

UNICORN 5.0

User Reference Manual





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Table of Contents

| 1. Introducing UNICORN | 9 |
|---|----|
| 1.1. About UNICORN | 10 |
| 1.2. About this manual | 14 |
| 1.3. About the UNICORN user documentation | 17 |
| 2. UNICORN concepts | 20 |
| 2.1. Concept definitions | 21 |
| 2.2. The UNICORN user interface | 24 |
| 2.2.1. UNICORN Manager | 25 |
| 2.2.2. The Method Editor module | 28 |
| 2.2.3. The System Control module | 32 |
| 2.2.4. The Evaluation module | 35 |
| 2.2.5. Search functions | 37 |
| 2.2.6. Help functions and manuals | 39 |
| 2.2.7. Snapshots | 41 |
| 2.3. Quick Start Guide | 44 |
| 3. General system operations | 45 |
| 3.1. Log on routines and log off routines | 46 |
| 3.2. How to create a new user | 50 |
| 3.3. How to assign user properties | 53 |
| 3.4. How to change your passwords and user attributes | 56 |
| 3.5. How to connect to the chromatography system | 58 |
| 3.6. How to back up and restore system data | |
| 3.7. How to set up a printer | 66 |
| 4. Files and folders in UNICORN | |
| 4.1. How to create folders | |
| 4.2. How to open and preview files | |
| 4.3. How to arrange and locate your files | |
| 4.4. How to copy, delete, rename and backup files and folders | 75 |
| 5. How to create a method | |
| 5.1. How to use the Method Wizard | 80 |
| 5.2. How to use the Method templates | 85 |
| 5.3. How to use Text instructions | 88 |
| 5.4. How to sign the method | 91 |
| 6. How to edit methods | 92 |
| 6.1. The Method Editor interface | 93 |

| 6.1.1. Method Editor module | 94 |
|---|-----|
| 6.1.2. Text Instructions editor | 95 |
| 6.2. Method blocks | 97 |
| 6.2.1. How to view method blocks | 98 |
| 6.2.2. How to call method blocks | 100 |
| 6.2.3. How to add method blocks | 101 |
| 6.2.4. How to delete method blocks | |
| 6.2.5. How to rename method blocks | 106 |
| 6.2.6. How to find, copy and move method blocks | |
| 6.2.7. How to import method blocks | |
| 6.3. Method instructions | 111 |
| 6.3.1. How to read method instructions | |
| 6.3.2. How to add method instructions | |
| 6.3.3. How to delete method instructions | |
| 6.3.4. How to change or move method instructions | |
| 6.4. How to use method variables | |
| 6.5. Run Setup | |
| 6.5.1. Overview of Run Setup | 123 |
| 6.5.2. The Variables tab | |
| 6.5.3. The Scouting tab | |
| 6.5.4. The Questions tab | |
| 6.5.5. The Gradient tab | |
| 6.5.6. The Notes tab | |
| 6.5.7. The Evaluation Procedures tab | |
| 6.5.8. The Reference Curves tab | 140 |
| 6.5.9. The Columns tab | |
| 6.5.10. The BufferPrep tab | |
| 6.5.11. The Method Information tab | 146 |
| 6.5.12. The Result Name tab | 147 |
| 6.5.13. The Frac-950 tab | 149 |
| 6.5.14. The Start Protocol tab | 151 |
| 6.5.15. How to export the values in the Run Setup | 153 |
| 6.6. How to use selected method instructions | |
| 6.6.1. Base instruction | 155 |
| 6.6.2. Instructions at the same breakpoint | 157 |
| 6.6.3 Block and method length | 158 |

| 6.6.4. Messages and Set_Marks | 160 |
|--|-----|
| 6.6.5. How to delay a method | 162 |
| 6.6.6. Linear flow rates | 163 |
| 6.6.7. Gradients and eluent concentrations | 164 |
| 6.7. Standard Watch conditions | 166 |
| 6.8. How to save or delete a method template | 171 |
| 6.9. How to print a method | 172 |
| 6.10. How to export a method | 174 |
| 7. Scouting | 175 |
| 7.1. How to set up a Scouting Scheme | 176 |
| 7.2. How to define different columns for scouting | 181 |
| 8. MethodQueues | 182 |
| 8.1. How to create a new MethodQueue | 183 |
| 8.2. How to edit a MethodQueue | 187 |
| 9. How to perform method runs | |
| 9.1. How to start a method run | 190 |
| 9.2. How to monitor a method run | 193 |
| 9.2.1. How to customize System Control panes | 194 |
| 9.2.2. The Run Data pane | |
| 9.2.3. The Curves pane | 199 |
| 9.2.4. The Flow Scheme pane | 204 |
| 9.2.5. The Logbook pane | 205 |
| 9.3. Manual system control | 207 |
| 9.3.1. The toolbar and status bar | |
| 9.3.2. Manual instructions | 212 |
| 9.3.3. Alarms and warnings | 215 |
| 9.4. How to perform a scouting run | 216 |
| 9.5. How to perform a MethodQueue run | 217 |
| 9.6. If the network connection fails | 219 |
| 10. How to view results | 220 |
| 10.1. How to open a result file | 221 |
| 10.2. How to use the File Navigator | 222 |
| 10.3. Basic presentation of chromatograms | 226 |
| 10.3.1. Introduction and temporary chromatograms | 227 |
| 10.3.2. The chromatogram window | 229 |
| 10.4. How to optimize the presentation of a chromatogram | |

| 10.4.1. How to make changes in the Chromatogram Layout dialog box | 235 |
|---|-----|
| 10.4.2. The Curve tab and Curve Names tab | 236 |
| 10.4.3. The Curve Style and Color tab | 238 |
| 10.4.4. How to change and fix the axes | 240 |
| 10.4.5. How to save and apply a layout | 242 |
| 10.4.6. How to show part of a curve | 244 |
| 10.4.7. How to change the size of Fraction, Injection and Logbook marks | |
| 10.5. How to print active chromatograms | 247 |
| 10.6. How to create and print reports | 249 |
| 10.6.1. How to create and print a customized report | 250 |
| 10.6.2. How to create and print a standard report | 264 |
| 10.6.3. How to edit an existing report format | 267 |
| 10.7. Run documentation | 270 |
| 11. How to edit results | 273 |
| 11.1. How to reduce noise and remove ghost peaks | 274 |
| 11.2. How to subtract a blank run curve | 275 |
| 11.3. How to add curves | 277 |
| 11.4. How to enter and edit text in the chromatogram | 278 |
| 11.5. How to pool fractions | 279 |
| 11.6. How to match protein activity to a curve | 285 |
| 11.7. How to rename chromatograms, curves and peak tables | 286 |
| 11.8. How to import and compare different runs | 287 |
| 11.8.1. How to use the Multifile Peak Compare wizard | 288 |
| 11.8.2. How to import and compare chromatograms | 302 |
| 11.8.3. How to import and compare curves | 305 |
| 11.8.4. How to stack and stretch curves | 312 |
| 11.8.5. How to produce a mirror image | 316 |
| 11.9. How to import and export results | 318 |
| 11.9.1. How to import results | 319 |
| 11.9.2. How to export results | 321 |
| 11.10. How to sign results electronically | 325 |
| 11.11. How to save results and exit the Evaluation module | 326 |
| 12. Evaluation | |
| 12.1. Peak integration | 328 |
| 12.1.1. Baseline calculation | 329 |
| 12.1.2. How to perform a peak integration | 330 |

| 12.1.3. How to optimize the baseline with a morphological algorithm | 336 |
|---|-----|
| 12.1.4. How to optimize the baseline with a classic algorithm | 340 |
| 12.1.5. How to edit the baseline manually | 348 |
| 12.1.6. How to edit the peaks | 351 |
| 12.1.7. How to integrate part of a curve and how to exclude or skim peaks | 358 |
| 12.1.8. Measurements | 363 |
| 12.2. Other evaluations | 365 |
| 12.2.1. Peak purity and peak identification | 366 |
| 12.2.2. How to find slope values | 369 |
| 12.2.3. How to simulate a peak fractionation | 372 |
| 12.2.4. How to create curves | 373 |
| 12.2.5. How to use the Fraction Histogram | 377 |
| 12.3. Automated evaluation procedures | 378 |
| 12.3.1. How to create a new procedure | 379 |
| 12.3.2. How to edit a procedure | 382 |
| 12.3.3. How to run a procedure | 384 |
| 12.3.4. How to rename and remove procedures | 387 |
| 13. The Analysis module | 388 |
| 13.1. General information about the module | 389 |
| 13.2. Quantitation overview | 393 |
| 13.2.1. General information about quantitation | 394 |
| 13.2.2. External standard quantitation | |
| 13.2.3. Internal standard quantitation | 400 |
| 13.2.4. Standard addition quantitation | 404 |
| 13.2.5. Recovery calculation | 406 |
| 13.2.6. General reliability factors for the quantitation techniques | 408 |
| 13.3. How to prepare for quantitation | 409 |
| 13.3.1. Preparations before quantitation | |
| 13.3.2. How to create a quantitation table | 411 |
| 13.3.3. How to edit and update a quantitation table | 422 |
| 13.4. How to quantitate the sample | 428 |
| 13.4.1. External and internal standard quantitation | |
| 13.4.2. Standard addition quantitation | |
| 13.4.3. How to calculate the recovery factor | |
| 13.5. Automated quantitation | 438 |
| 13.5.1. How to set up for automated quantitation | 439 |

| 13.5.2. How to perform automated quantitation | 441 |
|---|------------|
| 13.5.3. How to perform automated update | 442 |
| 13.6. How to measure molecular size | 448 |
| 13.6.1. Overview of molecular size determination | 449 |
| 13.6.2. How to determine the molecular size | 451 |
| 14. System settings | 458 |
| 14.1. General information about system settings | 459 |
| 14.2. Alarms | 461 |
| 14.3. Curves | 463 |
| 15. System maintenance and error reporting | 464 |
| 15.1. System maintenance functions | 465 |
| 15.2. How to generate problem reports | 469 |
| 15.2.1. How to generate a report from the UNICORN M | Лanager470 |
| 15.2.2. How to generate a report from the System Cont | rol473 |
| A. Troubleshooting | 476 |
| A.1. Logon | 477 |
| A.2. UNICORN access | 479 |
| A.3. Methods and method runs | 482 |
| A.4. Evaluation | 487 |
| A.5. ÄKTAdesign system specific problems | 488 |
| B. Evaluation functions and instructions | 489 |
| B.1. Smoothing algorithms | 490 |
| B.2. Baseline calculation theory | 493 |
| B.3. Peak table column components | 497 |
| B.4. Procedure instructions | 504 |
| C. Curve fit models and statistics | 520 |
| C.1. Curve fit models | 521 |
| C.2. Statistics | 525 |
| D. The Column list | 527 |
| D.1. How to edit the Column List | 528 |
| E. How to create and edit BufferPrep recipes | 536 |
| E.1. How to create a BufferPrep recipe | 537 |
| E.2. How to edit a BufferPrep recipe | 542 |
| F. Method examples | 545 |
| F.1. Simple equilibration | 546 |
| F.2. Equilibration with simple safeguard | 548 |

| F.3. Equilibration with extra safeguard | 549 |
|---|-----|
| F.4. Collection of absorbance peaks | |
| F.5. Collection of three absorbance peaks | 553 |
| F.6. Messages | 555 |
| F.7. Quality control procedure | 557 |

Introducing UNICORN 1

Introduction

This chapter contains:

- A general overview of the UNICORNTM system.
- Information about the user documentation for UNICORN and how to use it.

In this chapter

This chapter contains these sections.

| Topic | See |
|--------------------------------------|-----|
| About UNICORN | 1.1 |
| About this manual | 1.2 |
| About the UNICORN user documentation | 1.3 |

1.1 About UNICORN

Introduction

This section is a general overview of the UNICORN system.

What is UNICORN?

UNICORN is a complete package for control and supervision of chromatography systems. It consists of control software and a controller card for interfacing the controlling PC to the chromatography liquid handling module.

Liquid chromatography is used in separation processes, for analytical purposes or in the biochemical process industry.

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Operating environment

UNICORN runs on a PC under Microsoft® Windows® 2000 or Microsoft Windows XP. It is designed to run under English keyboard settings.

Note: Microsoft and Windows are registered trademarks of the Microsoft Corporation in the United States and/or in other countries.

Windows func-

Most Windows functions are also available in UNICORN, including

- cut and paste
- right-click short-cut menus

Note: Drag and drop is not available. File and folder handling in UNICORN also differs from the general Windows file manager standard.

Bar code reader

You can connect a bar code reader to the PC and use the reader to enter information instead of using the keyboard. This can be useful for example when entering information like batch IDs.

Compatible chromatography systems

UNICORN can be used with a number of systems including

- ÄKTATM design systems
- BioProcessTM systems

Note: All examples in this guide are based on an ÄKTAexplorer[™] 100 system that operates with the E100F400 strategy. If you use another system you may find that the descriptions and instructions do not match your system on every point. In that case you also need to refer to the user documentation for your specific chromatography system.

System networks

UNICORN can be installed on a stand-alone computer to control only a single, locally attached system. However, a stand-alone computer can control up to four separate systems. In a network installation each computer workstation can operate many systems regardless if they are locally connected or not. Each system can only be operated by one workstation at a time, but several may view the output data.



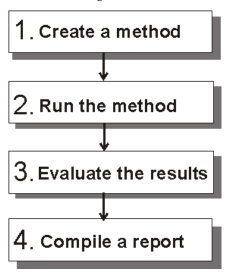
Software modules The UNICORN control software consists of four integrated modules:

| Module | Function |
|-----------------|---|
| UNICORN Manager | File handling and administration, e.g. definition of systems and user profile etc. |
| Method Editor | To create and edit methods for pre- programmed control of chromato- graphy systems. |
| System Control | To control and monitor the separation processes online, through method-based or manual control. |
| Evaluation | To evaluate and present stored results from separation processes. |

Note: All modules are active when the program is operating, and are not closed when they are minimized. A minimized System Control unit may control a process. All modules will normally open when the program is started. However, a user profile may be set up so that not all modules are available. Only the available modules will be displayed.

Work flow

The work flow in UNICORN can be divided into four distinct stages. Each stage is described in separate chapters in this manual. The flow chart below shows the work flow stages.



Help functions

An online help utility is included in the UNICORN software. The table below describes how to access the help utility.

| If you want to access | Then |
|-------------------------------|---|
| the general help utility. | open the Help menu in any of the software modules. |
| context-specific help topics. | click the Help button in the dialog box or press the F1 key on your keyboard. |
| the online manuals. | open the Help menu in any of the software modules and select Manuals . |

Security

The table below describes the main security functions in UNICORN:

| Feature | Function |
|-----------------|---|
| Access Security | Only authorized users can access UNICORN. Each user is assigned an access level, which defines the functions that the user is permitted to use. |

| Feature | Function |
|-----------------------|---|
| Connection Security | A running system can only be controlled from one connection. Systems may be locked with a password to prevent other, un-authorized users from changing parameters. |
| Data Security | Result files from an ongoing separation run can be saved automatically at preset intervals to minimize data loss if the system fails. The results are saved locally if the network communication fails. |
| Electronic Signatures | Method and result files can be signed electronically for enhanced security and accountability. |

1.2 About this manual

Introduction

This section is a general description of the manual, the contents and the pre-requisites for the examples and instructions that are presented in the User Reference Manual.

The purpose of the User Reference Manual

The purpose of the User Reference Manual is to present a comprehensive guide to the UNICORN system for a user either with previous experience of this system or from other, similar chromatography systems. The system is presented in detail, along with practical instructions of how to operate a model system.

Systems covered by this manual

This manual and the corresponding version of Getting Started with UNICORN covers the following systems:

- ÄKTAexplorer
- ÄKTApurifierTM
- ÄKTAFPLCTM
- ÄKTAbasicTM
- ÄKTApilotTM
- EttanTM LC



Note: Adapted versions of this manual are available for ÄKTAxpressTM, ÄKTA oligopilotTM and BioProcess systems.

The model system

For practical reasons the user documentation is based on a model system that consists of:

- ÄKTAexplorer 100
- Strategy **E100F400**
- Frac-950

Note: If you use another system you may find that the descriptions and instructions do not match your system on every point. In that case you also need to refer to the user documentation for your specific chromatography system.

Refer to other manuals

The User Reference Manual does not contain information about the installation procedure or network configuration. You will find this information in the Administration and Technical Manual.

Sometimes you may find it more convenient to refer to the Getting Started with UNICORN guide for a linear, step-by-step instruction how to perform a task.

Note: When you install the UNICORN software you choose which manuals you wish to install. You can also install the manuals after the program installation.

Document structure

The manual is divided into chapters. Each chapter starts with a brief overview that presents the contents and the headings for the sections that the chapter contains. Most sections begin with an introduction that summarizes the content. Some sections are divided into sub-sections.

A section is divided into blocks of information with separating lines. The blocks are identified by a label in the margin. This makes it easier for you to quickly scan a page to find the exact topic you are looking for.

Typographical representations

Menu commands, field names and other text items from the software are quoted exactly as they appear on the screen, in a bold typeface:

Example: Run Setup

Search paths are shown in a bold typeface with a separating colon between each level:

Example: View:Panes:Customize (i.e. the menu command Customize in the sub-menu Panes from the View-menu).

Text entries that UNICORN generates or that the user must type is represented by a monotype typeface:

Example: Connection change

Pre-requisites

The following pre-requisites must be fulfilled before you can use this manual the way it was intended:

- You need to have a general understanding of how your PC and Windows works. In most cases universal computer functions will not be explained.
- UNICORN must be installed and configured correctly on your computer.
- You need to understand the concepts of liquid chromatography. Terminology and functionalities will be explained only when they differ from normal practise.
- Before you try to operate a chromatography system based on the instructions in this manual you need to study and understand the safety information that is part of the system documentation.

1.3 **About the UNICORN user documentation**

Introduction

The user documentation for UNICORN is divided into three separate manuals. This section is an overview of the contents and the relationship between the manuals.

The manuals

The three manuals are:

- Getting Started with UNICORN
- UNICORN User Reference Manual (See 1.2 About this manual on page 14).
- UNICORN Administration and Technical Manual

User info about **Getting Started**

The questions and answers in the table below describe the features of the Getting Started manual.

| Question | Answer |
|---|--|
| Who should read Getting Started? | Users that are new to the UNICORN system and with limited experience from other chromatography systems. |
| What do I need before I start? | A basic knowledge of PC and Windows functions and an understanding of the concepts and terminology of liquid chromatography. |
| What are the contents of Getting Started? | Basic descriptions of UNICORN and its use, based on a model system. |
| How should I use Getting Started? | Read in front of your computer and test the instructions at the same time. |

User info about the User Reference Manual

The questions and answers in the table below describes the features of the User Reference Manual.

| Question | Answer |
|--|---|
| Who should read the User Reference Manual? | Users that are experienced with previous UNICORN system ver- sions. |
| | Users with vast experience from other chromatography systems. |

| Question | Answer |
|---|--|
| What do I need before I start? | Knowledge of PC and Windows functions and an understanding of the concepts and terminology of liquid chromatography. Preferably previous experience with UNICORN. |
| What are the contents of the User Reference Manual? | Detailed descriptions of UNICORN. Instructions on how to use the system, with suggested alternatives. Most instructions are based on a model system. |
| How should I use the User Reference Manual? | Depending on your previous experience you can either read whole chapters from the beginning to the end, or only selected sections for reference. |

User info about The Administration and Technical Manual

The questions and answers in the table below describes the features of the Administration and Technical Manual.

| Question | Answer |
|---|---|
| Who should read the Administration and Technical Manual? | System administrators. |
| What do I need before I start? | General knowledge of UNICORN. Knowledge of PC, Windows and general network administration functions. An understanding of the concepts and terminology of liquid chromatography. |
| What are the contents of the Administration and Technical Manual? | Detailed instructions of: How to install and maintain UNICORN in a network environment. How to create and administrate user profiles. Most instructions are based on a model system. |

| Question | Answer |
|---|---|
| How should I use the Administration and Technical Manual? | If you are an experienced administrator of previous UNICORN versions you can read selected sections for reference. If this is your first experience of UNICORN we recommend that you study the manual in detail. |

2 UNICORN concepts

Introduction

This chapter contains:

- Definitions and descriptions of some of the specific concepts that are presented in this manual and in other UNICORN manuals.
- An overview of the UNICORN user interface.
- A Quick Start Guide that can be used as a shortcut for experienced users that want to start right away.

Note: General concepts and common chromatography terminology are not explained here.

In this chapter

This chapter contains these sections.

| Topic | See |
|----------------------------|-----|
| Concept definitions | 2.1 |
| The UNICORN user interface | 2.2 |
| Quick Start Guide | 2.3 |

2.1 Concept definitions

Introduction

This chapter contains explanations and definitions of a number of UNICORN concepts that are used in this manual.

The concepts are organized in alphabetical order.

Alarms

Systems settings or method instructions specify acceptable limits for monitor signals during a separation run. An **Alarm** dialog box will be displayed on the screen if the monitored values exceed or fall below specified limits. The system will be paused.

Batch run

You can perform a **Batch run** of a number of result files in the **Evaluation** module. The files do not have to be open and the run operates in the background. The procedure is useful if you want to print a number of results with the same settings, or if you want to perform integration with the same parameter settings on many results.

BufferPrep

BufferPrep is a function to prepare a buffer of different pH and salt concentrations online from four stock solutions. This eliminates the need to manually prepare new buffers every time the pH needs to be changed.

Note: **BufferPrep** is only available for some ÄKTAdesign systems.

Chromatogram

A chromatogram is a collection of data represented by a number of curves that have been created during a separation run, including UV, conductivity, pH, fraction marks etc. The original raw data curves cannot be deleted or modified. They can be used as a basis for evaluation procedures and subsequent creation of new curves.

A chromatogram can also contain curves that have been created and saved during an evaluation session.

Curves

The monitor signals from the chromatography run are displayed graphically as curves.

Method

The program instructions for a chromatography run are defined in a **Method**. A Method can be divided into blocks that represent steps in the separation process. Each block consists of a series of instructions that request specific operations in the system.

MethodQueue

MethodQueues are used to link several methods together, on the same or on different systems.

Example: A **MethodQueue** can be set up to conduct a CIP study of a number of columns, through a controlled series of scouting runs.

Method Wizard

The **Method Wizard** is a user-friendly tool to create new methods. The **Wizard** takes the user step-by-step through the creation process.

Method Wizards are supplied with UNICORN installations for ÄKTAdesign systems.

Result files

UNICORN creates **Result files** when a method is run. The **Result files** contain:

• Run data from the monitors in the chromatography system.

Example: UV absorbance, flow rate, conductivity etc.

• Documentation from the run.

Example: Logbook entries, calibration settings, scouting parameters, text method etc.

• Saved results from evaluations of the run data.

Example: Peak integrations, simulated peak fractionations etc.

Scouting

Scouting is used to repeat a series of **Method runs** automatically with predetermined changes in the values for one or more **Variables**. A **Scouting Scheme** is defined as part of the method.

Scouting is used for optimizing chromatographic processes.

Strategy

Part of the UNICORN software is specific for the system that it is set up to operate. The system specific part is usually referred to as the **Strategy**. The **Strategy** defines available method and manual instructions, system settings, run data, curves and Method Wizards.



Note: The examples in this guide are generally based on the **E100F400** strategy.

Template

Templates are basic methods that can be used as a starting point for developing customized methods. The method variables in a suitable **Template** is adjusted to create a method for another application.

Variable

Instruction parameters and values at breakpoints in the **Method**may be defined as **Variables**. **Variables** makes it easy to adapt a method to a particular chromatography run.

- A framework **Method** with default parameters can be changed to create variants.
- A **Method** can be used in automatic **Method Scouting**, where one or more parameter **Variables** are changed systematically.

Warnings

Systems settings or method instructions specify acceptable limits for monitor signals during a separation run. A Warning dialog box may be displayed on the screen if a specified limit is exceeded. The system will still continue to run after a Warning.

2.2 The UNICORN user interface

Introduction

This section is an overview of the four UNICORN modules with descriptions of some of the elements of the user interface. The section also contains a description of the search functions in UNICORN.

Note: A user profile can be set up so that the user only has limited access to the modules described in this chapter. Only the available modules will open when the program is started.

In this section

This section contains these sub-sections.

| Topic | See |
|------------------------------|-------|
| UNICORN Manager | 2.2.1 |
| The Method Editor module | 2.2.2 |
| The System Control module | 2.2.3 |
| The Evaluation module | 2.2.4 |
| Search functions | 2.2.5 |
| Help functions and manuals | 2.2.6 |
| Snapshots | 2.2.7 |

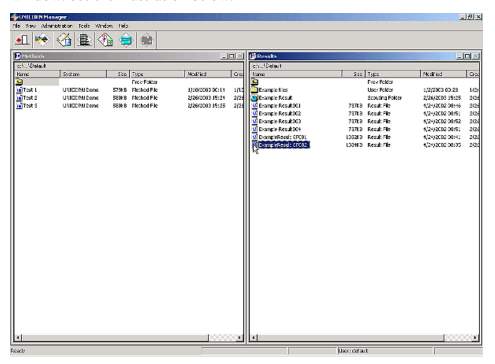
2.2.1 UNICORN Manager

Introduction

The **UNICORN Manager** is mainly used for file and folder administration.

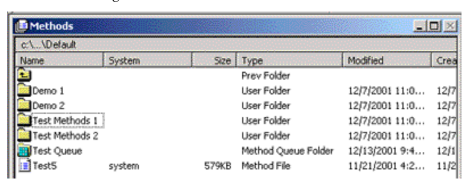
The UNICORN Manager windows

The module is divided into two windows, the **Methods** window and the **Results** window. See the illustration below:



The Methods window

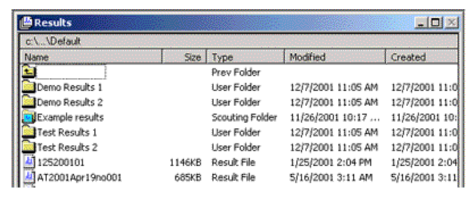
The **Methods** window contains all the saved methods, **MethodQueues** and all the folders containing methods that are available to the user. See the illustration below:



Note: The icons for **MethodQueue** folders are different from the regular folder icon.

The Results window

The **Results** window contains all the saved results and all the result folders.



Note: The icons for **Scouting** folders are different from the regular folder icon.

Toolbar icons in the UNICORN Manager

The table below describes the toolbar icons in the module.

| Icon | Function |
|-------------|---|
| ◆ [] | The Logon/Logoff icon is used to log on or log off the system. Note: The arrow in the Logoff icon points away from the door. |
| | The Instant Run icon immediately starts a run from a selected template or from a wizard. |
| | The New Method icon opens the Method Editor module and displays the New Method dialog box. |
| | The System Control icon activates the first connected System Control module and displays the Manual instruction dialog box. |
| | The Evaluation icon opens the Open Result dialog box. Select a result file and click OK to start the Evaluation module. |
| (| The MethodQueue icon opens the MethodQueue Editor. |
| (| The Existing MethodQueue icon opens the Running MethodQueue dialog box to display MethodQueues in progress. |

Limited access to the UNICORN Manager

Some user groups may be defined to have only a limited access to the **UNICORN Manager** functions. The available functions in the limited version are:

- Log off
- Change User Attributes
- Change Password
- Quit Program
- Help

There is also a **Cancel** button which minimizes the dialog box. The illustration below shows the limited access version of the **UNICORN Manager**.



Note: For more information about how to change passwords and user attributes please refer to 3.4 How to change your passwords and user attributes on page 56. For more information about how to log off and quit the program, please refer to 3.1 Log on routines and log off routines on page 46.

2.2.2 The Method Editor module

Introduction

The **Method Editor** module provides complete facilities for advanced editing of the methods.

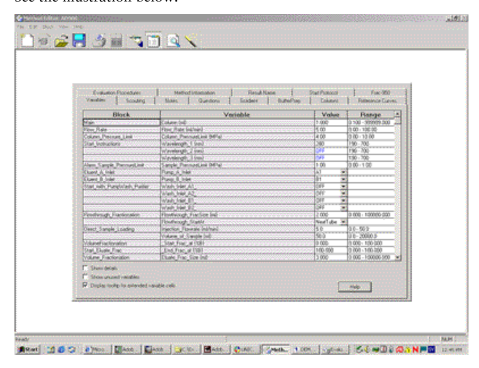
Two modes

The **Method Editor** interface operates in two modes:

- Run Setup
- Text Instructions

Run Setup

Run Setup is a dialog box with a number of tabs that define the method properties. See the illustration below:

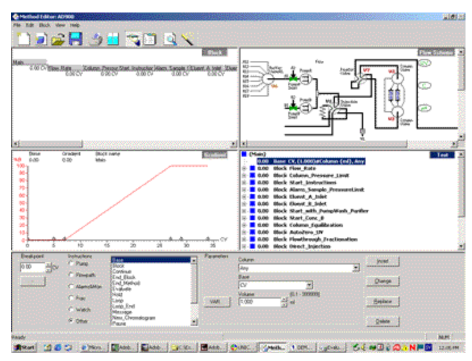


Text Instructions

Text Instructions are used for advanced editing. Up to five different display panes can be open at the same time:

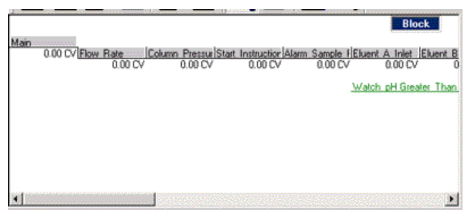
- The Block pane.
- The Flow Scheme pane.
- The **Gradient** pane.
- The **Text** pane.
- The **Instruction box** pane.

See the illustration below:



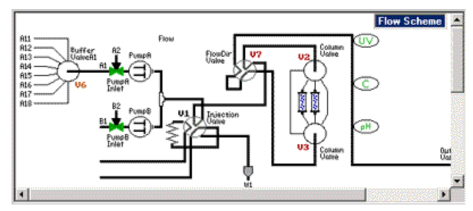
The Block pane

The **Block** pane contains a graphical representation of the method organized in blocks. See the illustration below:



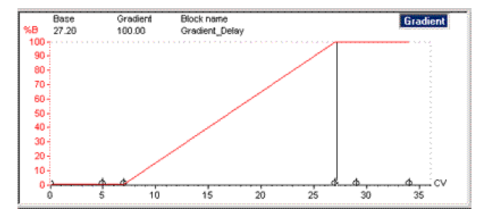
The Flow Scheme pane

The **Flow Scheme** pane displays the configuration of the system components. The pane is static and for information only. See the illustration below:



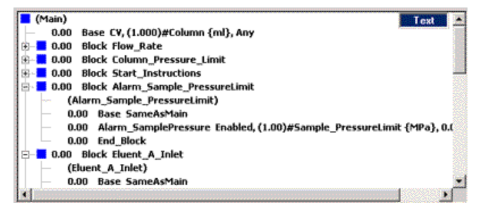
The Gradient pane

The **Gradient** pane provides a graphical overview of the block structure and eluent gradient in the current method. See the illustration below:



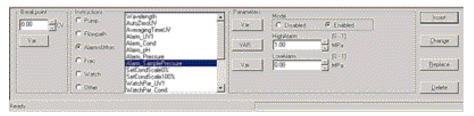
The Text pane

The **Text** pane displays the method as a list of text instructions. The instructions can be organized in blocks, denoted by blue square symbols. The blocks can be expanded to show the instructions within the block. See the illustration below:



The Instruction box pane

The Instruction box pane is used to enter, edit or delete instructions. See the illustration below:



Toolbar icons in the Method Editor

The table below describes the toolbar icons in the module.

| Icon | Function |
|----------|--|
| * | The New icon opens the New Method dialog box. The dialog box is used to create a new method. |
| | The New Block icon opens the New Block dialog box, which is used to add blocks to a method. |
| | The Open icon displays all available method files and method folders in the Open dialog box. |
| | The Save Method icon saves the edited method. |
| | The Print icon opens the Print dialog box. Select the method elements that you want to print. |
| | The Customise Panes icon opens the Customise Panes dialog box, which is used to select the panes that are open in Text Instructions mode. |
| * | The Text Instructions icon opens the Method Editor in Text Instructions mode. |
| H 00000 | The Run Setup icon opens the Method Editor in Run Setup mode. |
| | The Log Format icon opens the Log Format dialog box, which is used to display the accumulated time or volume for a method. |
| | The Method Wizard icon opens the Method Wizard , which is used to create new methods. |

2.2.3 The System Control module

Introduction

The **System Control** module is used to perform and monitor separation runs.

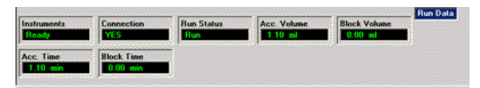
The System Control panes

The **System Control** module contains four different display panes that can be opened all at once or in any combination:

- The **Run Data** pane.
- The **Curves** pane.
- The Flow Scheme pane.
- The **Logbook** pane.

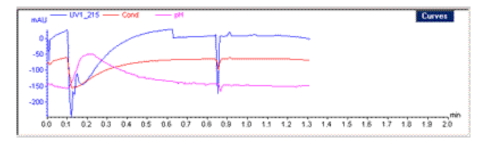
The Run Data pane

The **Run Data** pane displays the current values for the selected run parameters. The values are updated at regular intervals, which are defined in the system strategy. See the illustration below:



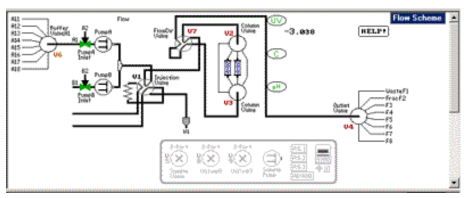
The Curves pane

The **Curves** pane displays monitor signal values graphically. See the illustration below:



The Flow Scheme pane

The **Flow Scheme** is a graphical representation of the chromatography system. During a run, the **Flow Scheme** displays open flow paths in color. Monitor signals can be displayed numerically. See the illustration below:



The Logbook pane

The **Logbook** pane displays all actions during a separation run, e.g. method start and end, base instruction, method instructions and manual instructions such as **Pause** or **Hold**. See the illustration below:



The Status bar

The **Status bar** in the bottom of the **System Control** module displays the current status of the separation run. See the illustration below:



The current system status is represented by the colored dot:

- A green dot represents a running system.
- A red dot represents a system in **Pause** state.
- A yellow dot represents a system in a **Hold** state.
- A white dot represents a system in an **End** state.

Toolbar icons in the System Control The table below describes the toolbar icons in the module:

| Icon | Function |
|------|---|
| Run | The Run icon opens the Run dialog box, which shows all available methods. If a method is loaded, Run Setup |
| | opens. |

| Icon | Function |
|----------|---|
| Hold | The Hold icon suspends execution of the method, while liquid is still pumped at the current flow rate and eluent concentration. |
| Pause | The function of the Pause icon depends on the strategy. The Pause icon suspends execution of the method and stops all pumps so that the system comes to a standstill. |
| Continue | The Continue icon resumes the execution of a paused or held method. |
| End | The End icon terminates the method execution and puts the system into an End state. |
| | The Customise Panes icon opens the Customise Panes dialog box, which is used to select the display panes that are open. |
| 11 0000 | The View Documentation icon opens the documentation pages. Run notes can be entered in the Notes page and settings can be changed. |
| Lite | The View Properties icon opens the Properties dialog box, which is used to control the data display in the System Control panes. |
| 1 | The Connect System icon is used to connect a system. |
| 11 | The Disconnect System icon is used to disconnect the system. |
| ₩ | The Take Control of the System icon is used to leave the view mode for the system and change into a control mode. |
| | The Leave Control of the System icon is used to leave the control mode for the system and change into a view mode. |

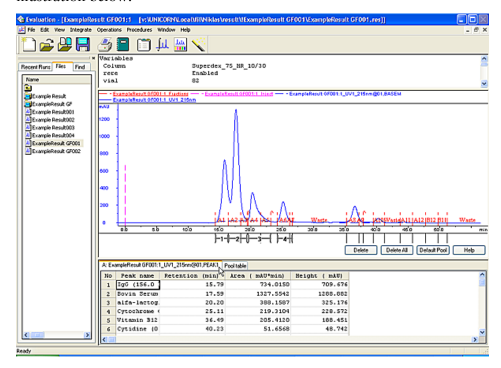
2.2.4 The Evaluation module

Introduction

The **Evaluation** module provides extensive facilities to present and to evaluate curve data.

The module window

Opened result files are displayed in the **Evaluation** module window. See the illustration below:



Toolbar icons in the Evaluation module The table below describes the toolbar icons in the module:

| Icon | Function |
|------|--|
| * | The New icon opens an empty chromatogram. |
| | The Open icon displays all available result files and result folders in the Open Result dialog box. |
| | The Open Curves to Compare icon opens the Open Curves to Compare dialog box, which is used to select and open curves for comparison. |
| | The Save icon saves the edited result file. |
| | The Print icon opens the Print Chromatograms dialog box. |

| Icon | Function | |
|-------|---|--|
| | The Report icon opens the Generate Report dialog box, which is used to select a report format. | |
| | The View Documentation icon opens the Documentation dialog box, which is used to view and edit the result documentation. | |
| Jul | The Peak Integrate icon opens the Integrate dialog box, which is used to select peaks to integrate in a modified peak table. | |
| Juli. | The Chromatogram Layout icon opens the Chromatogram Layout dialog box, which is used to select and format curves and display items in the chromatogram. | |
| | The Multifile Peak Compare icon opens the Multifile Peak Compare Wizard, which is used to compare peak data from different result files. | |

2.2.5 Search functions

Introduction

This section describes the general search functions that can be used to locate for example chromatograms, curves and text strings in UNICORN. These functions can be used in several program modules, dialog boxes and wizards.

Search the Folder list

The search will take place in the displayed folder only. To select another folder, click the **Browse** button and open the desired folder.

Search the Result list

- The search will take place in *all* result files within the selected folder as denoted by the asterisk (*). To select specific result file(s), click the **Browse** button and select the result file(s).
- You can use wildcard characters to search for chromatograms within result files with a specific name profile.
 - * represents any number of characters
 - ? represents any single character

Wildcard character examples:

iex will search files named "iex"

iex* will search all files with names that begin with "iex"

*iex will search all files with names that end with "iex"

?iex will search only 4-character names that end with iex

Search the Chromatogram list

The asterisk (*) indicates that all chromatograms within a result file will be selected. Click **Browse** to select one or several specific chromatograms.

Search the Curve name list

The UV curves are identified by number and sometimes wavelength. For example, UV1_280, UV2_280 and UV1_254 are all different curves. To search for all UV curves, select *UV* in the **Curve name** text field.

Searches for Sample ID

A **Sample ID** can be used as a search criteria if it has been defined as a variable. The **Sample ID** can be entered in searches for result files both in the **UNICORN Manager** and in the **Evaluation** module.

Find a text string

The **Find** command is used to search for text strings:



| Field | Description |
|-----------------------------|--|
| Find what | Type the text string you want to find. |
| Match whole word only | Select the check-box if you only want complete string matches, not partial matches. |
| Match case | Select the check-box if you only want matches which correspond according to upper-case and lower-case letters. |
| Search from top of document | Select the check-box to start the search from the top of the document, otherwise the search will start from the cursor position. |
| Direction | Choose whether to search upwards or downwards in the document. |

Commands

Use the commands below to find more occurrences of a text string after you have found the first one:

- Press F3 to search for the next occurrence of the string or right-click and choose
 Find next.
- Right-click and choose **Find previous** to search for a previous occurrence.

General information about searches

- The default setting is to search in all result files or chromatograms.
- User-entered search filters (to a maximum of 10) will be saved in the drop-down menus for both **Result** and **Chromatogram** selections. More than one string can be used as a search delimiter (insert ";" between strings), and search filters are automatically saved and stored within user profiles.
- Click **All** to return to the default setting to search in all result files or chromatograms.

Help functions and manuals 2.2.6

Introduction

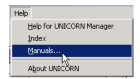
There are different ways to get help and instructions in the UNICORN application:

- From the **Help** menu in each module
- From the context-sensitive help in each dialog box
- By selecting the **Online Manual** from the **Help** menu
- By pressing the <F1> key
- By right-clicking an instruction in the **Method Editor** and selecting the **What's** This? menu item

The Help menu

- From the **Help** menu in each module you can access the **Help** file.
- From the Help menu of the UNICORN Manager module you can also access the installed manuals.

The illustration below shows the **Help** menu of the **UNICORN Manager** module:



The Help file

The table below describes how to open and use the Help file:

| Step | Action |
|------|---|
| 1 | Choose Help:Index . Result: The Help file is displayed |
| 2 | Type a word you want help on in the text box in the left pane. Result: The closest matches are displayed in the list. Select a match and click the Display button. Result: The associated help text is displayed in the right pane. |
| 3 | You can also click the Contents tab to view the contents of the Help file divided into sections. Click the plus signs to expand the tree structure. Click a topic to read the associated help text. |

Manuals

When UNICORN was installed, the administrator selected which manuals to install. Therefore the available manuals may be different on your system than in the illustration below.

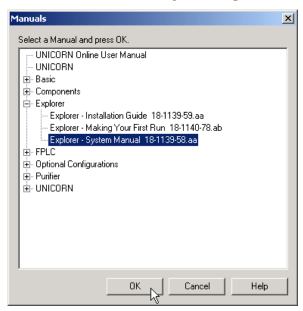
Note: Manuals can be added after the UNICORN installation. See the Administration and Technical manual for more information.

How to open a manual

To open a manual

• choose Help:Manuals in the UNICORN Manager module.

Result: The Manuals dialog box is opened.



• Select the manual and click the **OK** button.

Note: Some manuals are only available in PDF format.

Context-sensitive help

In each dialog box there is a **Help** button. If you press that button, either of the following will be displayed:

- A message box with relevant information, for example the dialog box options.
- The Help file, with relevant information displayed in the right pane.

2.2.7 Snapshots

Introduction

A **Snapshot** provides information about a method run at a certain point in time. It contains information about the values of all the variables at the selected point. Snapshot functionality is available in

- the **Method Editor**, where Snapshot instructions can be inserted in a method to be recorded during the method run.
- the **Evaluation** module, where you can take Snapshots from a result file using the Marker.
- the **System Control** module, where you can take Snapshots during a run using the Marker.

How to view recorded Snapshots

The table below describes how to view Snapshots which have been recorded during a method run using the **Snapshot** text instruction.

Note: How to insert the **Snapshot** text instruction in a method is described in **5.3** How to use Text instructions on page 90.

| Step | Action | |
|------|--|--|
| 1 | In the Evaluation module, | |
| | choose View:Documentation | |
| | or | |
| | • click the View Documentation icon. | |
| | Result: the Documentation dialog box is displayed. | |
| 2 | Select the Result Information tab. | |
| | • Select the Snapshots sub-tab. | |
| | <i>Result</i> : The recorded Snapshot information for a chromatogram is displayed in a list. | |
| 3 | You can | |
| | • select other chromatograms in the Select chromatogram dropdown box. | |
| | • select the Rows or Columns radio button to display each Snapshot as a row or a column. | |
| | • select the Time or Volume radio button depending on which quantity you want as a base. | |

| Step | Action | |
|------|---|--|
| 4 | To print the Snapshot information | |
| | click the Print button select the Snapshot check box in the Print dialog box. | |
| | • click OK . | |
| 5 | Click OK (or the Cancel button) to exit the Documentation dialog box. | |

How to take Snapshots in the Evaluation module

The table below describes how to take Snapshots in the **Evaluation** module:

| Step | Action | |
|------|--|--|
| | | |
| 1 | • Open a result file in the Evaluation module. | |
| | Right-click and select Marker in the menu. | |
| | Result: A vertical line indicating a certain point is displayed. | |
| 2 | Click the marker line and drag it to the desired point where you want to take a Snapshot. | |
| 3 | Right-click and select Snapshot in the menu. | |
| | Result: The Snapshot is displayed in the Snap Shot dialog box. | |
| | Snap Shot | |
| | Curve Retention Amplitude Unit | |
| | 01:125200101:1_UV1_280nm 29.41 11.69 mAU | |
| | 02: 125200101:1_UV2_250nm | |
| | 03: 125200101:1_UV3_0nm 29.41 0.00 mAU | |
| | 04: 125200101:1_Cond 29.41 0.47 m8/cm | |
| | 05: 125200101:1_Cond% 29.41 0.50 % | |
| | 06: 125200101:1_Conc 29.41 100.00 %B | |
| | 07:125200101:1_pH 29.41 5.89 | |
| | 09: 125200101:1_Flow 29.41 1.00 ml/min 10: 125200101:1_Temp 29.41 26.60 gC | |
| | 14:125200101:1_SampleFlow 29.41 0.00 ml/min | |
| | Save to File Print Close Help | |
| | | |
| 4 | • Click the Save to File button if you want to save the information as an Excel file (.xls) or a tabbed text file (.txt). | |
| | You can also copy the information to the clipboard: | |
| | - Click and drag the mouse in the table to select the information you want to copy. | |
| | - Press CTRL+C. | |
| | The information can now be pasted in a text editor. | |
| | Click the Print button if you want to print the information. | |
| | Click the Close button. | |

| Step | Action |
|------|---|
| 5 | Repeat steps 2 to 4 if you want to view more Snapshots. |

How to view Snapshots during a method run

The table below describes how to view Snapshots in the System Control module during a method run:

| Step | Action | |
|------|--|--|
| 1 | A method is running and the System Control is displayed: • Right-click in the Curves pane and select Marker in the menu. *Result: A vertical line is displayed. | |
| 2 | Click the marker line and drag it to the desired point where you want to take a Snapshot. | |
| 3 | Right-click in the Curves pane and select Snapshot in the menu. **Result: The Snapshot is displayed in the Snap Shot dialog box. Curve | |
| 4 | Click the Save to File button if you want to save the information as an Excel file (.xls) or a tabbed text file (.txt). You can also copy the information to the clipboard: Click and drag the mouse in the table to select the information you want to copy. Press CTRL+C. The information can now be pasted in a text editor. Click the Print button if you want to print the information. Click the Close button. | |
| 5 | Repeat steps 2 to 4 if you want to view more Snapshots. | |

2.3 Quick Start Guide

Introduction

This guide is intended for users who are fully familiar with the safety precautions and operating instructions that are described in all manuals, i.e. experienced users of previous versions of UNICORN. The instructions assume that all installations were made according to the instructions, that the model system is used and is connected.

Quick Start instructions

The table below describes the easiest way to create a method, run the system and generate a printed chromatogram. The procedure is based on an **Instant run**.

| Step | Action | |
|------|--|--|
| 1 | Click the Instant run icon in the UNICORN Manager module. | |
| | | |
| | Result: The Instant run dialog box opens. | |
| 2 | Select Wizard. | |
| | Select a system (if necessary). | |
| | Click the Run button. | |
| | Result: The Method Wizard opens in the System Control module. | |
| 3 | Go through all selections on the Method Wizard pages. Click the Next button to proceed from page to page. | |
| 4 | Click the Run button on the last page. | |
| | Result: The start protocol opens. | |
| 5 | Verify the method on the Variables page and change values as required. Click the Next button to proceed through several pages. | |
| 6 | Select Print_Chromatogram in the Evaluation procedures page. | |
| | Result: A printout will automatically be generated after the run. | |
| 7 | Click the Start button on the last page. | |
| | Result: The run starts. | |

3 **General system operations**

Introduction

This chapter describes how to start the program, assign user properties and set up the system.

Refer to the Administration and Technical Manual for installation and network configuration instructions.

In this chapter

This chapter contains these sections.

| Topic | See |
|--|-----|
| Log on and log off routines | 3.1 |
| How to create a new user | 3.2 |
| How to assign user properties | 3.3 |
| How to change your passwords and user attributes | 3.4 |
| How to connect to the chromatography system | 3.5 |
| How to back up and restore system data | 3.6 |
| How to set up a printer | 3.7 |

3.1 Log on routines and log off routines

Introduction

This section describes how to start and quit the UNICORN program, and how to log on and log off.

Username and password

Normally the system administrator defines the users and creates your first password. The program can also be set up so you can log on without a password.

Note: The first time after UNICORN has been installed, you may need to log on as a default user and create a user profile. This process is described in **3.2** How to create a new user on page 50.

How to start the program

Note: if UNICORN is already started by a previous user, proceed to How to log on.

There are two ways to start the program:

| If you start with | Then |
|--|--|
| a UNICORN icon on your desktop | double-click the icon |
| the Windows Start menu in Windows 2000 | locate the program under Programs:Unicorn and click the UNICORN logo |
| the Windows Start menu in Windows XP | locate the program under All programs:Unicorn and click the UNICORN logo |

How to log on

The table below describes how to log on to UNICORN.

| Step | Action |
|------|--|
| 1 | Select Tools:Logon in the UNICORN Manager module |
| | Or Click the Logon/Logoff icon in the UNICORN Manager module |
| | → |
| | Result: the Logon dialog box is displayed. |
| | <i>Note</i> : You do not have to perform this step if you start up UNICORN. When you start UNICORN the Logon dialog box is automatically displayed. |

| Step | Action |
|------|-------------------------------------|
| 2 | Select your username from the list. |
| 3 | Type your password (optional). |
| 4 | Click 0K . |

The four program modules

The program has four modules. When you start the program and log on you work in the **UNICORN Manager** module. UNICORN also automatically opens the **Method Editor**, the **System Control** and the **Evaluation** modules. These modules are minimized until you activate them. Up to four System Control module windows may open if UNICORN was set up to control more than one system at the installation.

Note: If the access rights are limited to only some modules, the other modules will not open.

Log off after you are finished

Always log off when you leave the computer to prevent others from accidentally changing or deleting your files, or disturbing your UNICORN runs. There are two ways to log off in the **UNICORN Manager**:

• Select Tools:Logoff

or

• Click the Logon/Logoff icon.



Note: In case your access to the **UNICORN Manager** is restricted you will still be able to log off.

Processes can run after log off

The process will continue even if you log off while a separation run is in progress. You can leave the process locked and set a password to protect it from interference. The table below describes how to log off and set a password for a running process.

| Step | Action |
|------|---|
| 1 | Select Tools:Logoff in the UNICORN Manager module. |
| | or |
| | Click the Logoff icon. |
| | Result: A confirmation box opens. |
| 2 | Click Yes to confirm that you want to log off. |
| | Result: The Leave Control of system dialog box opens. |
| 3 | Click the Locked radio button. |

| Step | Action |
|------|--|
| 4 | Type a password in the Password text box. |
| 5 | Click OK . |

Unlocked Log off

It is not recommended that you log off and leave a running system unlocked. This means that the run is in progress without a user that is responsible for the process.

Automated workstation lock or logout

The system administrator may set an automatic workstation lock or log off after a specified time for a user. If there are no keyboard entries or mouse movements within the time limit, the workstation will be locked or logged off.

Note: A locked workstation can be activated again only by the previous user if the regular log in password is entered. If another user wants to log on and use the workstation the previous user can be logged off without entering the correct password. The previous user's files will be closed and the new user will only have access to his own files. Automated logout will not happen while a **MethodQueue** or a **Scouting** scheme is operating.

How to log on and unlock the system

When you log on again after leaving the system locked with a process running or after an automated workstation lock, you will be asked to unlock the system.

| Step | Action |
|------|---|
| 1 | Log on to the system. Result: The System Unlock Confirmation dialog box opens. |
| 2 | Type your login password or the password that the system was locked with in the Password text box. |
| 3 | Click OK |

Note: You can connect in view mode only without providing the password.

Systems locked by other users

You can unlock a system that has been locked by another user if you have the correct password.

You may still be able to unlock a system even if you do not have the password. Any user with **Unlock locked systems** authorization can override another user's lock by entering his or her own logon password. However, it is recommended that this authorization is limited to only a few users.

How to quit UNICORN

UNICORN will still be open after you have logged off. To close the program you must log in again and quit UNICORN (you cannot quit the program if you are not logged in). The table below describes how to do this.

| Step | Action |
|------|---|
| 1 | Select the File:Quit Program menu command in the UNICORN Manager module. |
| | or |
| | • Click the close icon in the top right-hand corner of the program window. |
| | Result: A confirmation box opens. |
| 2 | Click Yes to confirm that you want to quit. |
| 3 | A Warning opens if you have any unsaved data in the Method Editor or Evaluation module. |
| | • Click Yes to continue to close the program. Your unsaved data will be lost when the program is closed. |
| | • Click No to return to the program and save your data. |
| 4 | The Leave Control of system dialog box opens. Select the locked or unlocked option as in the logoff procedure. |
| | <i>Note</i> : This step only happens when a system is connected. |
| 5 | Click OK. |

Note: Do not shut down Windows 2000/XP or turn off the computer if you quit UNICORN with a separation run in progress. If you are performing a **Scouting run** or a **MethodQueue run** you cannot quit the program at all.

In case your access to the **UNICORN Manager** is restricted you will still be able to quit the program.

3.2 How to create a new user

Introduction

This section describes how to create a new user and assign a home folder for the user's methods and results.

Default user

A default user is created when the system is installed. The default user has unrestricted access to all UNICORN functions. You log on with this profile when you access a newly installed system for the first time.

The table below describes how to log on as the default user.

| Step | Action |
|------|---|
| 1 | Select user default from the user drop-list. |
| 2 | Type password default if necessary. Note: The default user is the only user that is allowed to use the user name as password. |
| 3 | Click OK or press the Enter key. |



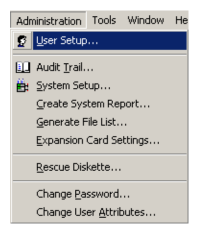
Note: We recommend that the default user is deleted when regular user profiles are created.

How to open User Setup

All user administration is performed in the **User Setup** dialog box in the **Main Menu** module. It is accessible only to authorized users (and the default user).

User Setup is found on the **Administration** menu.

• Choose Administration: User Setup.



The User Setup dialog box

The illustration below shows the User Setup dialog box.



How to create a new user

The table below describes how to create a new user.

| Step | Action |
|------|--|
| 1 | Click the New button in the User Setup dialog box. |
| | Result: The Create New User dialog box opens. |

| Step | Action |
|------|--|
| 2 | Enter a user name in the User name text box. |
| 3 | Enter the full name of the user in the Full name text box. |
| 4 | Enter the position of the user in the Position text box. |
| 5 | Select or create a Home folder: Select a Drive and a folder from the Name drop-list and proceed to step 9. or If you need to create a new home folder, proceed with step 6. |
| 6 | Click New. Result: the Create New Folder dialog box opens. |
| 7 | Select a Drive and type a folder name. |
| 8 | Click OK to create the folder and return to the Create New User dialog box. |
| 9 | Click OK . Result: The new user is created and added to the User Setup list. |
| 10 | Click Close. Or Click the New button and repeat steps 1 - 8 to create more users. |

Home folders

Each user must be assigned to a home folder. The **Default** folder can be used if you do not want to assign an individual home folder.

Note: If you create a home folder on the C: (local) drive it will not be accessible from other computers. If you select a network, make sure that is addressed by the same drive letter from all computers in the network.

3.3 How to assign user properties

Introduction

A user is assigned properties that define password rules, and the folders and chromatography systems that the user can access. This section describes how to assign properties.

How to open User properties

The user properties are defined in the **User Setup** dialog box in the **UNICORN Manager** module. The table below describes how to open **User Setup**.

| Step | Action |
|------|---|
| 1 | Select Administration:User Setup. |
| 2 | Select a user in the Users list. |
| 3 | Click the Edit button. Result: The User properties dialog box opens. |

The **User properties** dialog box is used to edit the user definition and assign properties for passwords, folder and system access, and available manual instructions.

How to edit the user definition

The table below describes how to edit the user definition in the **User properties** dialog box.

| Step | Action |
|------|--|
| 1 | Select the User item. |
| 2 | Select an access group from the Group drop-down box. <i>Note</i> : A pre-defined access group is assigned a certain level of access to UNICORN. |
| 3 | Select a folder from the Home folder drop-down box. |
| 4 | Click the check boxes to select Administrator Attributes. |
| 5 | Click OK to finalize or select another definition to edit. |

How to edit the user attributes

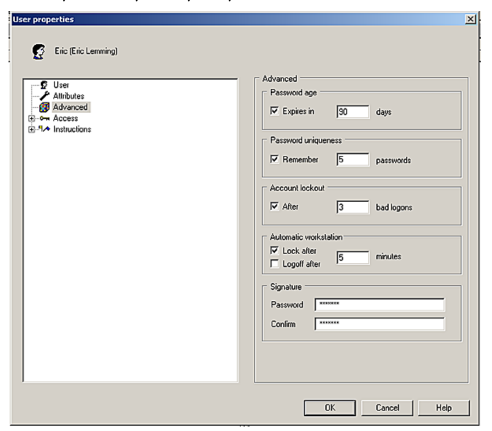
The table below describes how to edit the attributes in the **User Attributes** window pane.

| Step | Action |
|------|------------------------------------|
| 1 | Select the Attributes item. |

| Step | Action |
|------|---|
| 2 | Select applicable attribute items in the User Attributes pane: Use large toolbar icons Show unused variables Show variable details Default overwrite of baselines and peak tables Prompt for column before manual runs |
| 3 | Type which curve to display in the Quick view dialog box. |
| 4 | Select a size definition and type a value for the Fraction mark height. Select a size definition and type a value for the Injection mark height. Select a size definition and type a value for the Logbook mark height. |
| 5 | Click OK to finalize or select another definition to edit. |

The Advanced dialog page

The **Advanced** window pane is used to define password policies for the user. Normally this is only used by the system administrator.



Note: This dialog page is only available if a required password was selected when the software was installed.

How to define access to folders and systems

The **Access** dialog page is used to define the folders and systems that the user has access to. Click the check box for each selected folder and system.

Up to 20 folders can be set up to be shared. The user has access to all files and sub-folders in the selected folders. Only selected folders will be visible in the methods or results panels of the **UNICORN Manager** module.

Note: All users should have access to the **Failed** folder on each local station in a network installation. This will ensure that users can access results that were saved in the **Failed** folder in case of a network communication error.

How to define available manual instructions

The **Instructions** dialog page is used to define the manual instructions and system sounds that are available to the user as well as which monitors the user is allowed to calibrate. Click the check box for each selected instruction, sound or monitor.

Access groups

The level of access to UNICORN functions for each user is determined by the **Access group** that the user is assigned to. The access authorizations can be edited for each group, normally by the systems administrator. Refer to the Technical and Administration Manual if you need to edit an **Access group**.

Note: User access can be limited to only some UNICORN modules. If that is the case the unavailable modules will not be displayed. E.g. if the **UNICORN Manager** is unavailable you will only have access to a dialog box with the basic functions to change limited user attributes, passwords and to log out and quit the program.

3.4 How to change your passwords and user attributes

Introduction

Every user can change his or her passwords and some user attributes even if user administration is handled exclusively by the system administrator. The changes are made in the **UNICORN Manager**.

How to change passwords

The table below describes how to change your logon and signature passwords.

| Step | Action |
|------|---|
| 1 | Select Administration:Change Password. Result: The Change Password dialog box opens. |
| 2 | Type your old logon password in the Old text box. Note: Your passwords will only be shown as asterisks. |
| 3 | Type a new password in the New text box. |
| 4 | Repeat the new password exactly in the Confirm text box. |
| 5 | Repeat steps 2 to 4 in the Signature password section if necessary. |
| 6 | Click OK . |

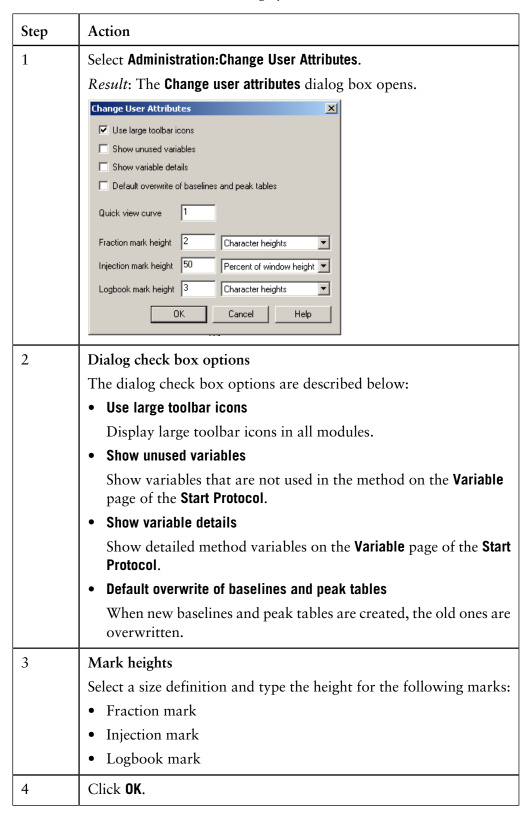
About passwords

The list below is a summary of facts and advice about UNICORN passwords:

- The system can be set up to operate without required passwords.
- The minimum number of password characters is set up at installation.
- Passwords can be any combination of letters and numbers.
- Passwords are case sensitive.
- Avoid using obvious passwords, e.g. your username, your telephone number, etc.
- The settings in the **User properties** determine the expiration for a password. Change passwords regularly even if your user profile is set up without password expiration.

How to change user attributes

The table below describes how to change your user attributes.



3.5 How to connect to the chromatography system

Introduction

A computer can have up to four chromatography systems connected at a time. This section describes how to connect to the systems, and different connection modes.

How to establish a connection

The table below describes how to connect a chromatography system that is locally connected to your computer.

| Step | Action |
|------|--|
| 1 | Open a System Control module. |
| | <i>Note</i> : Each UNICORN installation may have up to four System Control modules. The number of modules are selected when the software is installed. |
| 2 | Select the System:Connect menu command. |
| | or |
| | Click the Connect to system toolbar icon. |
| | |
| | Result: The System Connect dialog box opens. |
| 3 | Select the system you want to connect. |
| 4 | Click OK . |

Remote connections

Each computer workstation may have up to four chromatography systems connected locally. In a network installation you may connect a system that is physically connected to another computer, the local station. Your system is then a remote station.

The local station that is connected to the chromatography system must be logged on to the network and the UNICORN drivers must be running. However, the connection will work even if the UNICORN program is not running on the local station.

Network log on

Ensure that your workstation is logged on to the network before you start a chromatography system that is directly connected to the station. You can operate a local system without logging on to the network, but there are several disadvantages to this:

- Files stored on network drives are not accessible.
- Changes made to global files, e.g. user settings files, will apply only locally and will be lost the next time you log on to the network.
- Result files that are directed to a network drive will be stored in the **Failed** folder on the local station.

Connection modes

Several workstations can connect to a single chromatography system at the same time but only one workstation can be in control mode. The other connections are in view mode and the connected workstations can only monitor the system activity, but not issue any commands.

The system status is indicated on the status bar at the bottom of the **System Control** window. The table below describes the different connection modes, the corresponding status texts and some of the various actions you can take to change the connection mode.

| Connection mode | Status Text | Possible action to change connection mode |
|-----------------|------------------------|---|
| Not connected | Not Connected | Connect to a system. |
| Control mode. | Controlled By: default | Disconnect from or leave control of the system. (The system is controlled by you.) |
| View mode | Controlled By: Eric | No connection possible. (The system is controlled by another user.) |
| View mode | Locked By: Eric | Click the Connect to system icon and supply a password. (The system is locked by another user.) |
| View mode | System is available | Connect to the system. (The system has been left unlocked.) |

trol of a system

How to leave con- The table below describes how to leave control of a system so that it is available to be controlled by other users.

| Step | Action |
|------|---|
| 1 | Select System:Leave Control. |
| | or |
| | Click the Leave control of system icon. |
| | |
| | Result: The Leave Control of system dialog box opens. |
| 2 | Click the radio buttons to select to leave the system unlocked or locked. |
| 3 | Enter a password (if the system is to be locked). |
| 4 | Click OK. |

How to disconnect a system

The table below describes how to disconnect from a system.

| Step | Action |
|------|---|
| 1 | Select System:Disconnect. |
| | or |
| | Click the Disconnect from system icon. |
| | |
| 2 | Result: |
| | If the system is in view mode |
| | • the system is disconnected. |
| | If the system is in control mode |
| | • the Leave Control of System dialog box opens. |
| 3 | Select to leave the system locked or unlocked. |
| 4 | Click OK . |
| | Result: The system is disconnected. |

How to disconnect when quitting

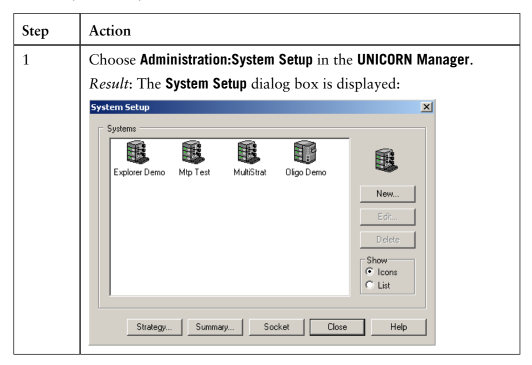
When you log off or quit from UNICORN you automatically disconnect all connected systems. A **Leave Control of System** dialog box will be opened for each system that was connected.

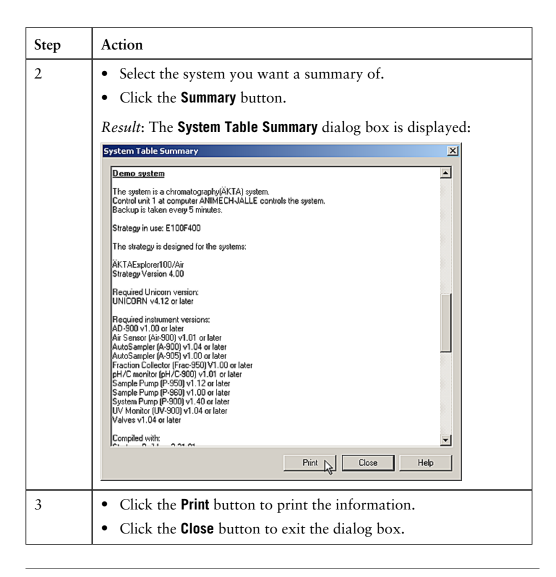
Note: If you disconnect from a system in control mode and re-connect to it, you may be connected in view mode. Another user may have taken control in the meantime.

How to view or print a system summary

You can view and print a total summary of a selected system from the **System Table Summary** dialog box.

The table below describes how to view and print an information summary of a selected systemthe systems:





3.6 How to back up and restore system data

Introduction

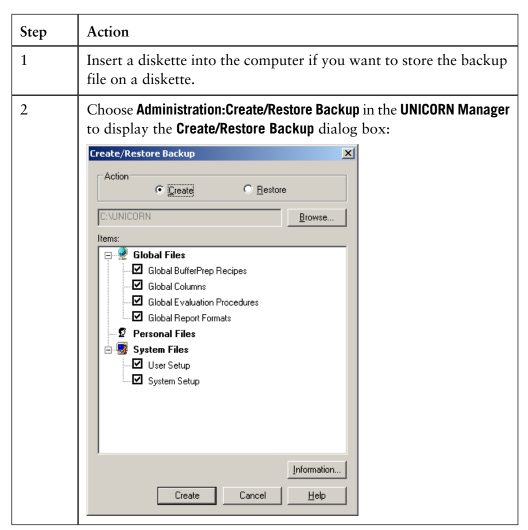
You can create a backup file with system information and store it on a diskette or another drive. The backup file will contain information about

- Global Files
- Personal Files
- System Files

Afterwards you can use the backup file to restore the system definitions in case they are corrupted.

How to create a backup file

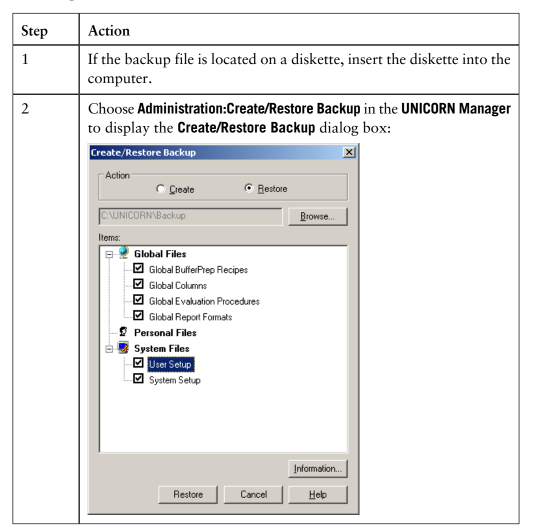
The table below describes how to create a backup file and store it for example on a rescue diskette:



| Step | Action |
|------|---|
| 3 | • In the Action field, make sure that the Create option is selected. |
| | • Click the Browse button to select where to store the backup file. |
| | . $Note$: Select A: \ to store the file on the diskette. |
| | • In the Items field, select which information to include on the backup file. |
| | • Click the Create button to create the backup file and store it in the selected location. |
| | <i>Note</i> : You can click the Information button to see which information files will be included in the backup file. |

How to restore the system data

The table below describes how to restore the system data from a backup file, located for example on a rescue diskette:



| Step | Action |
|------|--|
| 3 | In the Action field, select the Restore option. |
| | • Click the Browse button to select the folder where the backup file is located. <i>Note</i> : Select A:\ if the file is located on the diskette. |
| | • In the Items field, select which information to include from the backup file. |
| | Click the Restore button to restore the system definitions. |
| | <i>Note</i> : You can click the Information button to see which information files are included in the backup file. |

3.7 How to set up a printer

Introduction

UNICORN uses the default printer and printer settings that are installed on your computer. You can change your printer by changing the default Windows settings, but you can also set up a printer in UNICORN for the current working session.

How to set up a printer

The table below describes how to set up a printer in UNICORN.

| Step | Action |
|------|---|
| 1 | Select the File:Printer Setup menu command in the UNICORN Manager module. |
| | Result: The Print Setup dialog box opens. |
| 2 | Select a printer from the Name drop-down box. |
| 3 | Change all printer properties as necessary. |
| 4 | Click OK . |

Note: To save created reports electronically you can select to print the files in PDF-format. To be able to do this you must have a full version of AdobeTM AcrobatTM installed and select PDF Writer or DistillerTM in the **Printer Setup**.

4 Files and folders in UNICORN

Introduction

All UNICORN data is organized in files and folders. Files and folders are handled like in any other Windows application, with some exceptions. This chapter describes how to work with UNICORN files and folders, with the focus on the topics that are specific for UNICORN.

In this chapter

This chapter contains these sections.

| Topic | See |
|---|-----|
| How to create folders | 4.1 |
| How to open and preview files | 4.2 |
| How to arrange and locate your files | 4.3 |
| How to copy, delete, rename and back-up files and folders | 4.4 |

4.1 How to create folders

Introduction

This section describes how folders are organized in UNICORN and how to create a new user-specific folder for the user's methods and results.

UNICORN folders

The files and folders are displayed in the two UNICORN Manager module windows.

- All method files and corresponding folders are listed in the **Methods** window.
- The result files and folders are listed in the **Results** window.
- You can only see folders that you have access to.
- You can only see method files that are written for systems that you have access to.

How to create a user-specific folder

The table below describes how to create a user-specific folder.

| Step | Action |
|------|--|
| 1 | Select the window you want to create the folder in: Methods or Results . |
| | (Result: The window title bar is highlighted.) |
| 2 | Select File:New:Folder. |
| | or |
| | Right-click and select the New Folder shortcut. |
| | Result: The Create New Folder dialog box opens. |
| 3 | Type a name for the new folder. |
| 4 | Click OK. |

4.2 How to open and preview files

Introduction

This section describes how to open your saved method files and result files. You can also preview your result files to identify the correct file before you open it.

How to open a method file

You open a method file in the **UNICORN Manager** module. Click the file in the **Methods** window to select it and

• choose File:Open.

or

• right-click the file and choose **Open** from the short-cut menu.

or

• double-click the file.

Result: The file is opened for editing in the **Method Editor** module.

Note: A method file cannot be opened on two workstations simultaneously.

How to open a result file in UNICORN Manager

You can open a result file in the **UNICORN Manager** module. Click the file in the **Results** window to select it and

• choose File:Open.

or

• right-click the file and choose **Open** from the short-cut menu.

or

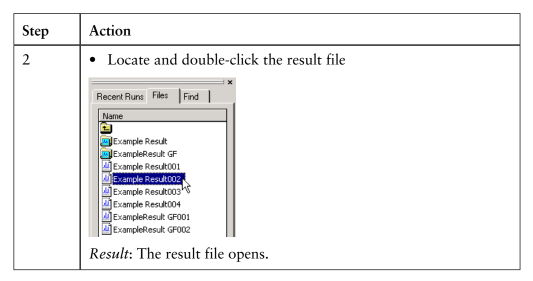
• double-click the file.

Result: The file is opened for editing in the **Evaluation** module.

How to open a result file in the Evaluation module

The table below describes how to open a result file from the **File Navigator** in the **Evaluation** module.

| Step | Action |
|------|-------------------------------|
| 1 | • Click the Files tab. |



Note: The **File Navigator** is opens by default in the **Evaluation** module. If it has been closed, select View:File Navigator in the Evaluation module.

Quick View

Quick View is a preview function for result files to make it easier to select the correct result file.

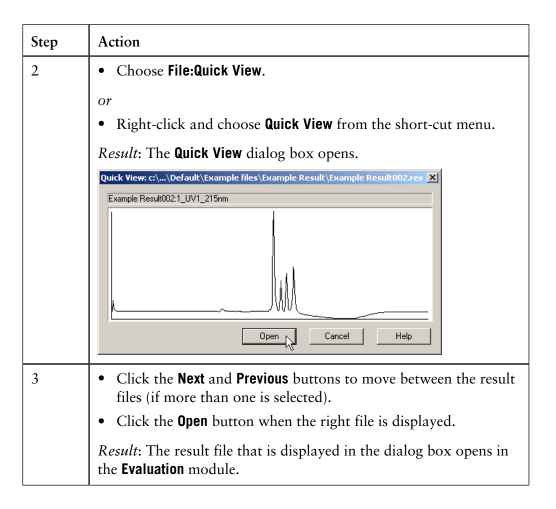
You can preview the first curve in the first chromatogram. You can also select to view another curve as default by selecting another curve number in your User **Attributes** settings, see 3.4 How to change user attributes on page 57.

Several files can be opened for comparison.

View

How to use Quick The table below describes how to preview result files in Quick View.

| Step | Action |
|------|---|
| 1 | Select one or more result files in the Result window of the UNICORN Manager . |



4.3 How to arrange and locate your files

Introduction

This section describes how to arrange the way the files are displayed in your UNICORN workspace and how to locate files through a search.

Different view modes

You can choose how the files and folders are displayed in the **UNICORN Manager** windows. The options are the standard Windows alternatives:

- Details
- List
- Large icons
- Small icons.

How to change the view mode

If you want to change the view you either:

• Select View and the option that you want,

or

 Right-click and select View and the option that you want from the shortcut menu.

Sort order in detailed view

The files can be sorted in a different order when a window is displayed in detailed view. The table below shows the options.

| Sorted by: | Order |
|------------|--|
| Name | Alphabetical order or reverse alphabetical order. |
| System | Alphabetical order or reverse alphabetical order (Method window only). |
| Size | Smallest or largest files first. |
| Туре | Alphabetical order of file extension type. |
| Modified | Most recently modified files first. |
| Created | Most recent creation dates first. |

How to change the sorting order

Select one of the methods below to change the sorting order:

• Select View:Sort and the option that you want,

or

• Right-click and select **Sort** and the option that you want from the short-cut menu.

or

• Click the column header for the option that you want to sort by (a second click on the same header will reverse the order).

Note: Only the currently active window is affected.

How to filter Method files

The files in the **Method** window can be filtered to show only methods for selected systems. You can also limit the displayed files by using standard Windows wildcard characters. The title bar of the **Method** window indicates if a filter has been activated.

The table below describes how to activate a filter.

| Step | Action |
|------|---|
| 1 | Select View:Filter. |
| | or |
| | Right-click and select Filter from the shortcut menu. |
| | Result: The Filter dialog box opens. |
| 2 | Click the check-boxes for the systems for which you want to show files. |
| 3 | Enter a file name specification (if necessary). |
| 4 | Click OK. |

How to find files

The table below describes how to perform a search for files.

| Step | Action |
|--------|---|
| Step 1 | Click either the Methods or Results window and: • Select the File:Find menu command. Or • Right-click and select Find from the shortcut menu. Result: The Find files dialog box opens. Result: The Find files dialog box opens. |
| 2 | Add search criteria to the dialog box, for example: • Type a name in the Name field. • Select a file type from the Type drop-down box. • Select if the search should include subfolders. • Select date limits in the Date drop-down boxes. • Type text strings to match Question or Answer texts. • Type a variable name and, if desired, a value. • Type a Batch ID. Note: You can search for a sample ID provided the sample ID is defined as a variable. |
| 3 | Click Find . Result: The search results are listed in the Found folders and files field. The search is limites to either methods or results and to the folder (including its subfolders) that is currently displayed. |
| 4 | Double-click a file in this list. Result: The dialog box is closed and the selected file is highlighted in the UNICORN Manager window. Note: If you click Close you will return to the UNICORN Manager window with no file highlighted regardless if you have selected one in the dialog box or not. |

4.4 How to copy, delete, rename and backup files and folders

Introduction

UNICORN has some file and folder handling functions that are slightly different from the general Windows functions. This section focuses on the differences.

Note: You need explicit authorization in your user profile to copy, move and delete files.

How to copy or move files and folders

There are some restrictions to how you can copy or move files and folders:

- Files and folders can only be copied or moved to folders that are specific to your user name.
- You can also copy files to and from the folders that you have access to on the network.
- Method files or folders cannot be copied to the **Results** window.
- Result files and folders cannot be copied to the **Methods** window.

If you copy a folder you will also at the same time copy all files and folders that it contains. The table below describes how to copy files and folders.

Note: Follow the same steps but select **Move** to move files and folders.

| Step | Action |
|------|--|
| 1 | Select one or more files and folders in either the Methods or Results window of the UNICORN Manager . |
| 2 | Select File:Copy. |
| | or |
| | Right-click and select Copy from the short-cut menu. |
| | Result: The Copy dialog box is opened. |
| 3 | Select a target folder or floppy disk drive. |
| 4 | Click OK . |

The function Copy to External

Use the function **Copy to External** when you need to copy files and folders outside of your own user folders. **Copy to External** should be used specifically when you need:

- to copy a method to another system (the method can then be connected to the appropriate system),
- to copy to a floppy disk drive. (The files are automatically compressed into a zip-file. The file will also automatically be spanned across several disks if necessary.)

How to Copy to External

The table below describes how to use the function **Copy to External**.

| Step | Action |
|------|--|
| 1 | Select the file you want to copy. |
| 2 | Select File:Copy to External. |
| | or |
| | Right-click and select Copy to External from the shortcut menu. |
| | Result: the Copy to External dialog box opens. |
| 3 | Select the destination drive and folder. |
| 4 | Click the Save button. |

The function Copy from Extern-

The function **Copy from External** can be used to import files and folders:

- If the files were saved using the function **Copy to External** they will automatically be decompressed.
- Copied method files must be connected to the same type of system they originally were created for. This is part of the **Copy from External** procedure.
- Method files that have been copied in and connected are displayed in the designated folder in the **Methods** window.

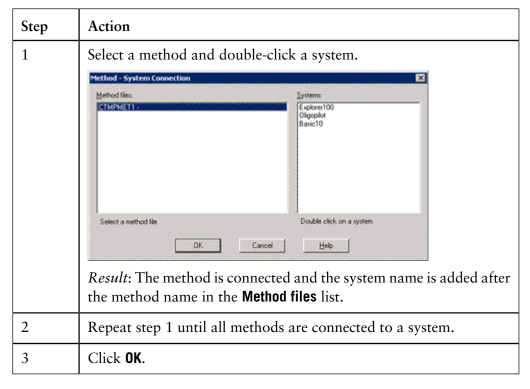
How to use Copy from External

The table below describes how to use the function **Copy from External**.

| Step | Action |
|------|--|
| 1 | Select a destination folder in the Methods or the Results window. |
| 2 | Select File:Copy from External. |
| | or |
| | Right-click and select Copy from External. |
| | Note: Do not select a file icon. |
| | Result: The Copy from External dialog box opens. |
| 3 | Select the files you want to copy. |
| 4 | Click Save. |
| | Result: |
| | • Result files are copied into the designated folder in the Results window. |
| | • If method files were selected, the Method-System Connection dialog box opens. |

How to connect a method to a system

The table below describes how to connect a method to a system.



How to rename files and folders

The table below describes how to rename files and folders in the **Methods** or **Results** windows in the **UNICORN Manager** module.

| Step | Action |
|------|--|
| 1 | Select the item that you want to rename. |
| 2 | Select File:Rename. |
| | or |
| | Right-click and select Rename from the shortcut menu. |
| | Result: The Rename dialog box opens. |
| 3 | Type a new name. |
| 4 | Click OK. |

How to delete files and folders

The table below describes how to delete files and folders in the **Methods** or **Results** windows in the **UNICORN Manager** module.

Note: Home folders cannot be deleted this way.

| Step | Action |
|------|--|
| 1 | Select the item that you want to delete. |

- 4 Files and folders in UNICORN
- 4.4 How to copy, delete, rename and backup files and folders

| Step | Action |
|------|--|
| 2 | Select File:Delete. |
| | Right-click and select Delete from the shortcut menu. |
| | • Press the Delete key. |
| 3 | Confirm the delete action in the confirmation dialog box |

Backup security

Backup copies should be taken regularly to avoid data loss in the event of hard disk failure or accidental deletion. You can use the function **Copy to External** to save your files on the network server.

Note: Amersham Biosciences cannot accept responsibility for the replacement of method programs that were lost as a result of computer failure or other incidents.

5 How to create a method

Introduction

Chromatography runs are programmed as Methods in UNICORN. Before you can proceed with a chromatography run you need either to use an existing method or create a new method. This chapter describes how to create new methods. It also contains instructions for signing a method.

In this chapter

This chapter contains these sections:

| Topic | See |
|---------------------------------|-----|
| How to use the Method Wizard | 5.1 |
| How to use the Method templates | 5.2 |
| How to use Text instructions | 5.3 |
| How to sign the method | 5.4 |

5.1 How to use the Method Wizard

Introduction

This section describes how to use a **Method Wizard** to create a new method. For most purposes customized methods can be created simply by setting appropriate values for the method variables.

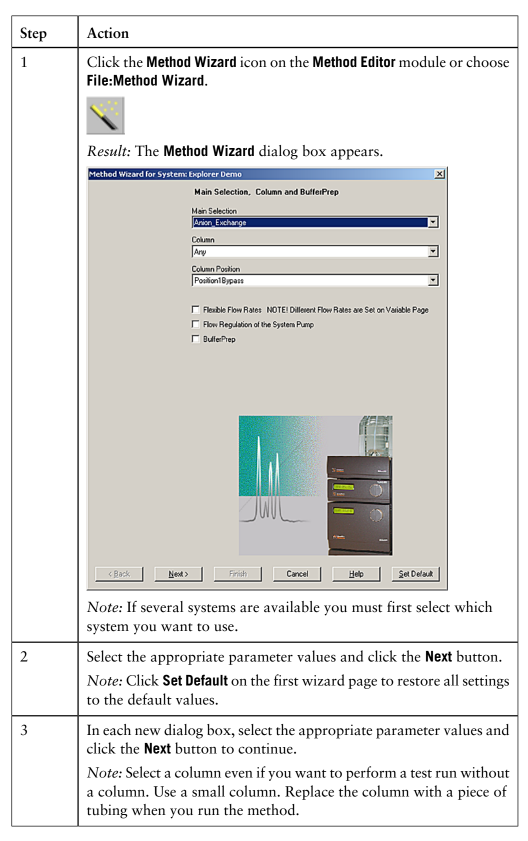
Note: Each method is written for a specific strategy. The function of the method cannot be guaranteed on systems having other strategies.

Are wizards always available?

Method Wizards are available for some **ÄKTAdesign** systems delivered with standard strategies. **Method Wizards** are not available for process systems.

How to create a new method

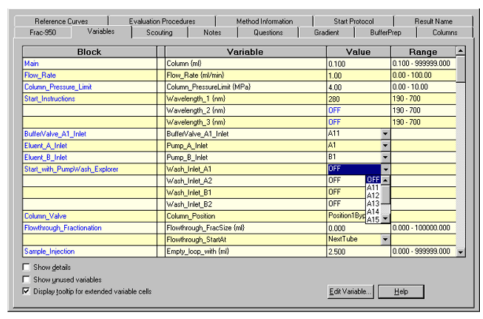
The table below describes how to create a method with the **Method Wizard**.



| Step | Action |
|------|--|
| 4 | Click the Finish button in the last dialog box. |
| | Result: The Run Setup opens. |

The Run Setup

The **Run Setup** consists of a number of tabs. Click on the appropriate tab at the top to select it.



The Variables tab

The method is represented by a number of blocks on the **Variables** tab. The blocks are typical steps in a chromatographic run.

Each block contains a number of **Method Variables** with suitable default values that can be changed to suit your application. Only the most commonly used variables are initially shown on the page. Click the **Show details** check box to display all variables in the method.

Default values for columns

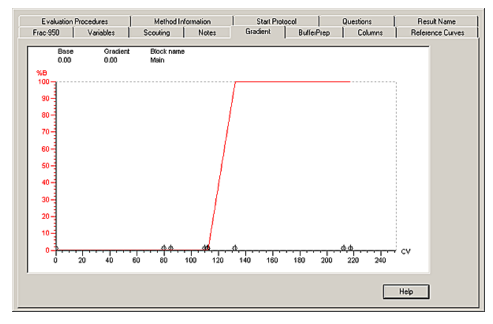
When you select a column, default values will be set for several parameters including the following:

- the correct column volume
- the recommended flow rate
- the correct pressure limit.

Note: If you exceed the recommended values for the selected column you will receive a warning when you save your method.



The **Gradient** tab shows the method graphically:



The length of each block is marked at the bottom of the graph.

- Click the X-axis to view the method in time, volume or column volumes.
- Drag the Y-axis to display a marker in the gradient. The **Base** value, **Gradient** value and **Block name** at the current marker position will be displayed in the upper part of the graph.

How to save the new method

A new method created from a **Wizard** is untitled, and must be saved under a method name before it can be used. The table below describes how to save a new method.

| Step | Action |
|------|---|
| 1 | Click the Save Method toolbar or choose File:Save. |
| 2 | If required, save the method in a folder other than the default home folder. |
| | • Enter a Method name for the method. The total path can be up to 256 characters long. The method name must be unique for the chosen system within the folder. |
| 3 | • If you have more than one system connected to the computer, choose the System for which the method is intended. The method can be run on any system that uses the same strategy. Remember that different systems may have different configurations and control capabilities. |
| | Choose the Technique for which the method was written. |

| Step | Action |
|------|--|
| 4 | Click OK . |
| | <i>Result:</i> The method is saved, but remains open in the Method Editor , so that you can continue editing if you wish. |

Note: You might want to sign your method. If you do so, you can choose to lock the method so that nobody will be able to change the method. See **5.4 How to sign the method** on page 91 for further instructions.

5.2 How to use the Method templates

Introduction

This section describes how to create methods based on an existing template.

Note: A custom system, for example a process system, requires that the users create their own templates by saving methods as templates. Each method is written for a specific strategy. The function of the method cannot be guaranteed on systems having other strategies.

How to create a new method

The table below describes how to create a method from the **UNICORN Manager** module.

Note: The **New Method** dialog box is also accessible from the **Method Editor** module using the same commands.

| Step | Action |
|------|---|
| 1 | Choose the File:New:Method menu command |
| | or |
| | • click the New Method icon. |
| | |
| | or |
| | • right-click in the Methods window and select New:Method from the shortcut menu. |
| | <i>Result</i> : The New Method dialog box opens in the Method Editor module. |
| 2 | • Select the system for which you want to create the method in the For system drop-down list. |
| | • Select Template in the Use field. |
| | • Select a chromatographic technique from the Technique dropdown list. |
| | Select a method template from the Template list. |
| | • Select a column from the For column list and click OK . |
| | <i>Result:</i> The method template will be opened as an untitled method in the Run Setup in the Method Editor . |
| | Note: If Any is selected in the For column list, you can use any column but must enter the column volume in the method on the Variables tab. It is recommended that a specific column is selected. |

Note: Only columns for the selected technique are displayed. If Any is selected as technique, all columns are displayed. Right-click in the textbox to open a list of the column categories to limit the number of displayed columns. If you type the beginning of a column name in the textbox UNICORN will automatically complete the column name.

If you do not find your specific column it can be added to the list. The column value, recommended flow rate, pressure limit and averaging time for the selected column will be automatically copied into the method, thus reducing the need to edit the method.

Method notes

Click the **Notes** and then the **Method Notes** tabs in the **Run Setup**. The notes describe important information about the template and how the system should be connected so that the method will work correctly.

Note: If your system does not correspond to the description on the **Method Notes** tab, either:

• rearrange the valves and tubing connections in accordance with the method notes description

or

• edit the method instructions in accordance with your system setup.

How to save the new method

A new method created from a method template is untitled, and must be saved under a method name before it can be used.

The table below describes how to save a new method.

| Step | Action |
|------|---|
| 1 | Click the Save Method toolbar icon or choose File:Save. |
| 2 | • If required, save the method in a folder other than the default home folder. |
| | • Enter a Method name for the method. The total path can be up to 256 characters long. The method name must be unique for the chosen system within the folder. |
| 3 | • If you have more than one system connected to the computer, choose the System for which the method is intended. The method can be run on any system that uses the same strategy. Remember that different systems may have different configurations and control capabilities. |
| | Choose the Technique for which the method was written. |
| | • Click OK . |
| | Result: The method is saved, but remains open in the Method Editor , so that you can continue editing if you wish. |

Note: You might want to sign your method. If you do so, you can choose to lock the method so that nobody else will be able to change the method. See 5.4 How to sign the method on page 91 for further instructions.

5.3 How to use Text instructions

Introduction

You can use the **Text Instructions** editor in the **Method Editor** to build your method step by step. You can also use the editor to modify instructions in methods created by wizards or based on templates.

Advanced editing facilities are available when you work directly in the **Text Instructions** editor. This section is a very brief description of this process. See 6 How to edit methods on page 92 for detailed instructions.

Note: Each method is written for a specific strategy. The function of the method cannot be guaranteed on systems having other strategies.

When do I use Text Instructions?

Use **Text Instructions** when you want:

- to change selected instructions in the method, for example the outlet valve position
- to add blocks or instructions, for example **Watch** instructions
- to change method instructions to adapt to non-standard system configurations
- to create new methods for applications not covered by the supplied templates or wizards.

How to edit Text Instructions

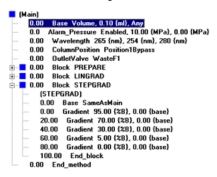
Open the **Text Instructions** editor by following the steps in the table below.

| Step | Action | |
|------|---|--|
| 1 | Select the Method Editor module and click the Text Instructions icon. | |
| | | |
| 2 | Click the Customise Panes icon and select Text and Instruction Box. | |
| | | |
| | Click 0K . | |
| 3 | Select instructions in the Instruction box in the lower part of the Method Editor , and use the Insert , Change , Replace or Delete buttor All text entries are shown in the Text pane. Applicable variables cabe edited for each selection. | |
| | The illustration below shows the Instruction box : | |
| | Part Part | |

Instructions can be organized in blocks

Individual text instructions can be grouped in blocks of instructions (marked by blue square symbols) for a specific functional use, e.g. to load a sample, to equilibrate a column etc. A block may contain other blocks or individual instructions.

This is an example of text instructions in the **Text** pane:



How to save the new method

A new method is untitled, and must be saved under a method name before it can be used.

The table below describes how to save a new method.

| Step | Action |
|------|---|
| 1 | Click the Save Method toolbar or choose File:Save. |
| 2 | If required, save the method in a folder other than the default home folder. |
| | • Enter a Method name for the method. The total path can be up to 256 characters long. The method name must be unique for the chosen system within the folder. |
| 3 | • If you have more than one system connected to the computer, choose the System for which the method is intended. The method can be run on any system that uses the same strategy. Remember that different systems may have different configurations and control capabilities. |
| | Choose the Technique for which the method was written. |
| | Click 0K . |
| | Result: The method is saved, but remains open in the Method Editor , so that you can continue editing if you wish. |

Note: You might want to sign your method. If you do so, you can choose to lock the method so that nobody will be able to change the method. See 5.4 How to sign the method on page 91 for further instructions.

How to display descriptions of instructions

A dedicated strategy is available for each system in the **ÄKTAdesign** platform. Although the majority of the instructions are general, some of them differ slightly between the individual strategies.

The list below describes two ways to display descriptions of the instructions in your particular strategy:

• Select the instruction in the **Instruction Box** of the **Method Editor** and press **<F1>**

or

 Right-click the instruction in the Text pane and choose the menu option What's This?

How to print descriptions of instructions

The table below describes how to print descriptions of the instructions in your particular strategy:

| Step | Action |
|------|---|
| 1 | Select File:Print in the Method Editor. |
| 2 | Select the Instruction set option to print the full set of instructions. Click OK. |

How to add a Snapshot

The **Snapshot** instruction can be used to record the curve values at a specific point in the method run. For example, a snapshot can be inserted to record the curve values immediately before an injection. The values are recorded in the result file and can be viewed in the **Snapshots** tab of the **Documentation** dialog box (See 10.7 **Run documentation** on page 270). Up to 500 snapshots can be recorded in each result file. The table below describes how to add a snapshot instruction to a method:

| Step | Action |
|------|---|
| 1 | • In the Text pane, select the instruction immediately before the position where you want to insert the Snapshot instruction. |
| 2 | Select Other in the Instructions field of the Instructions box. Select Snapshot in the instructions list. |
| 3 | Type a name in the Name text box in the Parameters field. • Click the Insert button. |

Note: Snapshots can also be taken in the **System Control** and **Evaluation** modules. However, these snapshots will only record the data for a specific moment. For more information about the **Snapshot** function see **2.2.7 Snapshots** on page 41.

5.4 How to sign the method

Instruction

If you sign the method, you can choose to lock it so that nobody will be able to change it.

The table below describes how to sign the method.

| Step | Action |
|------|---|
| 1 | Choose File:Sign Method in the Method Editor. |
| | Result: The Sign the Method dialog box is displayed. |
| 2 | Click the Signing tab and do the following: |
| | • Select a user in the User drop-down list box. In most instances, you will want to use the current user shown on the list. |
| | • In the Meaning field, provide a short text description explaining the meaning behind the signature (for example "Method now fully tested and approved"). |
| | • Type your signature password in the Password field. If desired, select the Lock box to lock the method permanently from further changes by other users. |
| | • If needed, view a list of all signatures associated with the current method on the View Signatures tab. |
| | Click OK on either the Signing or View Signatures tab. |

6 How to edit methods

Introduction

This chapter describes the complete facilities for editing methods in UNICORN. For many applications, suitable methods can be created by changing the default values in one of the wizard-generated methods supplied with UNICORN.

Use the more advanced editing facilities described here when you want

- to change selected instructions in the method, for example, change the outlet valve position
- to add blocks and instructions
- to change method instructions to adapt to non-standard system configurations.

In this chapter

This chapter contains these sections:

| Торіс | See |
|---|------|
| The Method Editor interface | 6.1 |
| Method blocks | 6.2 |
| Method instructions | 6.3 |
| How to use method variables | 6.4 |
| Run Setup | |
| How to use selected method instructions | 6.6 |
| Standard Watch conditions | 6.7 |
| How to save or delete a method template | |
| How to print a method | |
| How to export a method | 6.10 |

The Method Editor interface 6.1

Introduction

This section contains a general description of the Method Editor user interface and the editing operations that can be performed in the different parts of the module.

In this section

This section contains these topics:

| Topic | |
|--------------------------|-------|
| The Method Editor module | 6.1.1 |
| Text Instructions editor | |

6.1.1 Method Editor module

Two modes

The **Method Editor** interface operates in two modes:

- **Text Instructions** editor for entering and editing method instructions (see **6.1.2** Text Instructions editor on page 95)
- Run Setup for defining method properties (see 6.5 Run Setup on page 122).

How to open the Method Editor dialog boxes

The table below describes how to open the dialog boxes in the **Method Editor**:

| If you want to open | then |
|-------------------------------------|--|
| the Text Instructions editor | click the Text Instructions icon. |
| | choose View:Text Instructions. |
| the Run Setup | click the Run Setup icon. |
| | |
| | or |
| | choose View:Run Setup. |
| the Log Format | click the Log Format icon. |
| | |
| | or |
| | choose View:Log Format. |
| the Method Wizard | click the Method Wizard icon. |
| | |
| | or |
| | choose File:Method Wizard. |

Text Instructions editor 6.1.2

How to select panes to be dis-played You have a choice of four panes that can be open together with the Instruction box in the **Text Instructions** editor, all at once or one at a time.

Follow the steps in this table to select the panes to be displayed:

| Step | Action |
|------|---|
| 1 | In the Method Editor, choose View:Text Instructions |
| | or |
| | • click the Text Instructions icon. |
| | |
| 2 | Choose View:Panes:Customize (or select additional panes here) |
| | or |
| | click the Customize Panes icon. |
| | |
| 3 | Select panes |
| | Select panes in the dialog box and click the OK button. |
| | Customize Panes X |
| | ✓ Text ✓ Flow scheme |
| | ✓Instruction box ✓Block |
| | ⊈ Gradient |
| | OK Cancel <u>H</u> elp |
| | Deselect panes |
| | Deselect panes in the Customize Panes dialog box and click the OK button. |
| | or |
| | • right-click a window and select Hide . |

Method editing operations performed in the different panes

This table shows the method editing operations that can be performed in the different panes:

| The pane | Is used | See section |
|-----------------|---|--|
| Text | to display instructions to display and hide block instructions. to select current instruction. to edit instructions to cut, copy and paste instructions. to move instructions within a breakpoint. | 6.2.1 How to view method blocks on page 98 6.3 Method instructions on page 111 |
| Flow scheme | • for information only. This window is not updated according to system status and changes in the method. | 9.2.4 The Flow Scheme pane on page 204 |
| Instruction box | to specify break-points, instructions, parameters and variables. to insert, change and delete instructions. | 6.3.2 How to add method instructions on page 113 |
| Block | to select or display blocks. | 6.2 Method blocks on page 97 |
| Gradient | to display block duration and eluent gradient throughout the method. | 6.5.5 The Gradient tab on page 133 |

6.2 **Method blocks**

Introduction

This section contains a description of how to organize a method in blocks of instructions in order to make it more structured, and of how to work with method blocks.

In this section

This section contains these topics:

| Topic | See |
|--|-------|
| How to view method blocks | 6.2.1 |
| How to call method blocks | 6.2.2 |
| How to add method blocks | 6.2.3 |
| How to delete method blocks | 6.2.4 |
| How to rename method blocks | 6.2.5 |
| How to find, copy and move method blocks | 6.2.6 |
| How to import method blocks | 6.2.7 |

6.2.1 How to view method blocks

Instructions can be grouped into blocks

To view a method as a long list of individual text instructions can be confusing and inconvenient. Text instructions can therefore be grouped into blocks of instructions that define a specific functional use. For example, one block might contain the instructions necessary to equilibrate a column, and another block contains instructions to load a sample, etc.

The Text pane

In the **Text** pane of the **Method Editor**, the method is shown as a list of blocks, denoted by the blue square symbols. Note that a block can also contain sub-blocks.

The figure below shows the text instructions in blocks:

The table below describes how to view or hide the instructions:

| If you want | then |
|--------------------------|------------------------------|
| to view the instructions | click the "+" symbol |
| | or |
| | double-click the block name. |
| to hide the instructions | click the "-" symbol |
| | or |
| | double-click the block name. |

The Block pane

The organization of blocks in the method is shown graphically in the **Block** pane of the **Method Editor**.

Description

Each block is represented by a gray bar with the block name and the length of the block. The line is shifted down to indicate calls to other blocks.

Click on the line that represents a block in the **Block** window to expand the block in the **Text** pane and select the first instruction in the block.

Figure

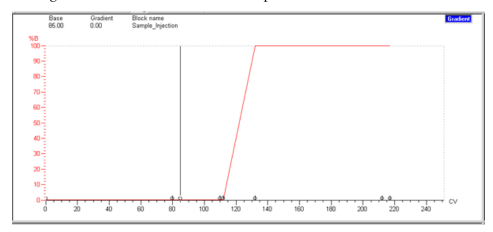
The figure below is an example with a **Watch** instruction to start the fraction collector which is active throughout the gradient elution block. **Loop** (to repeat a group of instructions) and **Hold_until** instructions are also indicated in the **Block** pane.



The Gradient pane

Blocks are represented in the **Gradient** pane of the **Method Editor** by marks on the X-axis. The marks show the length of each block. The name of the block in which the cursor line is currently placed is shown at the top of the pane.

The figure below describes the **Gradient** pane:



6.2.2 How to call method blocks

General description

To execute the instructions contained within a block in a method, the block must be called by the program. When a block is called, the instructions in the block are executed in the order that they are written until the block is finished or the **End_Block** instruction is executed. Any settings made in a block are valid throughout the method until the settings are changed.

Types of calls

There are two types of calls:

- Unconditional calls, which are made with a **Block** instruction.
- Conditional calls, which are made with a **Watch** instruction. This makes it possible to call a specified block or an instruction when a particular monitor signal meets a given condition. As long as the condition is not met, the block is not activated.

Watch instructions

Watch instructions are indicated by a green line that show the start and duration of the watch. These instructions can use various conditions to respond to absolute signal values or to rate of signal changes.

The breakpoint when the **Watch** instruction is issued determines when the watch begins, not when the block is activated. Once set, a watch remains active until the condition is met or a new **Watch** instruction is issued for the same monitor. The watch is cancelled automatically when the condition is met. A watch can also be turned off with the **Watch_off** instruction.

See F Method examples on page 545 for more details on Watch instructions.

6.2.3 How to add method blocks

Two ways to add method blocks

You can add method blocks to a method in two ways, using either

• the Instruction box of the Text Instructions editor,

or

• the New Block dialog box reached via the New Block icon.

Both these alternatives are described below.

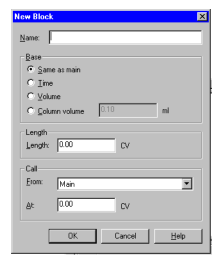
How to add blocks with the Instruction box

The table below describes how to add blocks with the **Instruction box**:

| Step | Action |
|------|--|
| 1 | In the Text pane of the Text Instructions editor, select the instruction or block that you want to precede the new block. |
| 2 | Select Other:Block in the Instruction box. |
| 3 | Enter a name for the block in the Block field. Click the Insert button. Result: The block is inserted after the block that was selected in step 1. |

The New Block dialog box

The illustration below shows the **New Block** dialog box that can be used when adding new method blocks:



How to add blocks with the New Block dialog box

The table below describes how to add blocks with the menu options of the **New Block** dialog box:

| Step | Action |
|------|---|
| 1 | Choose Block:New in the Method Editor |
| | or |
| | click the New Block icon. |
| | |
| | Result: The New Block dialog box is displayed. |
| 2 | Enter the relevant information in the New Block dialog box, and click OK . |
| | Result: The new block is added to the method, and placed last of all blocks. |
| | <i>Note</i> : The block can be placed in other positions by selecting something other than Main in the From droplist. |

The fields of the New Block dialog box

The table below describes the fields of the **New Block** dialog box:

| Field | Description |
|--------|--|
| Name | Block names can be up to 30 characters long, and can contain letters (A-Z), digits (0-9) and the underscore character. |
| | Block names must be unique within the method. The case of letters is retained but not significant (the names Start_Frac and START_FRAC are treated as identical). |
| Base | One of the following options can be selected: |
| | • SameAsMain : the new block will inherit the base from the Main block in the method. The corresponding Base instruction will be inserted in the block at breakpoint 0. |
| | Time: The block will be based on time. |
| | Volume: The block will be based on volume. |
| | Column volume: The block will be based on column volume. |
| Length | A block continues until the breakpoint for the End_Block instruction has been reached. |
| | An End_Block instruction will automatically be inserted in the block at the defined breakpoint. This field must not be left blank. |

| Field | Description |
|-------|---|
| Call | You can call the new block from an existing block (for example the Main block). |
| | Select values in the two fields: |
| | • From |
| | The block from which the newly created block should be called. |
| | • At |
| | The breakpoint at which the call is to be made. |
| | If you do not want to call the block (for example when the block being created is to be activated by a Watch instruction), choose the <unused></unused> line from the From drop-down list. Blocks using this line are placed last in the method in the Unused category. |
| | <i>Note</i> : You should not call a block from within itself. If you do, you will generate a potentially infinite loop that exceeds the maximum number of calls allowed in a method. A loop symbol is displayed at the beginning of the line if this occurs. |

6.2.4 How to delete method blocks

Four ways to delete blocks

There are four ways to delete blocks:

- To right-click a block and choose **Delete** from the shortcut menu
- To select a block and click Delete in the Instruction box
- To select a block and press the <Delete> key on the keyboard
- To select a block and use the Block:Delete Block command

Note: When you use any of the first three ways, the **Method Editor** dialog box will give you the option to transfer the block to the Unused section.

Delete options

The **Delete Block** dialog box is displayed when you delete a block with one of the first three options mentioned above.



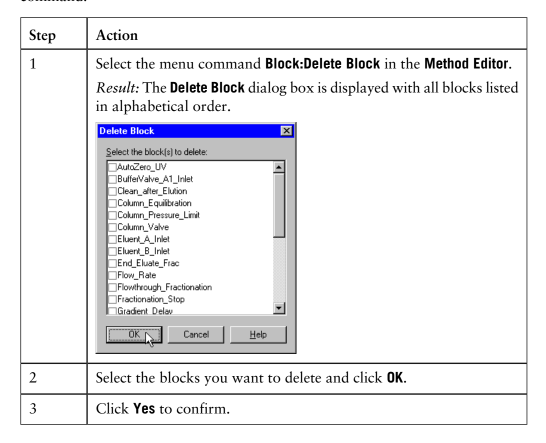
Options

Choose from the following options:

- **Delete:** The block is totally removed from the method. If the block is called several times in the method, all the blocks will be deleted. Blocks deleted in this fashion cannot be called again in the method.
 - *Note:* If the block contains sub-blocks, another dialog box is displayed, asking you if you want to delete the sub-blocks as well.
- Move: The block is deleted from the method and transferred to the Unused section. If the block is called several times in the method, however, only the row with the block currently marked in the Text pane will be deleted. In this case, the block will not be placed in the Unused section (since the block is still used in the method). Blocks deleted in this fashion can be called again in the method.

How to use the Block:Delete Block command

The table below describes how to delete a block using the Block:Delete Block command:



How to delete unused blocks

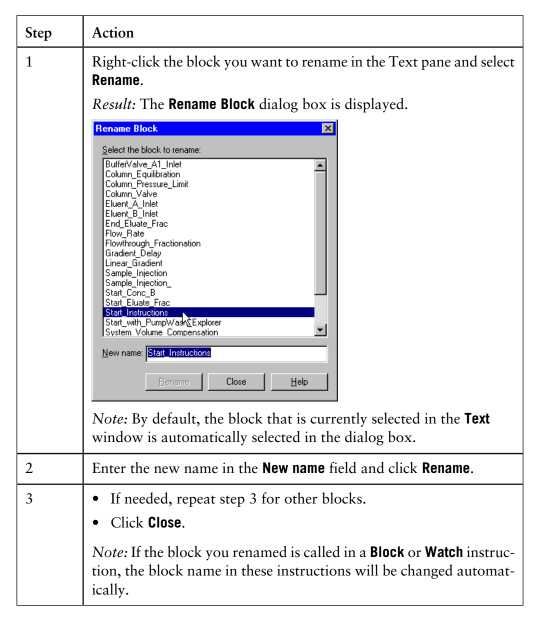
The table below describes how to delete an unused method block.

| Step | Action | |
|------|--|--|
| 1 | Highlight the method block. | |
| | Press the <delete> key</delete> | |
| | or | |
| | Right-click and choose Delete on the shortcut menu. | |
| | Result: The Delete Block dialog box opens. Note that the Move button is not available. | |
| 2 | Click the Delete button. | |
| | Result: The unused block is deleted and cannot be called upon again in the method. | |

6.2.5 How to rename method blocks

Instruction

The table below describes how to rename blocks:



6.2.6 How to find, copy and move method blocks

Introduction

By using the Edit options in the Method Editor, you can find, copy and paste and move blocks within a method.

How to find text strings in the method text

The table describes how to find text strings in the method text.

| Step | Action | |
|------|---|--|
| 1 | Choose Edit:Find in the Method Editor, | |
| | or | |
| | right-click an instruction or a block in the Text window and select Find . | |
| | Result: The Find dialog box is displayed. | |
| | Find Find what: Match whole word only Match gase Search from top of document Find what: Direction Cancel Cancel | |
| 2 | Enter the text you want to search for, search direction and case matching criteria. Click OK. | |

How to copy and paste a block

The table describes how to copy a block.

| Step | Action |
|------|---|
| 1 | Right-click the block you want to copy. |
| | Choose Copy. |
| 2 | Right-click the instruction line just above the point where you want the block to be pasted. Choose Paste. |
| | Result: A dialog box asks if you wish to rename the pasted block. |
| 3 | Click Yes to rename the block before insertion, or No to insert the copied block directly. |
| | <i>Result:</i> The pasted block is inserted with the same breakpoint value as the block or instruction selected for point of insertion. |

How to move a block

The table describes how to move a block.

| Step | Action |
|------|---|
| 1 | Right-click the block you want to move.Choose Cut. |
| 2 | Right-click the instruction line just above the point where you want the block to be pasted. Choose Paste. |
| | <i>Result:</i> The block is now removed from its original breakpoint and pasted at the new breakpoint. The pasted block is inserted with the same breakpoint value as the block or instruction selected for point of insertion. |

6.2.7 How to import method blocks

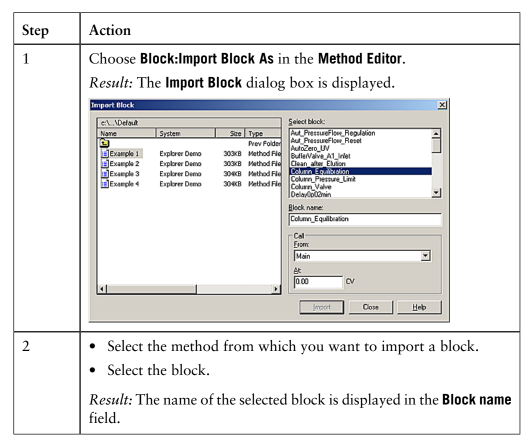
Introduction

You can import method blocks from other method files. You can also use this function to copy blocks within a method. In the latter case, it is important to note that it is the *saved* version of the method that will be copied, not changes that have been made after you last saved the method.

The block is imported exactly as it appears in the source method. If the base of the imported block is defined as **SameAsMain**, the block will inherit the main base in the new method, regardless of the base in the source method. Also, the imported block is inserted with the same breakpoint value as the block selected for point of insertion.

Instruction

The table below describes how to import method blocks:



| Step | Action | |
|------|---|--|
| 3 | In the Call field, do the following: | |
| | On the From drop-down list, select a block into which the block will be imported. | |
| | • In the At field, select the breakpoint value for the block to be imported. | |
| | Click the Import button. | |
| | <i>Note:</i> The imported block cannot have the same name as an existing block in the method. If the default name is not allowed for this reason, the Import button will be gray and locked. If this occurs, change the name of the imported block so that the Import button becomes available. | |
| 4 | Repeat steps 2 and 3 if needed. Click the Close button. | |

Method instructions 6.3

Introduction

This section describes how to work with the individual method instructions, in order to edit method blocks and methods.

In this section

This section contains these topics:

| Topic | See |
|---|-------|
| How to read method instructions | 6.3.1 |
| How to add method instructions | 6.3.2 |
| How to delete method instructions | 6.3.3 |
| How to change or move method instructions | 6.3.4 |

6.3.1 How to read method instructions

Description of instruction markings

Method instructions are displayed in the **Text** pane of the **Text Instructions Editor**. The table below explains the meaning of the markings:

| Marking | Explanation |
|------------------------------|--|
| Blue square beside text | Valid call instructions, that is, Block and Watch instructions to other blocks in the method. |
| Blue square with a red cross | Call instruction that contains one or more invalid instructions. |
| Bold text | Valid instructions. |
| Red dot | Instructions with invalid syntax. All such instructions must be deleted or changed before a method can be run. See 6.3.4 How to change or move method instructions on page 115. |
| | The instructions may be of the following types: |
| | Calls to blocks which are not defined in the method |
| | • Instructions that apply to a different system strategy (can occur if a method is written for one system and saved for another) |
| | • Instructions for components that have not been selected in the System Setup . |
| Normal text | Instructions that will not be executed because |
| | they are positioned after the end of a block or method |
| | or |
| | • they constitute a block to which there is no call. |
| Text with a loop symbol | When a block is called from within itself this will generate a potentially infinite loop, which might exceed the maximum number of calls allowed in a |
| ./5 | method. |

6.3.2 How to add method instructions

Instruction

The table below describes how to add a method instruction in the **Text Instructions Editor:**

| Step | Action | |
|------|--|--|
| 1 | Select a block in the Text pane, and display the instructions within the block. | |
| 2 | Select an instruction line in the block. Make sure that the selected instruction line is in the block, not the call to the block. | |
| 3 | Open the Instruction box if it is not already displayed (View: Panes). Do the following: Set the desired breakpoint in the Breakpoint field. Choose the instruction type and the instruction in the Instructions field. For basic help on each instruction, click the instruction and press <f1>.</f1> Type values for instruction parameters in the Parameters fields. If a scroll bar appears at the right side of the Parameters field, additional parameters are required. | |
| 4 | Click the Insert button. Result: The instruction will be inserted in the block at the position of the breakpoint of the new instruction, if there are no other instructions at that breakpoint immediately after the currently highlighted instruction, if the highlight is at the same breakpoint as the new instruction as the last instruction at the breakpoint, if there are several instructions at the same breakpoint and none of these is highlighted. Note: Instructions that are placed at the same breakpoint are executed simultaneously, with the exception of Block instructions which are executed in the sequence in which they are written. | |

Pause, Hold and Hold_until instructions

If you use ÄKTA systems, the Pause, Hold, and Hold_until instructions will stop execution at this breakpoint, that is, instructions following after Pause, Hold and Hold_until at the same breakpoint will not be executed until a Continue instruction is issued.

6.3.3 How to delete method instructions

Instruction

The table below describes how to delete method instructions in the **Text Instructions Editor**:

| Step | Action |
|------|---|
| 1 | Select the instruction in the Text pane. |
| 2 | Use one of the following alternatives: Right-click the instruction and choose Delete in the displayed menu, or |
| | press the Delete button in the Instruction box, or press the Delete key on your keyboard. |

End_Block instruction

If you delete the **End_Block** instruction, the block will end at the last instruction in the block. If a gradient is currently being formed, the gradient will continue into the next block.

How to suspend execution temporarily

An instruction that has been deleted can only be recovered by re-inserting the instruction. If you want to suspend execution of an instruction temporarily (for example during development work), you can replace the breakpoint with a value after the **End_Block** or **End_Method** instruction.

6.3.4 How to change or move method instructions

How to change an instruction

The table below describes how to change an instruction in the **Text** pane of the **Text Instructions Editor:**

| Step | Action | |
|------|--|--|
| 1 | Select the instruction. | |
| | <i>Result:</i> The instruction with its current parameters is displayed in the Instruction box . | |
| 2 | Make the required changes to the breakpoint or parameters | |
| | or | |
| | select a new instruction in the Instruction Box. | |
| 3 | Click the Change button | |
| | or | |
| | the Replace button. | |
| | <i>Note</i> : These buttons are equivalent unless changes are made to the breakpoint or the length of a gradient. See below. | |

Effects of the Change button and the Replace button on breakpoints

The table below describes the difference in function between the **Change** button and the Replace button when you change breakpoints:

| Button | Function |
|--------|--|
| Change | This button shifts all subsequent instructions in the block according to the change in the breakpoint. Change does not affect the relative order of instructions in the method. You cannot change the breakpoint of an instruction to earlier than the nearest previous breakpoint in a block. |
| | The illustration shows an example where Fractionation is changed from breakpoint 0 to 5: Change [Gradient] 0.00 Base SameAsMain 0.00 Gradient 100 (28) 20.00 (base) 20.00 Message "End of gradient", Screen, "No sound" 20.00 End, Block 20.00 End, Block 20.00 End, Block 20.00 End, Block 20.00 End, Block |

| Button | Function |
|---------|---|
| Replace | This button moves the selected instruction but does not change the breakpoint of any other instruction. Replace can change the relative order of instructions in the method. |
| | The illustration shows an example where Fractionation is changed from breakpoint 0 to 5: Replace |

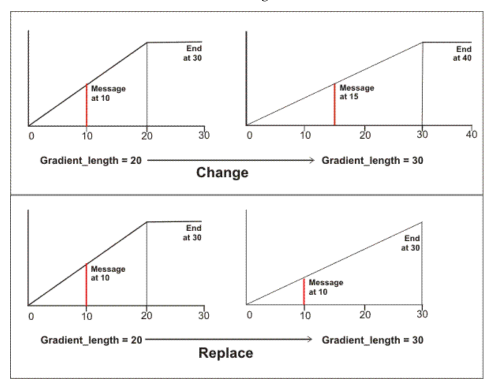
Effects of the Change button and the Replace button on gradient length

The **Length** parameter in the **Gradient** instruction affects the length of a gradient. Depending on which button you use, the change will have different results. The table below describes this:

| Command | Function |
|---------|---|
| Change | If this button is used to change the length of a gradient, the breakpoints for any instructions issued during the progress of the gradient will be adjusted proportionately so that they are always placed at the same relative position within the gradient. Instructions issued after the end of the gradient will be shifted by the amount of the change. Since the gradient works over time, any instruction that you want to insert after a gradient should be placed after the combined breakpoint and gradient length. Note: Moving the End_block instruction in a gradient |
| | block with the Change button does not affect the length of the gradient. |
| Replace | If this button is used to change the length of a gradient, other instructions are not affected. |

Illustration of the effects of the Change button vs. the Replace button on gradients

The illustration shows the different effects of the **Change** button and the **Replace** button on instructions within and after gradients:



How to move an instruction

Move an instruction within the same breakpoint

Select the instruction in the **Text** pane of the **Text Instructions Editor** and drag it to its new location to change the order of instructions within the same breakpoint in a block.

Move an instruction to another breakpoint

The table below describes how to move an instruction to another breakpoint:

| Step | Action |
|------|--|
| 1 | Select the instruction in the Text pane of the Text Instructions Editor. Choose Edit:Cut. |
| 2 | Select the instruction line just <i>above</i> the point where you want the cut instruction to be pasted. Choose Edit:Paste . |
| | <i>Result:</i> The instruction is now removed from its original breakpoint and pasted at the new breakpoint. The pasted instruction is inserted with the same breakpoint value as the instruction selected for point of insertion. |

6.4 How to use method variables

Introduction

Method variables can be used to edit suitable methods. Variables can be assigned to most instruction parameters including breakpoints. Variables also form the foundation for automatic method scouting.

Each parameter defined as a variable is also assigned a default value, which is used if no changes are made to variable values at the start of a run. Up to 500 variables can be defined in a single method.

All variables are listed on the **Variables** tab of the **Run Setup**, grouped according to the block in which they appear. See **6.5.2** The Variables tab on page 125.

Identifying variables

Parameters defined as variables can be identified in two ways:

• In the **Text** pane in **Text instructions**, the parameter is given as the default value in parentheses followed by the variable name. The illustration below shows an example of this:

```
0.00 Block Wash_Out_Unbound_Sample
[Wash_Out_Unbound_Sample]
0.00 Base SameAsMain
[2]#Wash_column_with End_block
0.00 Block VolumeFractionation
0.00 Block Linear_Gradient
```

 When the instruction is shown in the Instructions field of the Instruction box, the VAR button beside the parameter field is displayed in capital letters, that is VAR not Var.

The illustration below shows an example of the **Instruction box** where **UV1** and **UV2** are defined as variables and the **UV3** position is fixed.



When to change variable values

Variable values can be changed immediately before the start of a method run without using the **Method Editor**, allowing one method to be used for runs under a variety of conditions.

How to change variable values

To change default variable values, you can either

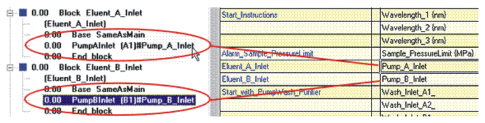
• edit the instruction in the **Instruction box**

or

• change the value in the **Variables** tab of **Run Setup**.

Changes made in the **Text** pane are automatically updated on the **Variables** tab and vice versa.

The figure below illustrates the relationship between variables in the **Text** pane and on the **Variables** tab of **Run Setup**:



Breakpoints or gradient lengths

If a breakpoint or gradient length is defined as a variable, changing the variable value in the **Variables** tab when the method run is started will shift other instruction breakpoints accordingly. This functionality is equivalent to using the **Change** button to alter a breakpoint or gradient length (see **6.3.4 How to change or move method instructions** on page 116 for how the **Change** button affects instructions within gradients).

How to define new variables

Only one variable that affects block length (breakpoint or gradient length) may be defined within each block. However, any number of parameters may be defined as variables within a block. The table below describes how to define a new variable.

| Step | Action |
|------|--|
| 1 | Select the instruction where you want to define the variable in the Text pane of Text instructions . |
| | <i>Result:</i> The parameters for the instruction are shown in the Instruction box . |
| 2 | Locate the breakpoint or the required parameter in the Instruction box . Oliver to Manufacture. |
| | Click the Var button. Result: The Variable Name Definition dialog box opens. |

| Step | Action |
|------|--|
| 3 | Enter a name for the variable. |
| | • Select the Visible in details only check box if you want to set the variable as a "details" variable. Detail variables only become visible on the Variables tab if the Show details check box is selected. This option is useful for hiding less important variables. |
| | • Click 0K . |
| | Result: The Var button changes to VAR to confirm the new variable. The variable is displayed in the Text pane. |

Variable names

Variables are defined with names that can be explicit descriptions of the variable function, for example **Sample_volume** and **Gradient_length**. Suitable choices of variable names can make the method easier to read and understand, and also help the operator in setting variable values at the start of a method run.

The names can be up to 32 characters long and the following characters can be used:

- Letters (A-Z)
- Digits (0-9)
- The underscore character (_)

The case of letters is retained, but not significant. The names **Flow_Rate** and **FLOW RATE** are treated as identical.

How to rename a variable

The table below describes how to rename a variable:

| Step | Action |
|------|--|
| 1 | Select the instruction that includes the variable you wish to rename in the Text pane of Text instructions . Result: The parameters for the instruction are shown in the Instruction box . |
| 2 | Locate the required parameter in the Parameters field. Click the VAR button. |
| 3 | Enter a new variable name in the dialog box and click OK . |

Note: Variables can also be renamed in the **Edit Variables** dialog box in the **Method Editor**. See **6.5.2** The Variables tab on page 126 for more information.

variable

How to remove a The table below describes how to remove a variable by converting it into a fixed value:

| Step | Action |
|------|--|
| 1 | In the Text pane of Text instructions , select the instruction with the variable you want to remove. |
| | <i>Result:</i> The parameters for the instruction are shown in the Instruction box . |
| 2 | Locate the required parameter in the Parameters field. Click the VAR button. |
| 3 | Click the Clear button to delete the variable. Click OK. |
| | Result: The VAR button changes to Var to confirm that the variable is removed. |

Note: Variables can also be deleted in the Edit Variables dialog box in the Method **Editor**. See **6.5.2** The Variables tab on page 126 for more information.

6.5 Run Setup

Introduction

Run Setup is a part of the **Method Editor**. It has several tabs for defining method properties. This section describes how to use the tabs and the information displayed on the tabs.

In this section

This section contains these topics:

| Topic | See |
|---|--------|
| Overview of Run Setup | 6.5.1 |
| The Variables tab | 6.5.2 |
| The Scouting tab | 6.5.3 |
| The Questions tab | 6.5.4 |
| The Gradient tab | 6.5.5 |
| The Notes tab | 6.5.6 |
| The Evaluation Procedures tab | 6.5.7 |
| The Reference Curves tab | 6.5.8 |
| The Columns tab | 6.5.9 |
| The BufferPrep tab | 6.5.10 |
| The Method Information tab | 6.5.11 |
| The Result Name tab | 6.5.12 |
| The Frac-950 tab | 6.5.13 |
| The Start Protocol tab | 6.5.14 |
| How to export the values in the Run Setup | 6.5.15 |

6.5.1 Overview of Run Setup

Introduction

To access **Run Setup**, either

• Click the Run Setup icon on the Method Editor toolbar,

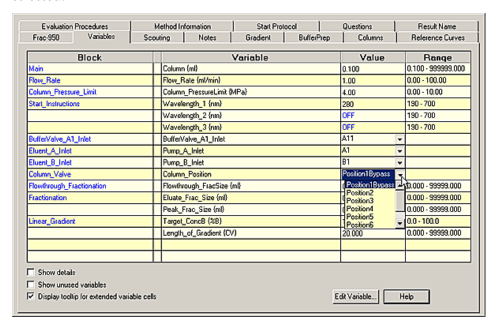


 α

• Select View: Run Setup.

Illustration of Run Setup

The illustration below shows an example of the **Run Setup** with the **Variables** tab selected:



The tabs

The table below contains brief descriptions of the tabs of **Run Setup**. If you want more detailed descriptions, see sections on the respective tabs:

| Tab | This tab |
|-----------|--|
| Frac-950 | allows the user to choose rack type and the fractionation order for the Frac-950 fraction collector. |
| Variables | lists all variables used in the method with their default values, organized by method block. |
| Scouting | shows the scouting scheme used for the method. The scouting scheme can also be set up from this tab. |
| Notes | shows the descriptive comments that form a part of the method documentation. |

| Tab | This tab |
|-----------------------|---|
| Gradient | provides a graphical overview of the block structure and eluent gradient tab in the current method. |
| BufferPrep | displays information about the selected buffer preparation recipe for the current method. |
| Columns | displays the columns used in the current method. |
| Reference curves | displays the reference curves that will appear in the System Control curve dialog box during the run of the current method. |
| Evaluation Procedures | shows the evaluation procedures that will run at the end of the current method. |
| Method Information | displays information about the method, such as method name, target system, and last date of change. |
| Start Protocol | determines which items of the Run Setup that are displayed at the start of the run. |
| Questions | displays the questions used in the method. Questions provide a means for entering run-specific information at the start of a run. Use this tab when you want to define questions. |
| Result name | specifies how the result files will be named for the results of a run, and where the result file will be saved. |

6.5.2 The Variables tab

Introduction

The **Variables** tab lists all variables used in the method with their default values, organized by method block. You can change the default values to create a variant of the method.

Note: The variables of a block are only displayed once on the **Variables** tab, even if the block is called several times in a method. **Variables** are displayed only if the method contains variables.

Check boxes

There are three check boxes on the **Variables** tab. The table below describes these boxes:

| Check box | Select this box if you want |
|---|---|
| Show details | detail variables to be shown. Detail variables are indicated by a D in the column immediately to the left of the Variable column. |
| Show unused variables | unused variables to be shown. Unused variables are indicated by a U in the column immediately to the left of the Variable column. |
| Display tooltip for extended variable cells | to display useful tips when you move the cursor to fields that can have several functions. |

Note: The options to show detail and unused variables can be set up as default options in the **Administration:Change User Attributes** settings in the **UNICORN Manager**.

How to change the default values

Enter new values in the appropriate fields to change the default variable values. For some variables, pre-set values are available on drop-down menus. Save the method when you have made your changes.

Note: The **Variables** box must be selected on the **Start Protocol** tab if you want to be able to change variable values at the start of a method.

Blue values

For variables with values shown in blue, the value input can be toggled between **OFF**, **INFINITE** or other single position values, and a variable range. To change the value, right-click the value cell.

Variables can also be changed in the Text Instructions Editor

Variables can be changed in the **Text Instructions Editor** as well as on the **Variables** tab of the **Method Editor**. Changed values will be displayed for the corresponding instructions in both windows.

How to delete or rename variables

The table below describes how to delete or rename a variable in the **Run Setup**.

| Step | Action |
|------|--|
| 1 | Click the Edit Variable button on the Run Setup Variables tab. |
| | or |
| | Choose the Edit:VariableMethod Editor menu option. |
| | Result: The Edit Variables dialog box opens. The variables are listed alphabetically. |
| 2 | Select the variable to edit. |
| 3 | Rename |
| | Type a new variable name in the New name text box. |
| | Click the Rename button. |
| | Result: The variable is renamed. |
| | Delete |
| | Click the Delete button. |
| | Confirm that you want to delete the variable. |
| | Result: The variable is deleted. |

How to change a variable into a detail variable

Detail variables are only shown if the **Show details** checkbox is selected on the **Variables** tab. The table below describes how to set up a detail variable.

| Step | Action |
|------|---|
| 1 | Click the Edit Variable button on the Run Setup Variables tab. |
| | Or Choose the Edit:VariableMethod Editor menu option. |
| | Result: The Edit Variables dialog box opens. The variables are listed alphabetically. |
| 2 | Select the variable to be changed. |
| 3 | Select the Set visible in details only checkbox. Click the Close button. |
| | Result: The variable is marked by the detail indicator D . |

How to change a detail variable into a regular variable

The table below describes how to change a detail variable into a regular variable.

| Step | Action |
|------|---|
| 1 | Click the Edit Variable button on the Run Setup Variables tab. |
| | Or Choose the Edit:VariableMethod Editor menu option. |
| | <i>Result</i> : The Edit Variables dialog box opens. The variables are listed alphabetically. |
| 2 | Select the variable to be changed. |
| 3 | De-select the Set visible in details only checkbox. Click the Close button. Result: The detail variable indicator D is removed. |

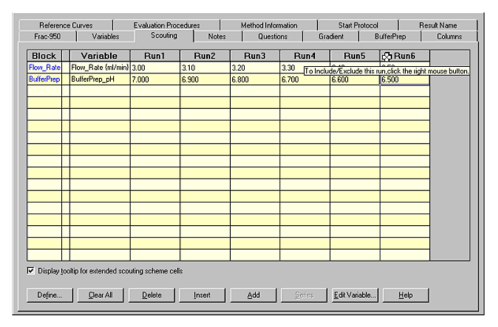
6.5.3 The Scouting tab

Introduction

A scouting scheme is a series of runs where chosen variable values are varied. You can define up to 99 runs in a scouting scheme. When a method is run with scouting, the method is automatically repeated for each selected run in the scouting scheme. Typically, scouting will vary one or more variables in a series of runs, for example, flow rate or elution gradient. See 7 Scouting on page 175 for instructions on how to set up a scouting scheme, and 9.4 How to perform a scouting run. on page 216 Note: The Scouting tab is available only if the method contains variables.

Example of a scouting scheme

The illustration below shows a scouting scheme for six flow rates and different pH values:



Note: The **Edit Variable...** button on the **Scouting** tab opens the same **Edit Variables** dialog box that can be accessed from the **Variables** tab.

6.5.4 The Questions tab

Introduction

The **Questions** tab of **Run Setup** is used for viewing and adding questions that the system asks a user at the start of a run. These questions provide a means for entering structured run-specific information. Method wizards and templates supplied with UNICORN are defined with a set of questions for sample, column and eluent identification.

Note: For questions to be shown in the start protocol, the **Questions** option must be checked on the **Start Protocol** tab of **Run Setup**.

Question status

Different types of questions have different status. The illustration below shows the **Question** field, an example of a question and the status alternatives that can be used:

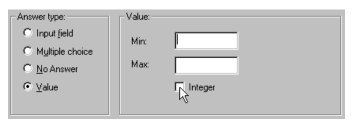


The table below explains the different alternatives:

| Question status | Explanation |
|-----------------|---|
| Mandatory | These questions must be answered before a method is started. |
| Authorized | These questions must be signed with the users signature password to unlock and continue the method. |
| Chromatogram | These questions will be printed with the answers on the same page as the chromatogram, if a question is chosen in an evaluation report. |

Answer type

A question has to be defined to accept one of four types of answers. The illustration below shows an example where the **Value** option has been selected. The appearance of the box to the right of the **Answer type** field depends on the answer type option selected:



The table below describes the different answer types:

| Answer type | This option |
|-----------------|---|
| Input field | accepts any alphanumerical input as the answer. Input field questions may have a default answer. |
| Multiple choice | allows the user to choose one of a defined set of answers. To allow a blank answer, enter a space in one of the predefined answers. |
| NoAnswer | is used to display important information or to split a question over more than one line by setting all but the last line in a question to No answer. (Normally, each question consists of one line only.) It is impossible to give an answer to questions with this option selected. |
| Value | accepts only numerical answers. Value questions must have specified maximum and minimum limits, and may be defined to accept only integer values. |

How to insert a question

The table below describes how to insert a question:

| Step | Action |
|------|--|
| 1 | If there are questions on the list, select the question that should be followed by the new question. |
| 2 | Enter the question text, status, answer type and answer option as required. |

| Step | Action |
|------|---|
| 3 | The Answer type determines what is displayed in the question definition field to the right of the Answer type field. For each answer type, do as follows: |
| | Input field |
| | Enter a default answer if required. |
| | Multiple choice |
| | Click in the text field under Alternatives. |
| | Enter the answer. |
| | Click the Add/Delete button. |
| | Result: The new alternative is added at the end of the list. |
| | • Repeat this procedure to add new alternatives. To remove an alternative, mark the alternative in the scroll list and click the Add/Delete button. |
| | No answer |
| | No action taken. |
| | Value |
| | Enter maximum and minimum limits. Select the Integer box if the question is to accept integers only as answers. |
| 4 | Click the Insert button. |
| | Result: The new question is added to the list. |

How to preview questions

The table below describes how to preview the questions as they will appear in the Start Protocol.

| Step | Action |
|------|--|
| 1 | Select a question. |
| | Click the Preview button. |
| | Result: The question is displayed. |
| 2 | Click the Edit button to return to the question editing mode. |

How to edit a question

The table below describes how to edit a question:

| Step | Action |
|------|---------------------------------------|
| 1 | Select the question you want to edit. |

| Step | Action |
|------|--|
| 2 | Change the text, status, type and answer as required |
| 3 | Click the Replace button. |

How to delete a question

Do one of the following to delete a question:

- Select a question and click the **Delete** button to remove the selected question.
- Click the **Delete all** button to delete all questions.

6.5.5 The Gradient tab

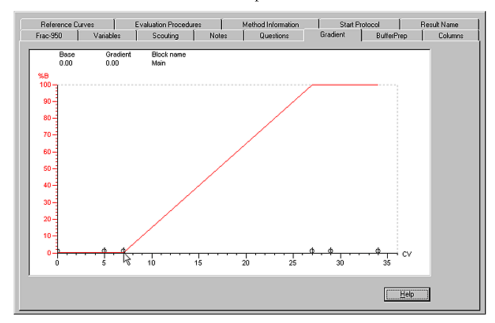
Introduction

The **Gradient** tab provides a graphical overview of the block structure and eluent gradient in the current method. The description of this tab can also serve as a description of the **Gradient** pane of the **Text Instructions**.

Note: For scouting runs, click **Run X** to see the gradient for each run.

Illustration

The illustration below shows an example of a **Gradient** tab:



How to zoom in on a selected region

The table below describes how to zoom in on a selected area of the **Gradient** tab:

| Step | Action |
|------|---|
| 1 | Press and hold the left mouse button and drag a rectangle on the screen to select the area you want to zoom in on. Release the mouse button. Result: The display is now zoomed in on the selected area. |
| 2 | Repeat the process for further magnification of selected areas. |

How to reduce the scale of the zoom function

To reduce the scale of the zoom function, right-click the tab and choose either:

- Undo Zoom to reverse each zoom-in action a step at a time, or
- **Reset Zoom** to reverse all of the zoom-in actions to the default scale setting.

How to use the vertical marker line

A vertical marker line can be dragged from the Y-axis with the mouse. As you drag the marker line, the current position is identified at the top of the tab in terms of the block name, X-position in the currently displayed base and eluent concentration in per cent of eluent B.

How to change the base shown on the X-axis

You can change the base shown on the **Gradient** X-axis. The alternatives are time, volume and column volumes. Changing the base for the display does not affect the base in the method instructions, which means that you can check how long a method will take simply by setting the axis scale to time, even if the method blocks are written in volume or column volume base.

The list below describes two ways to change the base shown on the X-axis:

• Click the X-axis to toggle between the base types.

or

- Right-click anywhere on the **Gradient** tab. *Result:* A sub-menu is displayed.
- Select **Base** and make the appropriate choice: **Time**, **Volume** or **CV**.

How to view hatch marks

You can display a hatched background on the **Gradient** tab. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Right-click anywhere on the Gradient tab. |
| | Result: A sub-menu is displayed. |
| 2 | Select Hatch. |
| | Result: The Gradient background is hatched. |
| 3 | To hide the hatch marks, repeat steps 1 and 2. |

6.5.6 The Notes tab

Introduction

Notes are descriptive comments that form part of the method documentation. Method templates are supplied with notes describing the system requirements for running the method. Read through these notes carefully before using a method.

Sub-tabs

There are four sub-tabs:

- **Method Notes**
- **Start Notes**
- Run Notes
- **Evaluation Notes**

Only the **Method Notes** can be edited from the **Method Editor**; the other notes are accessible at the respective stages in a run.

Recommended usage

We recommend that you use **Method Notes** to describe the system setup required by the method (for example eluent and sample inlets, outlets and column connections).

Use the **Start Notes** or **Run Notes** for run-specific information.

Note: **Method Notes** are saved with the method and apply to all runs made with the method.

How to write method notes

To write method notes in your own methods, place the cursor in the white area of the Notes tab and type the relevant text. Use standard Windows editing functions to edit the notes.

How to search for text strings

You can search for text strings in the method notes. The table below describes how to perform a search.

| Step | Action |
|------|--|
| 1 | Click the Find button. |
| | Result: The Find dialog box opens. |
| 2 | Type the text string in the Find what text box. |
| | Select search criteria and click 0K . |
| | Result: The located text string is highlighted in the text area. |

6.5.7 The Evaluation Procedures tab

Introduction

The **Evaluation Procedures** tab lists all evaluation procedures associated with the method. Evaluation procedures can be called automatically at the end of a method to evaluate and/or print the results.

Many UNICORN strategies are supplied with method templates or wizards that include a number of evaluation procedures. User defined procedures are created in the evaluation module and can be saved in method files (see 12.3 Automated evaluations procedures on page 378).

Changes in the Evaluation module

A procedure in a method will not be updated when a procedure with the same name is changed in the **Evaluation** module. The same applies to report formats saved in a procedure.

References to curves

Evaluation procedures that process chromatogram data rely on consistent identification of curves in the result file for correct operation. If you include evaluation procedures with a method, make sure that references to curves in the procedure will be valid when the procedure is executed at the end of the run (see 12.3 Automated evaluation procedures on page 378 for more details).

How to print evaluation results

If you use an evaluation procedure to print results automatically from a run controlled from a remote station in a network installation, the results will be printed on the printer currently set up on the local station, not on the remote station.

If you execute the procedure interactively from the **Evaluation** module on the remote station, the results will be printed on the printer set up on the remote station where you are working.

How to define and view evaluation procedures

Evaluation procedures are defined in the **Evaluation** module.

Procedures imported to a method can also be viewed and edited in the **Method Editor**. To do this, select the required procedure on the list and click the **Edit** button.

How to select procedures to run

To select procedures to run, select the procedure(s) that are to be executed at the end of the run. The procedures will be executed in the order they appear on the list.

How to import evaluation procedures

The table describes how to import global evaluation procedures:

Note: Procedures saved with one method file can be imported to another.

| C | . . |
|------|---|
| Step | Action |
| 1 | Select the Evaluation Procedures tab and click the Import button. |
| | Result: The Import dialog box is displayed. |
| 2 | Choose either option 1 or 2 below. |
| | Option 1: Select a global UNICORN procedure |
| | 1. Select a procedure on the Select list. |
| | Result: The evaluation procedure name is displayed in the Import as field. |
| | Option 2: Select a procedure from another method |
| | 1. Select a method, that contains a procedure, in the left part of the dialog box. |
| | <i>Result:</i> The procedures of the selected method will be displayed on the Select list. |
| | 2. Select the desired procedure on the Select list. |
| | Result: The method name is displayed in the Import as field. |
| | <i>Note:</i> Click Procedure List to return to the list of UNICORN's global evaluation procedures. |
| 3 | If desired, change the procedure name in the Import as field. |
| | <i>Note:</i> The imported evaluation procedure cannot have the same name as an existing evaluation procedure in the method. If the default name is not allowed for this reason, the Import button will be gray and disabled. When you change the name in the Import as field, the button will become available again. |
| 4 | Click the Import button. |
| | Result: The evaluation procedure is imported into the method. |
| 5 | Repeat steps 2 - 4 until you have imported all procedures. |
| 6 | Click the Close button. |

How to delete evaluation procedures

The table describes how to delete evaluation procedures from the method:

| Step | Action |
|------|--|
| 1 | Select the Evaluation Procedures tab. |
| 2 | Select the procedure(s) that you want to delete. |

| Step | Action |
|------|---|
| 3 | Click the Delete button and confirm the deletion when prompted. |
| | <i>Result:</i> The deleted procedures are immediately removed from the method file. |

How to rename evaluation procedures

The table describes how to rename evaluation procedures in a method.

| Step | Action |
|------|---|
| 1 | Select the Evaluation Procedures tab and click the Rename button. Result: The Rename dialog box is displayed. |
| 2 | Select a procedure from the list and change the name in the Rename item to field. Click Rename. |
| 3 | Repeat step 2 until you have renamed all procedures required. |
| 4 | Click the Close button. |

How to edit an evaluation procedure

The table describes how to edit evaluation procedures in a specific method:

| Step | Action |
|------|--|
| 1 | Select a procedure on the Evaluation Procedures tab and click the Edit button. |
| | <i>Result:</i> The Procedure Editor dialog box is displayed, with information about the selected procedure. |
| 2 | Enter the new parameter values in the appropriate place of the Parameters field, and click the Replace button. |
| | <i>Result:</i> The selected instruction in the evaluation procedure is updated in accordance with the new parameters assigned to it. |
| 3 | If needed, insert new instructions after the currently selected procedure instruction. Do the following: |
| | 1. Select an instruction type and instruction in the Instructions field. |
| | 2. Enter the appropriate parameter values in the Parameters field. |
| | 3. Click the Insert button. |
| | Result: The new instruction is added to the evaluation procedure. |
| 4 | To remove an instruction from the evaluation procedure, select it and click the Delete button. |

| Step | Action |
|------|---|
| 5 | Select File:Save as to save the edited procedure with a new name. Click the Close button. |
| 6 | Select File:Close from the menu in the Procedure Editor dialog box. <i>Result:</i> The Procedure Editor dialog box is closed and the procedure is saved automatically. |

6.5.8 The Reference Curves tab

Introduction

Reference curves are curves from existing result files that you can display in the **Curves** pane of **System Control** during a run.

How to choose and display reference curves

You can include up to five reference curves in a method. You choose which curves to display during the run with the **View:Properties:Curves** command in **System Control** (see **9.2.3 The Curves** pane on page 199). Reference curves are only displayed during the run. Reference curves are not saved in the result file.

How to add reference curves

The table below describes how to add a reference curve from a result file:

| Step | Action |
|------|--|
| 1 | Select the Reference Curves tab and click the Import button. |
| | Result: The Import Reference Curve dialog is displayed. |
| 2 | • In the left field, select the result file containing the curve to be added. |
| | Result: The Select list displays the available curves for the result file. |
| | Select the curve you want to add from the Select list. |
| 3 | If desired, change the curve name in the Import as field. |
| | Note: The curve name has to be changed if a reference curve with that name already exists.Click Import. |
| 4 | Repeat steps 2 and 3 if you want to add more curves. |
| 5 | Click the Close button to close the Import Reference Curve dialog box. |

How to delete reference curves

The table describes how to delete reference curves.

| Step | Action |
|------|--|
| 1 | Select the curves you want to delete. |
| 2 | Click the Delete button and confirm the action when prompted. |

Note: Deleting curves from the method does not affect the curves in the result file from which they were imported.

How to rename reference curves

The table below describes how to rename a reference curve in a method:

| Step | Action |
|------|---|
| 1 | Click the Rename button. |
| 2 | Select a curve from the list. Change the name in the Rename item to field. Click the Rename button. |
| 3 | Repeat steps 2 and 3 if you want to rename more reference curves. |
| 4 | Click the Close button. Result: The reference curve name is changed. |

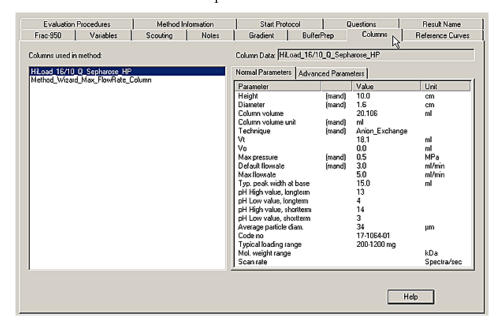
6.5.9 The Columns tab

Display of the column parameters

The **Columns** tab shows the parameters of the column selected for your method. The column parameters are displayed in the **Column Data** field. If you perform scouting runs with different columns, all of these will be listed. Select the appropriate column to display the parameters.

Illustration

The illustration shows an example of the **Columns** tab:



6.5.10 The BufferPrep tab

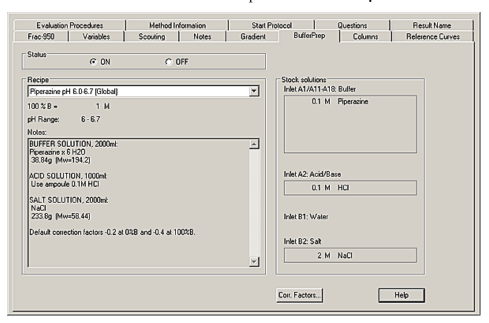
BufferPrep usage

BufferPrep allows a buffer of different pH and salt concentrations to be prepared on-line from four stock solutions. This removes the need to manually prepare new buffers every time the pH needs to be changed. Linear and step salt gradients can be run and pH can be used as a variable scouting parameter. **BufferPrep** is optimized for cation and anion exchange chromatography. For a complete description of **BufferPrep**, see the user manual for ÄKTAdesign systems.

Note: BufferPrep is only available for some ÄKTAdesign systems.

Illustration

The illustration below shows an example of the **BufferPrep** tab:



Stock solutions

The solutions and the inlets to which they should be connected are displayed to the right of the dialog box. Accuracy of preparation is essential. The four stock solutions consist of:

- a mix of buffering components (there can be up to five different buffering components enabling a broad pH range to be covered),
- an acid (HCl) or base (NaOH) for pH on-line titration,
- distilled water,
- an inert salt (for example NaCl) for salt gradient formation.

How to create a BufferPrep method

If a suitable template or wizard is not available, you can create a **BufferPrep** method yourself. The instruction **BufferPrep_pH** must be available at breakpoint zero at the beginning of the method. The method must not contain the instructions **PumpAlnlet** or **PumpBlnlet**.

The table shows one way to create a **BufferPrep** method.

| Step | Action |
|------|--|
| 1 | In the Text Instruction editor: Insert a BufferPrep-pH block at breakpoint zero at the beginning of the method. Define BufferPrep-pH as a variable. |
| 1 | Change to the Run Setup. Select the BufferPrep tab. Click the ON radio button in the Status field. Select a Recipe from the drop-down list box. There are two main alternatives: AIEX or CIEX, which are recipes covering a broad pH range, single buffer recipes for more narrow pH ranges. Result: All information relevant to the selected recipe will be displayed on the tab. |
| 2 | Prepare the required stock solutions. |
| 3 | Do one of the following: Select the Variables tab. Set the required pH for the method run in the variable BufferPrep_pH, or If you want to perform pH scouting, click the Scouting tab and select BufferPrep_pH as a scouting variable. Enter the pH values for the different runs. |

BufferPrep recipes

The recipe saved in the method (the one selected on the **BufferPrep** tab) cannot be edited, although fine tuning is possible. However, the recipes on the list of all **BufferPrep** recipes can be edited. New recipes can also be created (see E How to create and edit BufferPrep recipes on page 536).

How to fine tune the BufferPrep recipe with correction factors

In order to obtain high pH accuracy, the recipe can be fine tuned around a specific pH by setting correction factors. The table below describes how to fine tune the recipe with correction factors:

| Step | Action |
|------|--|
| 1 | In System Control, select Manual: Other. |
| 2 | Select BufferPrep Recipe and Recipe Name. Click the Execute button. |
| 3 | Set the pH in the instruction BufferPrep_pH in group Pump. Click the Execute button. |
| 4 | Set the flow rate to be used during the run in the Flow instruction. Click the Execute button. |
| 5 | Check the pH reading when stable. Allow at least 30 ml of eluent to pass through before expecting a steady pH reading. |
| 6 | Change to 100% B by setting the Gradient instruction in Manual:Pump to 100% for Target and 0 for Length. Click the Execute button. |
| 7 | Check the pH reading when stable at 100% B. |
| 8 | If the readings are acceptable at both 0% and 100%, the correction factors do not need to be changed. If the readings are not acceptable, click the Corr. Factors button in the BufferPrep tab in the method. |
| 9 | Enter the deviation at 0% and 100%. (e.g., if the pH is set to 7.0 and the actual pH is 7.1 enter 0.1. Enter -0.1 if the pH is 6.9). Note: If correction factors already exist, the measured pH deviation should be added to the old factors. |
| 10 | Save the method. |

Note: When changing the correction factors for the recipe selected in the method, the recipe with the same name on the list of all **BufferPrep** recipes is not affected. The changes will only apply in the specific method.

6.5.11 The Method Information tab

Introduction

The **Method Information** tab displays information about the method. This tab is for information only and cannot be edited.

There are three sub-tabs on this tab: Information, Signatures, and Method duration.

The Information sub-tab

The **Information** sub-tab displays

- method information such as method name, creation date, creator and date of last change,
- target system,
- strategy information such as strategy name, date and size.

The **Strategy Notes** button displays what systems, programs and file versions the strategy is designed for.

The Signatures sub-tab

The **Signatures** sub-tab has five information fields for all signatures. The table below describes the content of each field:

| Field | Description |
|-----------|--|
| Date | Date of the signature. |
| Meaning | Short description explaining the meaning behind the signature. |
| User Name | User name of the user who signed the method. |
| Full Name | Full name of the user who signed the method. |
| Position | Position of the user. |

The Method Duration sub-tab

The **Method Duration** sub-tab presents

- the estimated total time
- the estimated buffer volume required for the method.

If the method includes a scouting scheme, click the **Run** 'x' button to see values for the different scouting runs.

6.5.12 The Result Name tab

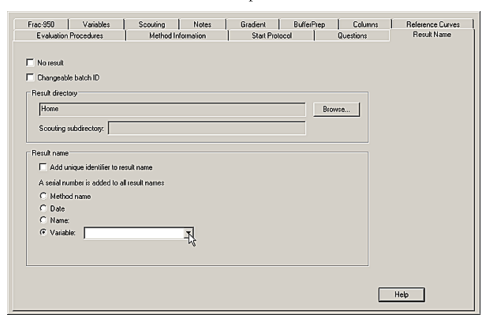
Introduction

The **Result Name** tab is used to specify:

- how the result files will be named for the results of a run
- where the result file will be saved
- the name of the special scouting folder where results from scouting runs will be stored.

Illustration

The illustration below shows an example of the **Result Name** tab:



Construction of the result file name

The result file name is constructed by one of the base options listed below. The serial number is changed automatically each time the method is run.

Base options of the result file name are:

- The **Method name** plus a 3-digit serial number,
- The **Date** of the run (in an 8-digit format determined by the country setting in Windows 2000 or XP) plus a 3-digit serial number,
- A freely specified **Name** (within the file naming restrictions of the operating system) plus a 3-digit serial number.
- A selected **Variable** (from the droplist) plus a 3-digit serial number.

Note: If a result names includes decimal points (e.g. numeric variables) or underscore characters, these characters will automatically be replaced by spaces. Points and underscores are not allowed in the result names.

Serial numbers and unique identifiers

If the result file folder already contains files with the same file name base, the serial number is changed automatically. For scouting runs, the 3-digit serial number will be the number of the executed run column in the scouting scheme.

A unique identifier can also be generated automatically, in addition to the serial number. The identifier is a string of numbers inserted between the result file name and the three-digit serial number.

• Select Add unique identifier to result name in the Result name field.

Batch ID for each test run

UNICORN will automatically issue a **Batch ID** to each method run. This ID is displayed before the **Base** in the logbook and can be used to identify individual runs. See illustration in 9.2.5 The Logbook pane on page 205. If **Changeable batch ID** is selected, another ID string can be typed in the **Start Protocol**.

Specify result name as changeable

The result name can be specified as changeable in the **Start Protocol** (see **6.5.14** The **Start Protocol** tab on page 152). In that case, the information you supply on the **Result Name** tab will be the suggested result name, but you can change this at the start of the run.

How to save the result files in a different folder

By default, result files are stored in the home folder of the user who starts the run. The table below describes how to change the folder where the result file will be stored:

| Step | Action |
|------|--|
| 1 | If the run contains information that is not important, you can save disk space by selecting the No result check box, thereby storing the result in the Temporary folder (named Manual Runs , where only the latest 10 result files are saved). If not, go to step 2. |
| 2 | Click the Browse button. |
| 3 | Double-click the required folder icon. Click the OK button. |

How to save scouting results

Scouting results will be saved in a special folder as specified by the result file path. To select a folder, type a name for the folder in the **Scouting** subdirectory field. Each time the scouting method is run, a new folder will be created with the name and a serial number (entering IEXSC will create folders IEXSC001, IEXSC002, etc.).

6.5.13 The Frac-950 tab

Introduction

The Frac-950 tab is used for defining options for Frac-950. The user can choose rack type and fractionation order.

How to set up fractionation a

The table describes how to set up the fractionation.

| Step | Action |
|------|--|
| 1 | Select rack type from the Rack drop-down list. |
| 2 | Select the order for fractionation by using the Fraction order radio buttons. |

Manual runs

In System Control, for manual runs, the Frac-950 tab cannot be used. Instead, use the manual fractionation instructions, starting with Man_.

• Choose Manual:Frac to open the Frac Instructions dialog box.

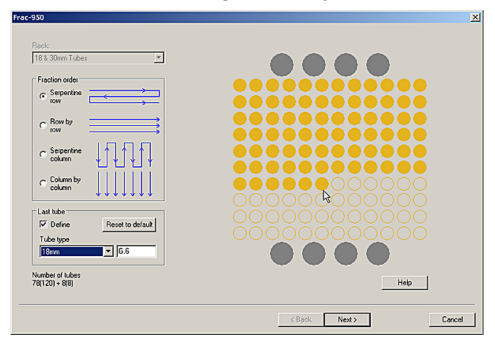
Total number of tubes

The total number of tubes sampled may differ if a last tube has been chosen. The Number of tubes equation in the bottom left corner of the Frac-950 dialog box of the Start Protocol shows the current number of available tubes chosen for fractionation, followed by the total possible number in parentheses.

How to select the last tube

You can select a position for the last tube to be used in the fractionation process. If the process attempts to go further than the selected last tube during a method run, an alarm will be executed.

The last tube position can only be selected on the **Frac-950** dialog box in the **Start Protocol** when you start a method run, or when you do an instant run. The illustration below shows an example of this dialog box:



The lower right box within the **Last tube** field shows the currently selected last tube. The table below describes how to re-define the last tube:

| Step | Action |
|------|---|
| 1 | In the Frac-950 dialog box, select the Define box in the Last tube field to select the last tube position. |
| 2 | Place the cursor over the appropriate tube (circle) within the tube matrix and click again. |

Note: When using different sized tubes in the same rack, the last tube can be set for both tube sizes. Use the **Tube type** drop-down list to choose the desired tube size, and then follow the procedures outlined above to select the last tube.

How to set the last tube to default setting

If you want to return to the default last tube position, click the **Reset to default** button of the **Frac-950** dialog box in the **Start Protocol** when you start a method run.

6.5.14 The Start Protocol tab

Introduction

The Start Protocol tab determines which items of the Run Setup are displayed at the start of a method run. Click the Start Protocol tab and select the items that you want to be displayed.

Checkboxes

The table below describes the check boxes of the **Start Protocol** tab:

| Checkbox | Displays |
|-----------------------|---|
| Frac-950 | the Frac-950 setup parameters, which can be changed. |
| Variables | values for method variables that can be changed at the start of the run. |
| | These values will override the default values for the particular run and be saved in the result file. The default values stored in the method are not affected. |
| Scouting | the scouting scheme which can be changed at the start of the run. Changes will override the default settings and values for the particular run and be saved in the result file. |
| Text Method | method instructions. They cannot be changed from this display. |
| Notes | the Notes tab. |
| Gradient | the gradient. |
| BufferPrep | the recipe selected in the method. The recipe cannot be changed during the start of a run. |
| Columns | the available column definitions. |
| Reference curves | the reference curves associated with the method. |
| Evaluation procedures | the evaluation procedures set to be executed at the end of the method. |
| Method information | the method information. |
| Settings | the settings. |
| Calibration | the monitor calibration settings. |
| Questions | questions defined in the method. You are recommended to always use this option, since the answers to questions can form an important part of the UNICORN run documentation. |

| Checkbox | Displays |
|-------------|--|
| Result name | the result name, which is changeable if this option has been selected. Click the Browse button to change the result folder. |
| | If the box is not selected, the result name will still be displayed, but you will not be able to change the name or folder. |

Scouting start pro- The table below describes the options in the **Scouting start protocol** field.

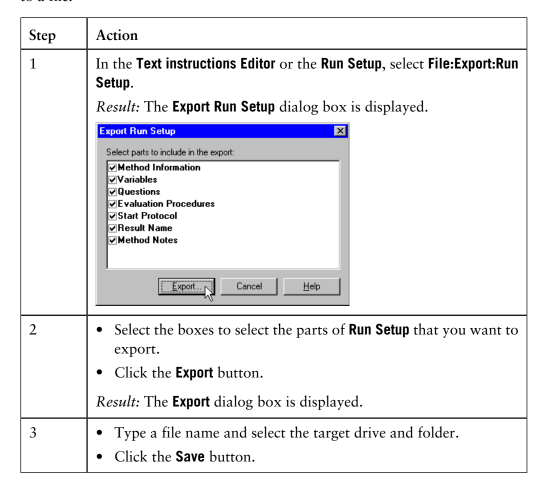
| Option | If you check this option |
|----------------|---|
| First run only | parameters for the scouting runs can be adjusted at the beginning of the first run only. After that, the runs will be performed automatically without operator intervention. |
| All runs | the Scouting start protocol will be displayed at the beginning of each run in the scouting scheme. |

6.5.15 How to export the values in the Run Setup

Instruction

You can easily export the values in the **Run Setup** to a file, and save it in ASCII format. This is useful when you want to enable others to read the methods without having access to UNICORN on their computers.

The table below describes how to export the values in the **Run Setup** and save them to a file.



6.6 How to use selected method instructions

Introduction

This section provides recommendations for how to use some common programming features in UNICORN methods. They are available from the **Instruction box** in the **Method Editor**.

In this section

This section contains these topics:

| Topic | See |
|-------------------------------------|-------|
| Base instruction | 6.6.1 |
| Instructions at the same breakpoint | 6.6.2 |
| Block and method length | 6.6.3 |
| Messages and Set_Marks | 6.6.4 |
| How to delay a method | 6.6.5 |
| Linear flow rates | 6.6.6 |
| Gradients and eluent concentrations | 6.6.7 |

6.6.1 **Base instruction**

Bases

Every method block must start with a **Base** instruction, defining the base for calculating breakpoints.

Different blocks can use different bases. The base can be one of the following:

- **volume** (the unit depends on the scale defined in the system strategy)
- time (minutes)
- **column volume**, CV (defined as a numerical value or taken from the column definition)
- SameAsMain (all blocks apart from the main block), which means that the block will inherit the base defined in the main block.

Method blocks that use a volume or column volume base

Make sure that the flow rate is not zero. Volume breakpoints are calculated from the flow rate of the pump, and the method will not progress if the flow rate is zero.

I use?

What base should Use the base that most closely suits the purpose of the block. Column volume is recommended as the base for most steps in a run. In some situations, however, it may be more suitable to use a time or volume base for individual blocks.

To change the base for an existing method

Be careful when changing the base for an existing method. Changing between time and volume bases can affect the relative duration of steps in the method if different steps use different flow rates.

Column parameter: named column

If a named column is selected for the **Column** parameter in the **Other:Base** instruction, the volume specified in the selected column definition will automatically be used for column volume in the method block. The column volume for base **CV** cannot then be changed in the instruction or defined as a variable. However, the **Column** parameter should be defined as a variable. Choosing a column definition also enables linear flow rate and column performance calculations.

Column parameter: Any

If the Column parameter in the Other:Base instruction is set to Any and the Base parameter is set to CV, the column volume is set numerically by the Volume parameter. The column volume may be defined as a variable, allowing the scale of the run to be decided when the method is actually run.

How to select columns for a template or wizard

In cases where a template or wizard-generated method and column are chosen, it is easy to select other columns for that method on the Variables tab in Run Setup. *Note*: This might not be possible for methods that you have created yourself.

How to select columns for a method not selected from a template

The table below describes how to select columns for a method, not selected from a template.

| Step | Action |
|------|---|
| 1 | In the Instruction box of the Text instruction dialog box, mark the Other:Base instruction. |
| 2 | Select the required column from the drop-down list for the Column parameter. Click the Var button to define the Column parameter as a variable. This is an optional but recommended step that will make it easy to change the column selection for different runs. |
| 3 | Enter a variable name and click 0K. Click Yes to confirm. |

Column definition

A column definition can be chosen and defined as a variable even if the base for the block is set to volume or time. Parameters in the column definition will then be used for linear flow rate and column performance calculations.

Recommendation

A selected column definition applies locally within the block for which it is selected, and is not transferred to other blocks. We strongly recommend that the column definition be selected for the main block.

Update parameters

If you want parameters (for example, flow, pressure and averaging time) to be updated when you change the column, you must define these as variables.

Pump:Methodbase instruction

Volume or column volume base is calculated from the flow rate of the **SystemPump** or the **SamplePump**, selected with the instruction **Pump:Methodbase**. If no **Pump:Methodbase** instruction is included in the method, the default setting **SystemPump** will be used.

6.6.2 Instructions at the same breakpoint

Description

Instructions placed at the same breakpoint in a block are executed simultaneously. **Exceptions**

Exceptions are successive **Block** instructions, which are executed in the sequence in which they are written. This can have important consequences in some situations.

The instruction sequence below shows an example of instructions with the same breakpoint, where the AutoZero_UV will start after the Wash block is completed.

| Breakpoint | Instruction |
|------------|--------------|
| 0.00 | Block WASH |
| 0.00 | AutoZero_UV |
| 0.00 | Block ELUATE |

6.6.3 Block and method length

General description

The time or volume of a method run is determined by the sum of the block lengths. In turn, the length of a block is determined by the breakpoint of the last instruction in the block.

Note: Depending on how conditional calls are used (see **6.7 Standard Watch conditions** on page 166), the overall method time or volume may vary according to watch events during the run.

Block length

A block in which all breakpoints are set to 0 will take no time or volume during a method run. The illustration below shows an example of this:

```
0.00 Block Initial_Eluent_Conditions_BP

(Initial_Eluent_Conditions_BP)

0.00 Base SameAsMain

0.00 BufferPrep_pH (7.000)#BufferPrep_pH

0.00 BufferValveA1 (A11)#Buffer_Inlet

0.00 Flow (5.00)#Flow_rate {ml/min}

0.00 Gradient (0.00)#Start_ConcB {%B}, 0.00 {base}

0.00 End_Block

0.00 End Block
```

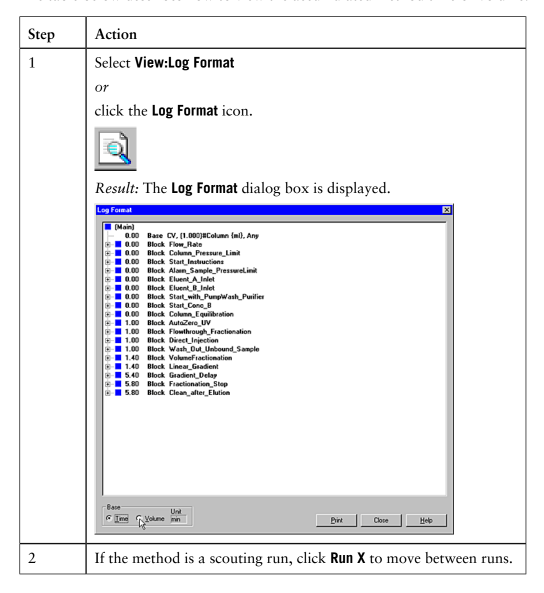
To extend the length of a block without performing any other operation, set the breakpoint of the **End_block** instruction appropriately, for example, as in the illustration below:

```
(Equilibration)
0.00 Base SameAsMain
4.00 End_Block
0.00 End_Block
```

How to view the accumulated method time or volume

The Log Format window in the **Method Editor** shows the accumulated method time or volume for the current method. The accumulated time/volume is an approximation and does not take into account time or volume for **Watch** blocks, **Wash** commands or programmed **Hold**. Also it does not compensate for splitter flow.

The table below describes how to view the accumulated method time or volume:



6.6.4 Messages and Set_Marks

When to use a message

Messages are used to inform the operator of the progress of the run. It is a good idea to issue messages at critical points in the method, for example, when **Watch** instructions are used for conditional events.

How to add a Message instruction

The **Message** instruction can be used to set up a message that will be displayed for the user during the execution of the method run. The message can be for information in a screen only, or it can require a signature before the user can control the system. The messages are all added to the logbook text. See **F.6 Appendix Messages** on page 555 for examples.

The table below describes how to add a **Message** instruction to the method.

| Step | Action |
|------|--|
| 1 | Select Other in the Instructions field of the Instructions box. |
| | Select Message in the instructions list. |
| 2 | Type a message in the Message text box in the Parameters field. |
| 3 | Select one of the display options on the Mode menu: |
| | Screen, i.e. only a text message is displayed. |
| | • Noscreen , i.e. the message will not be displayed but only inserted into the logbook. |
| | • Authorize , i.e. the message will require a signature from the user before the user can interact with the system again. |
| 4 | Select a sound on the Sound menu if desired. Click the Insert button. |

Note: If the **Message** instruction is inserted in a conditional block it will only be displayed if the conditions of the block (for example a **Watch**) is fulfilled.

Note: All messages are erased when the system reaches the **End** status. This also includes **Authorize** messages.

When to use a Set_Mark

Set_Mark instructions are useful text messages. They can be used

- to insert manual notes, for example, when a problem occurs in a run
- to highlight certain stages in a method.

Set_Marks differ from **Messages** in that they are inserted into the chromatogram at set points as well as into the logbook during a method run.

Example of a Set_Mark

The illustration below shows an example where **Set_Marks** are used to highlight the start and end of fractionation in a method:

```
(Volume_Fractionation)
0.00 Base SameAsMain
0.00 Set_Mark "Fractionation starts"
0.00 Set_Mark "Fractionation starts"
0.00 Fractionation (18mm)#TubeTypl√EluateFrac, (0)#Eluate_Frac_Size {ml}, (NextTube)#EluateFrac_StartAt, Volume
                0.00 End_block
0.00 End_Block
| Fractionation_Stop|
| 0.00 | Base SameAsMain |
| 0.00 | FractionationStop |
| 0.00 | Peak_FracStop |
| 0.00 | Set_Mark "Fractionation ends" |
| 0.00 | End_Block |
```

How to issue a Set_Mark

Set_Marks are issued from the Instructions box of the Text Instructions editor. The table below describes how to do this:

| Step | Action |
|------|---|
| 1 | Select Other:Set_Mark in the Instructions box. |
| 2 | Type the message in the Mark text field. |
| 3 | Click the Insert button. |
| | Result: A new line with the Set_Mark is added to the text instruction. |

6.6.5 How to delay a method

Introduction

A method can be programmed to be delayed at critical points. There are three instructions for this purpose: **Hold**, **Pause** and **Hold_Until**. These instructions are described below.

Hold

The **Hold** instruction suspends the execution of the method, but continues to pump eluent at the current flow rate and concentration settings. For example, this instruction is useful for giving the operator time to load a sample loop.

Resume the method

The method may be resumed if you click **Continue** on the **System Control** toolbar.

Pause

The **Pause** instruction suspends execution of the method and stops the pumps so that the system comes to a standstill. In ÄKTAdesign systems valves remain in the position they were in before the pause. The pause may be defined as indefinite or for a given number of minutes. This instruction is most useful for stopping the system in the event of an unexpected condition.

Resume the method

The method may be resumed if you click **Continue** on the **System Control** toolbar.

Hold_Until

The **Hold_Until** instruction is a special kind of **Watch** instruction. The method is put on hold until a specific condition is met (signal, test or value) or the time-out is reached. Thereafter the remaining instructions in the method are executed.

Instructions that share the same breakpoint as the **Hold_Until** instruction, but are placed after it in the method, will be executed after the **Hold_Until** conditions have been met.

6.6.6 **Linear flow rates**

Introduction

Linear flow rates (cm/h) can be specified for **Flow** instructions. The volume flow rate is calculated from a specified linear flow rate and the column diameter as given in the column definition.

How to use linear flow rates

The table describes how to use linear flow rates.

| Step | Action |
|------|--|
| 1 | Select a specific column on the Variables tab of the Run Setup , or Insert a column for the Base instruction of the block in the Text Instructions Editor . |
| 2 | In the Instruction box of the Text Instructions editor, select Flow and select the Linear Flow option as shown in the illustration below: Instructions |

Note: If the column is changed, you will be asked if the linear flow rate or the default flow rate should be used. If the linear flow rate cannot be used due to the max flow rate of the system or new column, you will be advised that the max flow rate will be used instead.

6.6.7 Gradients and eluent concentrations

Introduction

Gradient instructions are given in the **Text Instructions** editor of the **Method Editor**. This type of instruction defines gradients and immediate changes in eluent concentration.

Parameters of the Gradient

The table below shows the two parameters of the **Gradient** instruction:

| Parameter | Description |
|-----------|---|
| Target | Final eluent composition expressed in % eluent B. |
| Length | Duration of the gradient. |

Example of a Gradient instruction

The starting point for the **Gradient** is always the current eluent composition. The instruction can be read as follows: "form a **Gradient** to reach **Target** after **Length**".

Example of instruction

10.00 Gradient 50{%B}, 20{base}

The example instruction above forms a gradient to 50%B (**Target**) starting at breakpoint 10 with duration 20 method base units (**Length**). The example instruction will finish at breakpoint 30. If the current eluent concentration is greater than 50%, the gradient will be negative.

How to form a step gradient instruction

A step gradient is an immediate change in eluent composition. To form a step gradient, set the **Length** parameter to 0 in the **Gradient** instruction.

Example of instruction

10.00 Gradient 50{%B}, 0{base}

The example instruction above forms a step from the current eluent composition to 50%B at breakpoint 10. The method continues with 50%B.

Breakpoints for gradients

The breakpoint for a **Gradient** instruction defines the time or volume (according to method base) for the start of the gradient. A gradient with a non-zero duration occupies time and volume in the method, and breakpoints for other instructions may be set to occur before the gradient is completed. For most instructions, the instruction is simply carried out at the requested breakpoint, while the gradient is forming.

Instructions that affect gradients

The table below describes the instructions that affect the gradient:

| Instruction | Effect |
|-------------|--|
| Gradient | A new gradient will start at the requested breakpoint. Any remaining duration of the previous gradient is ignored. |
| Flow | The eluent flow rate will change at the requested breakpoint. If the current base is volume or column volume, the duration of the gradient will be changed. If the method base is time, the volume of the gradient will be changed. |
| End_Method | The whole method will stop, interrupting the gradient. |
| End_Block | The gradient formation will continue uninterrupted unless a new Gradient instruction is issued in the next block. For example, this means that a block can be called conditionally during gradient formation without interrupting the gradient. |

Gradients with variable length

For many purposes, it can be useful to define the length of the gradient as a variable. When this is done, breakpoints for instructions issued during or after the gradient in the same block are automatically shifted in proportion to the length of the gradient, with the same functionality as **Change** in the **Text Instructions** editor.

Instruction after a gradient

Any instruction that you want to insert after a gradient should be placed after the combined breakpoint and gradient length, since gradients function over time.

6.7 Standard Watch conditions

Introduction

Watch instructions allow the progress of a method run to be determined by the events during the method run, for example, start collecting fractions when the first peak eluates, or equilibrate the column until the eluent conductivity has reached a given value. This is facilitated by the **Watch** instructions.

The system strategy includes **Watch** instructions for each monitor defined in the system. These instructions are used to survey method runs, and instruct the system to call a specified block or an instruction when a particular monitor signal meets a given condition. As long as the condition is not met, the block is not activated.

Note: **Watch** instructions are shown in the **Instruction box** of the **Text Instructions** editor, indicated in the **Block** pane by a green line that shows the start and duration of the watch.

When is a Watch active?

The breakpoint when the **Watch** instruction is issued determines when the watch begins, not when the block is activated.

A watch is active from the point at which it is issued until

- the **Watch** condition is met
 - or
- a new watch is set for the same monitor *or*
- a **Watch_Off** instruction is issued for the monitor.

How to insert a Watch instruction

Watch instructions are inserted in the **Instruction box** of the **Text Instructions Editor**. The table below describes how to do this.

| Step | Action |
|------|---|
| 1 | In the Breakpoint field, select the appropriate breakpoint. This decides when the watch begins. |
| 2 | Select Watch in the Instructions field. Select a Watch instruction from the list. Select appropriate values under Test, Value and Action in the Parameters field. |
| 3 | Click the Insert button. Result: The new Watch instruction is inserted on the list of actions in the Text window. |

Test options in the Parameters field

The table below describes the **Watch** options that are available on the **Test** drop-down list of the Parameters field:

| Option | Explanation |
|---------------------|--|
| Greater_Than | The signal exceeds a certain value. |
| Less_Than | The signal falls below a specified value. |
| Slope_Greater_Than | The rate of change of the signal exceeds a specified value, expressed in monitor units/minute (for example, mAU/min). |
| Slope_Less_Than | The rate of change of the signal falls below a specified value, expressed in monitor units/minute (for example, mAU/min). |
| Less_Than_Or_Valley | The signal falls below a specified value or a valley is detected. A valley is detected only after a Peak_Max has been detected, and the valley is defined by a local minimum followed by an increase to 102% of the local minimum value plus the Delta_Peak value (see below). |
| Peak_Max | The signal falls to a specified fraction of the most recent peak maximum minus the Delta_Peak value. Factor=1 detects peak maximum. |
| Stable_Baseline | The signal is stable within the limits of the Delta_Base value for the period specified by the minutes parameter. |

Note: For slope values, use the Differentiate function in the Evaluation module to measure the slope of the test chromatogram. The Simulate Peak Fractionation technique can also be used to find the slope values.

Watch conditions for air sensors and AuxIn

Two Watch conditions are available for systems with air sensors, although they may be handled differently depending on the system. The table below describes the conditions and their explanations:

| Condition | Explanation |
|-----------|------------------|
| Equal O | No air detected. |

| Condition | Explanation |
|-----------|---------------|
| Equal 1 | Air detected. |

Note: To use the **Watch_AirSensor** instruction for air sensors, the **Alarm_AirSensor** setting must be disabled.

Actions when a Watch condition is met

The table below describes possible actions when a watch condition is met:

| Instruction | Effect |
|-------------|---|
| Block name | Calls the named block. |
| Pause, Hold | Pauses or holds the method. |
| Continue | Continues the method if paused or held. |
| End_block | Ends the current block and return to the point from which the block was called. |
| End_method | Ends the method. |
| Ready | Indicates that the next step in a MethodQueue may start. |

How to enter settings for Delta_Peak and Delta_Base

Permanent settings

Permanent settings for **Delta_Peak** and **Delta_Base** are entered with the **WatchPar** instruction (for example **WatchPar_UV**, **WatchPar_Cond**) under **System:Settings** in the **System Control** module (see the Administration and Technical Manual).

Temporary settings

Temporary settings that apply only for the duration of a given run can be entered in the Instructions field of the Instruction box in the Text Instructions editor. Select Alarms&Mon and then WatchPar.

The Delta_Peak setting

The **Delta_Peak** setting helps the software to detect valleys, peaks and peak maximum, and to ignore noise in the chromatogram.

The **Delta_Peak** value should be set

- large enough so that signal noise does not activate the conditions and
- small enough so that the condition is activated close to the valley or peak.

As a general guideline, set the value to 2-3 times the noise level and 5-10% of the smallest expected peak height. If you set a too high value you can prevent a new peak from being detected after a local minimum.

Use of the Delta_Peak setting

The **Delta_Peak** setting

• sets the threshold for signal increase after a local minimum that will be interpreted as a valley for the **Less_Than_Or_Valley** condition. A valley and a new peak are detected when the signal increases to 102% of the local minimum plus the **Delta_Peak** value.

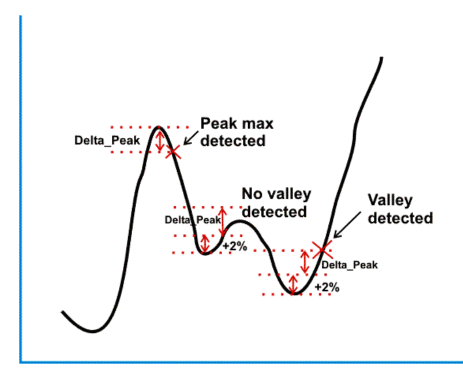
Note: A valley is detected only after a **Peak_Max** has been detected. *Example*:

If there is a local minimum at 0.05 AU and a **Delta_Peak** of 0.01 AU, a valley will be detected at:

 $(1.02 \times 0.05) + 0.01 = 0.111 \text{ AU}$

• sets the threshold for signal decrease after a local maximum that will activate the **Peak_Max** condition. **Peak_Max** is detected when the signal falls to the specified fraction of the most recent peak maximum minus the **Delta_Peak** value.

The figure below illustrates the **Delta_Peak** setting where **Peak_Max** is detected when the signal falls by **Delta_Peak** from a local maximum if the **Peak_Max** factor is set to 1:



The Delta_Base setting

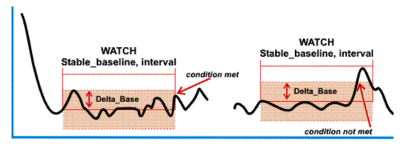
The **Delta_Base** setting helps the software to determine when the baseline is considered to be stable. In other words, it defines the permitted variation for the **Stable_Baseline** condition. For this condition to be activated, the signal may not vary by more than the **Delta_Base** value up or down over the time interval specified in the **Stable Baseline** condition in the **Watch** instruction.

Note: The **Delta_Base** setting affects the **Stable_Baseline** condition only.

The condition Watch Stable_Baseline

The condition **Watch Stable_Baseline** is met if the signal does not deviate by more than **±Delta_Base** from the baseline during the time interval specified for the watch. The baseline value is determined by the signal at the start of the watch. If the condition is not met, a new interval is started with a new baseline value defined by the signal level at the start of the new interval.

The illustration below shows an example of this:



6.8 How to save or delete a method template

How to save a method as a template

You can save a method that you have created yourself as a template if you have Edit global lists authorization (see the Administration and Technical Manual).

Recommendation

The templates for each system are common for all users. Be restrictive in saving methods as templates. We recommend that only methods that are useful for all users be saved as templates.

The table below describes how to save a method as a template:

| Step | Action |
|------|--|
| 1 | Choose File:Save as Template in the Method Editor. |
| | Result: The Save as Template dialog box is displayed. |
| 2 | Enter a name for the template in the Name field, |
| | choose an existing template name from the Templates list that shows the available templates within the chosen system. |
| 3 | • Select the system for which the template is intended in the For system field. |
| | Select the appropriate technique on the Technique list. |
| | Click 0K . |
| | Result: The method is saved as a template. |

How to delete a template

The table below describes how to delete a template:

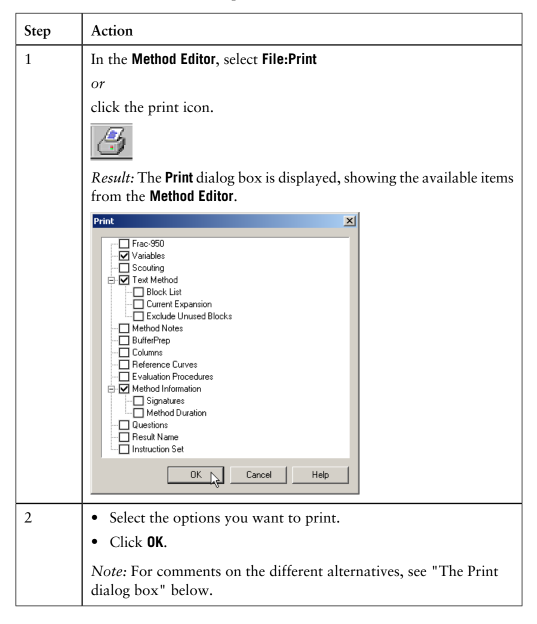
| Step | Action |
|------|---|
| 1 | Choose Edit:Delete template in the Method Editor. |
| 2 | Select the system and the template that you want to delete. Click the OK button and the Yes button to confirm. |

6.9 How to print a method

Instruction

You can print a copy of the method, including items from the method documentation, in **Run Setup** and the **Text Instructions** editor.

The table below describes how to print a method:



The Print dialog box

The table below describes some of the check box options in the **Print** dialog box:

| Check box | If you select this box |
|-------------|---|
| Text Method | all instructions will be printed, including those in unused blocks. |

| Check box | If you select this box |
|-------------------------------|--|
| Text Method:Current Expansion | the method will be printed according to the current expansion in the Text pane. (Only available from the Text Instructions editor.) |
| | (Only available from the Text instructions editor.) |
| Exclude Unused Blocks | only blocks that are used in the method will be printed. |
| Text Method: Block List | only the main method and a list of the blocks that are used in the method will be printed. |

6.10 How to export a method

Instruction

You can easily export a method to another file, and save it in another format, for instance .rtf. This is useful when you want to enable others to read the methods without having access to UNICORN on their computers.

The table below describes how to export a method and save it to another file:

| Step | Action |
|------|---|
| 1 | In the Text Instructions editor or the Run Setup, select File:Export:Method. Result: The Export Method dialog box is displayed. Export Method Options Main gnly Current expansion Export Cancel Help |
| 2 | Select whether the current method should be exported as a Method or as a Block list. Select the appropriate boxes in the Options field to define the level of detail in the information. Click the Export button. Result: The Export Method to file dialog box is displayed. |
| 3 | Enter a file name and select the target drive and folder. Click the Save button. |

Scouting 7

Introduction

Scouting is used to repeat a series of **Method runs** automatically with predetermined changes in the values for one or more Variables. A Scouting Scheme is defined as part of the method.

This chapter describes how to set up a **Scouting Scheme** and define columns. The chapter also provides some usage examples.

In this chapter

This chapter contains these topics:

| Topic | See |
|--|-----|
| How to set up a Scouting Scheme | 7.1 |
| How to define different columns for scouting | 7.2 |

7.1 How to set up a Scouting Scheme

Introduction

This section describes how to set up a method for scouting.

Any parameter can be scouted, provided that it can be defined as a variable in the method.

When to use scouting

Scouting is a facility for automatically repeating a run with systematic variation of one or more parameters. Some typical situations where scouting is useful are when you want:

- to screen for the best column
- to find the optimal pH
- to test column capacity (sample volume)
- to find the optimal flow rate for binding and elution
- to optimize gradient length and slope
- to optimize step gradients.

Variable values

The variables that appear in the scouting scheme are usually a subset of those on the **Variables** tab of the **Run Setup**. The values in the scouting scheme can only be set on the **Scouting** tab, while the default values in the method can be set either on the **Variables** tab or in the **Text instruction** pane.

Changing variable values in the scouting scheme does *not* change the values on the **Variables** tab or in the text instructions. Values for variables selected for scouting are grey on the **Variables** tab and cannot be changed there.

Any changes that you make to variable values when a scouting scheme is run are saved in the result file. Results from a scouting run are saved in a scouting folder.

Scouting tab buttons

There are seven buttons on the **Scouting** tab of the **Run Setup** plus the **Help** button. The table describes the functions of these buttons:

| Click the button | if you want |
|------------------|--|
| Define | to define new scouting variables. The Scouting Variables dialog box is displayed, and you can select variables to be used in the scouting series. Note: The variables that have been selected for scouting <i>cannot</i> be changed on the Variables tab. |
| Clear All | to clear all runs. This converts the scouting run to a non-scouting run so that it contains only the original method and values. |

| Click the button | if you want |
|------------------|--|
| Delete | to remove a run from the Scouting tab. Click on any variable in the run you want to remove, and then click the Delete button. |
| Insert | to insert a new scouting run before an existing run. Click on a run column and then click the Insert button. The new run will inherit the variable values from • the preceding run, or • from the default values in the method if the run is inserted at the beginning of the scouting series |
| Add | to add a scouting run if there are no runs previously in the scheme. Default values will be used for the first run. to add a scouting run after all other runs in the series. The new run inherits the values from the run that precedes the new run. |
| Series | to set up a series of runs with differing inputs. |
| Edit Variable | to rename or delete a variable, or change a variable into a detail variable. |

How to set up or edit a Scouting Scheme

The table below describes how to set up or edit a scouting scheme.

| Step | Action |
|------|---|
| 1 | Create a method. If you do not use a template or wizard, define appropriate variables in the method. |
| 2 | In the Run Setup, click the Scouting tab. |
| | Result: If no scouting variables have been previously defined, the Scouting Variables dialog box is displayed. If not, click the Define button. |

| Step | Action |
|------|--|
| 3 | Select the variables you want to scout. If you cannot find the variable you want, use the following options: |
| | • Show details to display variables created with the Visible in details only option. |
| | • Show unused variables to display all variables, including those that are not used in the method. |
| | Click 0K . |
| | Result: The selected scouting variables will appear in a column, with default valued inserted. |
| 4 | Make any required changes in the scouting variable values. |
| 5 | To add a new Run column, click the Add button to copy the values from the last run column, and then change variable values as required. |
| 6 | Repeat steps 3 to 5 as required until you have defined all the scouting runs you need. |
| 7 | To exclude scouting runs from the default scouting scheme, right- click the heading of the run. To include the scouting run again, right- click it again. |
| 8 | Click the Start Protocol tab in Run Setup. |
| | Select from the following options: |
| | • The Scouting box: Select this to display the Scouting page at the start of a run. This allows the operator to adjust the values for scouting variables before the method run starts. |
| | • The First run only button: Select this to display the start protocol before the first run only. The settings entered in the Start Protocol for the first run will apply throughout the run, and the scouting series will be performed automatically without user intervention. |
| | • The All runs button: Select this to display the Start Protocol before each run in the scheme. This gives the operator an opportunity to change variable values or fill the sample loop before each run. |
| | <i>Note:</i> The operator must then click the Start button before each run. |

How to set up series

The table below describes how to set up series.

| Step | Action |
|------|---|
| 1 | Select a cell on the Scouting tab, and click the Series button. Result: The Insert Series dialog box is displayed. |
| 2 | In the Insert Series dialog box, type the selected series values (within the specified range limits), separated by commas, and click OK. <i>Result:</i> A new set of runs is inserted on the Scouting tab with the values provided. |

How to delete or rename scouting variables

Scouting variables can be deleted or renamed in the scouting scheme in the same way as in the Variables tab. The table below describes how to delete or rename a variable in the **Scouting** tab.

| Step | Action |
|------|---|
| 1 | Click the Edit Variable button on the Scouting tab. |
| | Result: The Edit Variables dialog box opens. The variables are listed alphabetically. |
| 2 | Select the variable to edit. |
| 3 | Rename Type a new variable name in the New name text box. Click the Rename button. |
| | Result: The variable is renamed. Delete Click the Delete button. Confirm that you want to delete the variable. Result: The variable is deleted. |

How to change a scouting variable into a detail variable

Detail variables are indicated with a **D** to the left of the **Variable** column on the **Scouting** tab. The table below describes how to set up a detail variable.

| Step | Action |
|------|--|
| 1 | Click the Edit Variable button on the Scouting tab. |
| | Result: The Edit Variables dialog box opens. The variables are listed alphabetically. |
| 2 | Choose the variable to be changed. |

| Step | Action |
|------|--|
| 3 | Select the Set visible in details only checkbox. |
| | Click the Close button. |
| | Result: The variable is indicated by a D . |
| | Note: De-select the checkbox to make the variable fully visible again. |

How to copy contents to factorial design programs

The contents of the **Scouting** tab can be copied and pasted into a third-party factorial design program. Processed values can then be pasted back into the **Scouting** tab. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Select the text, etc., that you want to copy. |
| 2 | Press Ctrl+C. |
| 3 | Place the cursor where you want to insert the copied text. |
| 4 | Press Ctrl+V. |

7.2 How to define different columns for scouting

Instruction

You can define different columns for use in the various scouting runs. However, in selecting a different column, other variables may also be changed between runs. The table describes how to scout columns.

| Step | Action | |
|------|---|--|
| 1 | Choose a method with a column (not Any). Alternatively, you can have a method with CV as the main base and a column (not Any) selected as a variable called "column". | |
| 2 | Click the Define button on the Scouting tab. | |
| | Result: The Scouting Variables dialog box is displayed. | |
| 3 | Select Column and click OK. | |
| 4 | Click the Column drop-down menu item within the desired run. | |
| | Result: A menu is displayed. | |
| 5 | Select a column. | |
| | Result: The Column Value Update dialog box is displayed. | |
| 6 | The dialog box asks you whether you want to update the instructions with column default values. Select one of the following: | |
| | Yes | |
| | The method for the scouting run is updated with variable parameter values for the selected column, consisting of UV average time, pressure limit, flow rate, etc. These parameter values are added to the scouting variables on the Scouting tab. Note that the updated parameter values may differ from the values for the same variables in other scouting runs. | |
| | No | |
| | No changes are made. The method retains the parameter values corresponding to the column that were either originally selected during creation of the method, or included in an earlier version of the method on the Scouting tab for which the default values for that column were accepted. | |
| | <i>Note</i> : If the method contains a linear flow rate instruction, the user can keep the linear flow rate by selecting a check box in the dialog box. | |

8 MethodQueues

Introduction

MethodQueues provide a means for linking several methods together, on the same or different systems. For example, if a system wash procedure is programmed in a separate method, it can be linked in a **MethodQueue** to a series of different process methods, ensuring that the same wash procedure is used before every process. Alternatively, the product of a separation on one system might form the starting material for a separation on the next, allowing fully automated multi-step processing.

In this chapter

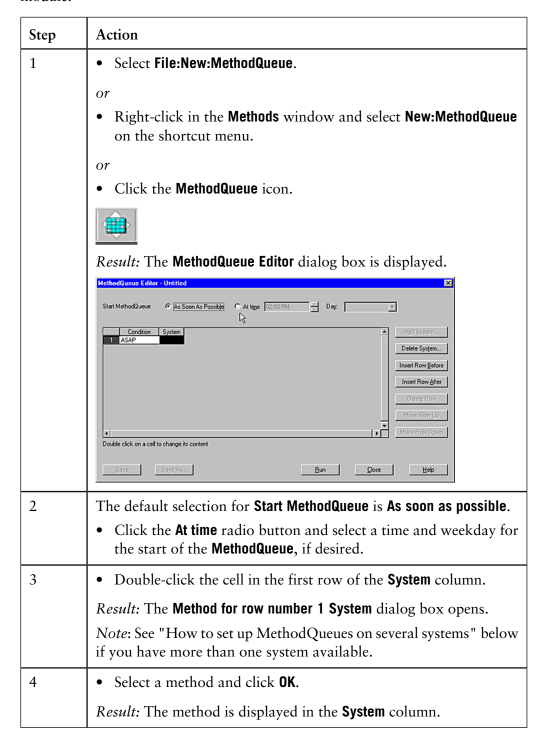
This chapter contains these topics:

| Topic | See |
|---------------------------------|-----|
| How to create a new MethodQueue | 8.1 |
| How to edit a MethodQueue | 8.2 |

8.1 How to create a new MethodQueue

Instruction

The table below describes how to create a **MethodQueue** in the **UNICORN Manager** module.



| Step | Action |
|------|---|
| 5 | Click the Insert Row After button and repeat steps 3 and 4 to add more methods to the MethodQueue. |
| | <i>Note:</i> The timing of MethodQueue steps performed on different systems can also be controlled by the Ready instruction in the method (see "Relative timing of steps" below). |
| | By default, each method step will start as soon as possible (ASAP) after the completion of the previous method step. Use the Condition cell of the chosen method to set another time interval for starting a selected step. |
| | • In the Conditions column, double-click the cell for the method to be delayed. |
| | Result: The Condition for row number X dialog box opens. |
| | Condition for row number 4 As soon as possible Wait Hours: Minutes: Previous Row Next Row Next Row Note: Use the Previous Row and Next Row buttons to select other |
| | methods for editing. |
| | • Click the Wait radio button, select the number of hours and minutes that the method is to be delayed and click OK . |
| | <i>Result</i> : The execution of the MethodQueue will be held for the selected number of hours and minutes and then resume. |
| | • Click the Save button to save the method. |
| | Result: The Save MethodQueue dialog box opens. • Type a file name and click the Save button. |

How to set up MethodQueues on several systems

If you have more than one system available, the **System** column will not be displayed at first in the **MethodQueue Editor**. The table below describes how to set up a **MethodQueue** for several systems.

| Step | Action | |
|------|--|--|
| 1 | • Click the MethodQueue icon. | |
| | Click the Add System button and select a system for the first MethodQueue step from the Add System dialog box. | |

| Step | Action |
|------|---|
| 2 | • Repeat this for each system when you want to use a different system in the MethodQueue . |
| | Result: Another system column will be added for each additional system. |

Relative timing of steps

The setting of the **Condition** dialog box (reached by double-clicking a **Condition** cell in the **MethodQueue Editor** dialog box), determines the relative timing of the steps of a **MethodQueue**. If successive methods are run on the same system, the timing set in **Condition** applies from the completion of one method to the start of the next.

If successive methods are run on different systems, you can use the **Ready** instruction in one method to trigger the start of the next method. In this way, you will be able to start the next method before the current method has ended. The **Condition** setting then applies from the **Ready** instruction to the start of the triggered method. This is useful in situations where a method on one system prepares the starting material for the next, and then continues to wash the system. See the example below:

| Instruction to System 1 | Instruction to System 2 |
|-------------------------|-------------------------|
| Apply sample | |
| Eluate | |
| READY | Apply sample |
| Wash | Eluate |

Unattended operation of the MethodQueue

The **Start Protocol** for each method step in the **MethodQueue** is displayed when the corresponding method is run. If you want the **MethodQueue** to operate unattended you must ensure that the methods do not include a **Start Protocol**.

See 5 How to create a method on page 79 for more information.

How to hold a method in queue while the system is busy The table below describes how you can create a **MethodQueue** if you try to start a new method run while the system is still busy with another method run.

| Step | Action |
|------|--|
| 1 | Right-click on the method in the UNICORN Manager module and select Run:system name on the shortcut menu. Result: The System Busy dialog box opens. |
| | Cannot start the method because the system is occupied. You have three options: Mait until the system is free and start the method again Add the method to a MethodQueue that will execute as soon as the system is free Add the Method to a MethodQueue and open the MethodQueue gditor NOTE! This method contains a start protocol that will be executed when the method starts. |
| 2 | Select the Add the method to a MethodQueue that will execute as soon as the system is free option. Click OK. |
| | Result: A MethodQueue will automatically be created in the default queue folder. The name of the MethodQueue will be the same as the method name, followed by a five-digit sequence number. |
| 3 | The method will be executed as soon as the system is free. Note: A warning note is displayed in the System Busy dialog box if the method includes a Start Protocol. The Start Protocol must be completed at the start of the method run before it can be executed. |

8.2 How to edit a MethodQueue

Method Queues are saved in a separate folder

MethodQueues are saved in a separate folder within the folder that you specified when you saved the **MethodQueue**. The **MethodQueue** folder is represented by a special icon in the **Methods** window of the **UNICORN Manager**.



A **MethodQueue** folder contains the **MethodQueue** definition and copies of all included methods.

How to edit a MethodQueue file

The **MethodQueue** files are *copies* of the original method files. If changes are made in the original method, these will not affect the method in the **MethodQueue**.

To avoid confusion between different versions of method files, make sure that **MethodQueue** definitions always contain updated methods. To implement changes in a **MethodQueue** method, do one of the following:

- Edit the method in the MethodQueue folder,
 or
- Edit the original method, then use the **MethodQueue** editor to update the **MethodQueue**, and replace the old method with the changed version.

Instruction

The table below describes how to edit an existing **MethodQueue**.

| Step | Action |
|------|---|
| 1 | Right-click the selected MethodQueue folder icon in the UNICORN Manager , and select Edit from the displayed menu. |
| | Result: The MethodQueue Editor dialog box is displayed. MethodQueue Editor - MQTest3 Start MethodQueue |

| Step | Action |
|------|---|
| 2 | Select a table row to edit and do the following as required: |
| | Double-click the System cell and select a new method from the Method for row dialog box. |
| | Double-click the Condition cell and edit the delay time for the method. |
| | • Click the Add System button to add a new system to the queue and use it for a MethodQueue step. |
| | • Click the Delete System button to remove a system and all associated methods from the MethodQueue . |
| | Click the Insert Row Before or Insert Row After buttons to add new rows before or after the selected row. |
| | Click the Delete Row button to remove the selected row. |
| | Click the Move Row Up or Move Row Down to move the selected row one step up or down in the queue. |
| 3 | Click the Save button. Click the Run button to execute the MethodQueue immediately or |
| | the Close button to close the dialog box. |

How to perform method runs 9

Introduction

This chapter describes how to perform and monitor different kinds of method runs from the System Control module. It also describes how to control the system with manual commands and instructions.

In this chapter

The chapter contains these sections:

| Topic | See |
|----------------------------------|-----|
| How to start a method run | 9.1 |
| How to monitor a method run | 9.2 |
| Manual system control | 9.3 |
| How to perform a scouting run | 9.4 |
| How to perform a MethodQueue run | 9.5 |
| If the network connection fails | 9.6 |

9.1 How to start a method run

Before you start

Before you start a method, make sure that

• the correct system is connected in control mode

Note: If the system is connected via a **CU-950 Advanced** unit, the Ethernet connection must not be broken during the start-up phase of the method run.

How to start from the UNICORN Manager

You can start a method from the UNICORN Manager in two ways:

- Select a method in the Methods window and select File:Run.
- Select a method, right-click and select **Run** from the displayed menu.

How to start from System Control

The table below describes how to start a method run from **System Control**:

| Step | Action |
|------|--|
| 1 | Select File:Run |
| | or |
| | click the Run button. |
| | Result: The Run dialog box is displayed. |
| | <i>Note</i> : The Run button will open the method that was used for the previous run, if a run has been performed since you logged on. |
| 2 | Select a method and double-click the method icon. |
| | <i>Result</i> : The method run starts. If the method includes a Start Protocol this must be completed before the actual method run starts. Se further instructions below. |

How to add methods to the File menu

For methods that are used frequently (for example column cleaning methods or routine separations), it may be convenient to define the methods as commands in the **File** menu.

The table below describes how to define a method as a command:

| Step | Action |
|------|--|
| 1 | Choose File:Menu in System Control and select the required method. |
| 2 | Click the Add button and click OK . |
| | Result: The method name will appear as a command in the File menu. If you choose the command, the method will start. |

How to start an instant run

You can start a method template or wizard directly if your system has defined templates or wizards.

To do this, either

• click the **Instant Run** icon in the **UNICORN Manager** toolbar



or

• select File:Instant Run in System Control.

How to use the Start Protocol

If the method is defined with a **Start Protocol**, this will be displayed before the method actually starts.

The table below describes how to use the **Start Protocol**:

| Step | Action |
|------|--|
| 1 | Start the method run. |
| | Work through the start protocol, answering questions as required. |
| | The start protocol items that can be displayed are described in 6.5.14 The start protocol tab on page 151. |
| | • As each screen is completed, click the Next button to move to the next screen or the Back button to return to the previous screen. |
| 2 | Click the Start button in the last window to start the run. |

Confirm/Sign authorization for the Start Protocol

If there are any questions in the **Start Protocol** that require authorized confirmation, you will be asked for a user name and password when you attempt to leave the screen containing the questions. Only users with **Confirm/Sign** authorization may authorize answers to such questions. Each question that requires an authorization must have a separate authorization.

How to start a method when the system is busy

If the system is busy with a method run in progress, you can still start a new method. You will have the option to place the method in a **MethodQueue**, which can be executed as soon as the system becomes available again. The table below describes how to do this.

| Step | Action |
|------|--|
| 1 | While a method run is in progress, right-click on the next method you want to run and select Run:System . |
| | Result: The System Busy dialog box opens. |

| Step | Action |
|------|---|
| 2 | Select the Add the method to a MethodQueue that will execute as soon as the system is free option. Click OK. |
| | Result: A MethodQueue will automatically be created in the default queue folder. The name of the MethodQueue will be the same as the method name, followed by a five-digit sequence number. |
| 3 | The method will be executed as soon as the system is free. Note: A warning note is displayed in the System Busy dialog box if the method includes a Start Protocol. The Start Protocol must be completed at the start of the method run before it can be executed. |

Note: See 8.2 How to edit a MethodQueue on page 187 for more information.

How to monitor a method run 9.2

Introduction

This section describes how to monitor a method run by using the **System Control** module and how to customize the different panes.

In this section

The table shows the topics that can be found in this section.

| Topic | See |
|---------------------------------------|-------|
| How to customize System Control panes | 9.2.1 |
| The Run Data pane | 9.2.2 |
| The Curves pane | 9.2.3 |
| The Flow Scheme pane | 9.2.4 |
| The Logbook pane | 9.2.5 |

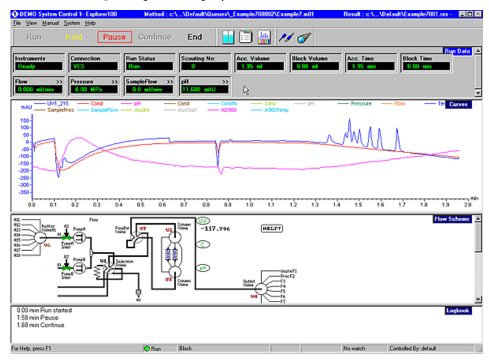
9.2.1 How to customize System Control panes

Introduction

The **System Control** module displays the status of the current system. On the Windows taskbar, there may be up to four **System Control** modules available that can be connected to different systems. Separate systems may be controlled and displayed independently of each other.

Illustration

The illustration shows the **System Control** module with the **Run Data**, **Curves**, **Flow scheme** and **Logbook** panes displayed.



How to select what panes to display

Each **System Control** module displays up to four panes for monitoring different aspects of the run. To select what panes to display, either

• click the Customize Panes icon,



01

• choose View:Panes.

How to customize **System Control** panes

Change the size

Select a split-bar and drag up and down to change the size of a specific pane.

Maximize, restore or hide

Right-click a pane and select the appropriate option to:

- maximize,
- restore

or

• hide the pane.

9.2.2 The Run Data pane

Description

The **Run Data** pane displays the current values for selected run parameters. The update interval is defined in the system strategy.

The figure below displays an example of the **Run Data** pane:



How to change the appearance of the pane

The appearance of the pane can be changed so that it includes more or fewer data displays. The table below describes how this is done:

| Step | Action |
|------|--|
| 1 | In System Control, select View:Properties |
| | or |
| | right-click on the pane and select Properties on the menu. |
| | Result: The Properties dialog box is displayed. |
| 2 | Select the Run Data Groups tab and, if desirable, do one or more of the following: |
| | Select an available group to be displayed in the list to the left. |
| | • <i>Edit</i> an available group: Select the group from the list on the left, and click the Edit Group button. Modify the included readings in the list to the right, and click OK . |
| | • <i>Create</i> a new group: Click the New group button and select the readings that you want to view from the list. Enter a name for the group, and click OK . |
| | • <i>Delete</i> a group: Click the Delete Group button and select a group in the Delete Layout dialog box, click OK and confirm the deletion. |
| 3 | Select the run data parameters that you want to display in the list to the right. |
| 4 | Click OK to view the selected items in the Run Data pane. The name of the selected layout replaces the default layout name Run Data . |

How to change text color or text background

The table describes how to change the text color or background in the displayed reading boxes.

| Step | Action |
|------|--|
| 1 | Right-click on the pane and select Properties. |
| | Result: The Properties dialog box is displayed. |
| 2 | Select the Run Data Color tab. |
| 3 | Click the Text or Background buttons. |
| | Select a new color, and click OK . |
| | Result: The color change is displayed in the test field. |
| 4 | Make further adjustments to the colors as appropriate. |
| 5 | Click OK to apply the changes. |

How to set the pressure units

If the **Pressure** reading box is displayed in the **Run Data** pane, you can set the displayed units. The table below describes how this is done:

| Step | Action |
|------|---|
| 1 | Right-click on the Pressure reading box to display the menu. |
| 2 | Select Set Unit and the appropriate unit (MPa , bar or psi). Result: The selected unit is displayed. |

How to view and select manual instructions

Some strategies directly link specific manual instructions to the reading boxes in the **Run Data** pane. This is indicated by a double arrow (>>). A particular reading box can have one or more instructions attached to it. In cases where there is more than one instruction, one of the instructions is the main instruction.

There are two ways to view the manual instructions:

Option 1:

• Double-click the reading box.

Result: The dialog box for manual instructions is displayed, showing the instruction, or main instruction if there is more than one.

Option 2:

- Right-click the reading box. Select **Instructions** in the displayed menu. Another menu shows the specific manual instruction(s).
- Click an instruction to select it.

Result: The dialog box for manual instructions is displayed in which you can execute the appropriate command.

- 9 How to perform method runs 9.2 How to monitor a method run 9.2.2 The Run Data pane

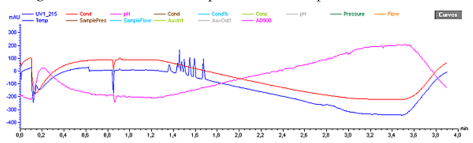
For more details on how to use manual instructions, please see 9.3.2 Manual instructions on page 212.

9.2.3 The Curves pane

Introduction

The **Curves** pane of the **System Control** module displays monitor signal values graphically.

The figure below shows an example of the **Curves** pane:



How to select curves to be displayed

You can decide which curves you want to display in the **Curves** pane. Curves will only be shown for components present in the chromatography system.

The table describes how to select the curves to be displayed on the screen.

| Step | Action |
|------|--|
| 1 | In System Control, select View:Properties. Result: The Properties dialog box is displayed. |
| 2 | Select the Curves tab. Note: The curves in the list are those for which Store is set to On in the system settings, together with any reference curves defined in the method. |
| 3 | In the Display curves list, select the curves you want to display. If you want all curves to be displayed, click the Select All button. If you do not want any curves to be displayed, click the Clear All button. Click OK . |

How to display a vertical marker line

The table below describes how to display a vertical marker line:

| Step | Action |
|------|---|
| 1 | Right-click the Curves pane and select Marker . |
| 2 | Drag the marker line with the mouse. Result: Where the line bisects the curve, the X-axis and Y-axis values are displayed at the top right corner of the pane. |

Note: Right-click and select **Snapshot** to record the marker position values. See 2.2.7 Snapshots on page 41 for more information about the Snapshot function.

How to set a reference point

When the vertical marker is displayed, you can set a reference point to display curve data. The table describes how to set a reference point:

| Step | Action |
|------|---|
| 1 | Display a Marker in the Curves pane. Right-click and select Set Marker Ref. Point to define a reference point for the marker position. |
| 2 | When the marker is moved from the reference point, the X-axis and Y-axis values for the new position are displayed together with: the new position in relation to the position of the reference point, the minimum, maximum and average values for the curve interval between the reference point and the new position. |

How to change the curve colors and styles

The **Curves** pane displays graphs for the selected curves in different colors, with any reference curves included with the method as dashed lines.

The table below describes how to change the curve colors and styles:

| Step | Action |
|------|--|
| 1 | Select View:Properties. |
| | Result: The Properties dialog box is displayed. |
| 2 | Select the Curve Style and Color tab. |
| 3 | Select a curve from the Curve list. Select an appropriate color and style. |

How to change the scale of the Yaxis

In most cases, the Y-axis is automatically scaled for each of the curves. Values on the Y-axis apply to the curve with the same color as the axis markings. To get the correct Y-axis, click the legend. The table below describes how to fix the scale of individual curves.

| Step | Action |
|------|--|
| 1 | Select View:Properties. |
| | Result: The Properties dialog box is displayed. • Select the Y-axis tab |
| | • Select the Y-axis tab. |

| Step | Action |
|------|---|
| 2 | Select the appropriate curve. Select Fixed and type a minimum and maximum range in the fields within the specified limits. |
| 3 | Repeat step 2 for other curves if needed. |
| 4 | Click OK . |

How to change the scale of the Xaxis

The table below describes how to change the scale of the X-axis:

| Step | Action |
|------|--|
| 1 | Select View:Properties. |
| | Result: The Properties dialog box is displayed.Select the X-axis tab. |
| 2 | Select the appropriate base, Time or Volume . |
| | <i>Note:</i> Curves are collected in time and recalculated for display in volume. Thus, the resolution of the two bases may appear slightly different. |
| 3 | Select the appropriate Axis scale : |
| | Total will show the curves as far as they have come in the run. |
| | • Window allows you to set the portion of the total pane to be displayed, either in minutes or ml depending on the selected base. |
| | • Adjust retention zero to injection sets the retention value to zero at the point of the first injection. |
| | • Click OK . |

How to switch between time and volume units

• Click the legend of the X-axis

or

• right-click and select Base Type

to switch the display between time and volume units. The run is controlled according to the time/volume base defined in the current block, regardless of the base in the curves display.

How to zoom in the Curves pane

The table below describes how to zoom in on a selected region of the curve pane:

| Step | Action |
|------|--|
| 1 | Press and hold the left mouse button and drag a rectangle out on the screen to encompass the area to be viewed. Release the mouse button. Result: The display is now zoomed in on the selected area. |
| 2 | Repeat the process for further magnification of selected areas. |

How to zoom out

To reduce the scale of the zoom, right-click in the **Curves** pane, and select one of the following options:

- **Undo Zoom**: reverses each zoom-in action a step at a time.
- **Reset Zoom:** reverses all zoom-in actions to the default scale.

How to select curve pressure units

If the **Pressure** curve is displayed in the **Curves** pane, you can set the displayed units. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Right-click in the Curves pane, and select Properties in the displayed menu. Result: The Properties dialog box is displayed. |
| 2 | Select the Y-Axis tab. |
| 3 | Select the Pressure curve and select the appropriate Pressure unit button. Click OK . |

How to edit text in the Curves pane

You can select the way that text is aligned for the **Logbook** and **Fraction** curves. You can also select to show only part of the **Logbook** information. The table below describes how to do this:

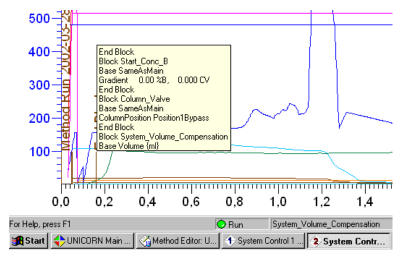
| Step | Action |
|------|--|
| 1 | Right-click in the Curves pane, and select Properties in the displayed menu. |
| | Result: The Properties dialog box is displayed. |
| 2 | Select the Curve Style and Color tab. |

| Step | Action |
|------|--|
| 3 | Select the following: |
| | Logbook or Fraction curve in the Curve list as appropriate. |
| | Select the appropriate Logbook text alignment or Fraction text alignment option: |
| | - Horizontal |
| | - Vertical |
| | - Fly over (displays the text if you place the mouse pointer over the generated mark). |
| 4 | To filter the type of Logbook information overlaid on the Curves pane, do the following: |
| | Click the Filter button. |
| | Result: The Filter Logbook dialog box is displayed. |
| | Select the appropriate check boxes and set the maximum block depth. |
| | Click OK to return to the Curve style and Color tab. |
| 5 | Click OK . |

How to view the complete logbook information

At some breakpoints there can be more logbook information than what is possible to conveniently display in the **Curves** pane. The additional information that is not displayed is indicated by an arrow point symbol by the break point.

• Hold the mouse cursor over the break point to display the complete information in a flyover text box, as shown in the illustration below.



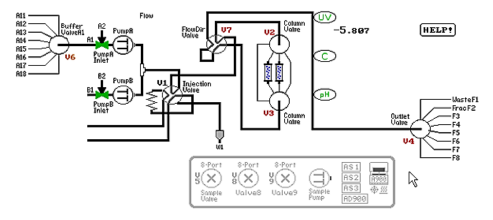
9.2.4 The Flow Scheme pane

Introduction

The flow scheme is a graphical representation of the chromatography system that shows the current status of the run. During a run, the flow scheme displays open flow path(s) in color and monitor signals with numerical displays.

Illustration

The illustration below shows an example of a flow scheme for a run:



How to stretch a flow scheme

The flow scheme can be stretched to fit the screen. To do this, right-click in the pane and select **Stretch** in the shortcut menu.

How to view and select flow scheme manual instructions

Some strategies link specific manual instructions directly to the components in the flow scheme pane. The components in the flow scheme that are associated with instructions are indicated with double arrows (>>). A particular component can have one or more instructions attached to it. In cases where there is more than one instruction, one of the instructions is the main instruction.

To display and select instructions:

double-click a component

or

• right-click a component, select **Instructions** and an instruction in the shortcut menu.

Result: The manual instructions dialog box for the selected instruction type opens.

9.2.5 The Logbook pane

Introduction

All actions (including method start and end, base instruction, method instructions and manual interventions such as **Pause** or **Hold**) and unexpected conditions such as warnings and alarms are logged for every run, with date, time and current user name where appropriate. The logbook thus provides a complete history of any given run. The log is saved in the result file.

Illustration

The illustration below shows an example of the **Logbook** pane:

```
0.00 min Method Run 3/26/2002_8:47:16 AM, Method : idtest, Result: v\_\Nklas\/idtest001.res
0.00 min Batch ID: 4E9820F4-380A-11D6-AC46-00D08728BCC0
0.00 min Base CV, 0.10 (m)
0.05 min Block Flow, Rate
0.05 min Base SameAsMain
0.05 min Block Bolock
0.05 min Block Column_Pressure_Limit
0.05 min Block Column_Pressure_Limit
0.05 min Block Column_Pressure_Enabled, 4.00 MPa, 0.00 MPa
0.05 min Alorm_Pressure Enabled, 4.00 MPa, 0.00 MPa
0.05 min Block Start_Instructions
0.05 min Block Start_Instructions
0.05 min Block Start_Instructions
0.05 min Block Start_Instructions
0.05 min Wavelength 280 nm, OFF, OFF
0.05 min Avereignit TimeUV 2.56 sec
0.05 min Avereignit TimeUV 2.56 sec
0.05 min Block Block
0.05 min Block Block
```

Note: The second logbook line is the **BatchID** that is automatically generated.

Autoscroll

The **Logbook** pane can autoscroll to display the latest entries. Right-click in the pane, and select **Autoscroll**. You can also select the **Autoscroll** option in the **Properties** dialog box (**View: Properties** and select the **Logbook** tab).

How to filter the logbook contents

You can choose to display only selected items in the logbook. The table below describes how to activate the filter.

| Step | Action |
|------|---|
| 1 | Right-click in the Logbook pane and choose Properties . |
| | Result: The Properties dialog box opens. |
| 2 | Choose the Logbook tab. |
| | • Select the items you want to display in the logbook (all items are selected by default). |
| | Click the OK button. |
| | Result: Only the selected items will be displayed in the logbook. The Logbook title in the upper right corner will show the text (Filter on) to indicate that not all items are visible. All items will still be logged in the result file. |

9 How to perform method runs

9.2 How to monitor a method run 9.2.5 The Logbook pane

How to find logbook text entries

The logbook can be searched for specific text entries. The table below describes the function:

| Step | Action |
|------|--|
| 1 | Right-click in the Logbook pane and choose Find . Result: The Find dialog box opens. |
| 2 | Type the text you want to locate. Select search criteria if necessary. Click 0K. Result: The located logbook entry is highlighted. |

9.3 **Manual system control**

Introduction

This section describes how to control the system with manual commands and instructions.

In this section

This section contains these topics:

| Topic | See |
|----------------------------|-----|
| The toolbar and status bar | |
| Manual instructions | |
| Alarms and warnings | |

9.3.1 The toolbar and status bar

Toolbar buttons

The toolbar at the top of the **System Control** module contains three sets of buttons:

- Manual Direct Commands buttons for starting and stopping the run
- **Windows** buttons to access dialog boxes for pane selection, documentation and layout properties
- System Access buttons to control the system connection.

Show and hide

The toolbars can be shown and hidden by choosing **View:Toolbars** and selecting the relevant boxes.

The figure below shows the toolbar:



Manual Direct Commands

The available **Manual Direct commands** buttons in **System Control** are dependent on the control status of the connection. The table below shows when each button is available:

| Control Status | Available buttons |
|----------------|----------------------|
| End | Run |
| Running | Hold, Pause, End |
| Manual | Run, Pause, End |
| Hold | Pause, Continue, End |
| Method pause | Hold, Continue, End |
| Manual pause | Run, Continue, End |

Direct command button functions

The table below describes the functions of the Manual direct command buttons.

| Button | Function |
|--------|--|
| Run | Opens the Run dialog box, which shows all available methods, as the first step in a method run. If a method is loaded, Run Setup opens. The run will start immediately if a start protocol isn't part of the method. |

| Button | Function |
|----------|---|
| Hold | Suspends execution of a method, but continues |
| | • to pump liquid at the current flow rate and eluent concentration settings. All settings remain unchanged. |
| | to increase accumulated time and volume. |
| | Method instructions are not executed until the Continue button is pressed. |
| Pause | Behavior of the Pause button is strategy-dependent. The Pause button suspends execution of a method and stops all pumps so that the system comes to a standstill. |
| | For ÄKTAdesign systems, valves remain in the position they were in before the pause. |
| | Accumulated time and volume is not increased during a Pause . |
| | Method instructions are not executed until Continue is pressed. |
| Continue | Resumes execution of a paused or held method. |
| End | Terminates method execution |
| | Puts the system into an End state. |
| | Note: You can choose to save the partial result or discard it. |

Note: The commands can also be found on the Manual menu.

Windows buttons The table below describes the functions of the Windows buttons:

| Button | Function |
|--------|---|
| | Opens a dialog box where you can choose which window panes to display. This button is equivalent to the menu command View:Panes . |
| | Opens the documentation pages. Run notes can be entered on the Notes tab and settings can be changed. Other tabs are displayed for information only. This button is equivalent to the menu command View:Documentation . |

| Button | Function |
|--------|--|
| Lili | Opens the properties pages. This button is equivalent to the menu command View:Properties . |

System Access buttons

There are two functions of the **System Access** buttons:

Disconnect/Connect system



The **Disconnect** button is used to disconnect the system and leave it in a locked or unlocked state.



The **Connect** button connects the system.

Leave/Take control of the system



The **Leave control** button leaves the system in a locked or unlocked state.



The **Take control** button takes control of the system.

Status bar, connection status

The status bar displays a message indicating the connection status of the window. The table below describes the different messages:

| Message | Connection status |
|------------------------------|---|
| Controlled by: <user></user> | The indicated user has a control mode connection to the system. Other users can establish a view mode connection. |

| Message | Connection status |
|--------------------------|--|
| Locked by: <user></user> | The indicated user has left the system in a locked state. Users who can supply the required password can unlock the system and establish a connection. The password is case sensitive. |
| | Note: It is possible to unlock with the "lock" password or with the UNICORN logon password. Anyone who uses the UNICORN logon password must have Unlock systems access rights. The "lock" password is the password entered by the user who locked the system. |
| System is available | Any user can establish a connection. |

status

Status bar, Watch The status bar displays a message indicating if a Watch is active in the method.

• Click the Active watch status message to open the Watch dialog box with information about the active Watch instruction.

9.3.2 Manual instructions

Introduction

The chromatography system can be controlled with manual instructions issued from the **Manual** menu in the **System Control** module. The available instruction options are dependent on the strategy.

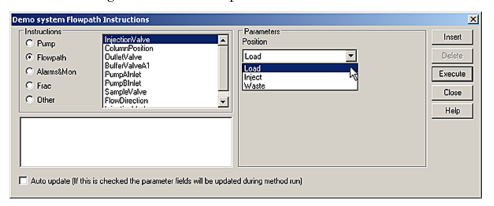
Manual instructions during a method run

Manual instructions can be issued while a method is running. A manual setting applies until the next method instruction of the same type is executed

Example: A manual **Flow** instruction will set the flow rate until the next **Flow** instruction in the method is executed. Manual instructions that you issue during a method are recorded in the logbook for the method run.

The manual instructions dialog box

The **Manual** menu in **System Control** opens a dialog box similar to the **Instruction box** in the **Method Editor**. The name of the connected system is displayed on the title bar of the dialog box. See an example in the illustration below:



Note: The parameter values will be updated continually during the run if the **Auto update** checkbox is selected.

Column protection

Your user attributes may include a requirement to always set pressure alarms.

| Step | Action |
|------|--|
| 1 | When you try to execute a pump instruction the Column protect mode dialog box opens. |
| 2 | Click the Yes button in the dialog box to select a column and retrieve the correct maximum pressure value. Click OK to close the column list. Click the Insert button to add the Alarm_Pressure instruction. |
| 3 | • If necessary, repeat step 2 to add an Alarm_SamplePressure instruction. |

How to use manual instructions

Manual instructions are entered in the same way as method instructions from the dialog box in the Method Editor. The table below describes how to add a manual instruction:

| Step | Action |
|------|--|
| 1 | Select an instruction group and a component in the Instructions field. Select instruction parameters in the Parameters field. |
| 2 | Click the Insert or Execute buttons as needed. (See the descriptions of the different functions below) |

manual instructions dialog box

The buttons of the The table below describes the functions of the manual instructions buttons:

| Button | Function |
|---------|---|
| Insert | This button places the current instruction in the list at the bottom left of the dialog box. |
| Delete | This button deletes the selected instruction from the current list only. One instruction can be deleted at a time. |
| Execute | This button |
| | • executes all instructions in the list at the same time |
| | or |
| | • executes the currently marked instruction if the list is empty. |
| | <i>Note:</i> Although all instructions are executed simultaneously, some (for example gradient and fraction instructions) may take some time to complete in the liquid handling module. |
| Close | If you click the Close button without first clicking the Execute button, commands in the list |
| | will not be executed |
| | will be deleted from the command list. |

How to save manual results

When you choose to run the system manually - as opposed to a **Method run** - the results are automatically stored in a folder called **Manual Runs**. The **Manual Runs** folder stores the ten most recent results from your manual runs. To save a result file from the **Manual Runs** folder more permanently, you need to move or copy it to another location.

An alternative way to save the results from a manual run is to record the results manually in a result file. The table below shows how to do this:

| Step | Action |
|------|--|
| 1 | Choose Manual:0ther. |
| | • Select the instruction Record On at the beginning of the run. |
| 2 | • Click the Execute button. |
| | Result: UNICORN will prompt for a result file name. |

9.3.3 Alarms and warnings

Introduction

Alarms and warnings are displayed regardless of the activity currently in progress in UNICORN. You will be notified of an exceeded limit in a running system even if you are developing a method, evaluating data or monitoring a method run on a different system. Warnings and alarms are also recorded in the logbook for the run.

Limits for monitor signals

The system settings determine the acceptable limits of monitor signals during a run. The limits can also be set for the current run by an instruction in the method. Limits set with a method instruction override the limits set in system settings. If these limits are exceeded in a run, a warning or alarm dialog box is displayed on the screen.

Effects of alarms and warnings

Alarms and warnings have different effects on the system:

- Warning: The run continues.
- *Alarm*: The system is paused.

In a network system

In a network installation, alarms and warnings are displayed on the controlling station and all stations viewing the system. An alarm can be acknowledged only from the computer connected in control mode. Alarms are displayed but cannot be acknowledged on computers connected in view mode.

9.4 How to perform a scouting run

More information on scouting runs

See 7.1 How to set up a scouting scheme on page 176 for information on how to set up scouting runs.

Instruction

The table below describes how to perform a scouting run:

| Step | Action |
|------|--|
| 1 | Start the method (see 9.1 How to start a method run on page 190). <i>Result:</i> The Start Protocol will display the scouting scheme as defined in the method (assuming that the Scouting box is selected on the Start Protocol tab of the Run Setup). |
| 2 | Check through the settings for the scouting scheme in the Scouting tab, and if required, do the following: • Change the scouting variable values. • Right-click the top of the Run column to toggle the run status between Run and Excluded . |
| 3 | Work through the rest of the Start Protocol . Click the Start button. |

Results of a scouting run

The results of a scouting run are saved in a special scouting folder as defined in the **Results** tab of the **Start Protocol**. Within the folder, each run is saved in a separate result file named according to the usual naming rules (see **6.5.12 The result name tab** on page 147).

If the **Start Protocol** is displayed for each run in a scouting scheme, you are able to change the result file name during scheme execution.

How to change scouting variables during a run

At any time during a run, you can click the **View Documentation** icon in **System Control** and change the scouting variables on the **Scouting** tab for runs which have not yet been started.



- Settings for the run that is currently in progress cannot be changed.
- You can add more scouting runs to the scheme as long as the last run has not been started.
- You can use this feature to adjust variables for scouting even if the start protocol is not displayed at the beginning of each run.

9.5 How to perform a MethodQueue run

Instruction

The table below describes how to run a MethodQueue:

| Step | Action |
|------|---|
| 1 | Make sure that all systems used in the MethodQueue are connected with control mode connections. |
| 2 | Select a MethodQueue in the Methods pane in the UNICORN Manager and • choose File:Run |
| | right-click the MethodQueue icon in the Methods pane and select Run from the shortcut menu. |
| | double-click the MethodQueue icon in the Methods pane and click the Run button in the MethodQueue Editor dialog box. |
| | Result: The MethodQueue will start in accordance with the conditions defined in the MethodQueue setup. |

See **8.1** How to create a new MethodQueue on page 183 for information about how to create a **MethodQueue**.

Unattended MethodQueue operation

The **Start Protocol** for the first and each subsequent method step in the **MethodQueue** is displayed when the corresponding method is run. If you require unattended **MethodQueue** operation after the start of the first method step, make sure that subsequent method steps do not include a **Start Protocol**.

Note: If the **Start Protocol** for a method in the queue is cancelled, the **MethodQueue** is paused. Select **MethodQueue:Display Running** in the **UNICORN Manager** and **Restart** or **End** the run in the displayed dialog box.

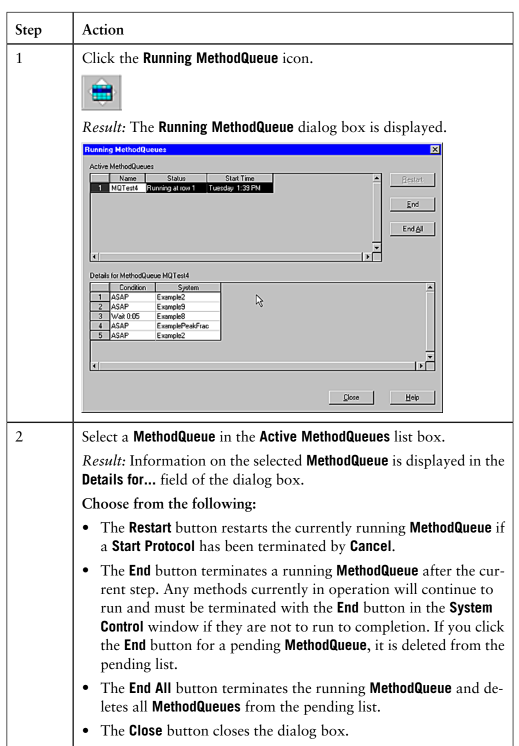
MethodQueues when the system is busy

You can choose to place a method in a **MethodQueue** if the system is already busy with another method run (See **8.1** How to create a new **MethodQueue** on page 183). In a similar manner you can also start a new **MethodQueue** while another **MethodQueue** is in progress. It will be placed in queue and executed when the first queue is completed.

How to display and edit pending and running MethodQueues

Definition: A pending **MethodQueue** is one for which **Run** has been requested but which has not yet started, either because the system is not available or because the setup time has not been reached.

The table describes how to display running and pending **MethodQueues**.



9.6 If the network connection fails

Results will be saved in the Failed folder

If the results of a method run are stored on a server or other location, and there is a network communication failure during a method run that has been started from a remote station, the method run will continue and the results will be saved in the **Failed** folder on the local station. A control mode connection can be established on the local station to control the running system. See the Administration and Technical Manual for more details.

How to view results

Introduction

A result file is automatically generated at the end of a method run and contains a complete record of the method run, including method, system settings, curve data and method run log. The **Evaluation** module offers extensive facilities for presentation and evaluation of curve data.

This chapter describes how to present the chromatograms and curves of your result file and how to create and print reports.

In this chapter

This chapter contains these sections:

| Topic | See |
|---|------|
| How to open a result file | 10.1 |
| How to use the File Navigator | 10.2 |
| Basic presentation of chromatograms | 10.3 |
| How to optimize the presentation of chromatograms | 10.4 |
| How to print active chromatograms | 10.5 |
| How to create and print reports | 10.6 |
| Run documentation | 10.7 |

10.1 How to open a result file

Introduction

All contents of the result files are opened in the **Evaluation** module. By default, the chromatograms in a run are shown as opened windows. The chromatogram window on top is the active window. There is also a minimized **Temporary** chromatogram window. See **10.3 Basic presentation of chromatograms** on page 226 for further information about chromatograms.

Note: It is not possible to open the same result file from two different locations simultaneously.

How to open a result from the UNICORN Manager

To open a result file from the **UNICORN Manager**, do one of the following:

• Double-click a result file in the **Results** window of the **UNICORN Manager**,

or

• Select a result file icon in the **Results** window of the **UNICORN Manager** and select **File:Open**,

or

• Click the **Evaluation** icon in the **UNICORN Manager**, open the **Evaluation** module and select a result file from the **Open Result** dialog box.



How to open a result in the Evaluation module

To open a result file in the **Evaluation** module:

- Do the following:
 - Select File:Open
 - Select a result file from the **Open Result** dialog box.

or

- Do the following:
 - Select View:File Navigator
 - Locate and select a result file from the **File Navigator**.

Note: See **10.2 How to use the File Navigator** on page 222 for detailed instructions on how to locate files and set up File Navigator preferences.

10.2 How to use the File Navigator

Introduction

The **File Navigator** can be used to locate and open result files in the **Evaluation** module. Recent runs are also listed based on the user preferences.

How to open the File Navigator

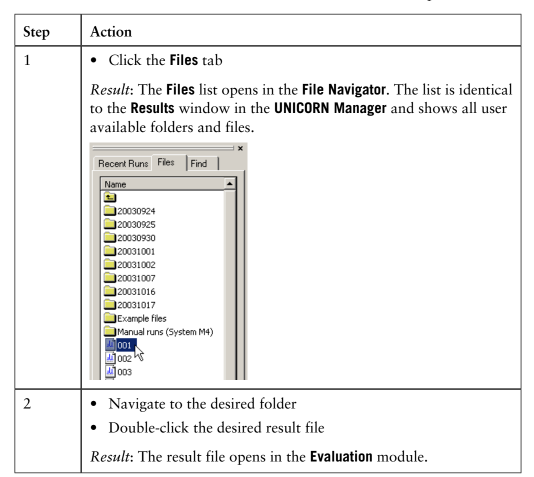
To open the **File Navigator**:

- Click the **Evaluation** module icon in the Windows task bar.
- Select View:File Navigator

Result: The **File Navigator** opens in the **Evaluation** module. The **File Navigator** can be resized and dragged to other positions in the module.

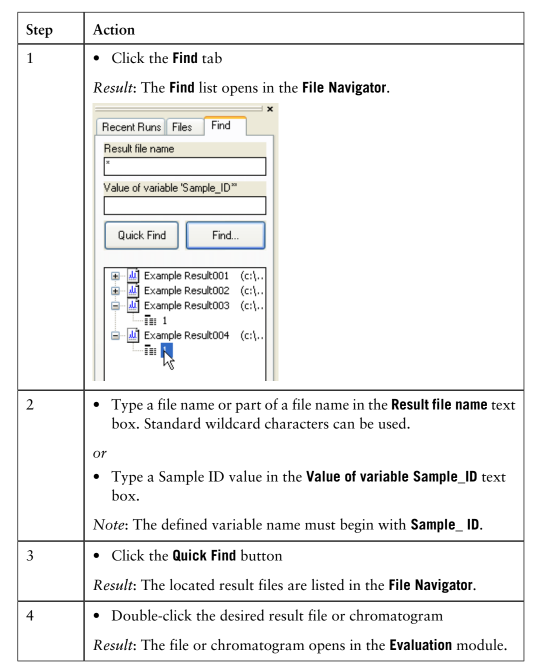
How to open files from the Files list

The table below describes how to use the **Files** list to locate and open a result file.



How to use Find to search for files

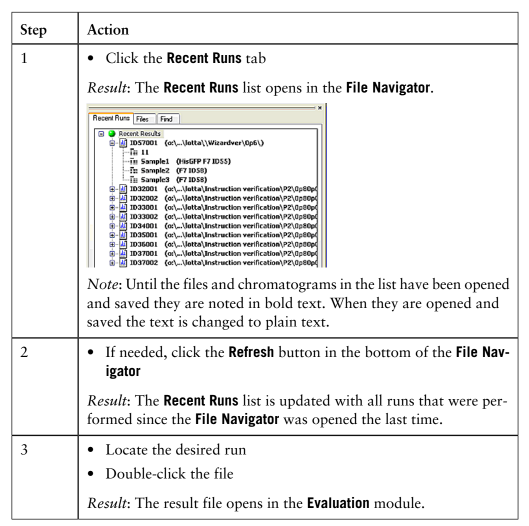
The **Find** function in the File Navigator is used to locate result files in the available folders. The table below describes how to use the **Find** function to locate and open a result file.



Note: Click the **Find** button to open the **Find Files** dialog box where more search functions are available. See **4.3** How to arrange and locate your files on page 74 for more information.

How to open a Recent Run

The **Recent Runs** list shows all the available recorded recent runs based on the selected user preferences. The table below describes how to use the **Recent Runs** list to locate and open a result file.

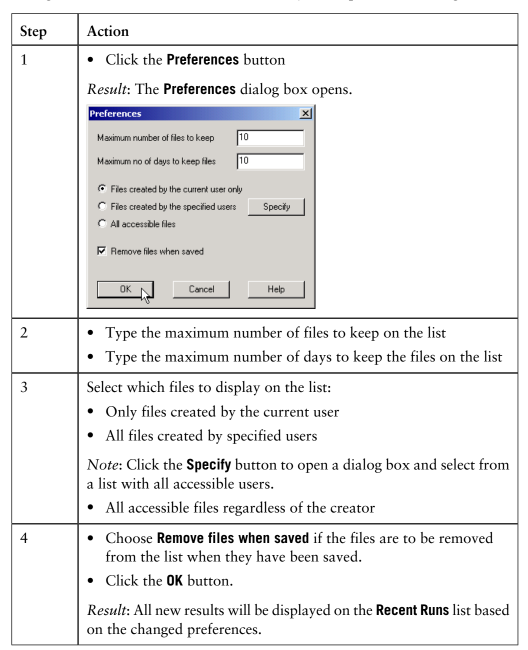


Note: Click the + signs to view or select individual chromatograms from the result files. Individual result files can be selected and removed from the list by clicking the **Remove** button. The **Remove** all button clears the whole list.

Note: **Remove** only clears the list, the files are not deleted.

How to set preferences for Recent Runs

The **File Navigator** will display **Recent Runs** based on the individual user preference settings. The table below describes how to adjust the preference settings:



How to close the File Navigator

To close the File Navigator:

• Click the small cross in the top right-hand corner of the File Navigator.

Result: The File Navigator closes.

10.3 Basic presentation of chromatograms

Introduction

This section describes how to access result files and optimize the presentation of a chromatogram and its curves via the **Chromatogram Layout** dialog box.

In this section

This section contains these topics:

| Topic | See |
|--|--------|
| Introduction and temporary chromatograms | 10.3.1 |
| The Chromatogram window | |

10.3.1 Introduction and temporary chromatograms

Contents of a chromatogram

Chromatograms can be viewed in the **Evaluation** module.

A chromatogram includes a number of curves that have been created during a method run, such as UV, conductivity, pH, fraction marks, etc. A chromatogram also contains the curves created and saved during an evaluation session. The original raw data curves cannot be deleted or modified, but they can be used as the basis for evaluation procedures and subsequent creation of new curves.

Temporary chromatograms

A **Temporary** chromatogram is essentially an empty chromatogram that is specific to the **Evaluation** module. It is also user-specific, so that all users have their own.

Information contained within a **Temporary** chromatogram is automatically saved from one evaluation session to the next, but is not saved within the result files.

How to copy curves into Temporary Curves can be copied into **Temporary** and comparisons or evaluations can be performed. This is particularly useful if you do not want to clutter up your original chromatograms with a large number of curves. It can also be used to keep blank run curves or curves to compare when you open different result files.

The table below describes how to copy curves into **Temporary**:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Select Edit:Copy:Curves. Result: The Copy Curve dialog box is displayed. |
| 3 | Select a source chromatogram and a curve to be copied in the Source Chromatogram fields. |
| 4 | Select Temporary as the target chromatogram and a position for the new curve in the Target Chromatogram fields. |
| 5 | Click the Copy button. Result: The curve is copied into the Temporary chromatogram. Click the Close button. |

How to clear a temporary chromatogram

The table below describes how to clear the contents of a temporary chromatogram:

| Step | Action | |
|------|--------------------------------|--|
| 1 | Open the relevant result file. | |

- 10 How to view results
- 10.3 Basic presentation of chromatograms 10.3.1 Introduction and temporary chromatograms

| Step | Action | |
|------|---|--|
| 2 | Select Edit:Clear Temporary Chromatogram. | |
| | Click the Yes button to confirm. | |

10.3.2 The chromatogram window

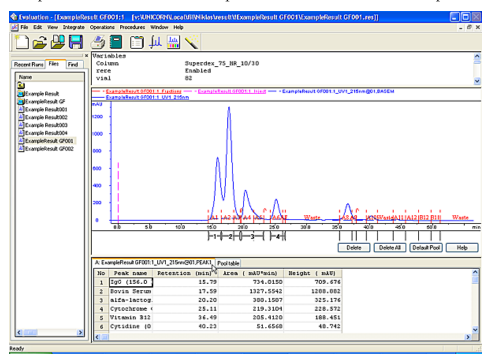
Main views

The chromatogram window is divided into four main views:

- File Navigator
- header information
- curves
- peak and pool tables

The displayed areas for the views can be adjusted by dragging the borders with the mouse cursor between the views.

The picture below shows an example of the window with all views present:



How to view header information

You can display header information at the top of a chromatogram, with details on variables, scouting variables, questions and/or notes. Header information cannot be displayed for imported chromatograms.

The table below describes how to display header information:

| Step | Action | |
|------|---|--|
| 1 | Open a result file. | |
| 2 | In the Evaluation module, select Edit:Chromatogram Layout | |
| | Result: The Chromatogram Layout dialog box is displayed. | |

| Step | Action | |
|------|--|--|
| 3 | Click the Header tab. | |
| | Select the options you want in the header. | |
| | Click 0K . | |
| 4 | • In the chromatogram window, place the cursor at the top of the curve window (just below the toolbar) until the window sizing tool appears. | |
| | Drag the cursor down to display the header window. | |

How to view peak table information

The table below describes how to display peak table information if the result has been integrated: been integrated:

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Choose Edit: Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box opens. |
| 3 | Click the Peak Table tab. |
| | Select a peak table in the Select peak table to display list. |
| | Select what peak table columns to display. |
| | Check if global peak table data should be displayed or not. |
| | • Click OK . |

How to view the Pool table

If fractions are pooled, the **Pool Table** is displayed in the same pane as the **Peak** Table.

• Click the **Pool Table** tab to display the **Pool Table** information.

See 11.5 How to pool fractions on page 279 for more information on how to create the Pool Table.

Run curves, default appearance and information

The first time a result file is opened and viewed, a default layout is applied to display all the original curves. The default layout can be changed by the user (see 10.4.5 How to save and apply a layout on page 242).

Information for each curve

Each curve is automatically assigned a default color and style, with default information about each curve displayed in the key above the curves. This information includes

- result file name
- chromatogram name
- curve name.

Choose the Y-axis scale

Each curve has a correspondingly colored Y-axis. To choose the appropriate Y-axis scale

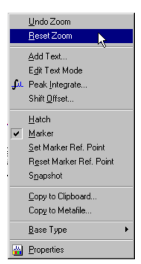
click on the Y-axis until the desired scale is displayed

or

click on the name of the curve.

Run curves, shortcut menu

When viewing curves in the **Evaluation** module, you can access a menu that provides a quick alternative to menu commands. Right-click the run curves view to display the menu shown in the picture below:



Optimizing the workspace

The chromatogram window can be minimized and maximized using ordinary Windows commands. The table below describes extra features to optimize the workspace:

| Use the command | if you want |
|----------------------|--|
| Window:Arrange icons | to arrange icons of minimized windows. |

| Use the command | if you want |
|-----------------|--|
| Window:Tile | to view several chromatogram windows side by side. |
| Window:Cascade | to stack the open windows like a deck of cards. |

How to display a vertical marker line

The table below describes how to display a vertical marker line:

| Step | Action |
|------|---|
| 1 | Right-click the Curves pane and select Marker . |
| 2 | Drag the marker line with the mouse. Result: Where the line bisects the curve, the X-axis and Y-axis values are displayed at the top right corner of the pane. |

Note: Right-click and select **Snapshot** to record the marker position values. See **2.2.7 Snapshots** on page 41 for more information about the **Snapshot** function.

How to set a reference point

The table describes how to set a reference point:

| Step | Action |
|------|---|
| 1 | Display a Marker in the Curves pane. Right-click and select Set Marker Ref. Point to define a reference point for the marker position. |
| 2 | When the marker is moved from the reference point, the X-axis and Y-axis values for the new position are displayed together with: the new position in relation to the reference point, the minimum, maximum and average values for the curve interval between the reference point and the new position. |

How to display the logbook overlay

The table below describes how to display the logbook entries as an overlay in the chromatogram.

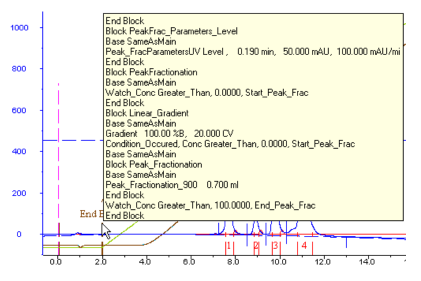
| Step | Action |
|------|---|
| 1 | Right-click in the chromatogram window and choose Properties on the shortcut menu. |
| | Result: The Chromatogram Layout dialog box opens. |
| 2 | Choose the Curve tab. |
| | Select the Logbook curve. |

| Step | Action |
|------|--|
| 3 | Choose the Curve Style and Color tab. Click the Filter button in the Logbook text alignment field. Result: The Filter Logbook dialog box opens. |
| 4 | Select all the logbook items you want to display and click OK. Click OK in the Chromatogram Layout dialog box. Result: The selected logbook items are displayed in the chromatogram window. |

How to view the complete logbook information

At some breakpoints there can be more logbook information than what is possible to conveniently display in the chromatogram window. The additional information that is not displayed is indicated by an arrow point symbol by the break point.

• Hold the mouse cursor over the break point to display the complete information in a flyover text box, as shown in the illustration below:



10.4 How to optimize the presentation of a chromatogram

Introduction

This section describes some of the ways you can optimize the presentation of a chromatogram.

In this section

This section contains these sub-sections:

| Topic | See |
|---|--------|
| How to make changes in the Chromatogram Layout dialog box | 10.4.1 |
| The Curve tab and Curve names tab | 10.4.2 |
| The Curve Style and Color tab | 10.4.3 |
| How to change and fix the axes | 10.4.4 |
| How to save and apply a layout | 10.4.5 |
| How to show part of a curve | 10.4.6 |
| How to change the size of Fraction , Injection and Logbook marks | 10.4.7 |

10.4.1 **How to make changes in the Chromatogram Layout** dialog box

Instruction

The **Chromatogram Layout** dialog box is used to make changes regarding chromatogram presentation. The main features of the Chromatogram Layout dialog box regarding chromatograms are described in the subsequent sections in this chapter. Features regarding peak tables are described in 12.1.2 How to perform a peak integration on page 330.

The table below describes how to make changes in the **Chromatogram Layout** dialog box:

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Right-click the chromatogram window and select Properties |
| | orChoose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. The view from which you activate the Properties command determines the tab that is displayed in the Chromatogram Layout dialog box. |
| 3 | Carry out the changes on the different tabs to get the desired layout for header, curves and peak table. |
| | Select Apply to all chromatograms if you want to apply changes made in the Chromatogram Layout dialog box to all open chromatograms. |
| | Click 0K . |

10.4.2 The Curve tab and Curve Names tab

The Curve tab

The **Curve** tab of the **Chromatogram Layout** dialog box contains a list of all the curves included in the chromatogram. Select the curves you want to display in the chromatogram, and click **OK**.

Curve name appearance

You select options for the curve name appearance on the **Curve Names** tab. This is an example of a default curve name:

Result:11_UV1_280

The table below describes the three components that make up the default curve name:

| Component | Description | Example |
|-------------------|--|---------|
| Result name | Name of the result. | Result |
| Chromatogram name | Number given automatically during a run or a name defined by the New_Chromatogram instruction. | 11 |
| Curve name | Curve type, for example detection of an eluted component. | UV1_280 |
| | In this example, the system uses a variable wavelength detector, so the wavelength (280) for the UV curve is also given. | |

How to choose curve name appearance

You can choose to view only part of the curve name. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Click the Curve Names tab. |

| Step | Action |
|------|---|
| 4 | Select the appropriate boxes for Curve name appearance. Select the appropriate Curve legend position. Click OK. |

Note: It is usually sufficient to select the **Curve Name** option if only one chromatogram is being evaluated. However, confusion can arise when more than one chromatogram is shown, so more complete names might be necessary.

10.4.3 The Curve Style and Color tab

Introduction

All curves within a chromatogram are represented by a default color and line style. Curves imported into the chromatogram or newly created curves are automatically assigned a color and line style.

Peak label settings

Peaks can be labeled on the **Curve Style and Color** tab of the **Chromatogram Layout** dialog box. Use a combination of the following labels:

- Retention (the default label)
- sequential Number
- user-defined Peak name.

Fraction text and Logbook text alignment settings

Both Fraction text and Logbook text can be set to the following alignment options:

- Vertical
- Horizontal
- **Fly Over**, which sets text labels as hidden text that appears only when the cursor is carefully positioned over a fraction mark.

How to change the color and style of a curve

The table below describes how to change the color and style of a curve:

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Click the Curve Style and Color tab. |
| 4 | Select the curve you want to change from the list. Select the desired color and style. Click OK. |

How to display and filter logbook information

The table below describes how to display and filter logbook curve information:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. |

| Step | Action |
|------|--|
| 3 | Click the Curve tab. |
| | Select the logbook curve. |
| 4 | Click the Curve Style and Color tab. |
| | Click the Filter button in the Logbook text alignment field. |
| | Result: The Filter Logbook dialog box is displayed. |
| 5 | Select the type of logbook information you want to show. |
| | Set the maximum block depth to show. |
| | Click 0K . |

How to display a hatched back-ground

The table below describes how to display a hatched background in the chromatogram window:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Click the Curve Style and Color tab. |
| | • Select the Hatch box. |
| | • If desired, select the Apply to all chromatograms box and click OK . |
| | Result: Hatch marks are displayed as a background. |

Note: You can also right-click in the **Chromatogram** window and select **Hatch**.

10.4.4 How to change and fix the axes

How to change and fix the Y-axis

The table below describes how to change and fix the Y-axis:

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Click the Y-Axis tab. |
| 4 | Select the appropriate curve from the list. Click the Fixed option. |
| 5 | Type the desired minimum and maximum values. Click the All with this unit button if you want other curves with the same Y-axis units as the current scaled curve to be similarly scaled. |
| | <i>Note</i> : The values will only be applied to existing curves. They will not be applied to new curves created after this function was last used. |
| | • Click the appropriate Pressure unit (MPa , psi , bar) option to change Y-axis units for pressure curves. |
| | <i>Note:</i> Default Pressure unit is From strategy , which is the unit defined in the original run strategy. |
| | • Click 0K . |

How to add a second Y-axis

The table below describes how to add a second Y-axis to the chromatogram.

| Step | Action |
|------|--|
| 1 | Choose Edit:Chromatogram Layout. Result: The Chromatogram Layout dialog box is displayed. |
| 2 | Click the Y-Axis tab. |
| 3 | Select the appropriate curve from the Right Axis droplist. Click the OK button. |

How to change and fix the X-axis

The table below describes how to change and fix the X-axis:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. |
| | Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Click the X-Axis tab. |
| 4 | Select the appropriate option in the Base field: • Time of retention • Volume • Column Volume |
| | Note: Some calculated curves, for example baselines, exist in only one base and might seem to disappear when the base is changed. Curves are collected in time and recalculated for display in volume. Thus, switching the base between Time and Volume can slightly alter the resolution. |
| 5 | Click the Fixed option in the Axis scale field to set the axis limits manually. Type the desired minimum and maximum values. If desired, de-select the Adjust retention zero to injection number checkbox. This checkbox is selected by default. The function sets the time/volume to zero at the injection mark, that is when the sample was injected. The time and volume before injection will become negative values. |
| | Click 0K . |

10.4.5 How to save and apply a layout

Introduction

All configurations that you make in the **Chromatogram Layout** dialog box can be saved as a layout. It is possible to apply saved layouts to other chromatograms. All saved layouts are user-specific.

How to save a layout

The table below describes how to save a layout:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Edit:Chromatogram Layout. Result: The Chromatogram Layout dialog box is displayed. |
| 3 | Make the appropriate layout configuration within the various tabs. View your changes Click OK if you want to return to the chromatogram window to see the applied affects of a given configuration. Return to the Chromatogram Layout dialog box to perform further changes. |
| 4 | Select the Layout Library tab. Click the Save current layout as button. Result: The Save Layout dialog box is displayed. |
| 5 | Type a name for the layout. If you want the current layout to be the new default layout, select the Save as default option. Click OK. Result: The new name is added to the Saved layouts list. Click OK. |

How to apply a layout

The table below describes how to apply a layout:

| Step | Action | |
|------|--|--|
| 1 | Select the Layout Library tab on the Chromatogram Layout dialog box. | |

| Step | Action |
|------|---|
| 2 | Select a layout from the Saved layouts list. |
| | Click the Apply selected layout button. |
| | Result: The layout is automatically applied to the active chromatogram window. |
| | • If the same layout is to be applied to all chromatograms on the Evaluation workspace, select the Apply to all chromatograms checkbox. |
| | • Click OK . |

10.4.6 How to show part of a curve

Introduction

You can select a part of a curve in order to examine details more closely.

You can

• use the zoom to magnify

01

• cut the axes.

It is also possible fix the axes, see 10.4.4 How to change and fix the axes on page 240.

How to use the zoom function

In the active chromatogram window, you can zoom in on a designated area of the chromatogram. This is the easiest and quickest way to enlarge different parts of a curve. The table below describes how to do this:

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Place the mouse pointer in any corner of the area you want to magnify. |
| | • Press and hold the left mouse button. A magnifying glass icon will be added to the mouse pointer arrow on the screen. |
| | Drag a box to cover the area to be magnified, and release the mouse button. |
| | <i>Result:</i> The selected region is now displayed in the entire chromatogram window, together with appropriate scales for the Y and X axes. |
| 3 | Use the arrow keys on the keyboard to move around in the chromatogram at the current zoom scale. |
| 4 | Undo zoom |
| | Right-click in the window and select Undo zoom to undo the last zoom step. |
| | Reset zoom |
| | Right-click in the window and select Reset zoom to reset all zoom steps at once. |

How to cut a curve and store as a new curve

The table below describes how to cut the curve between two values on the X-axis and store this part of the curve as a new curve:

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose Operations:Cut curve. Result: The Select Curve(s) to Operate On dialog box opens. |
| 3 | Select the curves to be operated on. Click OK. Result: The selected curves are shown in the Cut dialog box which contains two vertical cursor lines. |
| 4 | To select the region to be cut, either drag the two cursor lines to define the left and right limits of the cut area or type the desired left and right limit values in the Left limit and Right limit boxes. |
| | <i>Note:</i> The areas outside of the Left limit and Right limit will not be saved in the newly created cut curve. Thus, the X-axis of the new curve will not begin at zero unless this is designated as one of the limits. The original curve is not changed. |
| 5 | Click 0K . Result: The Save Cut Curves dialog box opens. |
| 6 | Select whether to save the new cut curve in the Source chromatogram, that is the current active chromatogram, or a New chromatogram (if you select this option, you can change the name of the chromatogram. Note that it is a recommendation not to use only numbers as names for chromatograms.). |
| | Click 0K. Result: If the destination of the cut curve was the source chromatogram, the cut curve is automatically displayed in the source chromatogram. If the destination of the cut curve was a new chromatogram, this will be represented as a new, open chromatogram window. |

10.4.7 How to change the size of Fraction, Injection and Logbook marks

Introduction

The sizes of **Fraction**, **Injection** and **Logbook** marks are all determined by your user settings. The settings are applied for all your chromatograms.

Instruction

The table below describes how to change the size of the **Fraction**, **Injection** and **Logbook** marks:

| Step | Action |
|------|--|
| 1 | Choose Administration:Change User Attributes in the UNICORN Manager module. |
| | Result: The Change user attributes dialog box opens. |
| 2 | Select the unit for the Fraction mark height: • Percent of window height • Character Heights • Pixels |
| | Type a new size value in the Fraction mark height box. |
| 3 | • Repeat step 2 for the Injection and Logbook marks if necessary. |
| | Click OK . |

10.5 How to print active chromatograms

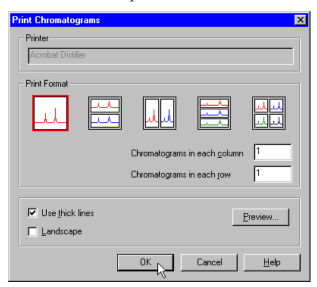
Introduction

This section describes how to print the chromatograms that are open in the **Evaluation** module.

The Print Chromatograms dialog box

This is an illustration of the Print Chromatograms dialog box.

Note: The selected print format is outlined in red.



Instruction

The table below describes how to print active chromatograms.

| Step | Action |
|------|--|
| 1 | Open all chromatograms that you want to print in the Evaluation module. |
| 2 | Select File:Print. or Click the Print toolbar icon. Result: The Print Chromatograms dialog box opens. |
| 3 | Select print format and layout options. |
| 4 | Click OK to print. or Proceed with step 5 to preview and edit the layout. |

| Step | Action |
|------|---|
| 5 | Click the Preview button. |
| | Result: The Customise Report window opens. |
| 6 | • Click the Edit Mode button to make changes, e.g. change the order of the chromatograms (see 10.6.1 How to create and print a customized report on page 250 for more information about how to edit). |
| | Click the Preview button to return to preview mode. |
| 7 | Select File:Print. |
| | or |
| | Click the Print toolbar icon. |
| | Result: The Print dialog box opens. |
| 8 | Select the print range and number of copies.Click OK. |

10.6 How to create and print reports

Introduction

The **Evaluation** module provides extensive tools to create detailed reports. This section describes how to create and print reports that are based either on a standard or a customized layout.

In this section

This section contains these topics.

| Topic | See |
|---|--------|
| How to create and print a customized report | 10.6.1 |
| How to create and print a standard report | 10.6.2 |
| How to edit an existing report format | 10.6.3 |

10.6.1 How to create and print a customized report

Introduction

You can choose from a variety of objects to include in a report, including chromatograms, methods, documentation, free text and more in the customized report interface. You can also place, align and size the objects as you please. This section describes how to create a customized report format.

Should you need to store store your reports in an electronic format you can save them as PDF files. This section also describes how to do this..

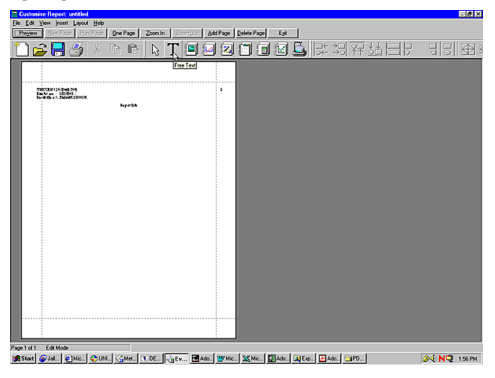
How to open the Report Editor in edit mode

The table below describes how to open the **Report Editor** in **Edit mode** to create a customized report format.

| Step | Action |
|------|--|
| 1 | Open a result file in the Evaluation module. |
| 2 | Select File:Report. |
| | or |
| | Click the Report icon. |
| | |
| | Result: The Generate Report dialog box opens. |
| 3 | Click the New button. |
| | Result: The Create New Report Format dialog box opens. |
| 4 | Select the Customised format and click OK. |
| | Result: The Report Editor opens in Edit mode. |

The Edit mode window

The illustration below shows the **Report Editor** window in **Edit mode** with a blank report open:



Toolbar button functions in the Report Editor

The table below describes the different functions of the Edit mode toolbar buttons in the **Report Editor**:

| Toolbar button | Function |
|--------------------|---|
| Preview/Edit | This button toggles between a print preview of the report and the Edit mode . |
| Next Page | This button displays the next page or pair of pages (where there are more than one page). |
| Prev Page | This button displays the previous page or pair of pages (where there are more than one page). |
| One Page/Two Pages | This button toggles between single page view and pairs of pages view, when there is more than one page. |
| Zoom In | This button increases the magnification of the view. |
| Zoom Out | This button decreases the magnification of the view. |
| Add Page | This button adds a blank page to the report. |
| Delete Page | This button deletes the current page from the report. |
| Exit | This button closes the Customize Report window. |

How to add and delete report pages

The table below describes how to add or delete report pages in the **Report Editor**:

| If you want | then |
|---|--|
| to add new pages, | click the Add Page toolbar button. |
| | Result: A new page is added after the last page. |
| to delete a page while in One Page mode, | select the page with Next Page or Prev Page, click the Delete Page toolbar button and confirm the deletion. |
| to delete a page in Two Page mode, | select the page with Next Page or Prev Page, click an object on the page, click the Delete Page toolbar button and confirm the deletion. |

How to change the page layout

The page layout is changed in the **Page Setup** dialog box. The table below describes how to set up the page layout:

| Step | Action |
|------|--|
| 1 | Double-click anywhere on the report page in the Report Editor (not on an object). Result: The Page Setup dialog box opens. |
| 2 | Type new values for the Margins if necessary. Select the appropriate Settings and Unit. |
| | Note: An extra Header tab will appear if you de-select the option to have the same header on all pages. The First Header tab is used for the first page header only, and the Header tab is used for all subsequent pages. Click the First Header tab. |
| 3 | Select all the items you want to include in the header from the Select Items list. Click the Font button to change the font for all items if necessary. |
| 4 | Type header text in the Free text box and click the Font button to alter the default font if necessary. Type the report title in the Report title box and click the Font button to alter the default font if necessary. |

| Step | Action |
|------|--|
| 5 | Select the Logo check box and click the Browse button if you want to locate and select a logo image file. |
| | Select the Alignment for the logo, if necessary. |
| | <i>Note</i> : The logo file must be in bitmap format (.bmp) and smaller than 64 kB. Larger logo files or files in other formats must be inserted as Picture objects. |
| 6 | If you want to have a line under or over the header, select the appropriate option in the Layout field. |
| 7 | • Repeat steps 3 to 6 on the Footer tab and the subsequent pages Header tab. |
| | Note: All Header and Footer tabs contain the same options. You can have all information in either the header or footer or split information between the header and footer as required. |
| | • Click 0K . |

How to add objects to the report

The table below describes how to add objects to the report. The various objects are described below this table.

| Step | Action |
|------|---|
| 1 | • Click the appropriate icon in the Report items toolbar. |
| | • Choose an object from the Insert menu. |
| | Insert Layout Help T Free text Text Method Chromatogram Documentation Evaluation log Quantitate and Mol. Size Frac 950 |
| 2 | Press and hold the left mouse button on the report page, and drag out a box to the size of the item you want to insert. |
| | <i>Note</i> : The mouse pointer shows a symbol for the type of item you have selected. |
| | Release the mouse button. |
| | Result: A Setup dialog box opens. The dialog is specific to the type of item that you want to insert. |

| Step | Action |
|------|--|
| 3 | Select the desired options and click 0K . |
| | Result: The object is inserted onto the page. |

Note: If you want to edit an object later, double-click the object box.

How to add free text

The table below describes how to add free text to the report:

| Step | Action |
|------|---|
| 1 | Click the Free Text icon. |
| | T |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the text. Release the button. |
| | Result: The Setup Free Text dialog box opens. |
| 2 | Type text in the edit field. |
| | Select if the text is to start on a new page. |
| | Select if the text box should be automatically sized. |
| | • Select if the text should appear in the same position on all pages, for example as header and footer text. |
| 3 | Click the Font button to change the default font. |
| | Result: The Font dialog box opens. |
| | Make the necessary changes and click 0K to return. |
| | • Click OK . |
| | Result: The text object is inserted onto the page. |

ture

How to add a pic- The Picture dialog box is useful to insert logos, pictures or other figures in the report. The table below describes how to add a picture object to the report:

| Step | Action |
|------|---|
| 1 | Press and hold the left mouse button on the report page and drag out a box to the size of the picture item. Release the mouse button. |
| | Click the Picture icon. |
| | |
| | Result: The Picture dialog box opens. |
| 2 | Click the Browse button to locate the desired picture file. |
| | Select the picture file and click the Open button. |
| | Note: The file formats .bmp, .emf, .jpg and .tif can be used. |
| | Result: A preview of the selected picture is displayed. |
| 3 | Select the desired Settings and click OK . |
| | Result: The picture is inserted onto the page. |

How to add a chromatogram or peak table

The table below describes how to add a chromatogram to the report. The layout can also be defined to include a peak or pool table if desired.

| Step | Action |
|------|--|
| 1 | Click the Chromatogram icon. |
| | |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the chromatogram. Release the mouse button. |
| | Result: The Setup Chromatogram dialog box opens. |
| | Selected chromatogram(s) Active chromatogram V |
| 2 | Select which chromatogram(s) to insert from the Selected chromatogram(s) droplist. • Active chromatogram inserts the chromatogram that currently is active in the Evaluation module. |
| | All chromatograms inserts all chromatograms that are open in the Evaluation module. |
| | • 1, 2etc. inserts the corresponding chromatogram. |
| 3 | Select the desired Settings . |
| | • If desired, change the Fonts . |
| | Note: Separate fonts can be selected for the Chromatogram, the Peak table and the Header text. |

| Step | Action |
|------|--|
| 4 | • Click the Define button in the Layout field if you want to re-define the layout of the chromatogram. |
| | Result: The Report Chromatogram Layout dialog box opens. |
| | Make the appropriate changes and click OK to return to the Setup Chromatogram dialog box. |
| | <i>Note</i> : The changes that you make will only affect the report and not the view of the chromatograms in the Evaluation module. |
| 5 | Click OK . |
| | Result: The chromatogram is inserted onto the page. |

Note: All curves can be de-selected in the **Report Chromatogram Layout** dialog box leaving only the selected peak table(s) in the report.

How to include a method

The table below describes how to include a method in the report:

| Step | Action |
|------|---|
| 1 | Click the Method icon. |
| | |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the item. Release the button. |
| | Result: The Setup Method dialog box opens. |
| 2 | Select the items to be included in the report: |
| | Main Method is the method on which the run was based. |
| | Blocks are the blocks that were used in the method. |
| 3 | Select the appropriate Settings . |
| | Note: Expand main displays the expanded method view. |
| | If desired, change the Fonts . |
| | Click OK . |
| | Result: The method object is inserted onto the page. |

mentation

How to add documentation to the report:

| Step | Action |
|------|--|
| 1 | Click the Documentation icon. |
| | |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the item. Release the button. |
| | Result: The Setup Documentation dialog box opens. |
| 2 | Select the items to be included in the report: |
| | Select All includes all items in the report. |
| | Clear All removes all selections. |
| 3 | If desired, change the Fonts . |
| | Select if the documentation should start on a new page. |
| | • If Select All, Logbook or Run summarySelect All or Logbook was selected, make the necessary changes to the Base and Logbook filter settings. |
| | • Click OK . |
| | Result: The selected documentation items are inserted into the report. |

How to add the **Evaluation Log**

The table below describes how to add the **Evaluation Log** to the report:

| Step | Action |
|------|---|
| 1 | Click the Evaluation Log icon. |
| | Press and hold the left mouse button on the report page and drag out a box to the size of the item. Release the mouse button. |
| | Result: The Setup Evaluation Log dialog box opens. |
| 2 | If desired, change the Fonts . |
| | Select if the Evaluation Log should start on a new page. |
| | Click 0K . |
| | Result: The Evaluation Log is inserted into the report. |

How to include Quantitate and Molecular Size data

The table below describes how to include **Quantitate** and **Molecular Size** data in the report.

Note: This option is only available if the **Analysis** module has been installed.

| Step | Action |
|------|---|
| 1 | Click the Quantitate and Mol Size icon. |
| | |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the item. Release the mouse button. |
| | Result: The Setup Quantitate dialog box opens. |
| 2 | If desired, change the Fonts . |
| | The default option is that the Quantitate and Molecular Size data |
| | will start on a new page. |
| | • Click OK . |
| | Result: The Quantitate and Molecular Size data is inserted into the |
| | report. |

How to include Frac-950 data

The table below describes how to include Frac-950 data in the report.

Note: This option is available only if a **Frac-950** has been installed and if the result file contains data from the **Frac-950**.

| Step | Action |
|------|---|
| 1 | Click the Frac-950 icon. |
| | |
| | • Press and hold the left mouse button on the report page and drag out a box to the size of the item. Release the mouse button. |
| | Result: The Setup Frac-950 dialog box opens. |
| 2 | If desired, change the Fonts . |
| | Select if the Frac-950 data should start on a new page. |
| | The Include rack layout option is selected by default. This will display the rack layout that was used in the run. |
| | Click OK . |
| | Result: The Frac-950 data is inserted into the report. |

How to move and resize objects freely

How to move and The table below describes how to select, move and resize objects freely:

| If you want | then |
|-----------------------------------|---|
| to select a single object, | • click the Select icon, |
| | B |
| | click the object of interest. |
| to select several ob- | • click the Select icon, |
| jects, | • press and hold the <ctrl> key while you click the objects.</ctrl> |
| to move the selected object(s), | click on the objects, hold down the left mouse button and drag the object(s) to the new position. |
| to resize the selected object(s), | click one of the object border anchors, either in the corners or in the middle of a border, and drag the box to the new size. |
| | Note: Some Text objects cannot be resized. |

Alignment toolbar icon functions

Objects can be placed in exact positions and sized in relation to other objects. The table below describes the function of the **Alignment** toolbar icons in the **Report Editor**:

| Tool- bar icon | Function |
|----------------------|---|
| □← | Align left Matches the left alignment of all selected objects to that of the highlighted object. |
| → □ | Align right Matches the right alignment of all selected objects to that of the highlighted object. |
| <u>↑</u> ↑ | Align top Matches the top alignment of all selected objects to that of the highlighted object. |
| <u>⋄</u> | Align bottom Matches the bottom alignment of all selected objects to that of the highlighted object. |

| Tool- bar icon | Function |
|----------------------|---|
| | Adjust to margins Stretches the selected object(s) to the left and right margins. |
| | Adjust to left margin Adjusts the selected object(s) to the left margin. |
| | Adjust to right margin Adjusts the selected object(s) to the right margin. |
| | Adjust to centre Adjusts the selected object(s) to the center of the page. |
| ** | Make same size Adjusts the selected objects to the same size as the highlighted reference object. |
| | Make same width Adjusts the selected objects to the same width as the highlighted reference object. |
| | Make same height Adjusts the selected objects to the same height as the highlighted reference object. |

Note: The **Make same size** and **Make same width** functions can only be used to resize the width of chromatograms, free text and picture objects.

How to print the report

The table below describes how to print the report:

| Step | Action |
|------|--|
| 1 | Choose File:Print. |
| | or |
| | Click the Print icon. |
| | 3 |
| | Result: The Print dialog box opens. |
| | Note: Printers are set up in the File menu of the UNICORN Manager. |

| Step | Action |
|------|------------------------------|
| 2 | Select the printing range. |
| | Select the number of copies. |
| | • Click OK . |

Note: You can also print the report from the Generate Report dialog box.

How to save the report in PDF format

The table below describes how to save the finished report as a PDF file:

| Step | Action |
|------|---|
| 1 | Click the UNICORN Manager icon on the Windows taskbar. |
| | Result: The UNICORN Manager opens. |
| | Choose File:Printer Setup. |
| | Result: The Print Setup dialog box opens. |
| 2 | • Select an Adobe Acrobat printer from the Printer Name list (e.g. Acrobat Distiller). |
| | • Click the Properties button and edit the document properties if needed. |
| | Select the appropriate paper size and orientation. |
| | • Click OK . |
| 3 | Click the Evaluation icon on the Windows taskbar. |
| | Result: The Evaluation module opens |
| 4 | Print the report as described in "How to print the report". |
| | Result: The report is created as a PDF file and saved in the location specified in your Acrobat settings. |

Note: You must have a full installation of Adobe Acrobat or a suitable printer driver to be able to do this.

How to save the report format

The table below describes how to save the finished report format:

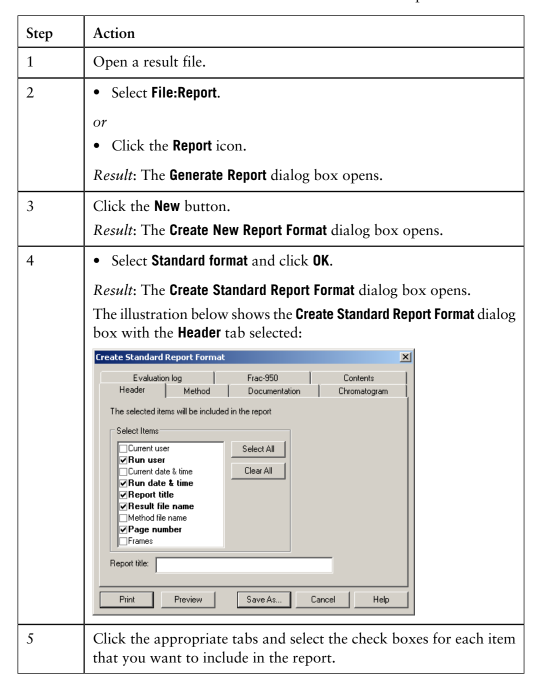
| Step | Action |
|------|--|
| 1 | Choose File:Save. |
| | or |
| | • Click the Save icon. |
| | |
| | Result: The Save Report Format dialog box opens. |
| 2 | Type a name for the format. |
| | Select if you want to save the format for global use. |
| | Select if you want to save the format as default. |
| | Note: The name for the default format will automatically be changed to DEFAULT.Click OK. |

10.6.2 How to create and print a standard report

How to create a Standard report

You can only select a number of pre-formatted items when you create a **Standard** report format. If you want to edit the layout in detail you must create a **Customized** report format. See **10.6.1 How to create and print a customized report** on page 250.

The table below describes how to create and save a **Standard** report format:

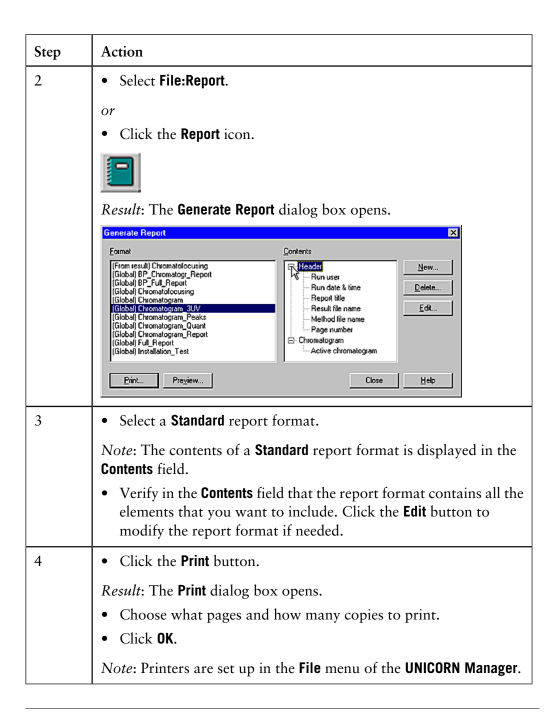


| Step | Action |
|------|---|
| 6 | Click the Chromatogram tab and select the chromatogram(s) you want to include. |
| | Select the Current option in the Layout field to apply the current layout in the Evaluation module. |
| | or |
| | Click the Define button in the Layout field to open the Curve tab in the Report Chromatogram Layout . |
| | - Select the curves that you want to include in the report and click 0K . |
| 7 | Click the Contents tab to see a list of all the selected items. |
| | Click the Preview button to see the entire report layout. |
| | Click the Close button to return. |
| | Click the Print button to print a test report. |
| 8 | Click the Save As button. |
| | Result: The Save Report Format dialog box opens. |
| | Type a name in the Report format name text box. |
| | - Select the Save as global format check box to make the format available to other users. |
| | - Select the Save as default report format check box if desired (The format is saved as DEFAULT). |
| | Click 0K . |
| | Result: The Generate Report dialog box opens again. The new report is saved and available in the Format list. |
| 9 | Click the Close button |
| | or |
| | Click the New button to create another Standard report. |

How to print a standard report

The table below describes how to print a ${\bf Standard}$ report format in the ${\bf Evaluation}$ module.

| Step | Action |
|------|---------------------|
| 1 | Open a result file. |



10.6.3 How to edit an existing report format

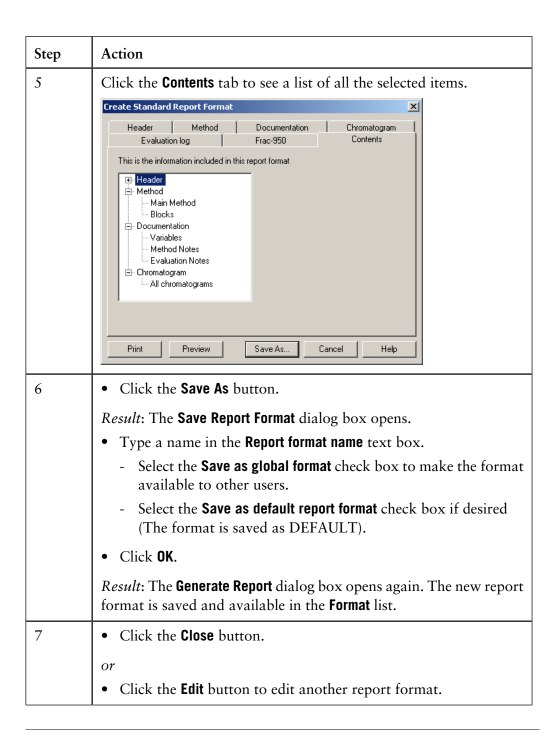
Introduction

This section describes how to edit an existing report format.

How to edit a standard report

The table below describes how to edit a standard report format in the **Evaluation** module.

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Select File:Report. |
| | or |
| | Click the Report icon. |
| | |
| | Result: The Generate Report dialog box opens. |
| 3 | Select a Standard report format to edit. |
| | Click the Edit button. |
| | Result: The Edit Report Format dialog box opens. |
| | Select Standard format and click OK. |
| | Result: The Edit Standard Report Format dialog box opens. |
| 4 | Click the appropriate tabs and select the check boxes for each item that you want to include in the report format. |
| | Note: See 10.6.2 How to create and print a standard report on page 264 for more information. |



How to edit a customized report

The table below describes how to edit a customized report format in the **Evaluation** module.

| Step | Action |
|------|---------------------|
| 1 | Open a result file. |

| Step | Action |
|------|--|
| 2 | Select File:Report. |
| | or |
| | Click the Report icon. |
| | |
| | Result: The Generate Report dialog box opens. |
| 3 | Select a Customized Report Format to edit. |
| | Click the Edit button. |
| | Result: The report format opens in the Report Editor. |
| 4 | Double-click the report item you want to edit. |
| | Make the desired changes in the dialog box. |
| | Continue to edit all items until the format is complete. |
| | Note: See 10.6.1 How to create and print a customized report on page 250 for more information. |
| 5 | Select File:Save As. |
| | Result: The Save Report Format dialog box opens. |
| | Type a name in the Report format name text box. |
| | - Select the Save as global format check box to make the format available to other users. |
| | - Select the Save as default report format check box if desired (The format is saved as DEFAULT). |
| | • Click OK . |
| | Result: The new report format is saved and available in the Format list. |

10.7 Run documentation

Introduction

The full documentation for a method run is stored in the result file. This section describes:

- some of the contents of the run documentation,
- how to view and print the run documentation,
- how to save the method from the run as a new method.

How to view and print the run documentation

The table below describes how to view and print the run documentation.

| Step | Action |
|------|--|
| 1 | Open a result file. |
| 2 | Choose View: Documentation in the Evaluation module. |
| | or |
| | Click the view Documentation icon. |
| | |
| | Result: The Documentation dialog box opens. |
| | See further information about some of the tabs below. |
| 3 | Click the Print button. |
| | Result: The Print dialog box opens. |
| | • Select the documentation items you want to print and click OK . |

The tabs of the Documentation dialog box

The table below describes the contents of some of the **Run Documentation** tabs.

| Documentation tab | Contents |
|-------------------|--|
| Variables | The Variables tab lists the parameters that were used during the method run. |
| Scouting | The Scouting tab displays the whole scouting scheme, with the values for the current result file displayed in yellow cells. |

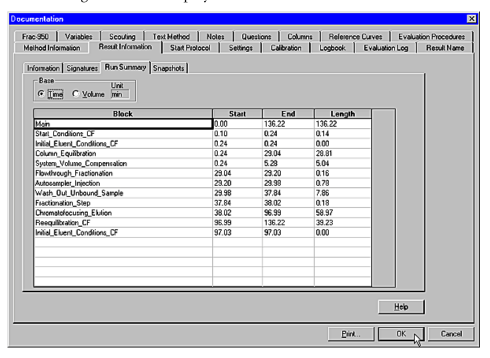
| Documentation tab | Contents |
|--------------------|---|
| Notes | The Notes tab displays the notes that you have made at various times during the method run. You are also able to type new comments on the Evaluation Notes sub-tab. |
| | <i>Note</i> : Click the Find button to search for a specific text string in the Notes . |
| Calibration | The Calibration tab displays the system calibrations and when and by whom they were made. |
| Logbook | The Logbook tab displays what happened during a method run. You can view information concerning alarms, the method, manual changes during the run, errors, etc. Note: Click the Find button to search for a specific text string in the Logbook . |
| Evaluation Log | The Evaluation Log lists all of the evaluation operations that you have performed for the result file during all sessions, including at the end of the method. |
| Method Information | The Method Information tab displays information about the method, such as the method name, the target system and the date of the last change. Information about the strategy includes name, date and size. There is also a sub-tab for Signatures . |
| Frac-950 | The Frac-950 tab displays the setup parameters for the fraction collector provided it is included in the strategy. |
| Result Information | See "The Result Information tab" in this section. |

The Result Information tab

The **Result Information** tab displays information about the result file, such as

- the result file name
- the system that was used
- the last date it was changed.

Information about the strategy includes name, date and size. The **Run Summary** sub-tab is a summary of the run expressed in volume or time per block. There is also a sub-tab for **Signatures** and a sub-tab where all **Snapshots** that have been taken during the run are displayed.



Save the method used for the run as a new method

You can save the method and the variables that were used for the run as a new method:

| Step | Action |
|------|---|
| 1 | Select the Text Method tab in the Documentation dialog box. |
| | Click the Save as button. |
| | Result: The Save As dialog box opens. |
| 2 | Select the appropriate destination folder. |
| | Type a name in the Method name text box. |
| | Select a system in the For System field. |
| | Select a technique in the Technique field. |
| | Click 0K . |
| | Result: The method is saved. |

11 How to edit results

Introduction

This chapter describes

- how to edit the results that are presented in the **Evaluation** module
- how to import and compare runs
- how to import and export results.

For more information about how to view results, see chapter 10 How to view results on page 220.

In this chapter

This chapter contains these sections:

| Topic | See |
|---|-------|
| How to reduce noise and remove ghost peaks | 11.1 |
| How to subtract a blank run curve | 11.2 |
| How to add curves | 11.3 |
| How to enter and edit text in the chromatogram | 11.4 |
| How to pool fractions | 11.5 |
| How to match protein activity to a curve | 11.6 |
| How to rename chromatograms, curves and peak tables | 11.7 |
| How to import and compare different runs | 11.8 |
| How to import and export results | 11.9 |
| How to sign results electronically | 11.10 |
| How to save results and exit the Evaluation module | 11.11 |

11 How to edit results

11.1 How to reduce noise and remove ghost peaks

Introduction

Sometimes the chromatograms contain curves with a noisy baseline. The noise can be caused by several factors, for example a dirty flow cell, air bubbles, electrical noise, dirty buffers, etc. The amount of noise can usually be reduced by taking proper precautions, for example filtration of buffers and instrument maintenance.

You can also use the smoothing function to reduce or remove background noise from a selected curve. Smoothing is always a compromise between noise removal and preservation of peak shape.

How to smooth a curve

The table below describes how to select a smoothing function and smooth a curve:

| Step | Action |
|------|--|
| 1 | Select Operations:Smooth. Result: The Smooth dialog box is displayed. |
| 2 | Select the curve to be smoothed and its target destination. |
| 3 | Select the Filter type to be applied. The options are: Moving average. Use this if you have noise along most of the curve. It affects peak height but not retention. There is little effect on the peak area. Autoregressive. Use this if you have periodic noise along the whole curve. It affects peak height and retention, although this has little effect on the peak area. Median. Use this if there is only one or a few noise spikes, for example caused by air bubbles, or if the noise is confined to only a small part of the curve. It can flatten peaks and affect peaks areas slightly, but does not affect retention. Savitzky-Golay. Use this to calculate the smoothing and differentiation of data by a least squares technique. |
| 4 | Select an appropriate smoothing parameter value from Light to Hard for the selected filter in the Filter Parameters field. Use the slider, or insert a value manually in the text field. The smoothing effect increases with increasing parameter values. • Click OK . |

Tip: Start with a low parameter value, for example the default value, and increase it until the best result is achieved. A useful strategy is to increase the parameter value by the default value for each try.

Note: By default, smoothed curves are given the suffix SMTH. The default curve name can be changed as needed.

11.2 How to subtract a blank run curve

Introduction

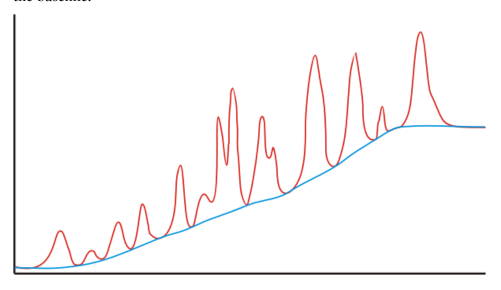
Subtracting a blank run curve is a frequently used function in presentations, especially if the curves have a drifting baseline or ghost peaks.

Ghost peaks

If the ghost peaks come from impurities in the eluents, all equilibration of the columns should be the same from method run to method run. If, for example, the equilibration volume with buffer A is larger before a blank run curve than before a separation, your ghost peaks might be higher in the blank run curve.

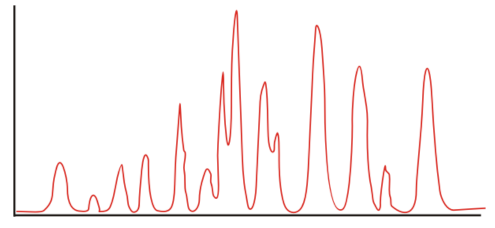
Example of a UV curve with baseline

The illustration below shows the UV curve with baseline prior to subtraction of the baseline:



Example of a UV curve after subtraction of the baseline

The illustration below shows the UV curve after subtraction of the baseline:



How to import a blank run curve

If a blank run curve was made, this might have been stored in another result file. If there is no blank run curve, you can create one with Integrate:Calculate baseline. The table below describes how to import the blank run curve:

| Step | Action |
|------|--|
| 1 | Ensure that the destination chromatogram has been opened and is the active window on the screen. |
| 2 | Choose File:Open:Curves. |
| | Result: The Open Curves dialog box is displayed. |
| 3 | Double-click the result file that contains the blank run curve. |
| | Result: The curves in the first chromatogram are displayed. |
| 4 | Select the appropriate chromatogram in the Chromatogram list . |
| | Result: The curves for that chromatogram are displayed on the Available list. |
| | • Select the curves that correspond to the blank run curve and click the Select button. |
| | Result: The selected curve is displayed on the Selected curves list. |
| 5 | If you want to remove a curve from the list, select it and click the Remove button. |
| | Click 0K to import the curve. |

Note: For more detailed information on how to import curves, chromatograms and other results see 11.8 How to import and compare different runs on page 287.

blank run curve

How to subtract a You can subtract the blank run curve or the baseline from the sample curve. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Select Operations:Subtract. |
| | Result: The Subtract dialog box is displayed. |
| 2 | Select the sample chromatogram and curve in the left field and the baseline or blank run curve to be subtracted in the middle field. Click OK . |

Note: All resulting curves from the subtract operation receive the SUB suffix by default. The default curve name can be changed as needed.

11.3 How to add curves

Introduction

In some method runs, several sequential chromatograms might have been created. This can occur, for example, when the instruction **New chromatogram** has been used in the method, thus creating different chromatograms during the run.

In order to view and evaluate the resultant curve of all the chromatogram parts, the curves must be added together. Usually, you have a number of chromatograms within the same result file and you want to add the curves. In some circumstances, curves might need to be imported from other result files.

Instruction

The table below describes how to add curves:

| Step | Action |
|------|--|
| 1 | Select and view the first chromatogram in the sequence. |
| 2 | Choose Operations:Add . Result: The Add dialog box is displayed. |
| 3 | Select the first curve in the desired sequence in the left field. Select the second curve in the sequence in the middle field. Click the OK button to add the two curves together in a new result curve. |
| 4 | Open the Add dialog box again. Select the result curve (.ADD) from the previous addition in the left field. Select the next curve in the sequence in the middle field. Click OK to add the two curves together in a new result curve. |
| 5 | Repeat steps 3 and 4 until all curves have been added together. The final curve should be the cumulative curve for the whole run. |

Note: All curves created using the **Add** operation receive the **ADD** suffix by default. The default curve name can be changed as needed. The original curves are distinguished in the chromatogram by underlined curve names.

11.4 How to enter and edit text in the chromatogram

How to enter text Text can be added to the chromatogram. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Right-click the curves view of the chromatogram window and select Add text from the menu. |
| | or |
| | Choose Edit:Text:Add. |
| 2 | Click where you want to insert text in the chromatogram. |
| | Result: A text box opens. |
| | Type the text. |
| | Click outside the text box to set the text. |

How to edit the text

The table below describes how to edit inserted text:

| Step | Action |
|------|--|
| 1 | Choose Edit:Text:Edit. Result: The Edit Texts tab of the Chromatogram Layout dialog box is displayed. |
| 2 | Select the text that you want to edit and make the appropriate changes in the Selected text field. Click the Change text button or the Delete text button. Use the Font and Set Orientation buttons if needed, and make the desired changes in the resulting dialog boxes. Click OK to apply the changes. |

Shortcut option

You can also right-click outside the text box and select Edit Text Mode from the shortcut menu. This activates all the text boxes in the chromatogram. The list below describes how to edit the text:

- Click the text and type the new text.
- Click outside the text box to set the text.

11.5 How to pool fractions

Introduction

Fractions are collected sequentially during a separation. Each fraction contains a set volume of sample. This section describes how to pool the information on several fractions into a new curve.

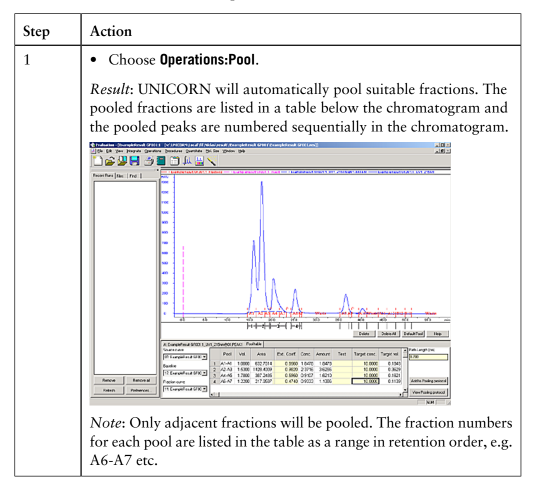
How to view the contents of a fraction

Each fraction is numbered according to its order in the sequence. The information is saved as a curve under the name **Fractions**.

 Select this curve on the Curve tab in the Chromatogram Layout dialog box to display the contents of each fraction in relation to the information displayed on the UV detection curve.

How to pool fractions

The table below describes how to pool fractions.



| Step | Action |
|------|---|
| 2 | The pooled fractions can be adjusted manually: |
| | To include or exclude adjacent fractions in a pool |
| | Click the numbered marker under the pool and drag the sideline. |
| | To add more pools |
| | • Click between the droplines under a fraction to create a new pool, and drag the sidelines to include more adjacent fractions. |
| | To delete pools |
| | • Click the numbered marker to select the pool and click the Delete button. Click the Delete All button to clear all pools. |
| | To restore the pools created by UNICORN |
| | Click the Default Pool button. |
| 3 | Other curves can be selected for the operation: |
| | • Select another source curve from the Source curve droplist and click the Default Pool button. |
| | or |
| | • Select another baseline curve from the Baseline droplist and click the Default Pool button. |
| | or |
| | • Select another fraction curve from the Fraction curve droplist and click the Default Pool button. |
| | <i>Result</i> : The pooled fractions in the list are replaced by the pooled fractions for the selected curve. |

How to create a pool fraction curve

The pooled fractions can be stored as a new curve.

Note: You must store the pooled fractions as a new curve in order to be able to proceed with other operations using the pooled fractions.

| Step | Action |
|------|--|
| 1 | Choose Operations:Create Pool Fraction curve. |
| | Result: The Create Pool Fraction Curve dialog box opens. |

| Step | Action |
|------|---|
| 2 | • Select a position where the curve will be stored from the Save curve in list. |
| | • If needed, type a new name in the Curve name text box. |
| | <i>Note</i>: The suggested curve name will have the default suffix POOL.Click the OK button. |
| | Result: The Pool Fraction curve is displayed in the chromatogram. |

How to show only the pooled fractions

The active chromatogram will now show both the original and the pooled fraction curves. The table below describes how to show only the pooled fractions.

| Step | Action |
|------|--|
| 1 | Choose Edit:Chromatogram Layout. |
| | or |
| | • Right-click in the chromatogram and choose Properties from the shortcut menu. |
| | Result: The Chromatogram Layout dialog box opens. |
| 2 | Select the Curve tab. |
| | De-select the check box for the original fraction curve (remove the check mark). |
| | Result: The original fraction curve is de-selected and is not displayed. |

How to calculate concentration and amount in the pools

Protein concentrations

The protein concentration in the fractions are calculated using the following formula:

Concentration [mg/ml] = A / (d * 1000 *Ext.Coeff.)

A = Average fraction absorbance = Area / Volume [mAu].

d = UV-cell path length [cm]

Ext.Coeff. = Protein coefficient at used wavelength. [1 g⁻¹ cm⁻¹]

Protein amounts

The total amount of protein found in the pool fraction is calculated using the following formula:

Amount [mg] = Concentration [mg/ml] * pooled fraction volume [ml]

How to calculate the concentrations and amounts:

- Type the UV path length expressed in centimeters in the Path Length text box.
- Type the extiction coefficient in the **Ext.Coeff.** table cell for each pool.

Result: The sample concentration and amount for each pool is calculated in the corresponding table cell.

How to determine a pool target volume

The **Target conc.** and **Target vol.** cells are used to calculate the pool volume at a specific concentration level. The result can then be used to determine if the pool needs to be concentrated further or diluted.

• Type the desired concentration level (mg/ml) in the **Target conc.** table cell.

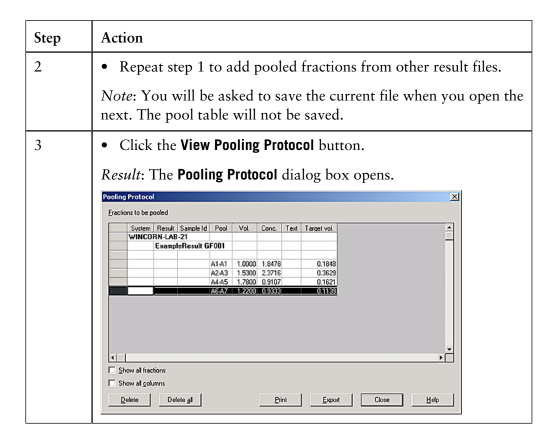
Result: The corresponding target volume is calculated in the **Target vol**. table cell using the following formula:

Target vol. = Conc. * (Vol./Target conc.)

How to use the Pooling Protocol

A protocol of the pooled fractions can be printed for use when handling the samples. The table below describes how to add pools to the **Pooling Protocol** and send the list to a printer or export the list to a file.

| Step | Action |
|------|--|
| 1 | Open a result file in the Evaluation module. |
| | Pool fractions as described in How to pool fractions above. |
| | Click the Add to Pooling Protocol button. |
| | <i>Result</i> : The pooled fractions from the active result file is added to the Pooling Protocol . |



| Step | Action |
|------|--|
| 4 | Click the Show all fractions checkbox to display the individual fractions instead of fraction ranges for the pools. Click the Show all columns checkbox to display all the information columns from the Pool table. |
| | Possible actions in the Pooling Protocol |
| | To delete a single pool |
| | select a pool and click the Delete button |
| | To clear the whole protocol |
| | • click the Delete all button. |
| | To print the protocol on the default Windows printer |
| | • click the Print button to print the protocol on the default Windows printer. |
| | To export the protocol |
| | • click the Export button to save the protocol in one of the following formats: |
| | - text (.txt) |
| | - Excel (.xls) |
| | - HTML (.htm) |
| | - XML (.xml) |
| | <i>Note</i> : The protocol is automatically saved for the user. The pooling protocol will be available again when the user starts UNICORN the next time. |
| 5 | Click the Close button to close the Pooling Protocol dialog box. |
| | Result: If the protocol was exported or only edited, the dialog box will close. If the protocol was printed, a dialog box will open asking if you want to delete the list and start a new. |

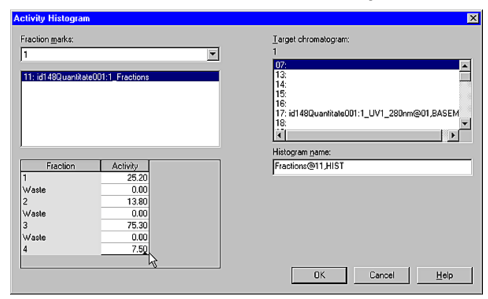
11.6 How to match protein activity to a curve

Introduction

You can compare data from the results of protein activity assays, such as ELISA, with the data contained in the UV curve. The activity curve and the UV curve can be compared in a combined presentation.

The Activity Histogram dialog box

The illustration below shows the **Activity Histogram** dialog box:



How to enter protein activity values for comparison The table below describes how to enter the values from a protein activity assay in a comparison histogram:

| Step | Action |
|------|---|
| 1 | Choose Operations:Activity Histogram. |
| | Result: The Activity Histogram dialog box opens. |
| 2 | By default, the fraction curve for the specific chromatogram is selected. |
| | If necessary, change the source and target chromatograms. |
| | All the component fractions of the fraction curve are listed in the Fraction/Activity field. |
| | Type an activity value for each fraction in the Activity column. Click OK. |

11.7 How to rename chromatograms, curves and peak tables

Instruction

The table below describes how to rename chromatograms, curves or peak tables in the **Evaluation** module:

| Step | Action |
|------|---|
| 1 | Choose Edit:Rename and the relevant option Chromatogram, Curve or Peak Table. |
| | Result: The Rename dialog box opens. |
| 2 | Select the appropriate object. |
| | Type a new name in the Name field. |
| | • Click OK . |

Note: The original raw data curves cannot be renamed. They will not be listed as options in the dialog box.

11.8 How to import and compare different runs

Introduction

This section describes

- how to make comparisons between curves or chromatograms from different
- how to present curves or chromatograms from different runs.
- how to compare curve parameters among curves from different runs
- how to view several chromatograms at the same time
- how to overlay curves from different runs in one chromatogram
- how to stack curves from different runs in one chromatogram
- how to stretch curves to make comparisons easier
- how to create mirror images

In this section

This section contains these sub-sections:

| Topic | See |
|--|--------|
| How to use the Multifile Peak Compare wizard | 11.8.1 |
| How to import and compare chromatograms | 11.8.2 |
| How to import and compare curves | 11.8.3 |
| How to stack and stretch curves | 11.8.4 |
| How to produce a mirror image | 11.8.5 |

11.8.1 How to use the Multifile Peak Compare wizard

Introduction

This section describes how to use the **Multifile Peak Compare** wizard to make comparisons between different results, for example, by comparing area, retention etc. The difference can be presented graphically or in a spreadsheet.

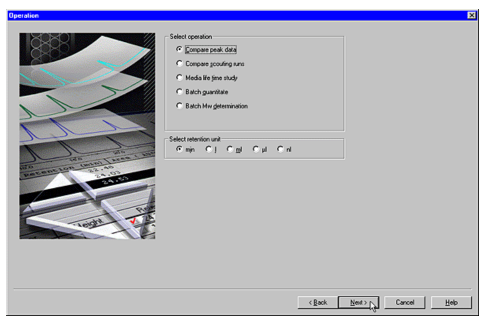
Step 1: How to select the Operation

The table below describes how to select the operation:

| Step | Action |
|------|--|
| 1 | In the Evaluation module, • choose File:Multifile Peak Compare:Start Wizard or • click the Multifile Peak Compare toolbar icon: |
| | Result: The Multifile Peak Compare wizard entry dialog box is displayed. |
| 2 | Click the Next button to display the Operation dialog box. |
| 3 | Select one of the available operations (see descriptions of the operations below this table) a retention unit. |
| | If you select Batch quantitate: |
| | Select a quantitation table in the Select quantitation table field. |
| | If you select Batch Mw determination: |
| | • Select a molecular size table in the Select mol. size table field. |
| | Click the Next button to proceed to the Data Selection dialog box. |

The Operation dialog box

The illustration below displays the **Operation** dialog box:



tions

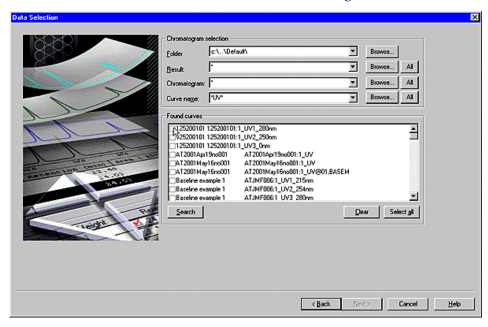
The operation op- The table below is a brief description of the operation options:

| Operation | Description |
|-----------------------|---|
| Compare peak data | This option is used to compare different results. |
| Compare scouting runs | This option is used to compare the results from scouting runs. The scouting variables can be displayed. |
| Media life time study | This option features different default values than the Compare peak data option, specially selected to measure changes in the column media. |
| Batch quantitate | This option is used to run several quantitations. This is an alternative to Quantitate:Calculate Amount and Conc. which is used to quantitate single results. A quantitation table must be created before this option can be used. This option is available only if the Analysis module has been installed. |

| Operation | Description |
|------------------------|---|
| Batch Mw determination | This option is used to batch run molecular size calculations. This is an alternative to Mol. Size:Calculate Mol.Size, which is used for single calculations. A molecular size table must be created before this option can be used. |
| | This option is available only if the Analysis module has been installed. |

The Data Selection dialog box

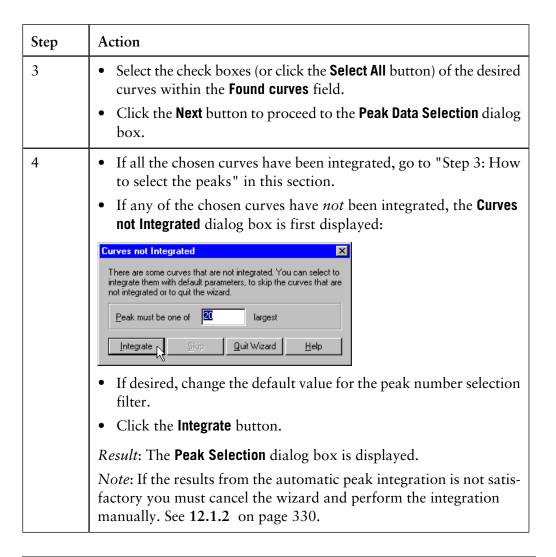
The illustration below shows the **Data Selection** dialog box.



Step 2: How to select data to compare

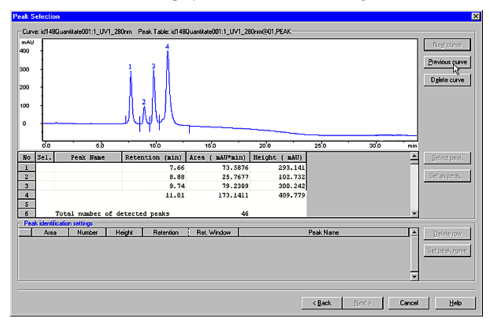
The table below describes how to select data to compare:

| Step | Action |
|------|--|
| 1 | Use the drop-down lists and Browse buttons in the Chromatogram selection field to specify the result files, chromatograms and curves for comparison. Click the All button if you want to select all available results, chromatograms or curves. |
| 2 | Click the Search button in the Found curves field. |
| | <i>Result</i> : A list of all curves that matched the search criteria is displayed in the Found curves field. |



The Peak Selection dialog box

The illustration below displays the **Peak Selection** dialog box:



- 11 How to edit results
- 11.8 How to import and compare different runs
- 11.8.1 How to use the Multifile Peak Compare wizard

Dialog box description

The dialog box displays the following properties for the first of the chosen curves:

- The integrated peak and the associated peak table
- The **Peak identification settings** table. Its purpose is to identify the peak parameter to be used in the comparison.

How to adjust improper peak integrations

The table below describes what to do if the peaks in the curve window do not appear to be integrated properly (for example if ghost peaks are labelled).

| Step | Action |
|------|---|
| 1 | Click the Cancel button to quit the wizard. |
| 2 | Perform a peak integration (see 12.1.2 How to perform a peak integration on page 330) and verify that the resulting curve is properly integrated. |
| 3 | Repeat the Multifile Peak Compare wizard operation. |

Step 3: How to select the peaks

The table below describes how select peaks in the **Peak Selection** dialog box:

| Step | Action |
|------|---|
| 1 | Choose a curve in the curve window: |
| | Double-click the peak, or click the peak once and then click the Select peak button. |
| | <i>Result</i> : The peak is assigned a letter (A, B, C) and the peak parameters are displayed in the Peak identification settings table. |

| Step | Action |
|--------|--|
| Step 2 | Set the desired peak identification criterion: • Click the desired parameter value in the Peak identification settings table. Example: If you have selected the highest peak in the curve and want to compare the highest peak among all curves, select the Height check box. In the illustration below, the initial (A) peak and the Height check box have been selected: Production Total makes of detected peaks in the curve and want to compare the highest peak among all curves, select the Height check box. In the illustration below, the initial (A) peak and the Height check box have been selected: Production Total makes of detected peaks in the curve and want to compare the highest peak in the curve and want to compare the highest peak among all curves, select the Height check box. In the illustration below, the initial (A) peak and the Height check box have been selected: Total makes of detected peaks in the curve and want to compare the highest peak in the curve and want to compar |
| 3 | If desired, you can assign a name to a chosen peak: Click the name of the row, for example A. Click the Set peak name button. Type a new name and click OK. Note: This can be useful when you compare multiple peak parameters and you wish to have peak names other than "Peak A", "Peak B", etc. to simplify peak identification and clarity f.ex. when comparing peak data between batch quantitated results. |
| 4 | Repeat steps 1-3 for other desired peaks in the current curve. |
| 5 | Use the Next curve and Previous curve buttons to navigate forward and backward among your selected curves and manually check the selections made by the software if necessary. |
| 6 | Other possible actions you can perform If the current curve does not prove useful for your comparison, click the Delete curve button to delete it from the comparison. Click the Back button to navigate back to the Data Selection dialog box and add new curves to your comparison. See also How to change the peak identification below. |

| Step | Action |
|------|--|
| 7 | When all peak selections and identification settings are complete, click the Next button to proceed to the Peak Data Selection dialog box. |

Note: Click and drag in the curve window to zoom into selected peaks to simplify accurate peak identification. Right-click and click the **Reset Zoom** button to reset the zoom to the full view.

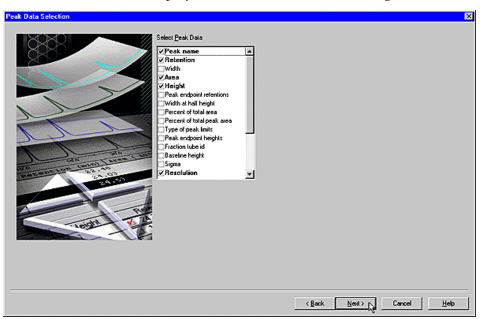
How to change the peak identification

In the **Peak identification settings table**, each column identifies a peak parameter to be compared among all peaks. If UNICORN has identified other peaks than the intended ones, you can change the peak identification manually. The table below describes how to change the identification:

| If you want to | then |
|--------------------------------------|--|
| remove a peak identification | click the desired peak in the curves window click the Set as peak button choose None in the Set As Peak dialog box click OK. |
| replace or add a peak identification | click a peak in the curves window click the Set as peak button choose a letter in the Set As Peak dialog box click OK. |
| remove a row from the table | select the row click the Delete row button. <i>Note</i>: If you click Delete row without first selecting a row, the first row (A) is deleted by default. |

Step 4: How to select the Peak Data

The illustration below displays the **Peak Data Selection** dialog box:

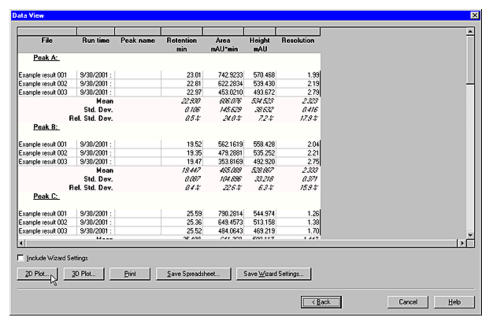


The table below describes how to select the peak data:

| Step | Action |
|------|---|
| 1 | In the Select Peak Data list, select the peak characteristics on the list that you want to include in your comparisons. If available, select the appropriate Scouting variables. |
| 2 | Click the Next button and proceed to step 5, How to use the Data |
| | View dialog box below. |
| | Note: If Media life time study was chosen in the Operation dialog box when the wizard was started, 2D Plot is selected in the Data View dialog box. |

Step 5: How to use the Data View dialog box

The **Data View** dialog box presents a comparison of the chosen data for the designated peak comparisons. The illustration below shows the dialog box:

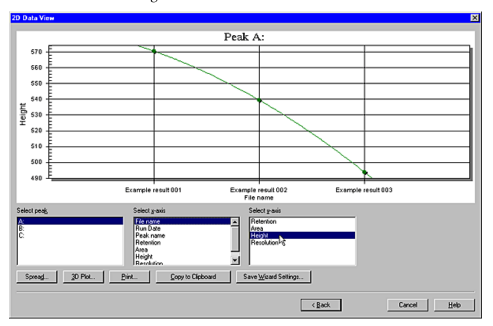


The table below describes how to use the command buttons of the dialog box:

| Command button | Function |
|----------------------|---|
| 2D Plot | Displays the data in 2-dimensional plot. See "How to use the 2D Data View dialog box" below. |
| 3D Plot | Displays the data in 3-dimensional plot. See "How to use the 3D Data View dialog box" below |
| Print | Prints the spreadsheet. |
| Save Spreadsheet | Allows you to save the data in different formats: • Excel (.xls) • Tabbed text (.txt) • FarPoint spread (.ss3) |
| Save Wizard Settings | See "How to save the Wizard Settings" below. |
| Cancel | Ends the Multifile Peak Compare wizard. |

How to use the 2D Data View

The **2D Data View** dialog box presents a two-dimensional plot of a selected peak. See also "How to use the 2D Data View shortcut menu" below. The illustration below shows the dialog box:



The list boxes

Use the list boxes to select which peak to plot and the units of the x- and y-axes.

The command buttons

The table below describes how to use the command buttons of the dialog box:

| Command button | Description |
|----------------------|--|
| Spread | Returns to the Data View dialog box. |
| 3D Plot | Displays the data in 3-dimensional plot. See "How to use the 3D Data View dialog box" below. |
| Print | Prints the spreadsheet. |
| Copy to Clipboard | Stores a figure for transfer to an external program. |
| Save Wizard Settings | See "How to save the Wizard Settings" below. |
| Cancel | Ends the Multifile Peak Compare wizard. |

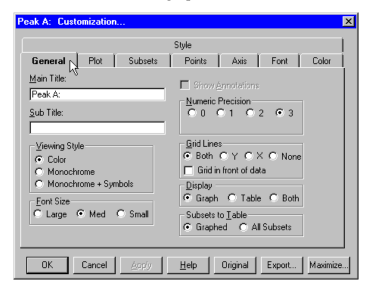
How to use the 2D Data View shortcut menu

Click the right mouse button in the plot area of the **2D Data View** dialog box to open the shortcut menu. See illustration below:



A wide array of plot presentation options can be found on the shortcut menu. Two of them are described below:

• Select **Customization Dialog** to open a dialog box which allows further customization of the graph:

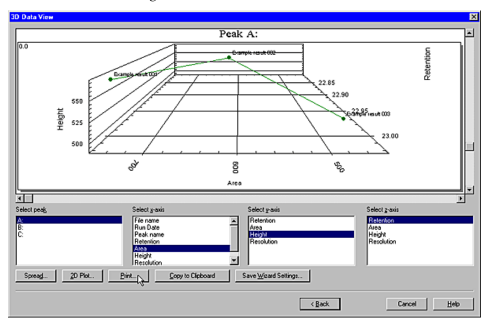


• Select **Export Dialog** to export the view.

Note: You can also click the **Export** button from the **Customization** dialog box.

How to use the 3D Data View dialog box

The **3D Data View** dialog box presents a three-dimensional plot of a selected peak. See also "How to use the 3D Data View shortcut menu" below. The illustration below shows the dialog box:



The list boxes

Use the list boxes to select which peak to plot and the units of the x-, y- and z-axes.

The command buttons

The table below describes how to use the command buttons of the dialog box:

| Command button | Function |
|----------------------|--|
| Spread | Returns to the Data View dialog box. |
| 2D Plot | Displays the data in 2-dimensional plot. See "How to use the 2D Data View dialog box" above. |
| Print | Prints the spreadsheet. |
| Copy to Clipboard | Stores a figure for transfer to an external program. |
| Save Wizard Settings | See "How to save the Wizard Settings" below. |
| Cancel | Ends the Multifile Peak Compare wizard. |

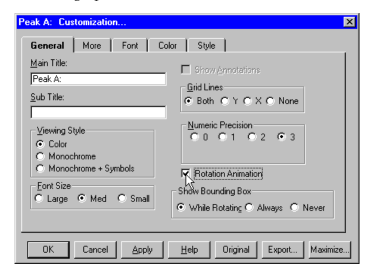
How to use the 3D Data View shortcut menu

Click the right mouse button in the plot area of the **3D Data View** dialog box to open the shortcut menu. See illustration below:



The **3D Data View** shortcut menu differs some from the **2D Data View** shortcut menu and allows the figure to be viewed by animated rotation. The shortcut menu displays different plot presentation options.

• Select **Customization Dialog** to open a dialog box that allows further customization of the graph:



• Select **Export Dialog** to export the view.

Note: You can also click the **Export** button from the **Customization** dialog box.

How to save the Wizard Settings

The wizard settings can be saved from either of these dialog boxes:

- The **Data View** dialog box
- The **2D Data View** dialog box
- The **3D Data View** dialog box

The table below describes how to save the wizard settings:

| Step | Action |
|------|---|
| 1 | Click the Save Wizard Settings button. |
| | Result: The Save Wizard Settings dialog box opens. |
| 2 | Type a name in the Wizard settings name field. |
| 3 | • If the settings are to be used by all users on the system, select the Global wizard settings check box. |
| | Click 0K . |
| | Click Cancel to close the wizard. |
| | <i>Note</i> : The Global wizard settings check box can also be used to toggle between lists of stored global and stored user settings. |

How to open the saved wizard settings

The table below describes how to open the saved wizard settings:

| Step | Action |
|------|--|
| 1 | Choose the File:Multifile Peak Compare:Start Wizard With Settings menu item. |
| | Result: The Select Wizard Settings dialog box opens. |
| 2 | Select the desired saved settings from the list. Click 0K. |
| | Result: The Multifile Peak Compare wizard opens with the saved settings. |
| | Note: The Global wizard settings check box is used to toggle between lists of stored global and stored user settings. |

11.8.2 How to import and compare chromatograms

Introduction

This section describes

- how to import chromatograms from other result files,
- how to compare with chromatograms in an already opened result file.

Commands to use

Two commands in the **Evaluation** module can be used to import chromatograms from result files into an already opened result file:

File:Open to compare

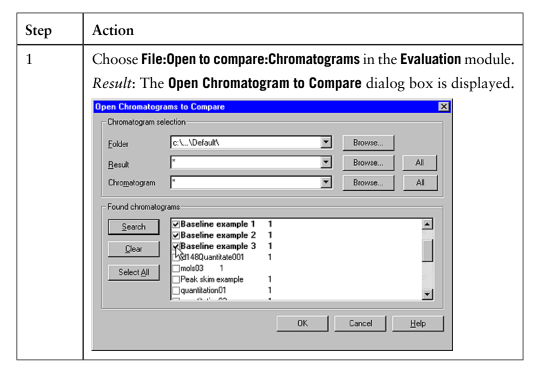
This is the preferred option when you search for many chromatograms in a *specific* folder based on defined selection criteria. See "How to import chromatograms with the command **File:Open to compare**" below.

• File:Open

This is the preferred option to import any individual chromatograms from result files in *different* folders. See "How to import chromatograms with the command **File:Open**" below.

How to import chromatograms with the command File:Open to compare

The table below describes how to import chromatograms with the **File:Open to compare** command. The search is performed at specific locations or with specific search criteria. This method is useful if you, for example, want to import chromatograms from all files of a scouting folder.



| Step | Action |
|------|---|
| 2 | • Click the Search button in the Found chromatograms field and a list of chromatograms will be displayed based on the designated search criteria. |
| | • A new search can be performed with new search criteria without erasing the first found chromatograms from the list. |
| | • Select the chromatograms that you want to import. If you click the Select All button, all the displayed chromatograms will be imported. |
| | • If you want to clear the list of displayed chromatograms, click the Clear button. |
| | Click 0K . |
| | <i>Result</i> : All the selected chromatograms are shown in the Evaluation workspace. |
| | <i>Note</i> : If the names of the imported chromatograms already are used they will be sequentially numbered for identification purposes. Up to 10 chromatograms can be made available at the same time in the Evaluation workspace. |

How to import chromatograms with the command File:Open

The table below describes how to import chromatograms one by one, using the command **File:Open:**

| Step | Action |
|------|--|
| 1 | Choose File:Open:Chromatogram in the Evaluation module. Result: The Open Chromatograms dialog box is displayed. |
| | |
| 2 | Double-click a result file to select it. |
| | Result: All the chromatograms contained in the result file will be displayed in the Available field. |
| 3 | Select the chromatogram(s) of interest and click the Select button. |
| | Result: Selected chromatograms are added to the Selected chromatograms list. |
| | <i>Note</i> : Chromatograms can be deselected with the Remove button. |

- 11 How to edit results
- 11.8 How to import and compare different runs
- 11.8.2 How to import and compare chromatograms

| Step | Action |
|------|---|
| 4 | Repeat steps 2-3 if you want to import chromatograms from other result files. Click 0K. |
| | <i>Note</i> : If the names of the imported chromatograms already are used they will be sequentially numbered for identification purposes. Up to 10 chromatograms can be made available at the same time in the Evaluation workspace. |

How to display and compare the imported chromatograms

The table below describes how to simultaneously display and compare the imported chromatograms:

| Step | Action |
|------|--|
| 1 | In the Evaluation module, select |
| | Window:Tile to display the chromatograms side by side. |
| | or |
| | Window:Cascade to display the chromatograms in layers. |
| | <i>Note</i> : Chromatogram windows can be individually sized and the presentation of the curves changed. |
| 2 | Display all chromatograms on the same scale |
| | Open the Chromatogram Layout dialog box for any chromatogram |
| | Make the changes to the chromatogram axes. |
| | Select the Apply to all chromatograms option. |

Note: Imported chromatograms cannot be shown with column volume as the X-axis base.

11.8.3 How to import and compare curves

Introduction

This section describes how to import or copy curves from different runs into one chromatogram for comparison.

Commands to use

Two commands can be used to import curves from result files into one chromatogram:

• File:Open to compare

This is the preferred option if you want to automatically search result files that are stored in the same folder to locate all curves of a specified type, for example, all UV curves. This is especially useful for comparison of curves from scouting runs. Moreover, the imported curves can be automatically overlaid, stacked or presented as mirror images. See "How to use File:Open to compare" below.

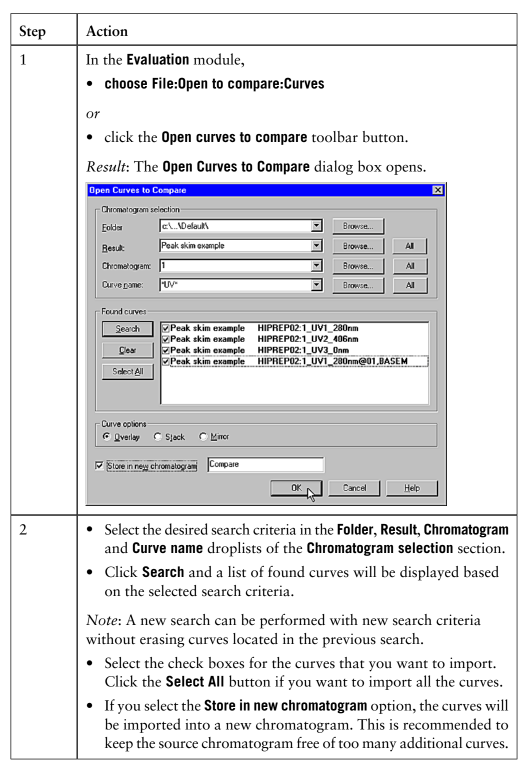
• File:Open:Curves

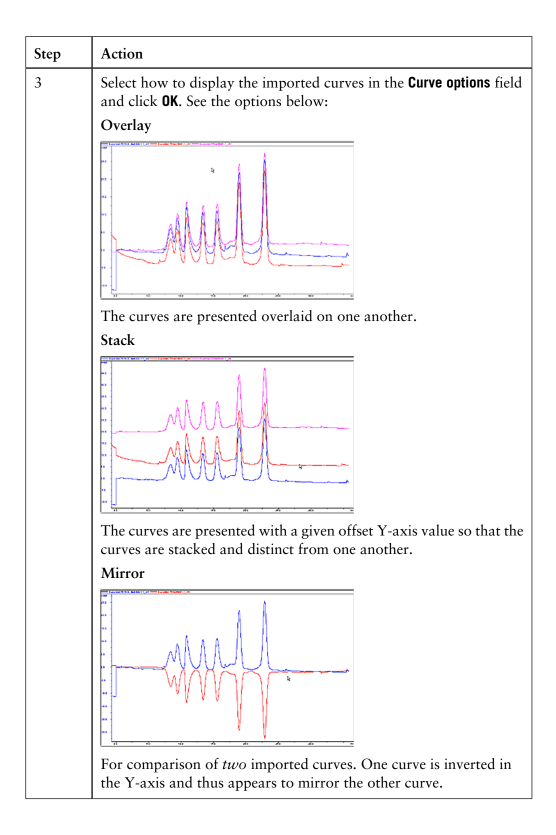
This is the preferred option to import individual curves. See " How to use File:Open:Curves" below.

Note: Original curves are underlined in the chromatogram, imported and created curves are not underlined.

How to use File:Open to compare

The table below describes how to import curves to a chromatogram with the command **File:Open to compare**:





| Step | Action |
|------|---|
| 4 | If you selected the Stack option in step 3, the Shift Curves by Offset dialog box is displayed: |
| | Shift Curves by Offset Offset: Unit: Selected curves will be shifted V01: Superdex 75 NCC test 3001:1_UV V02: Superdex 75test 4001:1_UV V03: Superdex 75test5001:1_UV V04: Superdex 75test5001:1_UV OK Cancel Help |
| | You can set the Offset value to increase or decrease the offset distance between the curves. Click OK. |
| | Result: Depending on your previous choices, the imported curves are now displayed in the source chromatogram or in a newly created chromatogram. Note: If curves with several different units have been selected, the curves with each different unit will be grouped together with separate offset from the other groups. |
| 5 | Change some comparison settings • Choose Edit:Chromatogram Layout to open the Chromatogram Layout dialog box. |
| | • Select or de-select the check boxes on the Curve tab to compare a different set of curves. |
| | On the Y-Axis tab, the curves can be scaled individually |
| | all with the same scale (click the All with this unit button). Click OK to display the curves. |
| 6 | If you stacked the curves and want to change the stack offset • choose Operations:Shift offset • type a new Offset value and click OK. |
| | Note: The individual curves can also be moved (see 11.8.4 How to stack and stretch curves on page 312). |

How to use File:Open:Curves

The table below describes how to import individual curves into an active chromatogram with the File:Open:Curves command:

| Step | Action |
|------|---|
| 1 | Make sure that the destination chromatogram for the imported curve(s) is active on the screen. |
| | Select File:Open:Curves in the Evaluation module. |
| | Result: The Open Curves dialog box is displayed. |
| 2 | Select curves in the Open curves dialog box |
| | • Select the folder and the result file in the upper part of the dialog box. |
| | Select a chromatogram on the Chromatogram drop-down list. Usually there is just one chromatogram. |
| | Result: The available curves are listed on the Available list. |
| | • Click the check boxes on the Available list for the curves that you want to import and click the Select button. |
| | Result: The selected curve(s) is displayed in the Selected curves list. To remove a curve from the Selected curves list, click the check box and then click the Remove button. |
| 3 | • Repeat step 2 if you want to import curves from other chromatograms. |
| | Click 0K when you have selected the curves you want. |
| 4 | Change some comparison settings |
| | Choose Edit:Chromatogram Layout to open the Chromatogram Layout dialog box. |
| | • Select or de-select the check boxes on the Curve tab to compare a different set of curves. |
| | On the Y-Axis tab, the curves can be scaled |
| | - individually |
| | - all with the same scale (click the All with this unit button). |
| | Click OK to display the curves. |

How to copy curves into one chromatogram

A practical way to compare curves is to create a chromatogram and copy curves from different chromatograms into the new chromatogram. The comparisons are then performed in the new chromatogram.

The table below describes how to copy curves into a chromatogram:

| Step | Action |
|------|--|
| 1 | Perform either A or B below: A. Create a new chromatogram • Choose File:New:Chromatogram to create a new chromatogram. B. Use the Temporary chromatogram • Choose Window:Temporary. |
| 2 | Open the source chromatogram(s) Choose File:Open:Chromatogram to open the chromatogram(s) that contains the curves you want to copy. Result: The Open Chromatogram dialog box opens. |
| 3 | Select the result file. Click the check box for the source chromatogram in the Available list. Click the Select button. Click OK. Result: The source chromatogram opens. |
| 4 | Copy the curves • Choose Edit:Copy:Curves. Result: The Copy Curve dialog box is displayed. |
| 5 | Select the source chromatogram and a curve of interest in the Source Chromatogram field. Select the target chromatogram (the one you created, or Temporary) in the Target Chromatogram field. Click the Copy button. Repeat this step for as many curves as you want, from the same or other chromatograms. Note: You can open more source chromatograms with the File:Open:Chromatogram command. Click the Close button when you have copied all curves. |

| Step | Action |
|------|---|
| 6 | Change some comparison settings |
| | • Make sure the target chromatogram is open and that its window is active. |
| | Choose Edit:Chromatogram Layout to display the Chromatogram Layout dialog box. |
| | • Select the curves that you want to view on the Curve tab and click OK . |
| | • The curves can be scaled individually or all with the same Y-axis scale. Use the All with this unit button on the Y-Axis tab to scale all curves with the same scale. |
| 7 | If you used the Temporary chromatogram |
| | • If you used the Temporary chromatogram you can perform evaluations in the Temporary chromatogram and transfer the final curves to other destination chromatograms. |
| | All of the contents in the Temporary chromatogram can be removed with Edit:Clear Temporary Chromatogram . |

Alternative way to copy curves

An alternative way to copy curves into one chromatogram is to

- create a new chromatogram by copying an existing chromatogram and saving it under a new name
- import more curves into the new chromatogram according to the instructions described above in this section.

11.8.4 How to stack and stretch curves

Functions

You can stack and stretch curves from different runs to better visualize the differences. To achieve this you can use the following functions:

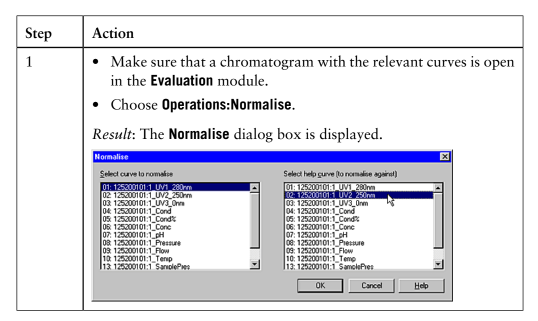
- Normalise
- Shift
- Multiply.

Note: All the functions require the curves to be present in one chromatogram.

How to use the Normalise function

The **Normalise** function provides the simplest method to align curves with respect to the X-axis or the Y-axis for easier visualization.

The table below describes how to use the **Normalise** function:



| Step | Action | | |
|------|--|--|--|
| 2 | • Select the curve you want to normalise in the left (Select curve normalise) field. | | |
| | Select the reference curve you want to normalise <i>against</i> in the right (Select help curve) field. <i>Example</i>: If you want to <i>stack</i> the curves, select the curve at the bottom of the stack as the reference curve. Click OK. | | |
| | Result: The Normalise window is displayed, where a box surrounds the curve selected to be normalised. | | |
| | 5-ye-day 75 ind 4001 1 _UVQQQ JACAM Superjay 75 HCC find 3001-1 _UV | | |
| | 150 - | | |
| | | | |
| | SD 100 - SD | | |

| Step | Action | |
|------|---|--|
| 3 | In the Normalise window, you can use the following command buttons: Size Allows the arrow keys to be used to stretch the selected curve | |
| | along its Y-axis or X-axis. This is useful for comparison of curves with, for example, different gradient lengths. | |
| | • Click the Size button and use the arrow keys to stretch the the curve either along its Y-axis or X-axis. | |
| | Move Allows the arrow keys to be used to move the selected curve to any position on the chromatogram. Axes are automatically rescaled to accommodate the new positioning. This function is useful for stacking curves. | |
| | • Click the Move button and use the arrow keys to move the curve into position. The curve can also be moved with the mouse pointer. Click the mouse button when the curve is in the correct position. | |
| | <i>Note</i> : The curve can also be moved and sized with the mouse pointer. | |
| | Normalise The curve to be normalised will be adjusted to the help curve. Thus, the height of the highest peak on both curves will be the same and will occur at the same retention point. | |
| | • Click the Normalise button. The curve to be normalised is automatically moved along the X-axis and stretched along the Y-axis. | |
| | • Click OK to save the new normalised curve. *Result: The Save Curve dialog box opens. | |
| 4 | Choose a curve position to save the curve in and click OK . | |
| 5 | Choose Edit:Chromatogram Layout to open the Chromatogram Layout dialog box. | |
| | Select the normalised curve for viewing on the Curve tab. Click OK. | |
| 6 | Repeat steps 1-5 for all curves you want to stack or stretch. | |

How to move a curve with the Shift function

If you want to position a curve more precisely, the **Shift** function should be used. The function is similar to **Normalise:Move** but each curve is repositioned by a precise value instead of by eye and the instruction is logged in the evaluation log. The table below describes how to use the **Shift** function:

| Step | Action | |
|------|---|--|
| 1 | Make sure that a chromatogram with the relevant curves is open in the Evaluation module. | |
| | Choose Operations:Shift. | |
| | Result: The Shift dialog box is displayed. | |
| 2 | Select the curve to be shifted in the Source chromatogram list. | |
| | • Select a curve position in the Target chromatogram list. | |
| | Type a new Curve name or accept the default. | |
| | Select the axis/axes along which the shift is to be made: along the X-axis (Shift retention) | |
| | | |
| | - along the Y-axis (Shift amplitude). | |
| | • Type the shift value(s). | |
| | Click OK . | |

How to stretch and shrink a curve with the Multiply function Curves can be stretched or shrunk on the x or y plane with the **Multiply** function. This function is similar to **Normalise:Size**, but each curve is repositioned with precise numbers instead of by eye and the instruction logged in the evaluation log. The table below describes how to use the **Multiply** function:

| Step | Action |
|------|---|
| 1 | • Make sure that a chromatogram with the relevant curves is open in the Evaluation module. |
| | Choose Operations:Multiply. |
| | Result: The Multiply dialog box is displayed. |
| 2 | Select the curve to be multiplied in the Source chromatogram list. |
| | Select a curve position in the Target chromatogram list. |
| | Type a new Curve name or accept the default. |
| | Select the axis/axes along which the multiplication is to be made: |
| | - along the X-axis (Multiply retention) |
| | - along the Y-axis (Multiply amplitude). |
| | Type the multiply value(s). |
| | Click OK . |

11.8.5 How to produce a mirror image

11.8.5 How to produce a mirror image

Instruction

A very useful way to compare the features of two curves is to produce a mirror image of one curve. The table below describes how to do this:

| Step | Action | |
|------|--|--|
| 1 | • Make sure that a chromatogram with the relevant curves is open in the Evaluation module. | |
| | Choose Operations:Multiply. | |
| | Result: The Multiply dialog box is displayed. | |
| 2 | Select the curve to be multiplied in the Source chromatogram list. Select a curve position in the Target chromatogram list. Type a new Curve name or accept the default. Select the Multiply amplitude check box. Type the multiply value -1. Click OK. | |
| | <i>Result</i> : The mirror image of the original curve is displayed in the active chromatogram window. | |

| Step | Action | |
|---|---|--|
| 3 | Shift the mirror image curve downwards | |
| | Shift the mirror image curve downwards for an improved presentation: | |
| | Choose Operations:Shift. | |
| | Result: The Shift dialog box is displayed. | |
| | • Select the curve to be shifted in the Source chromatogram list. | |
| | Select the same curve number in the Target chromatogram list box as in step 2. Select the Shift amplitude check box since the shift is to be made along the Y-axis. | |
| | | |
| | Type a shift value. | |
| | Click 0K . | |
| The illustration below shows the original curve and the mirridisplayed. | | |
| | 250 - 250 - 250 - 250 - 255 - 202 - 202 - 255 - 202 - | |
| 4 | If you want to display other curves in the active chromatogram window, • choose Edit:Chromatogram Layout to open the Chromatogram Layout dialog box • select the curves that you want to display • click OK. | |

11.9 How to import and export results

Introduction

Curves and data can be imported and exported in different formats. This section describes how to import and export results.

In this section

This section contains these topics:

| Topic | See |
|-----------------------|--------|
| How to import results | 11.9.1 |
| How to export results | |

11.9.1 **How to import results**

Introduction

This section describes how to import curves in different formats and how to import result data from SMART Manager, FPLCdirector™ or ÄKTAprime™.

Curve formats

You can import curve files in the following formats:

- AIA (.cdf)
- ASCII (text)
- Lotus 1-2-3 spreadsheet (.wks)

How to import curves

The table below describes how to import curves.

| Step | Action | |
|------|--|--|
| 1 | Choose File:Import:Curve. | |
| | Result: A menu with the available curve formats opens. | |
| 2 | Choose the correct curve format. | |
| | Result: The Choose File to Import From dialog box opens. | |
| 3 | Locate the file that contains the curve and double-click the file. | |
| | Result: The Import Curves dialog box opens. | |
| 4 | Select the curve(s) to import and click the OK button. | |
| | Result: The curves are opened in the Evaluation module. | |

How to import data from **SMART Manager** and FPLCdirector The table below describes how to import data from **SMART Manager** and **FPLC**director:

| Step | Action | |
|------|---|--|
| 1 | Choose File:Import:Result. | |
| | Result: A menu box with the available data sources opens. This box opens immediately after Import if no result file is open in the Evaluation module. | |
| 2 | Choose FPLCdirector or SMART. | |
| | Result: The Import FPLCdirector Result dialog box or the Import SMART Result dialog box opens. | |
| 3 | Locate and double-click the result file. | |
| | Result: The result file is opened in the Evaluation module. | |

- 11 How to edit results
- 11.9 How to import and export results
- 11.9.1 How to import results

Copy from a floppy disk

When you import **SMART** or **FPLCdirector** files from a floppy disk it is best to first copy the files to the hard disk and then import the files.

How to import data from AK-TAprime

The table below describes how to import **ÄKTAprime** data in the **Evaluation** module.

| Step | Action | |
|------|---|--|
| 1 | Choose File:Import:Result. Result: A menu box with the available data sources opens. This box opens immediately after Import if no result file is open in the Evaluation module. | |
| 2 | Choose ÄKTAprime. Result: The ÄKTAprime Data Collection wizard opens. | |
| 3 | Click the Next button and follow the instructions to connect ÄK-TAprime to the correct serial port on your computer. Click the Finish button to complete the setup. Result: All open result files are closed and UNICORN is set up to receive sample data from ÄKTAprime. | |

11.9.2 How to export results

Introduction

This section describes how to export curves in different formats and how to copy data and curves to the clipboard.

Data formats

You can export data in the following formats:

- AIA (.cdf)
- ASCII (.asc)
- Lotus 1-2-3 (.wks)
- Excel (.xls)
- XML (.xml)

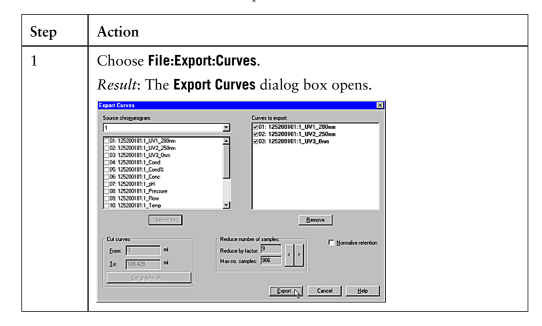
Export options

Select **File:Export** in the **Evaluation** module to export data from an open result file. The following export options are available:

- Curves
- Export curve to AIA
- · Peak table
- Method
- Documentation
- Evaluation log

How to export curves

The table below describes how to export curves in the **Evaluation** module.



| Step | Action | |
|------|---|--|
| 2 | Select the curve(s) you want to export. Enter parameters to limit the curve(s) if necessary. | |
| | | |
| | Click the Select button. | |
| | • Repeat Step 2 to select more curves. | |
| 3 | Click the Export button. | |
| | Result: The Export Curves to File dialog box opens. | |
| 4 | Select the export file format from the Save as type droplist. | |
| | ASCII files (*.asc) | |
| | • Lotus 1-2-3 files (*.wks) | |
| | Excel files (*.xls) | |
| | AIA files (*.cdf) | |
| 5 | Select a destination folder. | |
| | Type a file name and click 0K . | |

Note: Curves are exported as series of numerical coordinates that refers to the time/volume and signal respectively.

How to limit the exported curves

You can optimize the exported curves to only the parts that you want to focus on, in the **Export Curves** dialog box. The table below describes how to use these editing options.

| Dialog box option | Instruction |
|--------------------------|---|
| Cut curves | Enter retention values in the text boxes to limit the curve to only a portion of the original curve. |
| Cut graphically | This button opens the Export Cut dialog box. Move the vertical markers to the correct cutoff points. |
| Reduce number of samples | Adjust the factor value or the maximum number of samples. To reduce the number of samples by a factor of five means that only every fifth point will be sampled for export. |
| Normalise retention | Select the Normalise retention checkbox to have all exported curves normalized to a common X-axis. |

How to export curves in AIA format

The table below describes how to export curves in AIA format.

| Step | Action | |
|------|--|--|
| 1 | Select File:Export:Export curve to AIA. | |
| | Result: The Export curve in AIA format dialog box opens. | |
| 2 | Select the source chromatogram and the curve you want to export. Click the Export button. | |
| | Result: The Export Curves to File dialog box opens. | |
| 3 | Select a destination folder. Type a file name. Click OK. | |

How to export peak tables

The table below describes how to export peak tables.

| Step | Action | |
|------|---|--|
| 1 | Choose File:Export:Peak Table. Result: The Export Peak Table dialog box opens. | |
| | Result: The Export Fear Table dialog box opens. | |
| 2 | • Select the source chromatogram and the peak table you want to export. | |
| | Click the Export button. | |
| | Result: The Export Peak Table to File dialog box opens. | |
| 3 | Select the export file format from the Save as type drop-list. | |
| | ASCII files (*.asc) | |
| | • Lotus 1-2-3 files (*.wks) | |
| | Excel files (*.xls) | |
| | XML files (*.xml) | |
| 4 | Select a destination folder. | |
| | Type a file name. | |
| | • Click OK . | |

Note: Peak tables are exported as text strings in ASCII format and numerical values in the Lotus 1-2-3 formats. All possible columns in the peak table are exported.

- 11 How to edit results
- 11.9 How to import and export results
- 11.9.2 How to export results

How to export methods, documentation and evaluation logs

The table below shows how to export methods, documentation and evaluation logs:

| Step | Action | |
|------|---|--|
| 1 | Select the data you want to export. | |
| 2 | Select options in the dialog box. Click the Export button. | |
| 3 | Select a destination folder and type a file name. Click OK. | |

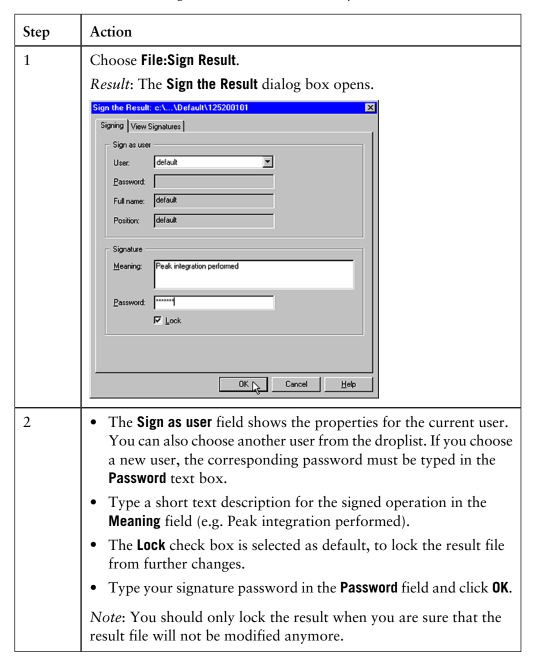
Copy to the clipboard

You can also use the **Windows** clipboard to copy the contents of the active window and paste it into other programs, e.g. **Microsoft Word**. Curves and documentation are copied as Windows enhanced metafiles (.emf) and peak tables are copied as text. Only the peak table columns that are selected in the spreadsheet will be copied.

11.10 How to sign results electronically

Instruction

Result files can be signed electronically to enhance data file security. The table below describes how to sign a result file electronically in the **Evaluation** module:



Signatures associated with the result

The **View Signatures** tab of the **Sign the Result** dialog box provides a list of all signatures associated with the current result. The information on this tab is for viewing purposes only and cannot be changed.

11.11 How to save results and exit the Evaluation module

Introduction

After you have finished the evaluation process, you can save all the changes you have made to the chromatograms, including newly created curves and chromatograms that you have imported and created.

How to delete unwanted curves

All the curves that you created during your manipulations will be saved in the chromatogram. If some of these curves are not be needed anymore, select **Edit:Delete:Curves** in the **Evaluation** module to remove the curves.

Note: The original curves that were created during the run can never be deleted.

How to save the results

You can either save your edited results in the original file or in a new result file. The table below describes how to save the results in the **Evaluation** module.

| If you want to save the edited results | then |
|--|---|
| in the original result file | select File:Save. or click the Save toolbar icon. |
| in a new result file | • select File:Save as. |

Note: The previous version of the result file will be overwritten if you save the changes. This cannot be reversed. However, the raw data curves remain unchanged.

How to exit the Evaluation module

The table below describes how to exit the **Evaluation** module:

| Step | Action | |
|------|---|--|
| 1 | Choose File:Exit. | |
| | Result: If there are unsaved changes, a dialog box opens with an option to save the changes before exit. | |
| 2 | Select Yes if you want to save the changes. | |
| | Result: The result file is closed in the Evaluation module and the UNICORN Manager module is displayed. | |

Evaluation 12

Introduction

This chapter describes:

- How to evaluate results with the focus on how to integrate peaks.
- How to automate evaluation operations.
- How to export data and curves.

In this chapter

This chapter contains these sections.

| Topic | See |
|---------------------------------|------|
| Peak integration | 12.1 |
| Other evaluations | |
| Automated evaluation procedures | |

12.1 Peak integration

Introduction

Peak integration is used to identify and measure a number of curve characteristics including peak areas, retention time and peak widths. This section describes:

- How to perform peak integrations.
- How to optimize peak integrations.

In this section

This section contains these sub-sections:

| Topic | See |
|---|--------|
| Baseline calculation | 12.1.1 |
| How to perform a peak integration | 12.1.2 |
| How to optimize the baseline with a morphological algorithm | 12.1.3 |
| How to optimize the baseline with a classic algorithm | 12.1.4 |
| How to edit the baseline manually | 12.1.5 |
| How to edit the peaks | 12.1.6 |
| How to integrate part of a curve and how to exclude or skim peaks | 12.1.7 |
| Measurements | 12.1.8 |

12.1.1 Baseline calculation

Introduction

The first step when you integrate peaks is to calculate a baseline. A correct baseline is crucial for accurate calculation of the peak areas. This section describes the options for how to calculate baselines in the **Integrate** dialog box.

Baseline options

UNICORN offers several options for how to create an accurate baseline:

- To use the automatic Calculate baseline function.
- To create a baseline based on a blank curve.
- To use a Zero baseline.
- To reuse an existing baseline.

The Calculate baseline function

The **Calculate baseline** instruction provides automatic calculation of the baseline. In most cases the measurement is very accurate. The calculation can be performed using the **Morphological** algorithm or the **Classical** algorithm.

Baselines based on a blank curve

A blank curve can be used as the baseline for peak integration.

• You can use a blank curve with the same chromatographic conditions as the corresponding sample.

or

• You can subtract the blank run from the source curve and then perform peak integration on the resulting curve with the **Calculate baseline** instruction.

Note: In addition to blank run curves, it is also possible to select any curve from the current chromatogram as the baseline, e.g. an edited baseline.

Zero baseline

To use a **Zero baseline** means that there is no baseline subtraction at all.

Reuse an existing baseline

To reuse an existing baseline for the selected curve is the default alternative whenever there is an existing baseline available. The option **Correlated baseline** is selected if this is the case.

12.1.2 How to perform a peak integration

How to perform a peak integration

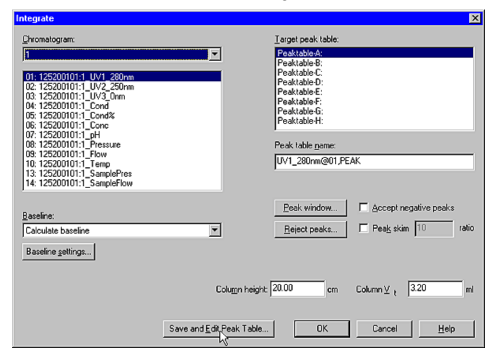
The table below describes how to perform a basic peak integration.

| Step | Action | |
|------|--|--|
| 1 | Open a result file in the Evaluation module. | |
| 2 | Choose Integrate:Peak Integrate. | |
| | or | |
| | Click the Peak Integrate toolbar icon. | |
| | Jul | |
| | Result: The Integrate dialog box opens. | |
| 3 | Select a source curve. | |
| | • Select a baseline or a calculation method from the Baseline list. | |
| | • Click OK to integrate with the default selections. or | |
| | | |
| | • Proceed with steps 4 to 6 to change the default selections. | |
| | Note: See also 12.1.3 How to optimize the baseline with a morpho logical algorithm on page 336 and 12.1.4 How to optimize the baseline with a classic algorithm on page 340. | |
| 4 | • Click the Baseline settings button to change the calculation algorithm in the Settings dialog box. The default algorithm is Morphological . | |
| | Change the selections or values. | |
| | Click OK | |
| 5 | Click the Peak window button to edit the peak window limits if necessary. | |
| | • Click the Reject peaks button to set the parameters for peak rejection if necessary. | |
| | • Edit the Column height or Column V values if necessary. | |

| Step | Action |
|------|--|
| 6 | Click OK to integrate and close the dialog box. |
| | or |
| | • Click Save and Edit Peak Table to save the integration and open the integrated curve for editing. |
| | - See 12.1.5 How to edit the baseline manually on page 348 |
| | - See 12.1.6 How to edit the peaks on page 351 |
| | - See 12.1.7 How to integrate part of a curve and how to exclude or skim peaks on page 358 |

Illustration

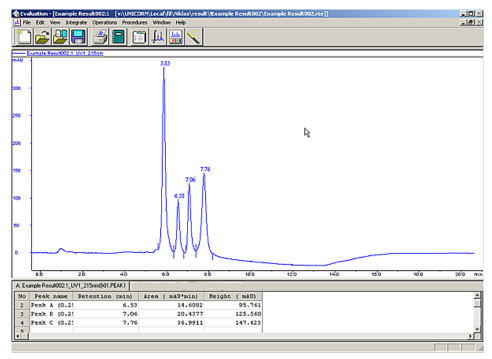
This is an illustration of the **Integrate** dialog box:



Peak integration results

The peak table is displayed underneath the active chromatogram. The start point and end point of each peak are marked by vertical marks, **drop-lines**, in the chromatogram. The peaks are automatically labelled according to what is selected in the **Curve Style and Color** tab of the **Chromatogram Layout** dialog box.

This is an illustration of the results after a peak integration:



Note: Peak tables can be copied from one chromatogram to another with the **Edit:Copy** command. However, to display the table you must right-click in the chromatogram, choose **Properties** and then select the new peak table on the **Peak Table** tab of the **Chromatogram Layout** dialog box.

How to display peak characteristics

The peak retention times and several other peak characteristics are calculated automatically. The table below describes how to display other peak characteristics.

| Step | Action |
|------|---|
| 1 | Right-click in the active chromatogram. Select Properties from the shortcut menu. Result: The Chromatogram Layout dialog box opens. |
| 2 | Click the Peak Table tab. |
| 3 | Select options from the Select peak table columns list. Click OK. Result: The selected items will be displayed in the peak table. |

How to filter peaks from view

Peaks can be removed from display in a peak table. The table below describes how to filter the peaks:

| Step | Action |
|------|---|
| 1 | Right-click in the active chromatogram or peak table. Select Properties from the shortcut menu. Result: The Chromatogram Layout dialog box opens. |
| 2 | Click the Peak Table tab. |
| 3 | Click the check boxes in the Filter Peaks field to select the filter criteria. Specify filter values. Click OK. |

To filter peaks vs. to reject peaks

The table below describes the major differences in the effect of filtering peaks compared to excluding the peaks by rejection.

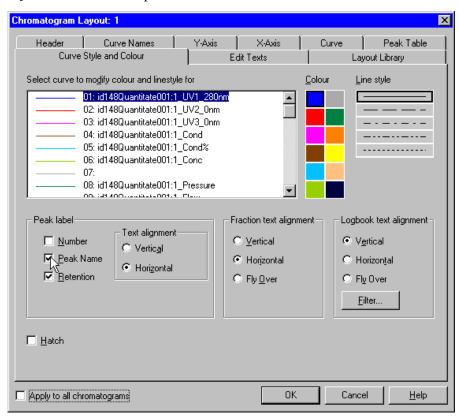
| Filter peaks | Reject peaks |
|---|---|
| excludes the peaks from display, | permanently excludes peaks from the integration, |
| does not exclude the peaks from the calculation of the total peak area, | excludes the peaks from the calculation of the total peak area, |
| can be reversed. | cannot be reversed. |

Peak labels

Peaks can be labelled with their retention, sequentially numbered, or be marked with specific identification names. See table below for an instruction on how to display peak labels.

The label type can be selected on the **Curve Style and Colour** tab in the **Chromatogram Layout** dialog box. De-select all label options to hide the labels, e.g. for presentations.

The illustration below shows the **Chromatogram Layout** dialog box with the **Curve Style and Colour** tab opened:



How to display peak labels

The table below describes how to display peak labels:

| Step | Action |
|------|---|
| 1 | Choose Edit:Chromatogram Layout. |
| | or |
| | Click the Chromatogram Layout icon. |
| | |
| | Result: The Chromatogram Layout dialog box opens. |
| 2 | Click the Curve Style and Colour tab. |

| Step | Action |
|------|---|
| 3 | Select one or more of the following labelling options in the Peak label field: |
| | Number |
| | Result: The peaks will be numbered sequentially. |
| | Peak Name |
| | Result: Peak names will be displayed. See 12.1.6 How to edit the peaks on page 351 for information about how to name the peaks. |
| | Retention |
| | <i>Result</i>: The retention volume or time will be displayed.Click 0K. |

12.1.3 How to optimize the baseline with a morphological algorithm

Introduction

The first choice when you want to optimize the peak integration is to change the baseline parameters. This section describes how to optimize the baseline with a morphological algorithm.

The Morphological algorithm

The **Morphological** algorithm can be described as a line that follows the chromatogram parallel to the X-axis. Data points for the baseline are created whenever the line touches the curve, and the points are joined at the end to create a baseline.

The **Morphological algorithm** gives the best result in curves with drifting baseline and peak clusters. The morphological baseline follows the curve faithfully, and a curve with a baseline at a more even level can be created by subtracting the morphological baseline.

The **Morphological algorithm** does *not* work well if there are negative peaks or if quantitative data from negative peaks are important in the run.

Note: The Morphological algorithm is the default baseline setting.

How to set a Morphological baseline

The table below describes how to choose a **Morphological algorithm** and define baseline settings.

| Step | Action |
|------|---|
| 1 | Select Integrate:Peak Integrate. Result: The Integrate dialog box opens. |
| 2 | Click the Baseline settings button in the Integrate dialog box. Result: The Settings dialog box opens. |
| 3 | Select the Morphological algorithm. Change the Baseline parameters if necessary. See more information about the parameters below this table. Click OK. |

Note: The same settings can be edited in the **Calculate Baseline** dialog box when a new baseline is created. Choose **Integrate:Calculate Baseline** to open the dialog box.

Morphological algorithm parameters

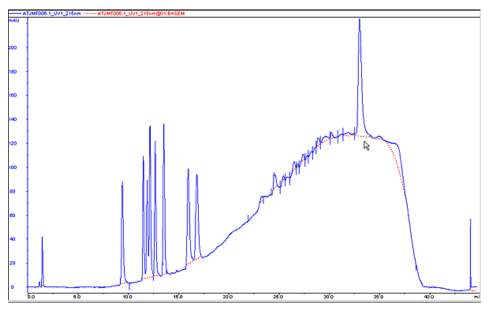
The parameters for the Morphological algorithm are:

- Structure width
- Noise window
- Minimum distance between points

Structure width

Structure width determines the length of the straight line that follows the chromatogram. The default value is set at the widest peak in the chromatogram multiplied by 1.5.

The illustration below is an example of how a morphological baseline follows the peaks at the different levels in the curve:



The correct structure width settings

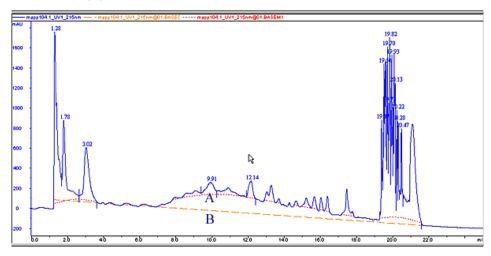
Too low settings

Too low **Structure width** settings can result in a baseline that reaches too high up in the peaks of the curve. Sometime a wider peak is not recognized because it contains a cluster of smaller peaks. The **Structure width** is then set to a value according to the largest width of the identified narrower peaks, and must be increased.

Too high settings

Too high **Structure width** settings mean that narrower peaks, especially in fluctuating curves, are not properly followed. This happens when an artifact in a curve is identified as the widest peak by the morphological algorithm, and then is used to set the default **Structure width** value.

The illustration below is an example of baselines using the default morphological algorithm settings (A) and a morphological algorithm with an increased **Structure** width value (B).



Noise window

Sometimes you get too many peaks after the peak integration, usually because noise on the baseline is erroneously detected as peaks.

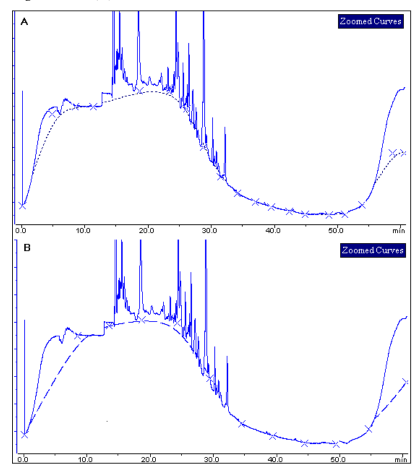
The solution to this is to increase the **Noise window** parameter. However, this can result in peak limits too high up on the peak slopes.

Note: You can also use the **Reject peaks** function in the **Integrate** dialog box to reduce the number of peaks based on the total number of accepted peaks or the minimum peak height.

Minimum distance between points

The **Minimum distance between points** is a measure of the distance between the data points used to generate a baseline. The largest number of data points is produced at the slopes of the curves. If you increase the **Minimum distance between points** value, fewer points will be collected on the slopes.

The illustration below is an example of a baseline (A) that is created with the **Minimum distance between points** parameter set at a low value. The number of data points is reduced when the **Minimum distance between points** parameter is set to a higher value (B).



12.1.4 How to optimize the baseline with a classic algorithm

Introduction

The first choice when you want to optimize the peak integration is to change the baseline parameters. This section describes how to optimize the baseline with a classical algorithm.

What is the Classic algorithm?

The **Classic algorithm** searches for all parts of the source curve that are longer than a defined minimum baseline segment and fall within limiting parameters. Together, the parameter values define the limits for a rectangular box. A part of the source curve must fit entirely inside this rectangular box to be identified as a baseline segment.

The **Classic algorithm** is particularly useful when you need to integrate curves with negative peaks and when quantitative data from negative peaks are important.

Classic algorithm parameters

The parameters for the Classic algorithm are:

- Shortest baseline segment
- Noise window
- Max baseline level
- Slope limit

See more information about the parameters below.

How to set a Classic baseline

The table below describes how to set a **Classic algorithm** and define a baseline.

| Step | Action |
|------|--|
| 1 | Click the Baseline settings button in the Integrate dialog box. |
| | Result: The Settings dialog box opens. |
| 2 | Select the Classic algorithm. |
| | Change the Baseline parameters. |
| | See more information about the parameters below this table. • Click OK . |

Note: The same settings can be edited in the **Calculate Baseline** dialog box when a new baseline is created. Choose **Integrate:Calculate Baseline** to open the dialog box.

Test your parameter changes

The best way to optimize the baseline is to change the baseline parameters step by step and then check the resulting baseline after each change. When the desired effect is accomplished it is best to go back and try a parameter value in between the two last settings to avoid an unnecessarily low or high value.

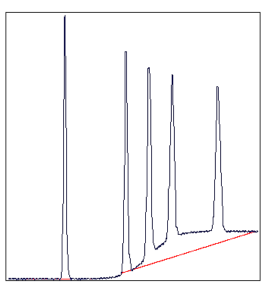
How much the values should be changed depends on the cause of the peak integration problem. The table below is a general guideline.

| Baseline parameter | Recommended initial change |
|---------------------------|---------------------------------|
| Shortest baseline segment | 20-50% |
| Noise window | 10-30% |
| Max baseline level | Usually not necessary to adjust |
| Slope limit | 25-50% |

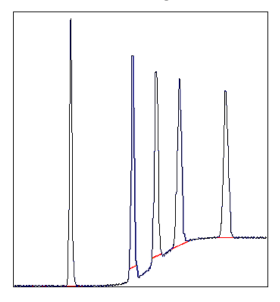
Note: If necessary, click the **Default** button to restore the default values.

Shortest baseline segment

If a too high **Shortest baseline segment** value is set, short curve segments between peaks in the middle of the chromatogram are not identified as baseline segments. The calculated baseline does not follow the source curve, see below:



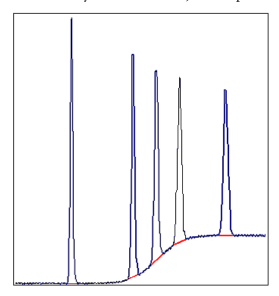
The **Shortest baseline segment** value is decreased by 50% in this example:



Slope limit

A changed **Slope limit** will often improve the baseline calculation. The **Slope limit** sets the maximum slope of the curve to define when a peak is recognized. A too high **Slope limit** will cause the up-slopes of the peaks to be recognized as baseline segments.

The example above was improved by the shorter baseline segments but the high slope of the short segments in the region between the second and the fourth peak still makes the baseline unacceptable. In the example below the **Slope limit** is increased by a factor of 2.5, which produces a correct baseline:

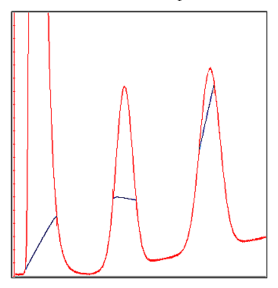


Too high slope limit

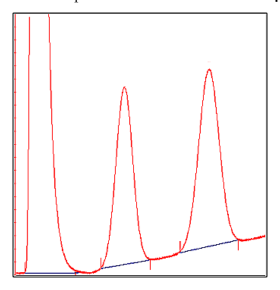
A too high **Slope limit** value can cause peak limits too high up on the peaks. This can be the case when the chromatogram includes a very large flow-through or solvent peak. The large peak affects the calculation of the default parameters and leads to too high values for the **Slope limit**.

Note: A too high value for the **Noise window** can have the same effect and be caused by the same situation, often also in combination with a high **Slope limit**.

Peak limits are defined on peaks in the example below due to the high Slope limit:



The example below has a much lower Slope limit, and a lower Noise window:

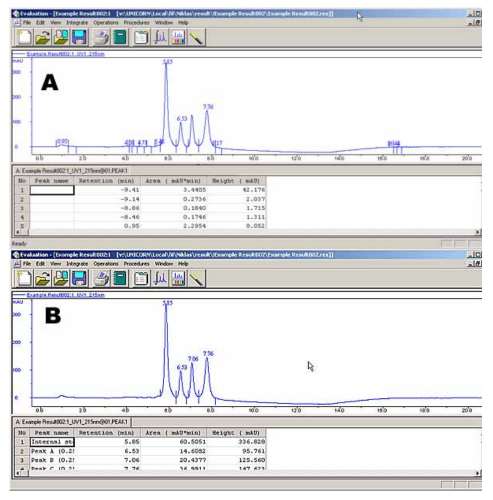


Noise window

Sometimes you get too many peaks after the peak integration, usually because noise on the baseline is erroneously detected as peaks.

The solution to this is to increase the **Noise window** parameter. However, this can result in peak limits too high up on the peak slopes.

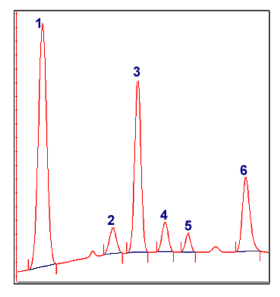
The illustration below is an example of noise detected as peaks (A) and the result of a second peak integration with an increased **Noise window** (B).



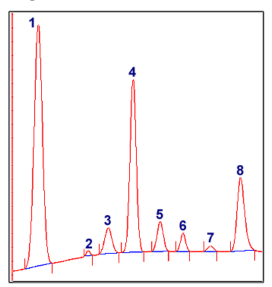
Note: You can also use the **Reject peaks** function in the **Integrate** dialog box to reduce the number of peaks based on the total number of accepted peaks or the minimum peak height.

Missing peaks

Sometimes obvious peaks are not detected in the peak integration. The probable cause is that the **Noise window** is set too high. See the illustration below:



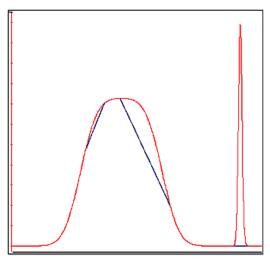
All peaks are detected if the **Noise window** is decreased, see example below:



Note: Missing peaks can also be caused by improper settings for **Reject peaks** in the **Integrate** dialog box, or **Filter peaks** in the **Chromatogram layout** dialog box.

When to change the Max baseline level

In rare cases the top of a broad, flat peak can be incorporated as a baseline segment. This is one of the very few situations where it is useful to change the **Max baseline level**. The illustration below is an example:



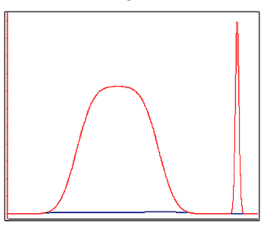
How to set the Max baseline level

The table below describes how to set the **Max baseline level**.

| Step | Action |
|------|--|
| 1 | Right-click in the chromatogram and select Marker. |
| | Result: A vertical line is set in the chromatogram. A text box in the top left corner of the chromatogram displays the X-axis and Y-axis values of the curve at the point where the vertical Marker line crosses the curve. |
| 2 | Move the Marker with your mouse. |
| | Measure the height of the peak you want to exclude from the baseline. |
| 3 | Choose Integrate:Calculate baseline. |
| 4 | Select the Classic checkbox as the Chosen algorithm. |
| | • Type a new value for Max baseline level . Set the level slightly lower than the value that you measured in step 2. |
| | Click 0K . |

Example of a correct baseline

The illustration below is an example of a correct baseline after the Max baseline level has been changed:



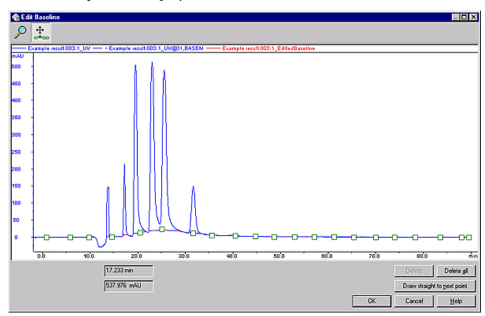
12.1.5 How to edit the baseline manually

The Edit Baseline dialog box

You can edit the baseline manually in the **Edit Baseline** dialog box in the **Evaluation** module:

• Select Integrate:Edit Baseline to display the dialog box.

The **Edit Baseline** dialog box displays the baseline and the curve it was calculated from. The baseline points are marked with green squares. Hold the cursor above the baseline point to display its coordinates. See the illustration below:



How to use the zoom function

The table below describes how to use the zoom function in the **Edit Baseline** dialog box.

| Step | Action |
|------|--|
| 1 | Click the Zoom icon. |
| | Result: The cursor is changed into a magnifying glass. |
| 2 | Press and hold the left mouse button. |
| | Drag the cursor over the area you want to zoom in on. |
| | Release the mouse button. |
| | Result: The area is enlarged. Right-click and select Reset zoom to restore the full view. |

How to edit and insert data points

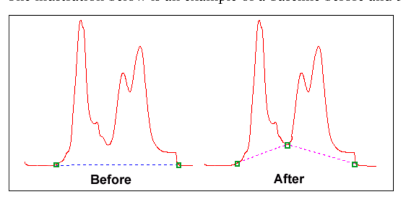
The table below describes how to edit and insert baseline data points:

| Step | Action |
|------|---|
| 1 | Select Integrate:Edit Baseline. |
| | Result: If there are more than one baseline available, the Select Baseline to Edit dialog box opens. If not, proceed to step 2. |
| | Select the baseline you want to edit from the list. |
| | • Click OK . |
| | Result: The Edit Baseline dialog box opens |
| 2 | Click the Set Curve Points icon. |
| | 0 · · · · · · · · · · · · · · · · · · · |
| | Result: The cursor is changed into a cross. |
| 3 | Add a data point |
| | Click the left mouse button to place a new baseline point in the chromatogram. |
| | Result: A new point is created, marked by a green square. The baseline curve is redrawn as a spline function based on the old and the new points. The baseline is guided by the points, but does not necessarily pass through them. |
| 4 | Delete a data point |
| | Double-click the data point. |
| | or |
| | Click the data point to select it and click the Delete button. |
| | or |
| | Right-click the data point and select Delete Point from the shortcut menu. |
| | Result: The data point is deleted and the curve is redrawn. |
| 5 | Move a data point |
| | Select the data point and drag it to a new position. |
| | Result: The baseline curve is redrawn. |
| 6 | Click OK . |
| | Result: The Save Edited Baseline dialog box opens. |

| Step | Action |
|------|--|
| 7 | Confirm the location and type a new name if necessary. Click 0K. |
| | Result: The new baseline is saved. |

Edited baseline

The illustration below is an example of a baseline before and after editing:



How to draw a straight line

The table below describes how to force a straight baseline between two points.

| Step | Action | |
|------|--|--|
| 1 | Select the first of the two points in the point list. | |
| 2 | Click the Draw straight to next point button. | |
| | Result: The baseline is drawn through the points as a straight line. | |

12.1.6 How to edit the peaks

Introduction

Once a peak table has been generated based on an appropriate baseline, it is possible to split or join peaks and to manually adjust the peak start and end points. The peaks will then be renumbered and the peak values will all be recalculated.

How to open the peak table for editing

The table below describes how open the peak table for editing. The editing options are described below this table:

| Step | Action |
|------|---|
| 1 | Select Integrate:Edit Peak Table. |
| | Result: If there are more than one peak table available, the Select Peak Table to Edit dialog box opens. The name of the baseline on which the peak table was based is displayed at the bottom of the panel. |
| 2 | Select the peak table from the list and click OK . |
| | Select one or more Help Curves to be displayed for reference if necessary. |
| | Result: The Edit Peak Table dialog box opens. |
| | <i>Note</i> : The Edit Peak Table dialog box will be opened immediately if you select Save and Edit Peak Table as the last step of the peak integration. |
| 3 | Perform the changes (described in the instructions below). |
| 4 | Click OK . |
| | Result: The Save Edited Peak Table dialog box opens. The dialog box displays a suggested name and location for the peak table. |
| 5 | Confirm the name and location and click 0K . |

How to adjust the baseline

The baseline can be adjusted graphically (see also 12.1.5 How to edit the baseline manually on page 348) in the **Edit Peak Table** dialog box. The table below describes this:

| Step | Action |
|------|---|
| 1 | Click the Set Curve Points icon. |
| | Result: The cursor is changed into a cross. |

| Step | Action |
|------|---|
| 2 | Perform the operations below as desired: |
| | Click to insert a new data point. |
| | Double-click on a data point or right-click the point and select Delete Point from the short-cut menu to delete the point. |
| | • Click a data point and drag the point to a new position to move the baseline. |
| | <i>Note</i> : Accept negative peaks must be selected before the peak integration if you want to be able to drag a data point to move the baseline above the curve. |

How to calculate a new baseline

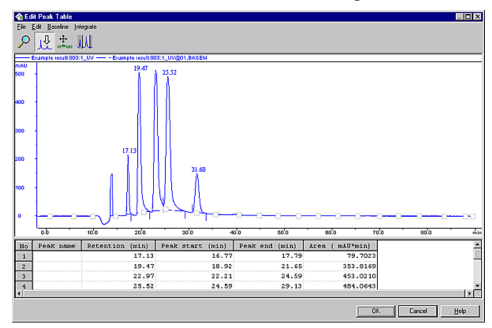
The baseline can be recalculated in the **Edit Peak Table** dialog box. The table below describes how to do this:

| Step | Action |
|------|---|
| 1 | Select Baseline:New:Calculate. |
| | or |
| | Right-click and select New Calculate from the shortcut menu. |
| | Result: The Settings dialog box opens. |
| 2 | Select an algorithm (Morphological is default). |
| 3 | Adjust the Baseline parameters as desired. |
| | or |
| | Click the Default Values button for the default values. |
| 4 | Click 0K . |
| | Result: The baseline is recalculated. |

Note: Select **Baseline:New:Zero Baseline** to replace the calculated baseline with a zero baseline.

The Edit Peak Table dialog box

The illustration below shows the **Edit Peak Table** dialog box.



How to delete a peak

The table below describes how to delete a peak in the **Edit Peak Table** dialog box:

| Step | Action |
|------|--|
| 1 | Click the Edit peaks icon. |
| | T. C. |
| | Click the peak in the curve or in the peak table to select the peak. |
| 2 | Right-click and select Delete Peaks from the shortcut menu. |
| | or |
| | Select Edit:Delete Peaks. |
| | Result: The peak is deleted and the remaining peaks are renumbered. |

to a peak

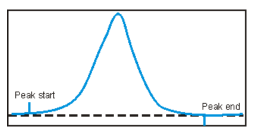
How to add color The table below describes how to add a fill color and a pattern to a peak in the **Edit Peak Table** dialog box:

| Step | Action |
|------|--|
| 1 | Click the Edit peaks icon. |
| | <u>\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\</u> |
| | Move the cursor over the peak you want to edit. |
| | Result: The cursor is changed into a larger arrow. |
| | Click to select the peak. |
| 2 | Right-click and select Fill Peak from the shortcut menu. |
| | or |
| | Select Edit:Fill Peak. |
| | Result: The Color and Pattern dialog box opens. |
| | Color Color Pattern OK Cancel |
| | • Select a color and a pattern. |
| | • Click OK . |
| | Result: The peak is filled according to the selections. |

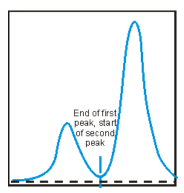
Note: The color and pattern selections will override the general Fill settings that can be selected for all peaks on the Peak Table tab in the Chromatogram Layout dialog box.

Peak start and end points

The beginning of each peak is marked with a drop-line above the curve, and the end of each peak is marked with a drop-line below the curve. The illustration below shows an example of start and end point drop-lines:



Where there are two peaks beside one another, the end of the first peak will be at the same point as the beginning of the next peak. Thus, there will be a drop-line below and above the curve at the same point. See the illustration below:



How to split a peak

It is possible to split the peak into two new peaks by inserting a drop-line. The table below describes how to split a peak in the **Edit Peak Table** dialog box:

| Step | Action |
|------|--|
| 1 | Click the Edit peaks icon. |
| | T. C. |
| | Click the peak in the curve or in the peak table to select the peak. |
| 2 | Right-click and select Split Peak from the shortcut menu. |
| | or |
| | Select Edit:Split Peaks. |
| | Result: A new drop-line is inserted at the middle point between the two existing drop-lines and the peak is split. |

Note: The area under each new peak will not be the same if the symmetry of the original peak was not perfect.

How to join peaks It is possible to join the areas of adjacent peaks if they are separated by a drop-line. The table below describes how to join adjacent peaks in the Edit Peak Table dialog box:

| Step | Action |
|------|---|
| 1 | Click the Edit peaks icon. |
| | T. C. |
| | Click the peak in the curve or in the peak table to select the peak. |
| 2 | Right-click and select Join Left or Join Right from the shortcut menu. |
| | or |
| | Select Edit:Join Left or Edit:Join Right. |
| | <i>Result</i> : The original intervening drop-line is removed and all peaks are renumbered. |

How to add peak names

The table below describes how to add names in the Edit Peak Table dialog box to identify the peaks:

| Step | Action |
|------|---|
| 1 | Click the Edit peaks icon. |
| | T. C. |
| | Click the peak in the curve or in the peak table to select the peak. |
| 2 | Right-click and select Peak Name from the shortcut menu. |
| | or |
| | Choose Edit:Peak name. |
| | or |
| | Double-click the peak in the peak table or the curve. |
| | Result: The Edit Peak Name dialog box opens. The number and retention of the selected peak is displayed. |
| 3 | Type a name in the Peak name textbox and click OK . |

How to adjust peak areas with drop-lines

The table below describes how to move the drop-lines to adjust the peak area in the **Edit Peak Table** dialog box.

| Step | Action |
|------|---|
| 1 | Click the Edit peaks icon. |
| | Click the peak in the curve or in the peak table to select the peak. |
| | Result: Two vertical bars become superimposed over the drop-lines that delimit the selected peak. The area between the bars is filled with a yellow fill pattern. |
| 2 | Drag the bars to define the new limits for the selected peak. *Result: The drop-lines are moved and the peak areas are automatically recalculated. |

Note: A drop-line can never be moved beyond another drop-line or beyond a point where the peak meets the baseline.

How to use the zoom function

The table below describes how to use the zoom function in the **Edit Peak Table** dialog box.

| Step | Action |
|------|--|
| 1 | Click the Zoom icon. |
| | |
| | Result: The cursor is changed into a magnifying glass. |
| 2 | Press and hold the left mouse button. |
| | Drag the cursor over the area you want to zoom in on. |
| | Release the mouse button. |
| | Result: The area is enlarged. Right-click and select Reset zoom to restore the full view. |

The Integrate menu

If needed you can use the selections on the **Integrate** menu to perform a peak integration in the **Edit Peak Table** dialog box. This is useful for example if you want to re-integrate the curve using different settings or integrate only part of a curve with different settings.

See 12.1.7 How to integrate part of a curve and how to exclude or skim peaks on page 358 for more information.

12.1.7 How to integrate part of a curve and how to exclude or skim peaks

Introduction

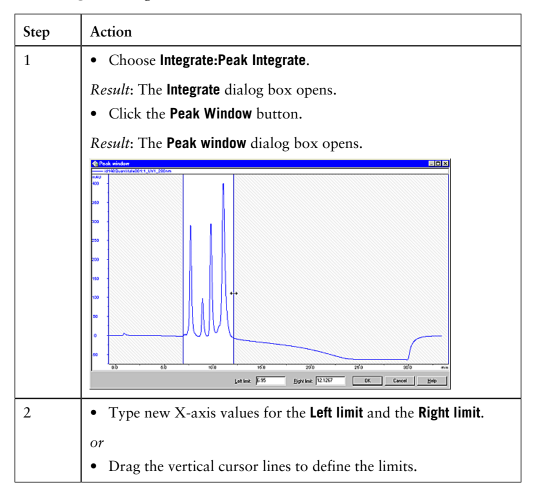
There are several possibilities to improve the results if the peak integration is unsatisfactory. This section describes:

- How to select only part of a curve for integration.
- How to exclude peaks.
- How to skim peaks.

These operations can be performed both in the **Integrate** dialog box in preparation for the peak integration, or in the **Edit Peak Table** dialog box to adjust an unsatisfactory peak integration. This section describes both alternatives.

How to select part of a curve

The table below describes how to select only a part of a curve for peak integration in the **Integrate** dialog box:



| Step | Action |
|------|--|
| 3 | Click OK . |
| | <i>Result</i> : The baseline will be calculated from the whole curve, but the calculation of the peak areas is only performed on the selected section. |

How to exclude peaks

You can define criteria to exclude peaks from integration. The table below describes how to define peaks to be excluded in the **Integrate** dialog box.

| Step | Action |
|------|---|
| 1 | Click the Reject peaks button. |
| | Result: The Reject Peaks dialog box opens. |
| 2 | Select the appropriate checkboxes and type values for height, width and area. |
| | Define how many of the largest peaks you want to include. |
| | • Click OK . |

How to include negative peaks

Select the **Accept negative peaks** checkbox of the **Integrate** dialog box to include negative peaks in the integration.

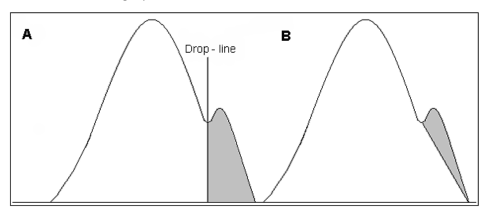
Result: The negative peaks will be reported as negative areas in the peak table. By default, negative peaks are not included in the integration.

Peak skimming vs. drop-lines

The area under a peak can be calculated either using separating drop-lines or peak skimming:

- **Drop-lines** are vertical marks that split two peaks at the valley. Drop-lines are used mostly for peaks of relatively similar size. When a peak has a shoulder, splitting with drop-lines will cause the first peak to lose too much of its area to the peak that forms its shoulder.
- The **Peak skim** option can be used to skim off the smaller peak with a straight line that starts in the valley between the peaks and ends at the other side of the smaller peak, at the point where the skim line and the curve slope are equal.

The illustration below is an example of how a drop-line (A) and a skimmed peak (B) affects the area under the main peak and the peak shoulder. The peak shoulder area is marked in gray:



How to skim peaks

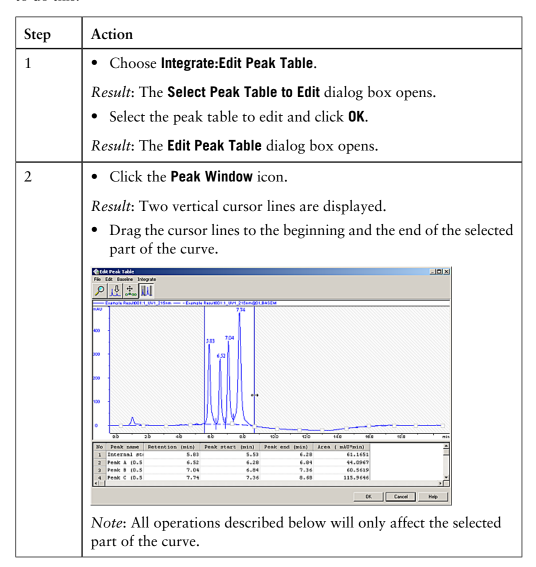
The table below describes how to select a ratio to skim peaks in the **Integrate** dialog box:

| Step | Action |
|------|--|
| 1 | Select the Peak skim checkbox. |
| 2 | Determine the ratio when peak skimming should be applied based on the relationship in the illustration below: |
| | $\frac{h_{p1} - h_{v}}{h_{p2} - h_{v}} > \text{skim ratio}$ $h_{p1} h_{v} h_{p2}$ $Note: \text{The default ratio value is } 10.$ |

| Step | Action |
|------|---------------------------------------|
| 3 | Type the ratio value in the text box. |

How to integrate part of a curve

Part of a curve can be selected in the **Edit Peak Table** dialog box and integrated with settings that differ from the rest of the curve. The table below describes how to do this.



| Step | Action |
|------|--|
| 3 | If desired, change the integration parameters: |
| | Reject peaks |
| | Choose Integrate:Settings. |
| | Result: The Reject Peaks dialog box opens. |
| | Change the settings as desired and click 0K . |
| | Skim peaks |
| | Choose Integrate:Peak Skim. |
| | Result: The Peak Skim dialog box opens. |
| | Select the Skim Peaks checkbox and type a ratio. |
| | • Click OK . |
| 4 | Choose Integrate:Peak Integrate. |
| | <i>Result</i> : The selected part of the curve is peak integrated based on the changed parameters. |

12.1.8 Measurements

Introduction

It is possible to determine the coordinates of any point on a curve and to obtain values for retention and peak height. This is a useful tool for many other functions, such as for measuring the parameters used in baseline calculations.

Measurement options

Coordinates can be obtained in two ways:

- Through direct measurement.
- From peak table data.

How to make direct measurements

The table below describes how to make direct measurements in a chromatogram:

| Step | Action |
|------|---|
| 1 | Right-click in the chromatogram and select Marker. Result: A vertical line is set in the chromatogram. A text box in the top left corner of the chromatogram displays the X-axis and Y-axis values of the curve at the point where the vertical Marker line crosses the curve. See the illustration below: |
| 2 | Move the Marker with your mouse to display the peak data. |
| 3 | Click the curve name legend above the chromatogram to change to another curve. Result: The Y-axis is changed to the one corresponding to the new curve. |
| 4 | Right-click and select Marker again to de-select the function. |

ence point

How to set a refer- The table describes how to set a reference point:

| Step | Action |
|------|--|
| 1 | Right-click in the chromatogram and select Set Marker Ref. Point to define a reference point for the marker position. |
| 2 | When the marker is moved from the reference point, the X-axis and Y-axis values for the new position are displayed together with: |
| | the new position in relation to the position of the reference point, the minimum, maximum and average values for the curve interval |
| | between the reference point and the new position. |

How to record a Snapshot

The table below describes how to record a **Snapshot** of the current curve values:

| Step | Action |
|------|---|
| 1 | Right-click in the chromatogram and select Snapshot from the shortcut menu. |
| | Result: The Snapshot dialog box opens. |
| 2 | The dialog box displays all the curve data that was current at the moment the snapshot was taken. |
| | Click the Save to file button to save the snapshot as an Excel file. |
| | Click the Print button to print the snapshot. |

How to select peak table data

The retention time and amplitude of any peak can be viewed directly in a peak table after an integration. This data and more is selected in the Chromatogram Layout dialog box. The table below describes how to select peak table data.

| Step | Action |
|------|---|
| 1 | Click the Chromatogram Layout icon. |
| | |
| | Result: The Chromatogram Layout dialog box opens. |
| 2 | Click the Peak Table tab. |
| 3 | Select the checkboxes on the Select peak table columns list for all items that you want to display in the table. Click OK. |

Other evaluations 12.2

Introduction

This section describes how the results can be used for other types of evaluations.

In this section

This section contains these sub-sections.

| Topic | See |
|-------------------------------------|--------|
| Peak purity and peak identification | 12.2.1 |
| How to find slope values | 12.2.2 |
| How to simulate peak fractionation | 12.2.3 |
| How to create curves | 12.2.4 |
| How to use the Fraction Histogram | |

12.2.1 Peak purity and peak identification

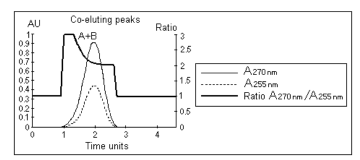
Introduction

Ratios between UV curves measured at different wavelengths give useful information about peak purity or peak identity.

Peak purity

The absorbance ratio can be used to check peak purity. If the peak is pure, the absorbance spectra are the same over the whole peak and the ratios should therefore remain constant. The peak is probably not pure if the absorbance ratio is not the same over the whole peak.

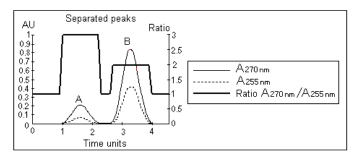
The illustration below shows a simulated chromatogram of two co-eluting components with differing absorbance spectra and a small difference in retention time:



Peak identification

The absorbance ratio can be used for peak identification. Different compounds have a specific ratio between absorbancies at different wavelengths.

The illustration below shows a simulated chromatogram of two components with differences in their absorbance spectra:



How to divide the curves

Both curves must have a baseline close to zero AU before they can be divided. This is achieved with baseline subtraction. The table below describes how to subtract the baseline from an earlier integration and divide the curves:

| Step | Action |
|------|--------------------------------------|
| 1 | Create a baseline for each UV curve. |

| Step | Action |
|------|--|
| 2 | Select Operations:Subtract. |
| | Result: The Subtract dialog box opens. |
| 3 | Select the UV curve in the first list of curves. |
| | Select its baseline in the second list of curves. |
| | • Click 0K . |
| | <i>Note</i> : You can also subtract corresponding blank runs if there are blank runs available. |
| 4 | Repeat steps 2 and 3 for the second UV curve. |
| 5 | Select Operations:Divide. |
| | Result: The Divide dialog box opens. |
| 6 | • Select the first result curve from the subtractions in the first list of curves. |
| | • Select the second result curve from the subtractions in the second list of curves. |
| 7 | Click the checkbox for Threshold and type values for each curve. This results in the following: |
| | • The quotient is set to 1.0 if either of the sample values is closer to zero than the threshold value. Very high quotient values are prevented if division is performed with values close to zero. Very low quotient values are also prevented. |
| | <i>Note</i> : Default Threshold values are entered by UNICORN. The values can be changed. |
| 8 | Click OK. |

How to filter the result curve

The resulting curve can be filtered to reduce noise and to remove ghost peaks. The table below describes how to filter the curve.

| Step | Action |
|------|---|
| 1 | Select Operations:Smooth. |
| | Result: The Smooth dialog box opens. |

| Step | Action |
|------|---|
| 2 | Select the Source Chromatogram. Select a Filter Type. |
| | Note: The Median filter is recommended to remove noise that appears as spikes or occurs in a small area of the curve.Click OK. |

12.2.2 How to find slope values

Introduction

With ÄKTAdesign systems it is possible to only collect peaks during fractionation. The way to find suitable slope values for a particular run is described in this section.

Where to use slope values

The slope values can be used in the **Method Editor**

- as StartSlope and EndSlope values in the Peak_FracParameters instruction.
- as parameters for the **Watch** instruction.

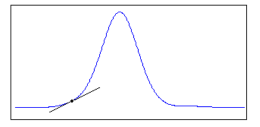
Using slope values for Watch instructions

Conditional **Watch** instructions can be set up to let the progress of a run be determined by the events during the run, e.g. start to collect fractions when the first peak emerges.

The slope of the curve can be set as a condition to satisfy a **Watch** condition in the method during the run. It is important to use accurate slope values for the specific **Watch** instruction parameter.

A sample run

You must first make a separation run with the sample you intend to purify. The result from this separation run is then used to find the slope values.



Retention scale

Time should be used as the X-axis scale for retention.

| Step | Action |
|------|-------------------------------------|
| 1 | Click the Chromatogram Layout icon. |
| 2 | • Click the X-axis tab. |
| | Select Time. |
| | • Click OK . |

How to differentiate the curve

The slope values are measured on a differentiated curve. The table below describes how to create a differentiated curve.

| Step | Action |
|------|---|
| 1 | Select Operations:Differentiate. |
| | Result: The Differentiate dialog box opens. |
| 2 | Select the UV curve you want in the Source chromatogram list. |
| | Click the First order radio button. |
| | • Click OK . |
| | Result: The differentiated curve opens in the chromatogram. |

How to measure the slope values

Sometimes the differentiated curve must be filtered to reduce noise and ghost peaks before the measurements. See section 12.2.1 Peak purity and peak identification on page 366.

The table below describes how to measure the slope values on the differentiated curve.

| Step | Action |
|------|---|
| 1 | Click the name of the differentiated curve (above the chromatogram window) to select the curve. |
| 2 | Use the zoom function to magnify the curve over an appropriate area. |
| 3 | Right-click and select Marker from the short-cut menu. Result: A vertical cursor bar opens in the chromatogram. |
| 4 | Place the Marker at the beginning of a peak where you want the Watch conditions to be fulfilled, i.e. where the slope becomes higher. |
| 5 | Read the actual slope value in the active Marker text box in the top left corner of the chromatogram window. |

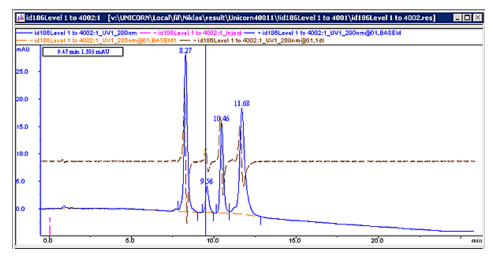
Note: The unit for the differentiated curve is mAU/min or AU/min. Any Y-axis value for the differentiated curve is the UV curve slope at the selected retention point.

Peak fractionation for ÄKTAdesign

If your system is an ÄKTAdesign system, measure the slope at the beginning and the end of the smallest, flattest peak of all the peaks of interest, and use these values.

Illustration: Slope value measurement

The illustration below shows a measurement of the slope limit after differentiation:



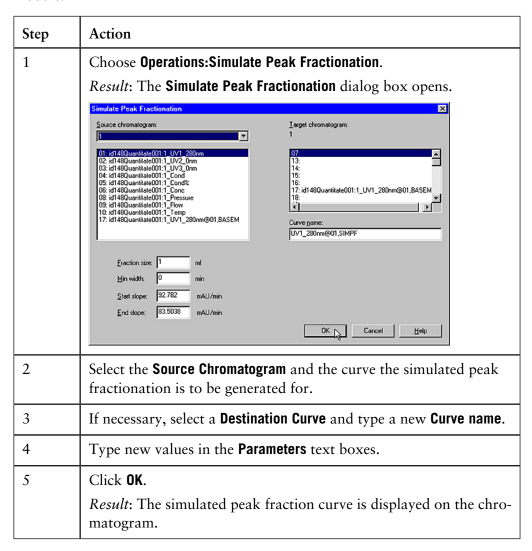
12.2.3 How to simulate a peak fractionation

Introduction

You can create a curve that simulates a peak fractionation to test the outcome before the actual peak fractionation is run. This section describes how this is done.

How to simulate a peak fractiona-

The table below describes how to simulate a peak fractionation in the **Evaluation** module.



12.2.4 How to create curves

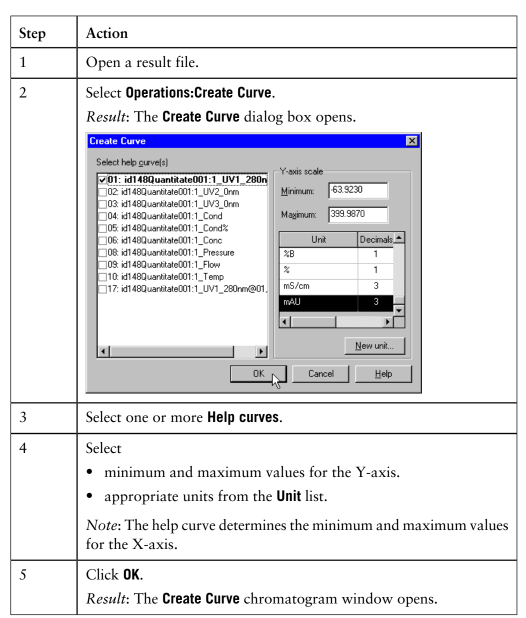
Introduction

You can draw a curve of your own in the **Evaluation** module. This section describes how this is done.

Note: The right to create and rename curves is defined in the user access rights and may be restricted.

How to create curves - step 1

The table below describes how to set up a chromatogram window to create a curve in the **Evaluation** module.



How to create new units

In the **Create Curve** dialog box you can also create new units for the curve. The table below describes how this is done.

| Step | Action |
|------|---|
| 1 | Click the New unit button. |
| | Result: The Create New Unit dialog box opens. |
| 2 | Type a new unit name and a number of decimal places. |
| 3 | Click OK . |
| | Result: The Create New Unit dialog box is closed. The new unit is now available in the Create Curve dialog box. |

How to create curves - step 2

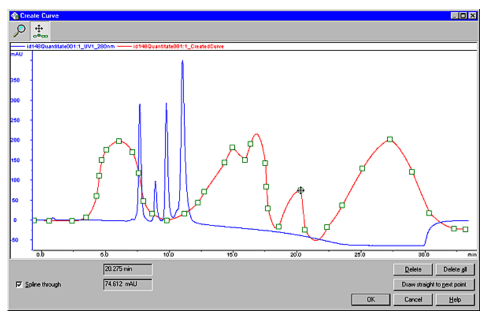
The new curve is created in the **Create Curve** window. The table below describes how to work in this window.

| Step | Action |
|------|---|
| 1 | Click the Set Curve Points icon. |
| | ⊕ □□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□ |
| 2 | Click to insert curve points in the chromatogram. |
| | Add more points to draw the curve. |
| | Result: A green square marks the new curve point. The curve is drawn from the previous point. Hold the cursor over the inserted point to see the coordinates displayed. |
| | Curve mode |
| | • The regular spline mode draws the curve as a smooth line near but not through every point. |
| | • Click the Spline through checkbox to draw the curve through all of the curve points. |
| 3 | Move a point |
| | Select the point and drag it to the new position. |
| | Result: The curve is redrawn. |

| Step | Action |
|------|---|
| 4 | Delete a curve point • Double-click the curve point. |
| | or |
| | • Select the point and click the Delete button. |
| | Select the point, right-click and choose Delete Point from the shortcut menu. |
| 5 | Click the Zoom icon to focus on details in the curve. |
| | Note: Right-click and select Reset zoom to return to the full view. |
| | Right-click in the chromatogram window and select Marker. |
| | • Position the Marker bar over peaks in the help curve to measure the coordinates. |
| | Result: The coordinates are displayed in the Marker text box in the top left corner of the chromatogram. |
| | <i>Note</i> : Click the Marker text box to display the coordinates for the created curve. Click again to return to the help curve coordinates. |
| 6 | Click OK . |
| | Result: The Save Curve dialog box opens. |
| 7 | Type a new name if desired and click OK . |

Curve example

The illustration below is an example of a curve created by using the **Draw Spline** command in the **Create Curve** chromatogram window.



How to force the curve through points

In cases where you have created a curve and not selected the **Spline through** option, you may want the curve to pass through some of the points that are outside the created curve. The table below describes how to force the curve through these points:

| Step | Action |
|------|--|
| 1 | Select the curve point immediately before the curve point you want to connect to. |
| 2 | Click the Draw straight to next point button. |
| | <i>Result</i> : The curve is adjusted so that it is drawn as a straight line between the two points. |

12.2.5 **How to use the Fraction Histogram**

Introduction

The Fraction Histogram dialog box in the Evaluation module can be used to create a curve for the average fraction absorbance.

How to create a Fraction Histogram

The table below describes how to create a **Fraction Histogram** curve.

| Step | Action |
|------|--|
| 1 | Select Operations:Fraction histogram. |
| | Result: The Fraction histogram dialog box opens. |
| 2 | Select the desired UV curve. |
| | <i>Note</i> : The fractions curve should already be selected on the middle list. If you have previous pooled fractions and created a pooled fraction curve, select the desired fraction curve. |
| 3 | Click 0K . |
| | <i>Result</i> : The average fraction absorbance values are displayed as a new curve in the chromatogram. |

12.3 Automated evaluation procedures

Introduction

An evaluation procedure is a recorded sequence of interactive operations in the **Evaluation** module, which can be executed for automated data evaluation and report generation. The concept is similar to the "macro" facilities in other programs. This section describes how to work with automated evaluation procedures.

In this section

This section contains these sub-sections.

| Topic | See |
|-------------------------------------|--------|
| How to create a new procedure | 12.3.1 |
| How to edit a procedure | 12.3.2 |
| How to run a procedure | 12.3.3 |
| How to rename and remove procedures | 12.3.4 |

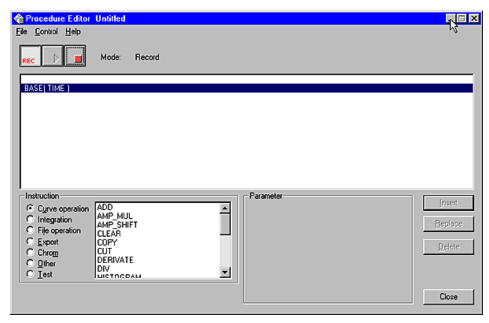
12.3.1 How to create a new procedure

Introduction

You can use the **Procedure Editor** to record or create a new procedure. The **Procedure Editor** can also be used to view and edit the instructions within a procedure. This section describes how to use the **Procedure Editor** to record new procedures.

The Procedure Editor dialog box

The illustration below shows the **Procedure Editor** in **Record mode**.



How to record a procedure

The table below describes how to record a new procedure.

| Step | Action |
|------|--|
| 1 | Open the result file in the Evaluation module. |
| 2 | Choose Procedures:Record On. |
| | Result: The Procedure Editor dialog box opens in record mode. |
| 3 | Minimize the Procedure Editor dialog box. |
| 4 | Perform the evaluation steps that the procedure is to contain. |
| | Result: The steps are recorded in the order that they are performed. |

| Step | Action |
|------|---|
| 5 | Stop the recording • Choose Procedures:Record Off. |
| | Restore the minimized Procedure Editor dialog box and click the Stop button. |
| | Restore the minimized Procedure Editor dialog box and select Control:End Record . |
| 6 | Choose File:Save or File:Save As in the dialog box. Result: The Save As dialog box opens. |
| 7 | Type a name for the new procedure in the Procedure name text box. Select the Global procedure checkbox if desired (see further information below). Click OK. Result: The procedure is saved and available for future use. |
| 8 | Click the Close button to close the dialog box. |

How to create a Global procedure

You can choose to save the new procedure as a **Global procedure**. This makes the procedure available to all users. The procedure will have **(Global)** before the name to designate that it is available to all users.

You must have **Edit global list(s)** authorization to be able to save **Global procedures**.

How to build a procedure with instructions

You can select instructions in the **Procedure Editor** dialog box to build a complete procedure step by step. The procedure instructions are described in **B.4 Procedure** instructions on page 504. The table below describes how to create a new procedure with instructions.

| Step | Action |
|------|---|
| 1 | Choose Procedures:Edit:New. |
| | Result: The Procedure Editor opens in Edit mode. |
| 2 | Select an instruction from the Instruction list. |
| | Type the appropriate parameters in the Parameter field. |
| | Click Insert. |

| Step | Action |
|------|---|
| 3 | Repeat step 2 until the procedure is complete. |
| 4 | Choose File:Save. |
| 5 | Type a procedure name and click OK . |
| 6 | Click the Close button in the Procedure Editor. |

12.3.2 How to edit a procedure

Introduction

Evaluation operations are represented by instructions in the **Procedure Editor** dialog box. The instructions can be modified to suit other specific evaluation needs and be saved for later use. This section describes how to use the **Procedure Editor** to edit a procedure.

How to edit a procedure

The table below describes how to edit an existing procedure:

| Step | Action |
|------|--|
| 1 | Select Procedures:Edit:Open. |
| | Result: The Open Procedure dialog box opens. |
| 2 | Select the procedure from the list and click OK . |
| | Result: The Procedure Editor opens in Edit Mode. |
| 3 | Select an instruction in the procedure window. |
| | <i>Result</i> : The instruction parameters are displayed in the Instruction and Parameter fields. A short definition of the selected instruction is displayed at the bottom left corner. |
| 4 | Type new values in the Parameter text boxes and click the Replace button. |
| | Result: The old parameters are replaced by the new parameters. |
| 5 | Add a new instruction |
| | • Select the instruction in the procedure immediately before where you want the new instruction. |
| | Select a type and an instruction in the Instruction field. |
| | Type parameter values in the Parameter field. |
| | Click the Insert button. |
| | <i>Result</i> : The new instruction is inserted after the selected instruction. |
| 6 | Remove an instruction |
| | Select an instruction in the procedure and click the Delete button to remove the instruction from the procedure. |
| 7 | Choose File:Save and click the Close button to close the dialog box. |

Descriptions of the procedure instructions

Appendix **B.4 Procedure instructions** on page 504 contains a list of procedure instructions with descriptions.

How to add instructions to a procedure when recording

If you start recording again you can add more instructions to a procedure that is already open in the **Procedure Editor**:

• The new instructions will be added to the end of the present procedure.

or

• The new instructions will be inserted after the selected instruction if an instruction has been selected.

Invalid instructions

The procedure will stop and display an error message if an instruction calls for an invalid operation when the procedure is run. Any subsequent instructions in the procedure will not be executed.

Address the right curves

Curves are identified only by their storage position. An instruction can become invalid if it addresses the wrong curve:

Example

- The instruction **ADD** (01,02,03) will try to add curve 01 to curve 02 and store the result in position 03.
- A curve in position 03 that is not a raw data curve will be overwritten.
- A raw data curve in position 03 cannot be overwritten and the procedure will be stopped at that point.

Default values for classic baseline instructions

When a classic or morphological algorithm is used to calculate a baseline, UNICORN will suggest default values for the four control parameters based on the appearance of the curve. To instruct UNICORN to use default values appropriate for the curve every time the procedure is run, choose the default setting in the appropriate fields for the parameters.

Example

• CALCULATE BASELINE (01, 06, XXX, XXX, XXX, XXX)

Can be changed to:

• CALCULATE_BASELINE (01, 06, DEFAULT, DEFAULT, DEFAULT, DEFAULT)

Global procedures

It is not advisable to edit existing global procedures. Open the global procedure instead and save a copy under a new name. Use this copy for editing purposes.

12.3.3 How to run a procedure

Introduction

You can run the saved procedures either for a specific chromatogram or as batch runs.

How to run a single procedure

The table below describes how to run a procedure for a specific chromatogram.

| Step | Action |
|------|---|
| 1 | Open a result file. |
| 2 | Select Procedures:Run. Result: The Run Procedure dialog box opens. |
| 3 | Select the procedure from the list and click OK . <i>Result</i> : The procedure is executed. |

Note: You can also open the procedure in the **Procedure Editor** dialog box and choose Control:Run or click the Play button.

Batch runs

It is possible to apply an evaluation procedure to a designated batch of result files if they are not open in the **Evaluation** module. An open file will not run and an error message will be displayed.

The batch run is performed in the background of the **Evaluation** module and the results of the run are not seen, with the exception of prints and documentation that are defined as steps in the procedure. For example, batch runs are useful

- to perform integration with the same parameter settings on many results,
- to print a number of results with the same settings.

batch run

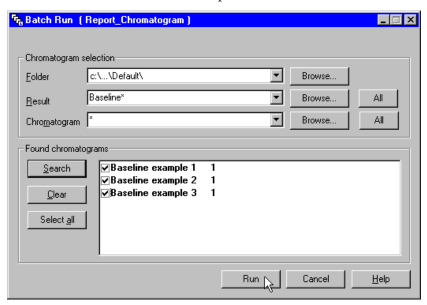
How to perform a The table below describes how to perform a batch run:

| Step | Action |
|------|--|
| 1 | Choose Procedures: Batch run. Result: The Open Procedure dialog box opens. |
| 2 | Select the procedure from the list and click OK . Result: The Batch Run dialog box opens. |

| Step | Action | |
|------|--|--|
| 3 | Use the Browse button to find and select the folder to search for result files and chromatograms. | |
| | <i>Note</i> : The search will only be performed in the selected folder. You can use standard wildcard characters and define restricting search criteria for the Result and Chromatogram fields. Up to 10 user-defined search filters can be saved in the drop-menus. | |
| 4 | Click the Search button. | |
| | Result: A list of found chromatograms is displayed. | |
| 5 | Select the chromatograms you want to perform the run on. | |
| | The Select All button selects all chromatograms. | |
| | The Clear button removes all chromatograms from the list. | |
| 6 | Click the Run button. | |
| | Result: The batch run is performed and any created curve or peak table will automatically be saved in each result file. | |

The Batch Run dialog box

The illustration below is an example of search results in the **Batch Run** dialog box:



How to batch-run reports

Evaluation procedures combined with batch runs can be a useful tool to produce printed documentation simultaneously for many result files, e.g. for a number of scouting runs. The table below describes how to create a procedure to batch-run reports.

| Step | Action | |
|------|--|--|
| 1 | Choose Procedures:Record On to record a procedure. | |
| 2 | Choose File:Report. Result: The Generate Report dialog box opens. | |
| 3 | Choose a report format. | |
| 4 | Click the Print button as the final instruction. | |
| 5 | Choose Procedures:Record Off. | |
| 6 | Save the procedure. Note: A printing procedure can also be saved with a method to produce automatic prints at the end of a run. | |

Note: When for example a batch run is performed, the latest version of the procedure will be used. However, procedures that are saved with a method are not affected if the original procedure is edited at a later time.

How to add procedures to the menu

You can add up to 15 created evaluation procedures to the **Procedures** menu in the **Evaluation** module. The table below describes how to add procedures to the menu:

| Step | Action |
|------|--|
| 1 | Select Procedures:Menu. |
| | Result: The Edit Procedures Menu dialog box opens. |
| 2 | Select the checkboxes of the procedures you want to display on the menu. Click OK. |
| | Result: The selected procedures are included on the Procedures menu. |

Remove a procedure

Open the **Edit Procedures Menu** dialog box and select the checkbox again to de-select and remove a procedure from the menu.

12.3.4 How to rename and remove procedures

Introduction

The procedures that you have created can be renamed or removed from the list of available procedures. This section describes how this is done.

How to rename a procedure

The table below describes how to rename a procedure.

| Step | Action |
|------|---|
| 1 | Choose Procedures:Edit:Rename. Result: The Rename Procedure dialog box opens. |
| 2 | Select a procedure. Result: The procedure name is displayed in the New name text box. |
| 3 | Type the new name. |
| 4 | Click OK. Result: The procedure name is changed. |

How to delete a procedure

The table below describes how to delete a procedure.

| Step | Action |
|------|---|
| 1 | Choose Procedures:Edit:Delete. |
| | Result: The Delete Procedure(s) dialog box opens. |
| 2 | Select a procedure. |
| 3 | Click 0K . |
| | Click the Yes button to confirm. |
| | Result: The procedure is deleted. |

Global procedures

It is not advisable to edit existing global procedures. Open the global procedure instead and save a copy under a new name. Use this copy for editing purposes.

13 The Analysis module

Introduction

This chapter describes how to use the **Analysis** module. This module is an optional feature that must be ordered separately and installed after the regular UNICORN installation.

The **Analysis** module is accessed in the **Evaluation** module. The **Analysis** module uses functions in the **Evaluation** module that are presented in the previous chapters. It is recommended that you are familiar with the contents of those chapters before you begin with this chapter.

In this chapter

This chapter contains these sections.

| Topic | See |
|--------------------------------------|------|
| General information about the module | 13.1 |
| Quantitation overview | 13.2 |
| How to prepare for quantitation | 13.3 |
| How to quantitate the sample | 13.4 |
| Automated quantitation | 13.5 |
| How to measure molecular size | 13.6 |

13.1 General information about the module

Introduction

This section is an overview of the **Analysis** module including:

- Definitions of terminology that will be used in this chapter.
- A description of how to install the **Analysis** module.
- A description of the new procedure instructions that become available when the **Analysis** module is installed.

Module functions

The **Analysis** module is an optional extra module that adds functionality to the regular UNICORN **Evaluation** module. Basically the **Analysis** module is used:

- to determine the absolute quantity or concentration of a component.
- to determine the molecular size of a component.

Module menus

The **Analysis** module is accessed in the **Evaluation** module. After the installation, two new **Evaluation** module menus are added:

- Quantitate.
- Mol.Size.

Note: The menus are only available when a result file is open in the **Evaluation** module.

Quantitate

The **Quantitate** function provides a wide range of techniques for quantitative analysis:

- External standard quantitation
- Internal standard quantitation
- Standard addition
- Recovery calculations

Quantitate uses peak data from standard runs to produce calibration curves which can then be used to evaluate the amount and concentration of components in a sample.

Molecular Size

The **Molecular Size** (**Mol.Size**) function determines the molecular size of components in a sample. The function uses a molecular size curve prepared from one or more standards.

Term definitions

The table below lists definitions for some terminology that is used in this chapter.

| Term | Definition |
|----------------------|---|
| Amount | This specifically refers to the injected amount. In most cases, the word "amount" is used as an abbreviation for "concentration or amount". Both concentration and injected amount can be used to produce the calibration curve. When analyzing the sample, both amount and concentration are calculated. |
| Calibration curve | The relationship between amount and peak size of a component. The relationship can be shown as a curve and as a mathematical expression. |
| Level | A known amount or concentration of a standard. The levels are numbered 1-20 in decreasing or increasing order of concentration. |
| Molecular size curve | The relationship between molecular size and retention volume for a number of components. The relationship can be shown as a curve and as a mathematical expression. |
| Molecular size table | All necessary data required to determine the molecular size of one or several components in a sample. The molecular size table contains the molecular size curve. |
| Peak size | Used generally as a common term for "peak area or peak height". |
| Peak table | The result of a peak integration presented in tabular form. The peak table can include, for example, height, area and retention volume. After the analysis, the peak table contains the values for concentration, amount (and molecular size). |

| Term | Definition |
|--------------------|--|
| Quantitation table | All necessary data required to quantitate one or several components in a sample. The quantitation table contains calibration curve(s) and peak identification settings. |
| Sample | A sample with an unknown concentration of the component(s) of interest. The concentration is determined by Quantitation . |
| | For molecular size calculations, the sample contains a component or several components of unknown molecular size. |
| Sample run | A chromatographic sample run of a sample to be analyzed. |
| Spiking | The addition of a known quantity of the component of interest to the sample prior to the sample preparation for the run. |
| Standard | A defined concentration of one or several components. The concentration does not have to be the same for all components in the standard. One or several standards are used to produce a calibration curve. |
| | For molecular size calculations, the standard contains components of known molecular size. |
| Standard run | A chromatographic standard run of a specific concentration level of a standard. |

How to install the Analysis module

The table below describes how to install the Analysis module.

| Step | Action |
|------|---|
| 1 | Close all other applications. |
| 2 | Insert the installation CD in the CD drive. |
| 3 | Open My Computer. |

| Step | Action |
|------|---|
| 4 | Double-click the CD drive icon. |
| | Result: The file window opens. |
| 5 | Double-click Setup.exe. |
| 6 | Follow the instructions on the screen. |
| 7 | Remove the CD after the installation is complete. |

Note: See the license agreement for information on the legal aspects of the installation.

The Analysis module in a network

One or several computers in a network may have the Analysis module installed. The module does not need to be installed on all network computers that run UNICORN. All installations must be made in accordance with the license agreement.

13.2 Quantitation overview

Introduction

Quantitation is used to determine the amount or concentration of components in a sample. This section is an overview over quantitation in general and the four quantitation techniques that the **Analysis** module provides. The section also contains information about the reliability of quantitation.

In this section

This section contains these sub-sections.

| Topic | See |
|---|--------|
| General information about quantitation | 13.2.1 |
| External standard quantitation | 13.2.2 |
| Internal standard quantitation | 13.2.3 |
| Standard addition quantitation | 13.2.4 |
| Recovery calculation | 13.2.5 |
| General reliability factors for the quantitation techniques | 13.2.6 |

13.2.1 General information about quantitation

Introduction

This section is a brief presentation of the quantitation techniques that the **Analysis** module provides. The section also contains an outline of the steps in a quantitation and the procedure instructions for quantitation that are added when the **Analysis** module is installed.

About quantitation

Most quantitation techniques use peak integration data from standards to produce calibration curves. These curves show the relationship between the amount of the components of interest and the peak sizes at different concentration levels of the standard. The relationship can be linear, quadratic or point-to-point. Quantitation is usually based on a number of test runs using a standard at several concentration levels.

The amount and concentration of the component(s) of interest in the sample are then determined from the peak size of the component using the calibration curve.

Note: Quantitation should only be performed on chromatograms that have been integrated and saved. Time is the recommended base unit for quantitation and it must be used for all integrations.

Quantitation steps

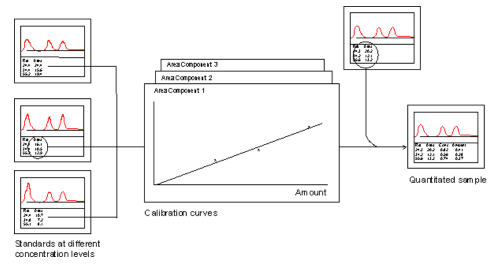
The table below describes the general steps in quantitation. The steps are described in detail in the sections about the different quantitation techniques.

| Step | Action |
|------|--|
| 1 | Run the different concentration levels of the standard. |
| 2 | Integrate the curves to produce peak tables.Check the integration. |
| 3 | Identify the components for which calibration curves will be produced. |
| 4 | Enter the known concentrations for the different standards to produce a calibration curve for each selected component. |
| 5 | Run the sample and integrate the curve. |
| 6 | Let the program calculate the concentration and amount of the components of interest in the sample. |

Note: The steps above do not apply to **Standard addition**. See "Standard addition quantitation" below.

Illustration of the work flow

The quantitation work flow is illustrated below.



The four quantitation techniques

The Analysis module provides four different quantitation techniques:

- External standard quantitation
- Internal standard quantitation
- Standard addition quantitation
- Recovery calculation

Each technique is described below.

External standard quantitation

One or several component(s) of interest are run to produce a calibration curve. The amount and concentration of the component in the sample is then determined from the calibration curve. This technique is fairly simple and usually produces accurate results.

Internal standard quantitation

Peak areas of the components of interest are related to the peak area of an internal standard added in a fixed amount to each concentration level of the standard and to the sample. This technique reduces errors that are caused by changes occurring between the separation runs and is therefore the technique that can produce the highest precision if a suitable internal standard can be selected.

Standard addition quantitation

The sample is spiked with a known amount of the component of interest. The areas of the spiked and unspiked sample are then compared and the amount in the unspiked sample is determined. No calibration curves from standards are used. Only one component can be quantitated. Compared to other techniques, results can be obtained more quickly when you are performing a small number of sample runs with standard addition. However, the precision is limited.

Recovery calculation

Recovery is used to determine the losses that can occur during the sample preparation process. The sample is spiked with a known amount of the component of interest. The amount in the spiked sample is then determined from a calibration curve and is compared with the amount in an unspiked sample. The recovery can only be determined for one component each time.

Analysis procedure instructions

The table below describes the new procedure instructions for quantitation that become available when the **Analysis** module is installed.

| Instruction | Description |
|-------------|---|
| QUANTITATE | The instruction calculates the concentration and amounts in the sample from a quantitation table. |
| | Amount and concentration columns will be added to the peak table. |
| UPDATE | The instruction updates a quantitation table with new data from one standard concentration level. |
| | The default Limit (+/-) value of 12.5% will be used. The quantitation table will not be updated if the peak area or peak height of the new and the previous results differ more than the Limit value. |
| | <i>Note</i> : Either peak area or height is selected for the Limit value. |

Default values

The DEFAULT value for the injection value will be taken from the injection volume reported by the **Autosampler A-900** from the method. DEFAULT can only be used when the injection is performed by the autosampler.

The DEFAULT value for the concentration level for the standard will be taken from the level entered in the **QuantitationData** instruction in the method.

13.2.2 External standard quantitation

General information

External standard quantitation is based on the use of a standard prepared in a number of concentration levels. A run is performed for each concentration level and calibration curves are produced to show the relationship between amount and peak size for each component. The calibration curves are used to quantitate the components in the sample.

Note: The standard should contain known amounts of all the components that are to be quantitated in the sample.

How to improve quantitation

External standard quantitation can be based on the use of a single standard concentration level, but the calibration curve is then limited to a linear through-the-origin relationship. The use of a number of different concentration levels of the standard broadens the range of the calibration curve. It also allows the development of non-linear calibration curves and improves precision. Multiple runs at each level improve precision further.

The description in this section is based on the use of a standard

- that contains two components,
- which is run at three different concentration levels.

How to perform External standard quantitation

The table below describes briefly how **External standard quantitation** is performed (based on the use of a standard which contains two components and which is run at three different concentration levels).

| Action |
|---|
| Perform a run for each standard level. |
| Integrate the curves to produce a peak table for each run. Standard 3 levels |
| I m 2 |

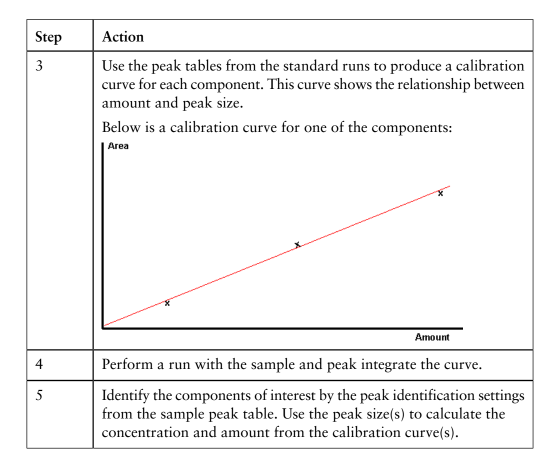
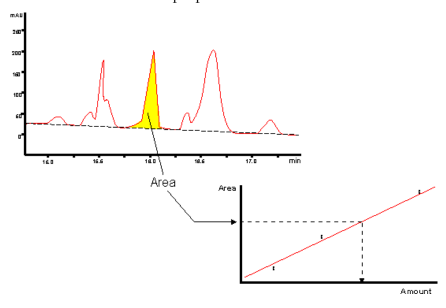


Illustration - how to use the calibration curve The illustration below describes how the calibration curve is used to determine the amount based on the sample peak area.



Reliability

External standard quantitation normally produces accurate results and is fairly simple. The following reliability factors are specific to the technique.

- Precision is limited by changes that may take place between the runs, for example column degradation and mobile phase variations.
- There is no compensation for losses of sample during the sample preparation process prior to analysis.

13.2.3 Internal standard quantitation

General information

Internal standard quantitation uses peak tables prepared from the standard, similar to the **External standard quantitation**. However, a fixed quantity of an additional component is added to every separation run, including the sample. The peak sizes of the standards and the sample are then related to the peak size of the internal standards to compensate for any changes that may have occurred between the runs.

General assump-

The internal standard technique relies on the assumption that any changes in the injected amount of the component(s) of interest, e.g. due to sample preparation losses, correspond to equal changes in the injected amount of the internal standard component.

Advantages

Internal standard quantitation reduces errors that are caused by changes in the system between successive runs with the sample and the standard concentration levels. For example, there may be unpredictable losses during the sample preparation procedure or unintentional changes in the amounts that are injected.

What is a suitable internal standard?

A suitable internal standard must meet the following conditions:

- It must be well separated from the components in the sample (not just from the components of interest).
- It must *not* be present naturally in the sample(s).
- It must have similar chemical properties to the component(s) of interest.

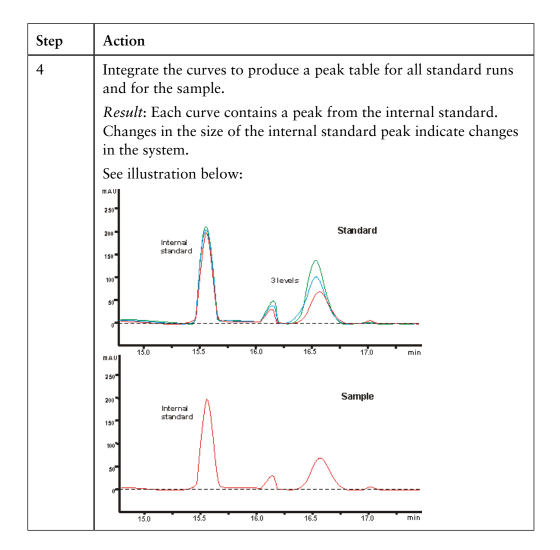
To be able to compensate for losses during the sample preparation, all the standard concentration levels must be subjected to the same sample preparation procedure as the samples.

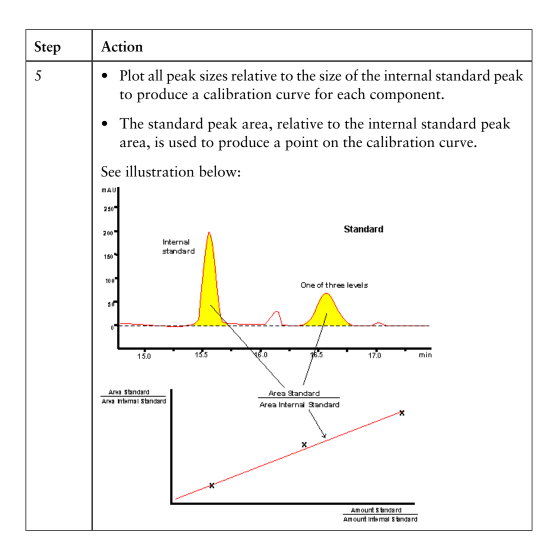
Note: If there are several components of interest, they must all be chemically similar.

How to perform Internal standard quantitation

The table below describes briefly how **Internal standard quantitation** is performed.

| Step | Action |
|------|---|
| 1 | Prepare a series of concentration levels from the standard. |
| 2 | Add an additional component, the internal standard, in the same concentration to all the standards and to the sample. |
| 3 | Perform a run for each standard and the sample. |





| Step | Action |
|------|--|
| 6 | • Prepare data from the sample in the same way as the data from the standard runs to produce peak sizes relative to the internal standard peak size. |
| | • The resulting relative value is applied to the calibration curve to determine the amount and concentration of the component of interest. |
| | See illustration below: |
| | Internal stan dard Sample Sample 150 150 150 150 150 150 150 15 |
| | Area internal Standard X Amount component of interest Amount Internal Standard |

Reliability

Internal standard quantitation is potentially the most reliable of the quantitation techniques. However, if the internal standard component is not selected carefully, the reliability will probably be worse than with the external standard technique. There are some specific factors that can affect the reliability:

- There is an increased risk of overlap when the extra component (the internal standard) is added if the sample contains many peaks.
- The addition of the internal standard must be accurate in both the standards and samples, otherwise, the precision of the quantitation will be reduced dramatically.

13.2.4 Standard addition quantitation

General informa-

Standard addition quantitation is a simple way to obtain measurements of amount in your sample (concentration is not calculated). It requires only a first sample run and a second sample run which has been spiked with a known amount of the component of interest. The technique is straight-forward and relatively fast when you are running only a few samples. Standard addition can be useful when you want to use the internal standard technique but do not have a suitable internal standard.

Disadvantages

The disadvantages of Standard addition quantitation are

- its limited precision compared to the external and internal standard techniques
- its lack of ability to measure more than one component.

How to perform Standard addition quantitation

The table below describes briefly how **Standard addition quantitation** is performed.

| Step | Action |
|------|---|
| 1 | Perform a sample run. |
| 2 | Perform a second run with a sample that has been spiked with a known quantity of the component of interest prior to the sample preparation. See illustration below: Sample (unspiked) Sample (unspiked) Sample with Standard addition (spiked) Area corresponding to the added amount so |
| | 15.0 15.5 16.0 16.5 17.0 min |

| Step | Action |
|------|---|
| 3 | Perform peak integration on both curves in the Evaluation module to produce a peak table for both the spiked and the unspiked sample. <i>Result</i> : The difference in peak area between the spiked and the unspiked sample represents the peak area from the added amount. |
| 4 | With the assumption of a linear proportionality between the peak area and amount, and with the added amount known, the software calculates the amount of the component of interest in the sample: Unspiked sample amount = Amount added × Peak area from unspiked sample (Peak area spiked sample - Peak area unspiked sample) |

Reliability

Standard addition is the least precise of the quantitation techniques since it is restricted to a single concentration level and the amount in the sample is calculated by extrapolation. Below are factors that determine if standard addition can be used with reliable results:

- The component of interest must be completely resolved from all other components in the chromatogram. Overlapping peaks will produce unreliable results.
- The peak integration parameters (baseline settings) must be correctly selected. The default settings will be satisfactory in many cases, but the integration results have to be checked for all chromatograms.
- The standard addition technique assumes a linear through-the-origin relationship between the amount of component and peak size. This is a good approximation for small quantities under normal conditions.
- Standard addition has no way of compensating for changes that are made between the runs. However, if losses during sample preparation are constant between the two runs, they may be accounted for by spiking the sample prior to the sample preparation.
- A spike amount which is of the same order of magnitude as the sample must be used to maximize precision.
- All the runs must be performed consecutively to reduce systematic errors and thereby maximize precision.

13.2.5 Recovery calculation

General informa-

Recovery calculation is used to determine losses that can occur during the sample preparation process. Recovery can also be used to determine the recovery factor of a preparative purification or a chromatographic process. The recovery factor can only be determined for a single component.

A calibration curve is produced using a concentration series of an external standard. The calibration range must cover the amount in both the sample and the spiked sample. Two runs are performed, one with the sample and a second with the sample that was spiked prior to the sample preparation with a known amount of the component of interest. Quantitation of the data from the two sample runs allows the recovery factor of the sample preparation to be calculated.

Note: The recovery is measured as the recovery for the sample preparation, not for the separation during the chromatographic analysis.

The recovery factor

The recovery factor can be used to manually compensate for losses during sample preparation. The apparent amount in a sample is divided by the recovery factor to obtain the corrected amount.

How to perform Recovery calculation

The table below describes briefly how **Recovery calculation** is performed.

| Step | Action |
|------|---|
| 1 | Perform a run with each level of the standard. |
| 2 | Peak integrate the curves to produce a peak table for each level. |
| 3 | Use the data from the peak tables to produce a calibration curve. Note: This is the same process that is used in the External standard quantitation. |
| 4 | Spike a portion of the sample with a known amount of the component of interest prior to the sample preparation. |
| 5 | Run both the spiked and an unspiked sample. |
| 6 | Peak integrate both samples to produce peak tables for the unspiked sample and the spiked sample. |

| Step | Action |
|------|--|
| 7 | The amounts for unspiked and spiked sample are calculated from the calibration curve. The difference between these amounts provides the apparent amount of the addition. See illustration below: |
| | Sample (unspiked) |
| | 150 |
| | 50 |
| | 15.0 15.5 16.0 16.5 17.0 min |
| | Sample with addition Area corresponding (spiked) to the sample |
| | Area corresponding to the addition |
| | 15.0 15.5 16.0 16.5 17.0 min |
| 8 | The ratio of this apparent amount compared to the amount actually added to the sample determines the recovery of the system. Recovery factor = Apparent amount added* Actual amount added = Amount of spiked sample - Amount of unspiked sample |
| | Example: If 2 mg of the component of interest had been added to the sample and quantitation indicated an apparent amount added of 1.6 mg, the recovery factor would then be 0.8. |

Reliability

Below are some specific factors that determine if the recovery factor result is reliable:

- A spike amount that is of the same order of magnitude as the sample must be used to maximize the precision.
- It is assumed that the recovery is the same for both the sample and the spiked sample. However, if the recovery varies according to the amount of the component of interest, the results are unreliable.

13.2.6 General reliability factors for the quantitation techniques

Reliability factors

The following factors are valid for all quantitation techniques, except for **Standard** addition:

- Quantitation requires that the components of interest are completely resolved from all other components in the chromatogram. Overlapping peaks will produce unreliable results.
- The peak integration parameters (baseline settings) must be correctly selected. The default settings will be satisfactory in many cases, but the integration results have to be checked for all chromatograms.
- All integrations must be performed using the same X-axis base unit. For highest reliability, time is the recommended unit.
- The concentration levels of the standard have to be accurately prepared. Errors in the amount or concentration values will lead to unpredictable results.
- Self-imposed limitations, such as the use of a small number of concentration levels of the standard, also limits precision.
- Precision is improved by the appropriate choice of the concentration range of the standard. The range should extend across the presumed amount in the sample.
- Use of the most appropriate curve model will maximize precision.
- Accuracy is improved if several runs are performed at each level.
- All the runs should be performed consecutively to reduce systematic errors and thereby maximize precision.

Further information

Refer to statistical reference books for more detailed information about quantitative analysis. An example is "Statistics for Analytical Chemistry", 3rd Edition 1993, J.C. Miller and J.N. MIller, Ellis Horwood PTR Prentice Hall.

13.3 How to prepare for quantitation

Introduction

This section describes how to use peak data from standards to prepare quantitation tables and calibration curves for use with External standard, Internal standard and Recovery quantitation.

In this section

This section contains these sub-sections.

| Topic | See |
|---|--------|
| Preparations before quantitation | 13.3.1 |
| How to create a quantitation table | 13.3.2 |
| How to edit and update a quantitation table | 13.3.3 |

13.3.1 **Preparations before quantitation**

Description

The table below describes the preparations before the quantitation.

| Step | Action |
|------|--|
| 1 | Create a method to be used for all the standard runs. The method and the injection volume must be the same for all the runs. |
| 2 | Perform at least one run for each concentration level of the standard. |
| 3 | Peak integrate the curves to produce a peak table for each of the standard curves. |
| | <i>Note</i> : When integrated, all standards must use the same X-axis base unit. Time is the recommended unit for the highest reliability. |
| 4 | Check that each integration is correct and consistent. |
| 5 | Select File:Save to save all the peak tables. |

Concentration levels

The standard series should include standard concentrations that extend beyond the lower and upper limits of the sample amount. If an internal standard is used, the internal standard must be added in the same concentration in all standards.

Methods created from a wizard

If the method is created from a wizard for ÄKTAdesign systems, you may select the correct standard concentration level in the variable Quantitation_Type. You can also set the level after the run has been performed. Each level is an alias for a specific concentration of the standard.

The list below describes how the levels are applied:

- Level 1 should be selected for the standard with the highest or lowest concentration.
- The levels must be set in consecutive order of changing concentration of the standard.
- All runs with the same concentration must be given the same level.

Reject irrelevant peaks

If many small irrelevant peaks are detected, it may be an advantage to re-integrate after adjusting the Reject peaks criteria. The number of largest peaks to detect has a default value of 20 and it may be helpful if this is set to a smaller value.

13.3.2 How to create a quantitation table

Introduction

The quantitation table contains all the necessary data, such as the calibration curves, that are needed to quantitate one or several components in a sample. This section describes how the quantitation table is created.

How quantitation tables are created

Quantitation tables are created in the same way for both external standard quantitation and for recovery calculations. They both use absolute values of standard peak data.

For quantitation with internal standard, the peak sizes relative to the size of the internal standard peak are used to create a calibration curve.

Four process steps

The creation of the quantitation table can be divided into four steps:

- 1. Standard data input
- 2. Component selection and definition
- 3. Peak identification
- 4. Calibration curve and quantitation table creation

Step 1 - How to input the standard data

The table below describes how to input the standard data in the **Evaluation** module.

| Step | Action |
|------|--|
| 1 | Select Quantitate:Edit Quantitation Table:New on the menu bar. |
| | <i>Result</i> : The New Quantitation Table dialog box opens with the name of the active chromatogram displayed in the Source chromatogram field. |

| Step | Action |
|------|---|
| 2 | Double-click a result file in the Select peak table list if you want to select a source chromatogram from another result file. |
| | Standards expressed in Ural Yolume unit Image Image |
| | If desired, the standard can be expressed in Concentration instead of in Amount . • Click the Concentration checkbox and edit the injection volume in the Inj. volume field. |
| | in the Inj. volume field. Note: The software will always calculate both amount and concentration for the sample. |
| | • Highlight the standard peak table of level 1 on the Peak table(s) list and click the Select button. |
| | Note: This should be the table for the highest or lowest concentration of the standard. Result: The peak table is added to the Level/Peak table(s) list. |
| 3 | The level is automatically copied onto the list if it already was set in the method. If so, continue with step 4. If a level has <i>not</i> been set, the Select Level dialog box opens. Select |
| 4 | 1 on the Level menu and click OK . |
| 4 | • Click another result file in the Results field and select the new source chromatogram. |
| | Result: The peak tables associated with this chromatogram are displayed on the Peak table(s) list. |

| Step | Action |
|------|---|
| 5 | • Repeat steps 3 and 4 until all the standard peak tables have been selected. |
| | <i>Note</i> : Increase the level number for each new standard concentration in consecutive order of decreasing or increasing concentration. |
| | • Click the Current button at any time to return to the chromatogram that was active before you activated Quantitate . |
| | Highlight unwanted tables on the list and click Remove . |
| | Click OK to finish the selection. |
| | Result: The Define Component(s) dialog box opens. |
| | Continue to "Step 2, How to select and define components" below this table. |

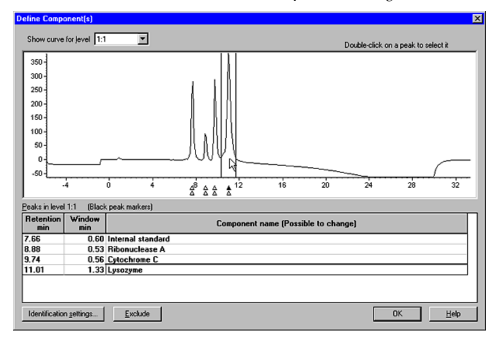
Standard concentration levels

It is useful to think of each level as an alias for a specific concentration of the standard. You can incorporate up to 10 peak tables at each level, prepared from runs repeated at the same concentration. **Quantitate** will later allocate each with an incrementing suffix, e.g. 1:1, 1:2 etc.

The Define Component(s) dialog box

The components that will be used to produce the calibration curves are selected in the **Define Component(s)** dialog box. **Quantitate** must be able to identify these components on all levels. This dialog box is used to set the criteria by which peaks are identified.

The illustration below shows the **Define Component(s)** dialog box.



Examine the components

The **Define Component(s)** dialog box initially displays the components from level 1:1, that is the peak table from the highest or lowest concentration of the standard. The **Show curve for level** list is used to examine the curve for each standard run. The size of the components are reduced or increased progressively as you select levels further down on the list, which reflects the decreasing or increasing concentration of the standard.

If an internal standard has been incorporated, its peak remains about the same size on each level.

Peaks detected during the peak integration

Each component peak that was detected during the peak integration, i.e. that is present in the peak table, is identified by a lower triangle (black in level 1:1, green in other levels). There may be different peaks detected for different levels. Upper triangles will later identify the peaks that are selected for quantitation.

Step 2 - How to select and define components

The table below describes how to select and define the components.

| Step | Action |
|------|---|
| 1 | Select level 1:1 in the Show curve for level list and click a peak. <i>Result</i> : The peak is highlighted in the table. |
| 2 | Double-click the peak. |
| | or |
| | Click the Include button. |
| | <i>Result</i> : The peak is selected for quantitation, marked with an upper triangle and " component no. " is listed as the Component name . The selected peak is affected on all levels. |
| | <i>Note</i> : More than one peak can be selected to produce calibration curves for several components. |
| 3 | Highlight the component name and type a new name. |
| 4 | Double-click the internal standard peak (if applicable) and type a new name. |
| 5 | Continue to "Step 3, How to identify the peaks" below this table. |

The Define component(s) peak table columns

The peak table within the **Define Component(s)** dialog box has three columns:

- The (absolute) **Retention** value of the component in level 1:1.
- The width of each component's window. If you change the width of the window by adjusting the cursor lines, this is reflected in the **Window** column.
- The **Component name**, with the currently selected component highlighted.

| Retention min | Window | Component name (Possible to change) |
|------------------|--------|-------------------------------------|
| 7.66 | 0.60 | Internal standard |
| 8.88 | 0.53 | Ribonuclease A |
| 9.74 | 0.56 | Cytochrome C |
| 11.01 | 1.33 | Lysozyme |

Step 3 - How to identify the peaks

Description

When a component is selected, vertical cursor lines show the current identification window. The software uses this window to search for peaks on other levels and in the sample runs. A peak found in the window is assumed to be the component of interest. You can change the limits by dragging a limit cursor line. Both cursor lines move symmetrically so that the limits center on the component peak.

The window should be set wide enough to include peaks on the other levels despite minor variations in retention volumes. However, the window should also be narrow enough to exclude unwanted peaks that will interfere with the quantitation.

Instruction

The table below describes how to adjust the window width for the best results.

| Step | Action |
|------|---|
| 1 | Drag the cursor lines to set the window to a suitable width. |
| 2 | • Use the Show curve for level menu to display all levels and check that the width is suitable (the window width is the same on all levels). |
| | Click the lower green or black triangle to display the actual retention for a peak. |
| 3 | Repeat steps 1 and 2 for all selected peaks. |
| | Note: Overlapping windows are not allowed. |
| 4 | If necessary, click the Identification settings button to edit the settings. See "How to adjust the identification settings" below this table. |
| 5 | Click the OK button to accept the default identification settings. |
| | Result: The Quantitation table dialog box opens. |
| 6 | Continue to "Step 4, How to create a calibration curve and a quantitation table" below this table. |

Identification settings

The criteria by which peaks are identified are set in the **Identification Settings** dialog box. The criteria are valid for all the selected peaks in the **Define Component(s)** dialog box. These settings also affect the information provided in the peak table in the dialog box.

How to adjust the identification settings

Description

By default, peaks are identified by their absolute retention values and the highest peak maximum within the window. In most cases, it is not necessary to change these default settings. Peak identification by absolute retention works well when there has been little or no drift in retention between successive runs of the standard. Quantitate will find corresponding peaks in these successive runs providing any drift in retention does not move a peak outside the peak window.

Instruction

If you have drifting retention that makes peak identification difficult you can choose to identify peaks according to their position relative to a reference peak. The table below describes how to adjust the identification settings in the **Define Component(s)** dialog box.

| Step | Action |
|------|---|
| 1 | Identify a component peak that can be used as the reference. Note: Choose a peak that is well separated from any other peaks. This enables the window to be set relatively wide and the system can accommodate a larger drift in retention value. |
| 2 | Click Identification Settings. Result: The Identification Settings dialog box opens. Identification Settings Peak identification Identify peaks on: Absolute retention Window width as Relative Absolute Reference peak for relative retention OK Cancel Help See "How to identify peaks within a window" below. |
| 3 | Select Relative retention on the Identify peaks on droplist. (See "Absolute and Relative window width" below) |
| 4 | Scroll down the Component menu and select the component to be used as the reference peak. |

| Step | Action |
|------|--|
| 5 | Type the window width for the reference peak (an absolute value). |
| | Note: Set the width fairly wide to accommodate a larger drift in the retention value. Make sure that there are no other large peaks within the window. • Click 0K . |
| | Result: A column for the relative retention is added in the peak table, Ret/Ref. The column displays the value of each component relative to the retention value of the reference component. This reference component is marked Ref. in the Window% column. The Window% column shows the window width for each peak expressed as a percentage of its relative retention value. |

How to identify peaks within a window

Quantitate must be advised of how the peaks are to be identified if any of the windows includes more than one peak. The second droplist in the **Peak identification** field of the **Identification Settings** dialog box offers the following options:

- Highest peak maximum (default).
- **Closest to retention**, i.e. closest to the center of the window (see the retention column in the peak table.)
- Maximum peak area.

Examine the nature of the peaks enclosed by the window and select the option that differs between the wanted and the unwanted peaks. Use **Closest to retention** if there are large peaks from components that are not going to be quantitated.

Note: The selection applies to all peaks, even the internal standard and reference if used.

Absolute and Relative window width

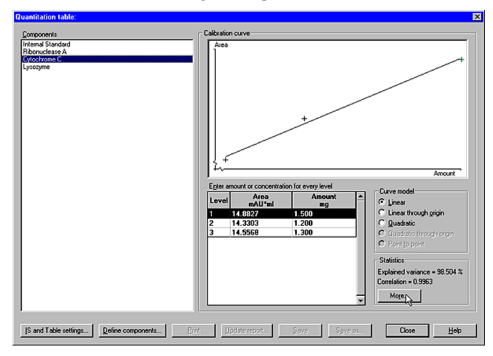
When the **Peak identification** is set to **Absolute retention**, the peak window width can be displayed as **Absolute** or **Relative**. Select the appropriate button in the **Identification Settings** dialog box.

- Select **Absolute** to show the window width of each peak in minutes (or the base volume unit).
- Select **Relative** to display the width of each component as a percentage of its retention.

If **Peak identification** is set to **Relative** retention, **Window** is set automatically to **Relative** except for the reference peak.

Step 4 - How to create a calibration curve and a quantitation table

When the component selection and identification settings are completed (see Step 3), the **Quantitation table** dialog box is opened:



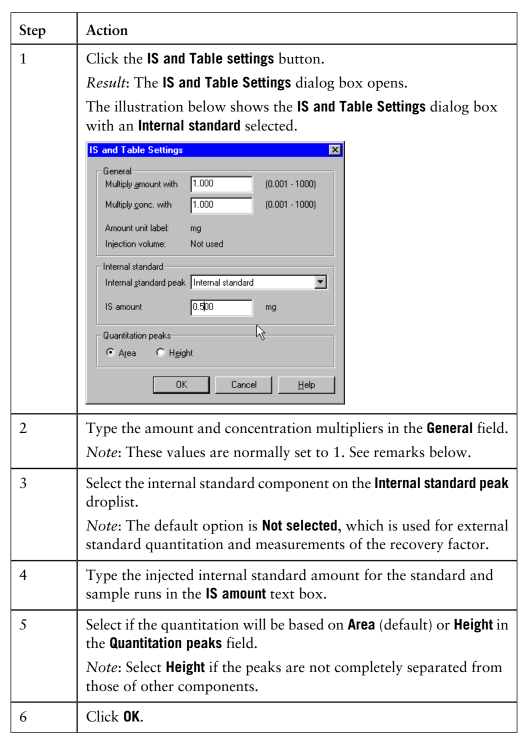
The table below describes how to enter data for the standards and create a quantitation table and a calibration curve.

| Step | Action |
|------|---|
| 1 | Click the IS and Table settings button if you want to use an internal standard or base the calibration curve on peak height (see "How to select an Internal Standard" below this table). |
| 2 | Verify that the selected components in the Components list are correct. If an internal standard is used, the related component is labelled (IS). |
| | - If relative retention has been used, the reference component is labelled (Ref). |
| | Click the Define components button to change the components. |

| Step | Action |
|------|---|
| 3 | Select the first component at the top of the Components list. |
| | <i>Note</i> : Do not select an internal standard component (if available) as the amount for this has already been entered and does not change between the levels. |
| | Highlight the Amount/Concentration for Level 1. |
| | • Type the amount or concentration of the component in the standard at this level. |
| | <i>Note</i> : This is the amount corresponding to the injected volume, not the total amount used when the standard level was prepared. |
| | Repeat this for the other levels for this component. |
| 4 | Click the Curve model radio button for the best curve model: • Linear (recommended). • Linear through origin. |
| | Quadratic. |
| | Quadratic through origin. |
| | Point to point. |
| | Result: The curve is displayed in the Calibration curve window. Each component level is labelled with crosses. If more than one run has been performed for any level, all points in that level will be shown. The average of these points is calculated and this value is used to produce the calibration curve. |
| 5 | Repeat steps 3 and 4 for all the remaining components. |
| | <i>Result</i> : The quantitation table is complete with a calibration curve for each component. |
| 6 | Save the quantitation table and click Close . |
| | or |
| | Click the Save as button. |
| | Result: The Save quantitation table dialog box opens. |
| | Note: The Save button is used to save updates in an existing quantitation table. However, this will overwrite the original table. You might prefer to use Save as and create a new name for the edited table to preserve the original. |
| 7 | Specify if the table is to be globally accessible to any user or restricted to your personal user ID. The default is global. |
| | Type a name in the Quantitation table name field. |
| | Click the 0K button. |

How to select an Internal Standard

The table below describes how to select an internal standard in the **Quantitation table** dialog box.

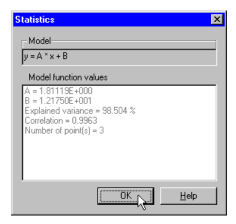


Note: The amount and concentration of the sample are multiplied by the multiplier values when the calibration curve is applied to a sample. Change the default values if you want to determine the amount or concentration in the starting volume of the sample instead of in the injected volume of the sample.

Quantitation statistics

The **Statistics** field in the **Quantitation table** dialog box displays the **Correlation** and **Explained variance** values when available.

Click the **More** button to open the **Statistics** dialog box for a complete display of available data.



Statistical reference values

- The correlation (only available for linear models) should be as close as possible to 1.00.
- The explained variance value should be as close as possible to 100%.

Note that the value is usually rather high even for poor models. A value of 90% indicates a very poor model.

The explained variance is not shown for curve models that are drawn through the origin.

Note: If the point-to-point curve model is selected, no statistics are available.

13.3.3 How to edit and update a quantitation table

How to open an existing table

The table below describes how to open an existing quantitation table for editing in the **Evaluation** module.

| Step | Action |
|------|---|
| 1 | Select Quantitate:Edit Quantitation Table:Open. |
| | Result: The Open quantitation table dialog box opens. |
| 2 | Select a quantitation table from the Quantitation table(s) list. Note: By default the list will show the quantitation tables that are globally available. Click the Personal radio button to display the tables that are restricted to your own user ID. |
| 3 | Click OK . Result: The Quantitation table dialog box opens. |

Note: **Quantitate** includes an update function that can be used to add new peak size data to an existing quantitation table in a simplified way. This function does *not* allow you to redefine components in the **Define Component(s)** dialog box.

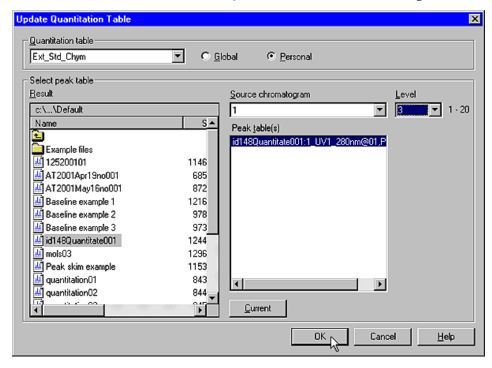
The update function

The update function can be used to add new peak size data to an existing quantitation table. This enables precision to be improved through the use of data from a number of standard runs. It also simplifies the process of renewing the calibration curves before each analysis.

Note: The injection volume must always be the same for the new run as it was for the previous standard runs.

The Update Quantitation Table dialog box

The illustration below shows the **Update Quantitation Table** dialog box.



How to prepare the calibration curve for updating

The table below describes how to open the function and prepare the calibration curve for updating.

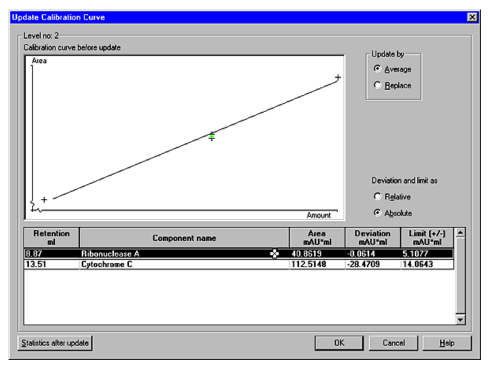
| Step | Action |
|------|--|
| 1 | Perform a peak integration for the new run and save the result. |
| 2 | Select Quantitate:Edit Quantitation Table:Update. Result: The Update Quantitation Table dialog box opens. |
| 3 | Select the Personal radio button if the table is located in your personal folder. Select the quantitation table that is to be updated in the Quantitation table field. |
| 4 | Double-click the result file in the Select peak table list to access the new data. Click the Current button if you want to use the result file that is open in the Evaluation module. |
| 5 | Select the chromatogram on the Source chromatogram list. Select the peak table that contains the new data in the Peak table(s) list. |

| Step | Action |
|------|--|
| 6 | Select the level you wish to update on the Level list. |
| | • If the selected quantitation table is based on concentration, verify or edit the Inj. Volume field. |
| | • Click OK . |
| | Result: The Update Calibration Curve dialog box opens. |
| | See "How to update a calibration curve" below. |

The Update Calibration Curve dialog box

Data on the selected components for the curve to be updated are shown in the **Component name** table. When a component is highlighted, its calibration curve is displayed above in the **Calibration curve before update** field.

The calibration curve to be updated is shown without taking the new point into consideration. A new point is shown either in green or red. If it is green, the area falls within the set **Limit (+/-)** value and this point will be used for calculation of the new calibration curve, instead of the old point. If it is red, it falls outside this range.



How to update a calibration curve

Peak size deviation

The **Deviation** column of the **Update Calibration Curve** dialog box shows how much the peak size for the proposed new point differs from the existing size. The **Limit** (+/-) column displays the set limit for the deviation. The default value is +/- 12.5% of the existing peak size. You can edit the **Limit** (+/-) value. Use the **Deviation and limit as** radio buttons to specify if both of these columns are expressed in **Absolute** or **Relative** (%) units.

Instruction

The table below describes how to use the **Update Calibration Curve** dialog box for calibration curve updates.

| Step | Action |
|------|---|
| 1 | Choose to update by Average or by Replace . The same selection applies to all components. |
| | See explanations for the options below this table. |
| 2 | Select each component table rows in turn and check that the new point falls within acceptable limits. |
| 3 | Click the Statistics after update button. |
| | Result: The Statistics after update dialog box opens. |
| 4 | Use the statistical data to check the curve model. |
| | <i>Note</i> : The old non-updated calibration curve is still shown, but the statistics apply to the data after the update. If the new point is red, the statistics shown will be those for the old curve. |
| | Click OK to close the Statistics after update dialog box. |
| 5 | Repeat steps 2-4 for each component. |
| 6 | Click OK. |
| | Result: The Update report dialog box opens. This report provides a summary of the proposed update so that you can assess its viability. See illustration below. |
| 7 | Click the Print button for a print-out of the Update report . |
| | and/or |
| | Click Save or Save as to save the updated table. |

Update by Average

The **Average** option means that the average area value is calculated from the old point (representing the average of the old points at this level) together with the new point. The green point represents the new average value and not the position of the point from the new peak table.

Update by **Average** may be used if you want to increase the precision of the calibration curve by performing several runs at each level.

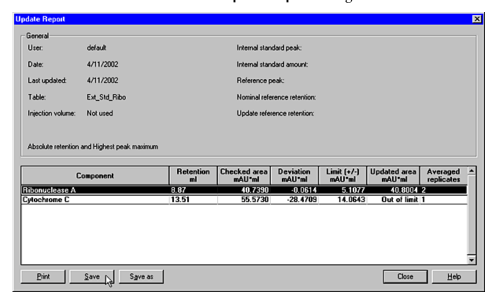
Update by Replace

The **Replace** option means that the old point (representing the average of the old points at this level) will be replaced with the new point shown in green. The data for the old point can then not be recovered.

Update by **Replace** may be used to simplify the process of renewing the calibration curve before each analysis. Instead of manually producing a new quantitation table, you may renew an existing table by running all standard levels again and updating the table with **Replace**. The old data will then be deleted.

The Update Report dialog box

The illustration below shows the **Update Report** dialog box.



Features

The list below describes some features of the dialog box.

- Components that will not be updated are shown in the column **Updated area** (or **Updated ratio** if an internal standard is used) with the text Out of limit.
- The column **Averaged replicates** shows the number of points used to calculate the average area value. After each update by **Average**, the number is increased by one. After an update by **Replace**, the number will be one.
- **Nominal reference retention** shows the retention for the reference peak in level 1:1.
- **Update reference retention** shows the retention for the reference peak in the new peak table.

How to rename a quantitation table

The table below describes how to rename an existing quantitation table.

| Step | Action |
|------|---|
| 1 | Select Quantitate:Edit Quantitation Table:Rename. |
| | Result: The Rename quantitation table dialog box opens. |
| 2 | • Select Personal to display the quantitation tables that are restricted to your own user ID, if needed. |
| | • Select the quantitation table you wish to rename on the Quantitation table(s) list. |
| | Click in the Quantitation table name text box and type a new name. |
| | Click the Rename button. |
| | Click the Close button. |

Note: You must have **Edit global list(s)** rights to be able to rename a global quantitation table.

How to delete a quantitation table

The table below describes how to delete an existing quantitation table.

| Step | Action |
|------|---|
| 1 | Select Quantitate:Edit Quantitation Table:Delete. |
| | Result: The Delete quantitation table dialog box opens. |
| 2 | • Select Personal to display the quantitation tables that are restricted to your own user ID, if needed. |
| | • Select the quantitation table you wish to delete on the Quantitation table(s) list. |
| | Click the Delete button. |
| | Click the Yes button to confirm. |
| | Click the Close button. |

Note: You must have **Edit global list(s)** rights to be able to delete a global quantitation table.

13.4 How to quantitate the sample

Introduction

This section describes how to use calibration curves to quantitate samples. Calibration curves are applicable to external and internal standard quantitation and to recovery factor measurement. Standard addition measurements are also described.

In this section

This section contains these sub-sections.

| Topic | See |
|---|-----|
| External and internal standard quantitation | |
| Standard addition quantitation | |
| How to calculate the recovery factor | |

13.4.1 External and internal standard quantitation

Introduction

This section describes how to perform quantitation in the **Evaluation** module using either an external standard or an internal standard.

The processes involved in both external standard and internal standard quantitation of a sample are very similar. The procedural differences mainly concern the creation of the quantitation tables. A quantitation table is specific to either external standard or internal standard quantitation.

Method for the sample runs

The method that is used for the sample runs must be the same as for the standard runs. If the method is created from a wizard or a template for ÄKTAdesign systems, select **Sample** in the variable **Quantitation_Type** on the **Variables** tab in the **Run Setup**.

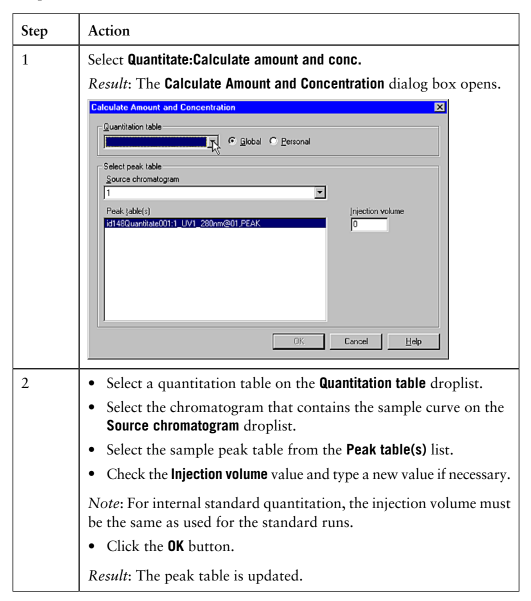
How to prepare for the quantitation

The table below describes briefly how to prepare for the quantitation.

| Step | Action |
|------|--|
| 1 | Prepare a quantitation table for the components of interest. See 13.3.2 How to create a quantitation table on page 411 for further information. |
| 2 | Perform a sample run. Note: If internal standard quantitation is used, the internal standard must have been added to the sample prior to the sample preparation procedure. The injected amount must be the same as on the standard levels. |
| 3 | Open the sample result file and peak integrate the sample curve to produce a peak table. Note: The sample curve must use the same X-axis base unit as the standards during the integration. Time is the recommended unit for highest reliability. |
| 4 | Select File:Save to save the peak table. |

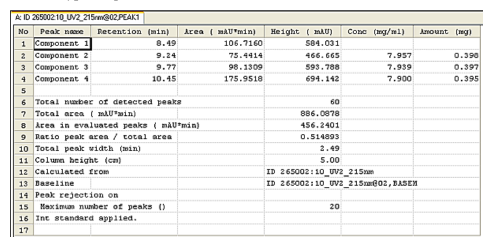
How to calculate the amount and concentration

The table below describes how to calculate the amount and concentration in the sample.



How to view the quantitation results

The results of the quantitation are shown in the **Concentration** and **Amount** peak table columns of the **Evaluation** module. The **Peak Names** are shown in the table and the type of quantitation is also listed. See illustration below:



The quantitation table used for the quantitation

When the result file is saved, it includes the quantitation table that was used for the quantitation. You can view the table that was used by selecting **Quantitate:Edit Quantitation Table:View Current**.

If the amount cannot be calculated

If the amount cannot be calculated, one of the following signs is shown in the peak table **Amount** column:

| Sign | Function |
|------|---|
| > | This means that the value is higher than the highest value in the calibration curve, i.e. outside the valid range of the calibration curve. |
| < | This means that the value is lower than the lowest value in the calibration curve, i.e. outside the valid range of the calibration curve. |
| - | This means that the value cannot be calculated. For example, this sign might indicate that the peak could not be identified. |

Standard addition quantitation 13.4.2

addition

Stages in standard Standard addition is performed in five stages:

| Stage | Description |
|-------|---|
| 1 | Perform two runs. |
| 2 | Copy the curves into one result file. |
| 3 | Integrate the curves to produce the peak tables. |
| 4 | Select the component to be used. |
| 5 | Evaluate the amount of a component in the sample. |

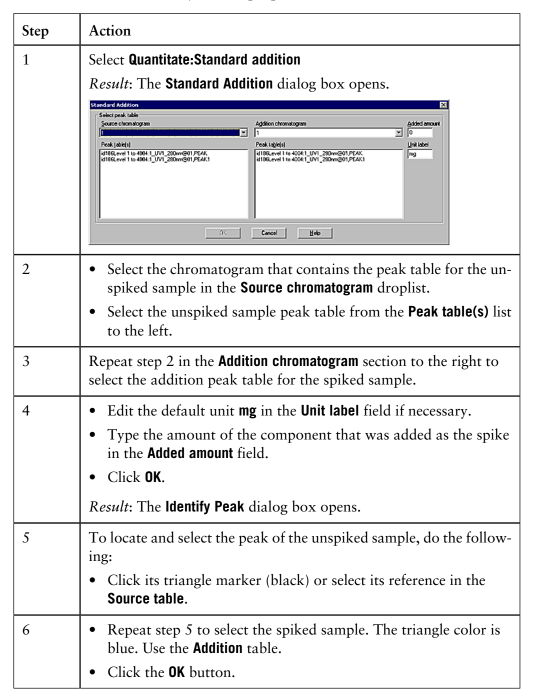
How to prepare for the Standard addition quantitation

The table below briefly describes how to prepare for the quantitation.

| Step | Action |
|------|---|
| 1 | Perform a sample run with the unspiked sample and a run with the spiked sample. |
| 2 | Open one of the two result files. Use File:Open:Curves to copy the second curve to the opened result file. |
| 3 | Peak integrate the sample curves to produce the peak tables for the unspiked and the spiked samples. Note: The sample curves must use the same X-axis base unit. Time is the recommended unit for highest reliability. |
| 4 | Check that the integrations are correct.Optimize the peak integration if necessary. |
| 5 | Select File:Save to save the peak table. |

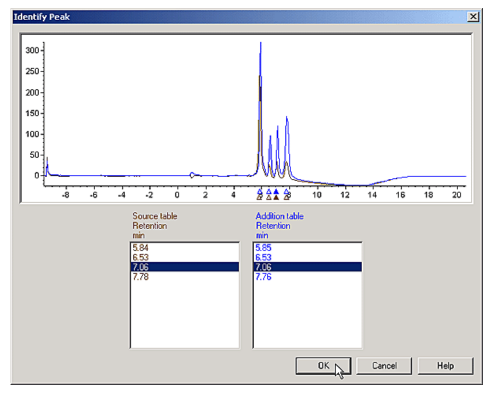
How to select the component and identify the sample peaks

The table below describes how to select the component to be used for the standard addition and how to identify the sample peaks.



The Identify Peak dialog box

The illustration shows the **Identify Peak** dialog box, described in the table above.



How to view the quantitation results

The amount of the component of interest is displayed in the peak table **Amount** columns of the **Evaluation** module.

| No | Retention (min) | Area (mAU*min) | Height (mAU) | Amount (mg) |
|----|------------------------------------|-----------------|---------------|-------------|
| 1 | 5.84 | 52.9263 | 282.754 | |
| 2 | 6.53 | 4.8881 | 30.365 | |
| 3 | 7.06 | 6.4036 | 38.687 | 0.913 |
| 4 | 7.78 | 11.3658 | 42.744 | |
| 5 | | | | μž |
| 6 | | | | 220 |
| 7 | 7 Total area (mAU*min) | | | 92.9166 |
| 8 | Area in evaluated peaks (mAU*min) | | | 75.5839 |
| 9 | Ratio peak area / | total area | | 0.813459 |

13.4.3 How to calculate the recovery factor

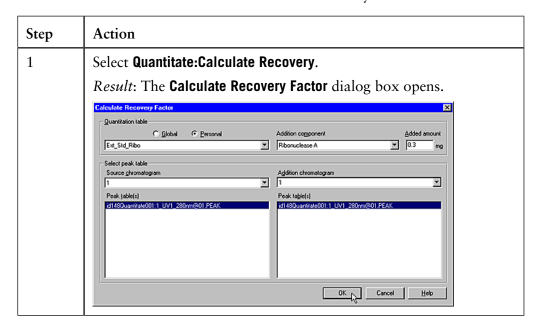
How to prepare for the quantitation

The table below briefly describes how to prepare for the quantitation.

| Step | Action |
|------|---|
| 1 | Prepare a quantitation table for the components of interest. Note: An external standard quantitation must be used. Internal standard quantitation tables cannot be used. |
| 2 | Perform a sample run with the unspiked sample and a run with the spiked sample. |
| 3 | Peak integrate the sample curves to produce the peak tables for the unspiked and the spiked samples. Note: The sample curves must use the same X-axis base unit as the standards during the integration. Time is the recommended unit for highest reliability. |
| 4 | Check that the integration is correct.Optimize the integration if necessary. |
| 5 | Open one of the sample result files. Use File:Open:Peak Tables to copy the other peak table to that result file. |
| 6 | Select File:Save to save the result. |

How to calculate the recovery

The table below describes how to calculate the recovery factor.



| Step | Action |
|------|---|
| 2 | Select Global or Personal quantitation tables. Select a quantitation table on the Quantitation table droplist. Note: Only external standard quantitation tables will be shown. Select the chromatogram that contains the unspiked sample peak table on the Source chromatogram droplist. |
| | • Select the unspiked sample peak table from the Peak table(s) list to the left. |
| 3 | • Repeat step 2 to select the peak table for the spiked sample on the Addition chromatogram fields. |
| | • Select the component that was added prior to the sample preparation on the Addition component droplist. |
| | • Type the injected amount of this component in the Added amount field. |
| | Click the OK button. |

How to view the recovery factor calculation results

The recovery factor calculated by the software is placed at the bottom of the peak table in the **Evaluation** module. You need to scroll to the end of the table to see it.

| No | Retention (min) | Area (mAU*min) | Area/Peak area ((time) % | Height (mAU) |
|----|---------------------|-----------------|--------------------------|-------------------------------|
| | Recovery Factor () | | | 0.989 |
| | Component name | | | Chymotrypsinogen A |
| | Calculated from | | | id 18501:1_UV1_280nm |
| | Baseline | | | id 18501:1_UV1_280nm@01,BASEC |
| | Peak rejection on | | | |
| | Maximum number of | peaks () | | 20 |
| | Recovery applied. | | | |
| | Current peak filte: | r settings | | |
| | Maximum number of | peaks () | | 20 |

Note: The checkbox **Do not show global peak table data** must be de-selected in the **Peak Table** tab of the **Chromatogram Layout** dialog box.

If the recovery cannot be calculated

If the recovery cannot be calculated, one of the following signs is shown in the peak table **Amount** column:

| Sign | Function |
|------|---|
| > | This means that one of the amounts/concentrations is higher than the highest value in the calibration curve, i.e. outside the valid range of the calibration curve. |
| < | This means that one of the amounts/concentrations is lower than the lowest value in the calibration curve, i.e. outside the valid range of the calibration curve. |

| Sign | Function |
|------|---|
| - | This means that the recovery factor cannot be calculated. For example, this sign might indicate that the peak could not be identified in both runs. |

13.5 Automated quantitation

Introduction

Some method wizards designed for quantitation are available for ÄKTAdesign systems supplied with **Autosampler A-900** or **A-905**. These can be used to quantitate a sample automatically or to update a quantitation table.

The procedures described in this chapter are designed for use with the systems mentioned above.

In this section

This section contains these sub-sections.

| Topic | See |
|--|--------|
| How to set up for automated quantitation | 13.5.1 |
| How to perform automated quantitation | |
| How to perform automated update | 13.5.3 |

13.5.1 How to set up for automated quantitation

Introduction

This section describes how to create a quantitation table for automated quantitation.

Basic conditions for the quantitation table

A quantitation table must be produced from standards before samples can be quantitated. The list below describes the basic conditions for the quantitation table:

- The same method must be used for all standard and sample runs.
- Each level is an alias for a specific concentration of the standard.
- All runs with the same concentration must be assigned the same level.
- Level 1 must be selected for the standard with the highest or lowest concentration.
- The levels must be set in order of decreasing or increasing concentration of the standard.

How to prepare the quantitation table

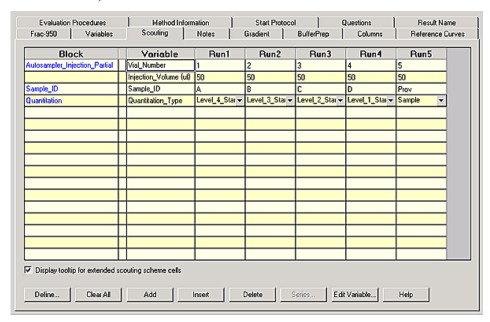
The table below describes how to prepare the quantitation table for automated quantitation.

| Step | Action |
|------|--|
| 1 | Use the Method Wizard to create a method. Select Autosampler from the Injection Technique droplist in the Sample Injection dialog box. |
| 2 | Proceed with the following dialog boxes in the Method Wizard and click the Finish button on the last dialog box. Result: The Run Setup opens. |
| 3 | Click the Scouting tab. (See illustration below) Select the Quantitation_Type variable from the Scouting Variables dialog box. Select other scouting variables of interest, e.g. Sample_ID, Vial_Number etc. Click OK. |
| 4 | Double-click the Quantitation_Type variable table cell. Select the correct standard concentration level. Note: This corresponds to the text instruction QuantitationData. You can also set this level after the run has been completed. For more information about scouting see 7.1 How to set up a scouting scheme on page 176. |

| Step | Action |
|------|---|
| 5 | Click the Evaluation Procedures tab. |
| | Select the Integrate_and_Print procedure. |
| | <i>Result</i> : This procedure will automatically use default baseline settings and integrate the first UV curve. |
| 6 | Save the method. |
| 7 | Perform all the standard runs. |
| 8 | In the Evaluation module, select Quantitate:Edit Quantitation Table:New . |
| 9 | Create a quantitation table manually from the standard runs. See 13.3.2 How to create a quantitation table on page 411. |

The Scouting tab

The illustration below shows the **Scouting** tab in the **Run Setup**, used to enter standard data, before the standard concentration level is defined.



13.5.2 How to perform automated quantitation

Instruction

The table below describes how to set up sample runs to perform automated quantitation.

| Step | Action |
|------|--|
| 1 | Select File:Open in the Method Editor module. Select a method that has been used for standard runs in the Open dialog box. (See 13.5.1 How to set up for automated quantitation on page 439) Click OK. |
| 2 | Click the Scouting tab in the Run Setup. Click the Clear All button to clear the scouting scheme. Double-click each Quantitation_Type table cell and select Sample for all sample runs. |
| 3 | Click the Evaluation Procedures tab. • Select only the Quantitate_Sample procedure. (Click the Import button to import the Quantitate_Sample procedure if it isn't displayed on the list) *Result: This procedure automatically integrates the first UV curve with default baseline settings and uses the selected quantitation table to quantitate the sample. The amounts and concentrations are then printed. |
| 4 | Click the Quantitate button. Result: The Quantitation table dialog box opens. |
| 5 | Select the quantitation table from the Global or Personal folder and click OK . *Result: The quantitation table is copied into the Quantitate_Sample procedure. *Note: The procedure cannot be executed if a quantitation table has not been selected. Time must have been selected as the X-axis base unit. |
| 6 | Save the method with a new name. |
| 7 | Perform the run(s). Result: The amount and concentration of the components in the samples will be printed automatically after each run. |

13.5.3 How to perform automated update

Introduction

This section describes how to update quantitation tables automatically, also in scouting runs. See also 13.3.3 How to edit and update a quantitation table on page 422.

How to perform automated update with the Replace option The table below describes how to automatically update a quantitation table with the **Replace** option (default).

| Step | Action |
|------|---|
| 1 | Open a method in the Method Editor . |
| 2 | Click the Scouting tab in the Run Setup. Click the Clear All button to clear the scouting scheme. Double-click each Quantitation_Type table cell and select the correct concentration level for the standards. |
| 3 | Click the Evaluation Procedures tab. • Select the Update_Quantitation procedure. (Click the Import button to import the Update_Quantitation procedure if it isn't displayed on the list) |
| | Result: This procedure automatically integrates the first UV curve with default baseline settings and updates the selected quantitation table with the new standard. An update report is then printed. |
| 4 | Click the Quantitate button. Result: The Quantitation table dialog box opens. |
| 5 | Select the quantitation table from the Global or Personal folder. Time must be selected as the X-axis base unit. Result: The quantitation table is copied into the Update_Quantitation procedure. Note: You can only perform one run at each level since the old points in the quantitation table will be replaced after each run. |
| 6 | Save the method with a new name. |
| 7 | Perform the run(s). Result: The quantitation table will be updated automatically after each run. |

Note: The quantitation table will not be updated if the peak area or peak height of the new and the previous results differ more than the **Limit** value. The **Limit** value is defined either for peak area or height.

How to perform automated update with the Average option

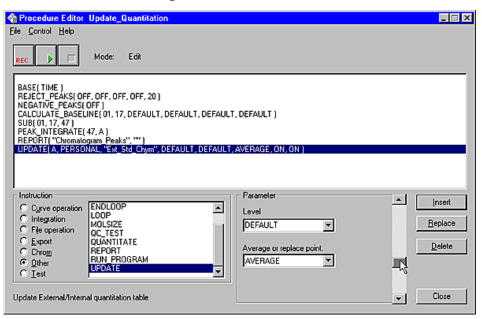
The table below describes how to automatically update a quantitation table with the **Average** option.

| Step | Action |
|------|---|
| 1 | Open a method in the Method Editor . |
| 2 | Click the Scouting tab in the Run Setup dialog box. Click the Clear All button to clear the scouting scheme. Double-click each Quantitation_Type table cell and select the correct concentration level for the standards. |
| 3 | Click the Evaluation Procedures tab. Select the Update_Quantitation procedure. Click the Quantitate button. Result: The Quantitation table dialog box opens. |
| 4 | Select the quantitation table from the Global or Personal folder and click OK . *Result: The quantitation table is copied into the Update_Quantitation procedure. |
| 5 | Click the Edit button on the Evaluation Procedures tab. *Result: The Procedure Editor dialog box opens. See illustration below. |
| 6 | Select the existing UPDATE instruction. |
| 7 | Use the scroll bar in the Parameter field to locate the Average or replace point droplist. • Select the AVERAGE option. • Click the Replace button to the right of the scroll bar. |
| 8 | Select File:Close in the Procedure Editor dialog box to return to the Run Setup. |
| 9 | Save the method and perform the runs. *Result: The quantitation table will be updated automatically after each run. New average values will be calculated from the old points together with the new points. |

Note: The quantitation table will not be updated if the peak area or peak height of the new and the previous results differ more than the **Limit** value. The **Limit** value is defined either for peak area or height.

The Procedure Editor dialog box

The **Procedure Editor** dialog box is illustrated below:



How to perform automated update in scouting runs - step 1

It is possible to run both standards and samples in the same scouting run and continuously update a previously created quantitation table with new values. The table below describes how to set up the evaluation procedures for the updates.

| Step | Action |
|------|--|
| 1 | Open the same method that was used to create the quantitation table from the standard runs and open the Run Setup . |
| 2 | Click the Evaluation Procedures tab. |
| 3 | Select Update_Quantitation and click Quantitate. Result: The Quantitation table dialog box opens. |
| 4 | Select the quantitation table and click OK . |
| 5 | Deselect the Update_Quantitation procedure on the Evaluation Procedures tab. |
| 6 | Repeat steps 3 to 5 for the Quantitate_Sample procedure. |
| | <i>Note</i> : Make sure that both procedures are deselected after this is completed. Otherwise they will be run twice. |
| 7 | Proceed with the instructions how to edit the instructions (see table below). |

How to perform automated update in scouting runs - step 2

The table below describes how to edit the text instructions.

| Step | Action |
|------|--|
| 1 | Click the Text Instructions icon. |
| 2 | Select the last instruction in the method in the Text pane. |
| 3 | Click the Other radio button in the Instructions field of the Instruction box. Select Evaluate on the Instructions list. |
| 4 | Select Update_Quantitation in the Procedure droplist of the Parameters field. |
| | Best point District Pump Base Bibble Pump Bibble Pump Bibble Pump Bibble Pump Pump |
| | Click the Var button in the Parameters field. |
| | Result: The Variable Name Definition dialog box opens. |
| 5 | Type a variable name, for example Procedure and click OK . |
| | <i>Result</i> : The Evaluate instruction is inserted in the method. By defining the evaluation procedure as a variable, different procedures can be selected on the Scouting tab for different scouting runs. |
| 6 | Proceed with the instructions on how to set up the scouting runs for the standards (see table below). |

How to perform automated update in scouting runs - step 3

The table below describes how to set up the scouting runs for the standards.

| Step | Action |
|------|---|
| 1 | Select View:Run Setup and click the Scouting tab. |
| 2 | Click the Define button. |
| | Result: The Scouting Variables dialog box opens. |

| Step | Action |
|------|---|
| 3 | Edit the scouting variables list to include: • Procedure • Vial_Number • Injection_volume • Sample_ID • Quantitation_Type Note: The Procedure variable will appear at the beginning of the list of variables, even though the Evaluate instruction is inserted at the end of the method. |
| 4 | Set up all the standard runs in the scouting scheme: Select the Update_Quantitation procedure. Ensure that Quantitation_Type is set to the correct standard level for each run. Result: The quantitation table will now be updated with new values after each run. Since the runs will be performed with the Replace (the default selection) option, you can only perform one run at each level. |
| 5 | Proceed with the instructions on how to set up the scouting runs for the samples in step 4 (see table below). |

Note: The quantitation table will not be updated if the peak area or peak height of the new and the previous results differ more than the **Limit** value. The **Limit** value is defined either for peak area or height.

How to perform automated update in scouting runs - step 4

The table below describes how to set up the scouting runs for the samples.

| Step | Action |
|------|---|
| 1 | Select the Quantitate_Sample procedure. |
| 2 | Select Sample in the variable Quantitation_Type for all the sample runs. |
| | Result: All samples will be run automatically and the amount and concentration of the components of interest will be printed after each run. |
| | <i>Note</i> : The result files will include an additional chromatogram (labelled 12) that contains a small part of the curves collected during the execution of the evaluation procedure. |

How to change the scouting runs to be updated by average

The table below describes how to change the scouting runs so that the quantitation table is updated by average after each standard run.

| Step | Action |
|------|--|
| 1 | Click the Evaluation Procedures tab in the Run Setup Editor. Click the Import button. |
| | Result: The Import dialog box opens. |
| 2 | Select the current method from the Method file menu. Highlight Update_Quantitation in the Select field. Type a new name, e.g. Update_Average in the Import as text box. Click the Import button. Click the Close button. Result: The new procedure is added to the list of Evaluation Procedures. |
| 3 | Select the new procedure and click the Edit button. *Result: The Procedure Editor dialog box opens. |
| 4 | Highlight the existing Update instruction. |
| 5 | Use the scroll bar in the Parameter field to locate the Average or replace point droplist. Select the AVERAGE option. Click the Replace button. |
| 6 | Select File:Close in the Procedure Editor dialog box to return to the Run Setup. |
| 7 | Select the Update_Average procedure and click the Quantitate button. *Result: The Quantitation table dialog box opens. |
| 8 | Select the quantitation table and click 0K . |
| 9 | Deselect all the procedures on the Evaluation Procedures tab, otherwise they will be run twice. |
| 10 | Click the Scouting tab. Select the Update_Average procedure for the second and all following runs at each standard level concentration. Note: The Update_Quantitation procedure (Update by Replace) should still be used for the first run at each level. |

13.6 How to measure molecular size

Introduction

The molecular size of components in a sample can be determined by size exclusion chromatography. A molecular size calibration curve must first be created with components of known molecular size. The retention is inversely related to the molecular size.

This section describes how to measure the molecular size.

In this section

This section contains these sub-sections.

| Topic | See |
|--|--------|
| Overview of molecular size determination | 13.6.1 |
| How to determine the molecular size | 13.6.2 |

13.6.1 Overview of molecular size determination

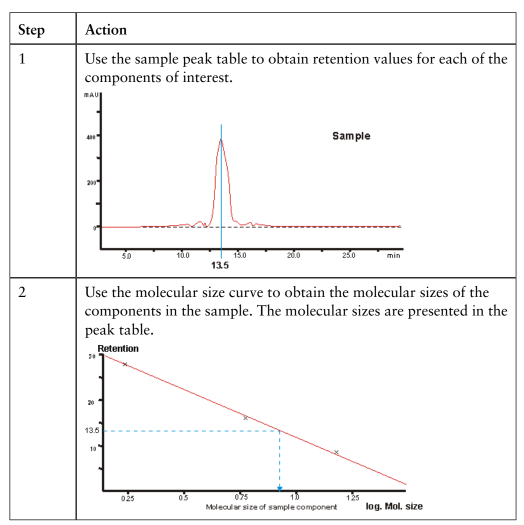
How to create a molecular size curve

The table below is a brief description of how to create a molecular size curve.

| Step | Action |
|------|--|
| 1 | Perform a run with one or more standards to create a standard curve. Note: The standards should contain a number of components of known molecular size and these should extend beyond the size limits that are expected in the test sample. |
| 2 | Peak integrate the standard curve to produce a peak table. |
| 3 | Use the peak table from the standard to produce a molecular size table. Each peak is represented by a retention value. |
| 4 | Select the relevant peaks and input data for the corresponding molecular sizes. **Result: The software plots these values as a molecular size curve. This curve has an inverse relationship between the logarithm of the molecular size and retention. **Retention** **Retention* |

How to calculate the molecular size in the sample

The table below is a brief description of how to use the molecular size table to calculate the molecular size of the components in the sample.



13.6.2 How to determine the molecular size

Introduction

This section describes the technique for measuring molecular size in detail.

Before you start

Before you create the molecular size curve, you need to do the following:

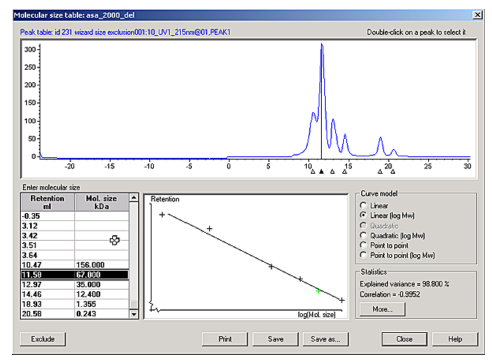
- Perform chromatographic runs with an appropriate standard with components of known molecular size. The standard should contain components of sizes that extend over the range that is expected in the sample. If you are using many components, it may be better to split them into two or more standard runs.
- Peak integrate the curves to produce peak tables. The standard curves must all use the same X-axis base unit during the integration. Volume is the recommended unit for molecular size determination.
- Save the results.

The Molecular size table dialog box

This dialog box is used to select the peaks that will be used to produce the molecular size curve. Each curve and its peak table name is color coded. All the available peaks for all the curves are listed together in the **Retention/Mol.size** table.

Triangles show that a peak has been selected. The name of its source peak table is shown above the curve window. This is useful when you wish to know which peak has been selected of two closely spaced peaks from different peak tables.

The illustration below shows the Molecular size table dialog box.



save a molecular size table

How to create and The table below describes how to create and save a molecular size table in the **Evaluation** module.

| Step | Action |
|------|---|
| 1 | Open a result file and select Mol. Size:Edit Mol. Size Table:New. Result: The New Molecular Size Table dialog box opens. |
| | New Molecular Size Table Urit Molecular size unit label: InDa Select peak table(s) C-\Default Name Example files 1 122001 or 16001 A 72001 Ay 16x001 A 8 seriene example 3 If 1880 unit sheet(s) Example files 1 20001 or 16x001 A 8 seriene example 2 B 8 seriene example 2 B 8 seriene example 3 If 1880 unit sheet(s) If 1890 unit sheet(s) Example files If 1890 unit sheet(s) If 1890 uni |
| 2 | Double-click the result file in the Select peak table(s) list. Select the source chromatogram on the Source chromatogram droplist. |
| 3 | Highlight a peak table that was prepared from the standard in the source Peak table(s) list and click the Select button. |
| 4 | Repeat step 3 to select more peak tables. <i>Note</i> : The runs must all have been made under the same conditions. |
| 5 | To deselect a table, highlight the table in the Peak table(s) list to the right and click the Remove button. |
| 6 | Repeat steps 2 to 4 to select peak tables from other result files. |
| 7 | Type the appropriate size measurement unit in the Molecular size unit label field (default kDa). Click OK when the Peak table(s) list to the right contains all the required peak tables. |
| | Result: The Molecular size table dialog box opens. |
| 8 | Use one of the following ways to select a peak: Click the peak in the curve. Click the peak entry in the Retention/Mol. size table. |
| 9 | Double-click in the Mol. size column cell and type the known molecular size from the standard. |

| Step | Action | |
|------|---|--|
| 10 | Repeat step 8 and 9 for all components of known molecular size. | |
| 11 | To remove unwanted entries, click the peak entry in the table and click the Exclude button. | |
| 12 | Select the appropriate curve model in the Curve model field (see "The molecular size curve" below). | |
| 13 | Click the Save as button. Result: The Save molecular size table dialog box opens. | |
| 14 | Choose if the table is to be globally accessible to any user or restricted to your personal user ID. The default is global. Type a name in the Molecular size table name field. Click OK. | |

The molecular size curve

The molecular size curve shows the relationship between molecular size and the corresponding retention. The curve is plotted from the **Retention/Mol. size** data that you have typed in the table as described above. Before this can be done, a curve model is needed, which describes the relationship between molecular size and retention. Each of the peaks selected is represented by a point in this curve, which is drawn according to the best fit that can be achieved using the selected model. Select one of the available models in the **Curve model** field:

- Linear
- **Linear (logMw)** (Theoretically, this is the best choice.)
- Quadratic
- Quadratic (logMw)
- Point to point
- Point to point (logMw)

Molecular size Statistics

With the exception for the two point-to-point models, the molecular size curves can be expressed as mathematical expressions. The expressions and related items can be viewed in the **Statistics** dialog box.

• Click the **More** button in the **Statistics** field of the **Molecular size table** to open the dialog box.

The expression is shown at the top of the window, followed by the values for the constant that it contains.

Statistical reference values

- The correlation value (only for linear models) should be as close to -1.00 as possible.
- The explained variance value should be as close to 100% as possible.

Note: Explained variance values are usually high. A value of 90% indicates a very poor model.

How to open an existing table

The table below describes how to open an already existing molecular size table for editing in the **Evaluation** module.

| Step | Action | |
|------|--|--|
| 1 | Select Mol. Size:Edit Mol. Size Table:Open. | |
| | Result: The Open molecular size table dialog box opens. | |
| 2 | • Select a molecular size table from the Molecular size table(s) li | |
| | <i>Note</i> : By default the list will show the molecular size tables that are globally available. Click the Personal radio button to display the tables that are restricted to your own user ID. | |
| | • Click 0K . | |
| | Result: The Molecular size table dialog box opens. | |

How to rename a molecular size table

The table below describes how to rename an existing molecular size table.

| Step | Action | |
|------|---|--|
| 1 | Select Mol. Size:Edit Mol. Size Table:Rename. | |
| | Result: The Rename molecular size table dialog box opens. | |

| Step | Action | |
|---|--|--|
| • Select Personal to display the tables that are restricted to own user ID, if needed. | | |
| | • Select the molecular size table you wish to rename in the Molecular size table(s) list. | |
| | • Click in the Molecular size table name text box and type a new name. | |
| | Click the Rename button. | |

Note: You must have **Edit global list(s)** rights to be able to rename global tables.

How to delete a molecular size table

The table below describes how to delete an existing molecular size table.

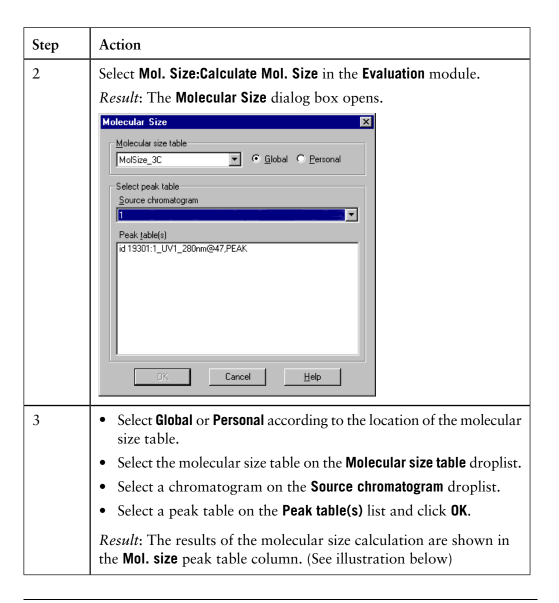
| Step | Action | |
|------|--|--|
| 1 | Select Mol. Size:Edit Mol. Size Table:Delete. | |
| | Result: The Delete molecular size table dialog box opens. | |
| 2 | • Select Personal to display the tables that are restricted to your own user ID, if needed. | |
| | • Select the molecular size table you wish to delete in the Molecular size table(s) list. | |
| | Click the Delete button. | |
| | Click the Yes button to confirm. | |

Note: You must have **Edit global list(s)** rights to be able to delete global tables.

How to calculate the molecular size

The table below describes how the molecular size curve is used to calculate the molecular sizes of the components in the sample.

| Step | Action | |
|------|---|--|
| 1 | Perform a sample run and peak integrate the curve to produce a peak table. | |
| | <i>Note</i> : The sample curve must use the same X-axis base unit as the standards. Use volume for molecular size calculations. | |



The Mol. size peak table column

The illustration below shows the Mol. size peak table column.

| | No | Retention | (min) | Mol. | size | (kDa) |
|---|----|-----------|-------|------|------|-------|
| ı | 8 | | 6.42 | | | > |
| l | 9 | | 7.90 | | | 38.22 |
| l | 10 | | 8.42 | | | 31.15 |
| l | 11 | | 9.59 | | | 19.57 |
| l | 12 | | 10.43 | | | 14.05 |
| L | 13 | | 11.71 | | | < |

When the result file is saved, it includes the molecular size table that was used for the molecular size determination. You can view the table that was used by selecting Mol. Size:Edit Mol. Size Table:View Current.

If the molecular size cannot be calculated

If the molecular size cannot be calculated, one of the following signs is shown in the peak table **Mol. size** column:

| Sign | Function | |
|------|---|--|
| > | This means that the molecular size is larger than the largest size in the molecular size curve, i.e. outside the valid range of the curve. | |
| < | This means that the molecular size is smaller than the smallest size in the molecular size curve, i.e. outside the valid range of the curve | |
| - | This means that the retention value is negative. | |

Molecular size procedure instruction

The table below describes the new procedure instruction for molecular size measurement that becomes available when the **Analysis** module is installed.

| Instruction | Description |
|-------------|---|
| MOLSIZE | The instruction calculates the molecular sizes from a molecular size curve. A Mol.size column will be added to the peak table. |

14 System settings

Introduction

This chapter describes some of the general system settings.

In this chapter

This chapter contains these sections:

| Topic | See |
|---|------|
| General information about system settings | 14.1 |
| Alarms | 14.2 |
| Curves | 14.3 |

14.1 General information about system settings

System settings

The system settings

- define settings for alarms and warnings
- select the data that will be stored in result files

When to change the system settings

Each system has a set of default settings.

• Changes to the default settings should be made when the system is installed.

Certain system settings may need to be adjusted in the following cases:

- If system components are changed: e.g. the alarm and warning limits
- For specific separation runs: e.g. the monitor and curve settings.

Note: Only the settings for the selected components will be shown for strategies where you select the system components.

How to change the default settings

The table below describes the two different ways to change the default system settings.

| Change | Effect |
|---|---|
| To assign a new value to a parameter within a method. | The specific change is valid only until End in the method. After End the parameter returns to its default setting. <i>Note</i> : Only some parameters can be changed in the method. |
| To assign a new value to the system setting. | The new value is valid for all runs and remains until you change the value again or return the setting to its default value. See "How to assign a new value to a system setting" below. Like the default values, the new value can be changed temporarily in a method. |

Note: You must have **System settings** authorization to assign a new value to an actual system setting.

How to assign a new value to a system setting

The table below describes how to assign a new value to a system setting in the **System Control** module.

| Step | Action | |
|------|--|--|
| 1 | Select System:Settings. Result: The Instructions dialog box for the connected system opens. The illustration below shows the dialog box opened with the Alarms group of settings selected. System Alarms Instructions Instructi | |
| 2 | Click the radio button to select one of the following instruction groups: • Alarms • Specials • Monitors • Curves Result: The instructions for the group are displayed. The parameters are listed below each instruction. The title bar of the dialog box shows the selected instruction group. | |
| 3 | Select a parameter from the list. Change the setting value in the Parameters field. Result: The parameter is updated with the new value in the list. | |
| 4 | Click the Set Selected Parameter To Strategy Default Value button to return to the default value (if necessary). Result: The default setting that was defined in the system strategy is restored. Only the selected parameters will be restored. | |
| 5 | Click OK. | |

Limits for monitor signals in methods

If the system strategy allows, limits for certain monitor signals can be set in the method. These limits will only work locally in the method and override the global settings as long as the method is in operation. This feature can be used to set the pH warning threshold to one value during the process operation and another during the system cleaning.

14.2 Alarms

Introduction

This section is a description of the **Alarms** system settings.

Alarms and Warnings

The **Alarms** settings define the upper and lower **Alarm** and **Warning** limits for process monitor signals.

The table below describes the difference between Alarms and Warnings.

| If the signal exceeds | then |
|-----------------------|--|
| the Alarm limits | an alarm sounds |
| | an alarm message is displayed |
| | • the process is paused (i.e. the method execution is suspended and all pumps are stopped) |
| | the alarm is noted in the logbook. |
| | The situation must be acknowledged and corrected before the process can be continued. |
| the Warning limits | a warning message is displayed |
| | the process continues |
| | the warning is noted in the logbook. |

Note: The message text in an **Alarm** dialog box and the corresponding text in the logbook are both color-coded in red. **Warning** texts are color-coded in orange both in the dialog box and in the logbook. The text in the logbook is changed into black when the **Alarm** or **Warning** is acknowledged.

Note: The **Alarms** are not active unless the mode is set to **Enabled**.

Alarms in a network

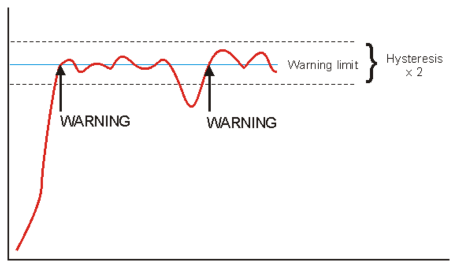
Alarms and warning messages are displayed on all stations with a connection to the concerned system. This is regardless of the activity that is currently performed in UNICORN and regardless of the identity and access rights of the current user.

Alarms and warnings can only be acknowledged from the station that is connected in control mode.

The hysteresis setting

The hysteresis setting (not available for ÄKTAdesign systems) for a warning determines to which extent the signal can oscillate up or down from the warning limit threshold without re-activating a warning.

After the signal has activated a warning, the warning will not be repeated as long as the signal remains within a window defined by the hysteresis setting above and below the warning limit. This prevents repeated warnings from noisy or oscillating signals close to the warning boundary.



Note: Hysteresis is only relevant for warnings, since an alarm puts the system into **Pause** mode at the first alarm.

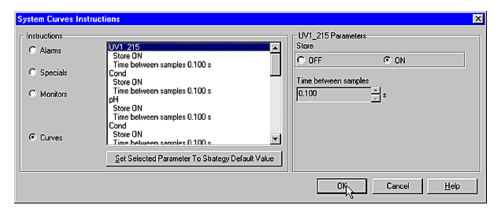
14.3 Curves

Introduction

This section is a short description of the **Curves** system settings.

The Instructions dialog box

The illustration below shows the **Instructions** dialog box with the **Curves** instructions selected.



Curve settings

The curve settings determine which monitor signals that will be stored as curves in the result file. Verify that **Store:ON** is set in the **Instructions** dialog box for all signals that are to be stored.

Warning: If a curve is set to **Store:OFF**, data from the specific monitor cannot be displayed in the curves window during a process run. The data will not be recorded in any way.

Store and Time between samples

The table below describes the function of the two curve settings.

| Setting | Function |
|----------------------|---|
| Store (OFF/ON) | This setting determines whether the curve data is stored or not. |
| Time between samples | This setting determines with which frequency curve data is recorded. It does not affect the reading frequency of the actual monitor. Default value is the shortest possible time between samples. |

15 System maintenance and error reporting

Introduction

This chapter describes the system maintenance and error reporting functions.

In this chapter

This chapter contains these sections:

| Topic | See |
|---------------------------------|-----|
| System maintenance functions | |
| How to generate problem reports | |

15.1 System maintenance functions

Introduction

Some strategies support the capacity to view system information for the components in a chromatography unit. The system information can be used to issue maintenance warnings for the components. This is featured in the strategies for the lab-scale ÄKTAdesign family.

This section describes the system maintenance functions.

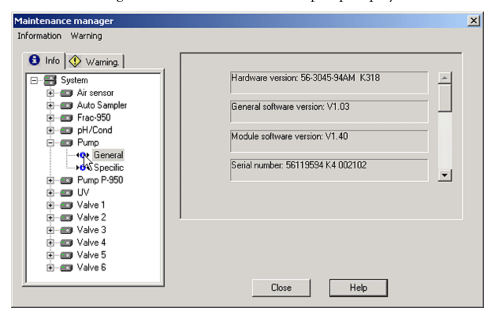
How to open the Maintenance manager

The system maintenance functions are controlled in the **Maintenance manager** dialog box in the **System Control** module.

• Select System:Maintenance.

Result: The **Maintenance manager** dialog box opens with the **Info** tab selected. The connected chromatography system is scanned for its components. After a while the components are displayed.

The illustration below shows the **Maintenance manager** dialog box with the **Info** tab selected and general information about the pump displayed:



How to display component information

Click a component in the list to display the component information.

You can choose two different views:

- **General**, e.g. serial number, version number etc.
- Specific, e.g. how many hours a pump has been run etc.

How to set up a maintenance warning

The table below describes how to set up a maintenance warning.

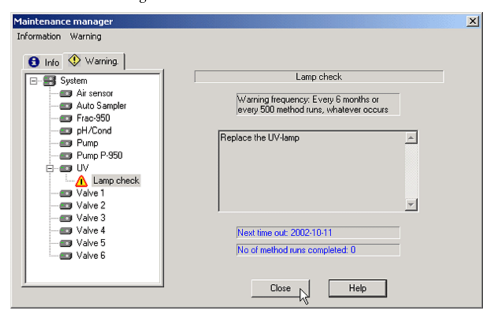
| Step | Action |
|------|---|
| 1 | Click the Warning tab. |
| 2 | Select a component. • Choose Warning:New. |
| | Right-click the component and select the New option on the shortcut menu. |
| 3 | Type the appropriate value in the Periodicity field. Type a warning text in the Pop up text box. Type a name for the warning type in the Name text box. Click the Save button. |
| 4 | Repeat steps 2 and 3 to set up more warnings. |
| 5 | Click the Close button. |

How to view the warning parameters and counters

The component that has been set up for a maintenance warning is marked by an icon and the name of the warning.

• Select the warning to display the parameters.

Counters show the remaining time or number of operations before the next maintenance warning. See the illustration below:



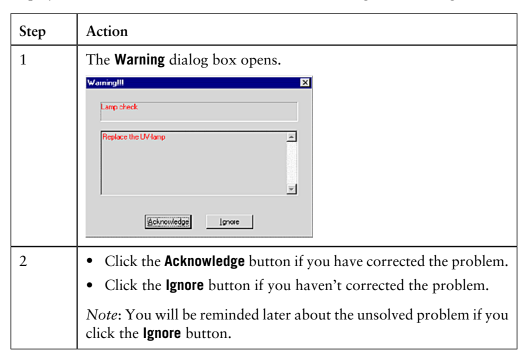
How to reset the counters

The table below describes how to reset the maintenance warning counters.

| Step | Action | |
|------|---|--|
| 1 | Select System:Maintenance in the System Control module to open the Maintenance manager dialog box. Click the Warning tab. | |
| 2 | Select the warning you want to reset on the component list. Choose Warning:Edit. Or Right-click and select Edit on the shortcut menu. Result: The Maintenance manager dialog box changes into edit mode and the text boxes are activated. | |
| 3 | Type new text if necessary. Click the Reset button. Result: The Reset parameters dialog box opens. | |
| | No of method runs completed: 0 Next time out: 2002-10-11 Reset OK Cancel Help | |
| 4 | Click one or both of the Reset buttons to reset the counters. Click OK. | |

How to acknowledge a warning

Once a specific **Periodicity** parameter has been reached, a warning message will be displayed. The table below describes how to acknowledge the warning.



15.2 How to generate problem reports

Introduction

UNICORN contains a Generate Report Wizard for registration of errors or problems that you have detected or that occur during a run. The Generate Report Wizard takes you through the steps to generate your report.

There are two ways of accessing the **Generate Report Wizard**:

- From the UNICORN Manager
- From the System Control.

In this section

This section contains these sub-sections:

| Topic | See |
|---|--------|
| How to generate a report from the UNICORN Manager | 15.2.1 |
| How to generate a report from the System Control | 15.2.2 |

15.2.1 How to generate a report from the UNICORN Manager

Introduction

The **Generate Report Wizard** is used to generate problem reports. This section describes how to generate a problem report from the **UNICORN Manager**.

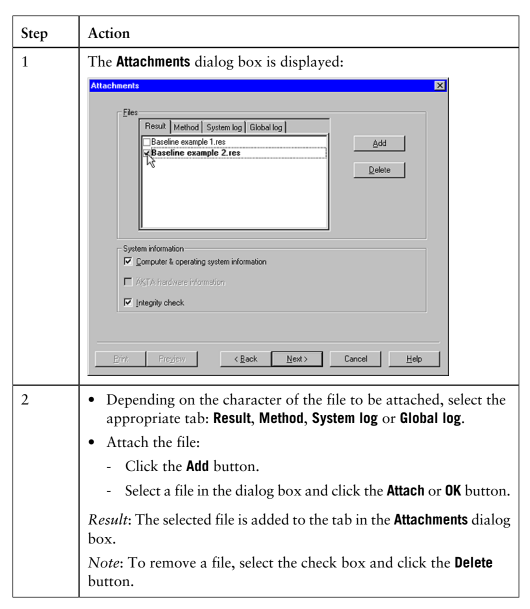
Step 1: How to create the report

The table below describes how to create a report with the **Generate Report Wizard**.

| Step | Action |
|------|--|
| 1 | Select Administration:Create System Report in the UNICORN Manager module. |
| 2 | The first step is a Welcome screen. |
| | • Click the Next button. |
| | <i>Result</i> : The Systems dialog box opens with a list of the available systems for the logged-on user. |
| | • Select a system for which the report is to be generated and click the Next button. |
| | Result: The Description dialog box opens. |
| 3 | Add the following information in the dialog box: |
| | a short description of the problem |
| | the circumstances under which the problem occurs |
| | • the consequences of the problem. |
| | Click the Next button. |
| | Result: The Reproducibility dialog box opens. |
| 4 | Specify whether the problem is reproducible or not. Select one of these alternatives: |
| | • Yes |
| | (Provide a short description in the text box of how the problem can be reproduced.) |
| | • No |
| | Unknown. |
| | Click the Next button to proceed to attach example files (see table below). |

tach a file

Step 2: How to at- You can attach result files, method files and/or log files to the problem report. The table below describes how to attach a file:



| Step | Action | |
|------|---|--|
| 3 | To include more information in the report, select the appropriate check boxes in the System information field. By default, all options are checked. | |
| | Computer & operating system information | |
| | A summary of the computer and operating system information, for example type of processor, processor speed, RAM, hard disk capacity and printer. | |
| | ÄKTA hardware information | |
| | A summary of the specific ÄKTAdesign hardware, for example the instrument and PROM version for every instrument that is connected. | |
| | Integrity check | |
| | When UNICORN is installed a checksum calculation is performed on the stationary files (*.dll and *.exe) for the system. An integrity check means that a new checksum calculation is performed for the same files in their folders. This new calculated value is compared with the checksum value obtained during installation. The results of the comparison are presented in the report and any deviations are included. | |
| | Click the Next button. | |
| | Result: The Generate report dialog box is displayed. | |
| 4 | Proceed to Step 3: How to generate and save the report below. | |

Step 3: How to generate and save the report

The table below describes how to generate and save the report:

| Step | Action |
|------|--|
| 1 | By default, the report is saved in the folder Unicorn\Reports. If you want to save the report at another location, select a folder in the tree structure. |
| 2 | You also have these options: • Click the Preview button to open the report in Notepad. • Click the Print button to print the report without any preview. |
| 3 | Click the Finish button to generate and save the report. |

15.2.2 How to generate a report from the System Control

Introduction

The **Generate Report Wizard** is used to generate problem reports. When an error message appears in **System Control**, you can activate the report wizard from the error message dialog box. The **Generate Report Wizard** can also be activated anytime if you choose **System:Report**.

Step 1: How to create the report

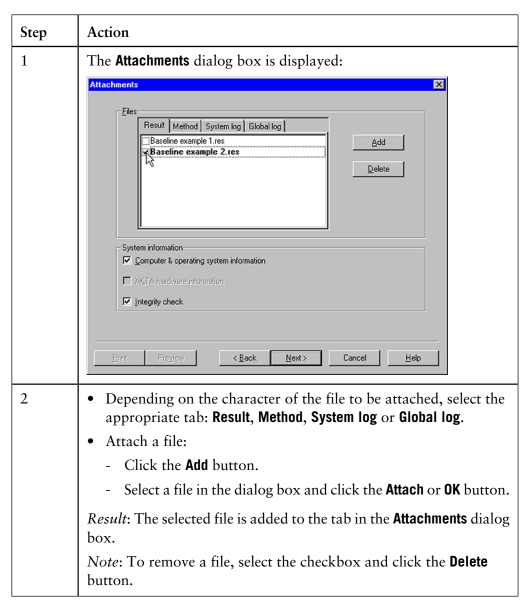
When an error message appears in **System Control**, follow the instructions in this table to activate the **Generate Report Wizard** and create a report:

| Step | Action | |
|------|---|--|
| 1 | • Click the Report button in the error message dialog box. | |
| | or | |
| | Choose System:Report. | |
| 2 | The first step is a Welcome screen. | |
| | Click the Next button. | |
| | <i>Result</i> : The Description dialog box is displayed and shows a list of the problems/errors that have occurred. All the problems/errors that have occurred, together with help texts, are automatically recorded and included in the report. | |
| | • If you select a specific error in the Description dialog box, the appropriate help text is shown in the error message box. | |
| 3 | Add the following information in the Description dialog box: | |
| | A short description of the problem. | |
| | The circumstances under which the problem occurs. | |
| | The consequences of the problem. | |
| | Click the Next button. | |
| | Result: The Reproducibility dialog box opens. | |
| 4 | Specify whether the problem is reproducible or not. Select one of these alternatives: | |
| | • Yes | |
| | (Provide a short description in the text box of how the problem can be reproduced.) | |
| | • No | |
| | Unknown. | |
| | Click the Next button to proceed to attach example files (see table below). | |

Step 2: How to attach a file

You can attach method files and/or log files to the problem report.

The table below describes how to attach a file:



| Step | Action | |
|------|--|--|
| 3 | To include more information in the report, select the appropriate check boxes in the System information field. By default, all options are checked. | |
| | Computer & operating system information | |
| | A summary of the computer and operating system information, for example type of processor, processor speed, RAM, hard disk capacity and printer. | |
| | ÄKTA hardware information | |
| | A summary of the specific ÄKTAdesign hardware, for example the instrument and PROM version for every instrument that is connected. | |
| | Integrity check | |
| | When UNICORN is installed a checksum calculation is performed on the stationary files (*.dll and *.exe) for the system. An integrity check means that a new checksum calculation is performed for the same files in their folders. This new calculated value is compared with the checksum value obtained during installation. The results of the comparison are presented in the report and any deviations are included. • Click the Next button. | |
| | | |
| | Result: The Generate report dialog box is displayed. | |
| 4 | Go to step 3 below. | |

Step 3: How to generate and save the report

The table below describes how to generate and save the report.

| Step | Action |
|------|--|
| 1 | By default, the report is saved in the folder: Unicorn\Reports. If you want to save the report in another location, select a folder in the tree structure. |
| 2 | You also have these options: Click the Preview button to open the report in Notepad. Click the Print button to print the report without any preview. |
| 3 | Click the Finish button to generate and save the report. |

A Troubleshooting

Introduction

This appendix describes different problems which may arise in UNICORN and how to solve the problems.

In this appendix

This appendix contains these sections:

| Topic | See |
|-------------------------------------|-----|
| Logon | A.1 |
| UNICORN access | A.2 |
| Methods and method runs | A.3 |
| Evaluation | |
| ÄKTAdesign system specific problems | |

A.1 Logon

In this section

This section describes how to solve the following log on problems:

- Unable to log on to UNICORN
- Error message "Strategy file error".

Unable to log on to UNICORN

The table below describes some log on problems and their solutions:

| Problem description | Solution |
|--|---|
| You have forgotten your password. | Ask the system administrator to supply a new password. |
| You cannot log on although you use your correct username and password. Reason: The file USERS30.MPM in the folder \UNICORN\SERVER\FIL could be corrupt. | Restore the file USERS30.MPM from the latest back-up copy or reinstall the default user. |
| No user names: Remote station Both these conditions must apply: The User name drop-down box in the Logon dialog box is empty. You are trying to log on from a remote station in a network installation. | Make sure that the computer is logged on to the network before you start UNICORN. Note: A remote station accesses the user list directly from the network server. |
| No user names: Local station The user list on a local station in a network installation is not up to date. | Make sure that the computer is logged on to the network before starting UNICORN. Note: The user list is stored locally on a local station, and is updated automatically from the network server if the computer is logged on to the network. |

Error message "Strategy file error"

The table below describes some problems and their solutions:

| Problem description | Solution |
|---|---|
| Stand-alone installation If you receive the error message "Strategy file error" in a <i>stand-alone</i> installation, the strategy file is probably corrupt. | Reinstall the strategy as described in the Administration and technical manual "Install selected software components after the initial installation". |
| Network installation In a network installation, the error message "Strategy file error" may appear if you try to create a method for a system not physically connected to the computer. | Make sure that the computer is logged on to the network before UNICORN is started, so that the strategy file on the server disk is accessible. |

A.2 UNICORN access

In this section

This section describes how to solve the following UNICORN access problems:

- Unable to access certain UNICORN functions
- Connection problems
 - Connections are not available
 - System is not available
 - Error message in a network installation
 - You cannot control the system
- Run data Connection in System Control displays a "NO [1]", "NO [2]" or "NO [3]".

Unable to access certain UNICORN functions

The table below describes an access problem and its solution:

| Problem description | Solution |
|---|---|
| UNICORN functions to which you do not have access appear grey in the menu and cannot be used. | Choose Administration:User Setup in the UNICORN Manager to change the user profile. |
| | <i>Note</i> : Contact the system administrator if you are not authorized to change your user profile. |

Connection prob- The table below describes some connection problems and their solutions:

| Problem description | Solution |
|---|--|
| The connections are not available. | Check the connection between the PC and the chromatography system. Check that the power to the chromatography system is turned on. |
| The connections are not available even though the connection between PC and chromatography system appears to be correct the power is turned on. | Quit UNICORN. Shut down and switch off the computer. Switch off the chromatography system. Restart the entire system. |

| Problem description | Solution |
|--|---|
| A system is not available when you attempt to establish a connection. | Check that you have access rights to the system. Access rights are not automatically assigned for a newly defined system. |
| You receive the error message "Cannot connect to system" in a network installation. | Check that the local computer to which the system is connected is turned on and logged on to the network. Check that the computer where you try to establish a connection is logged on to the network. Check that the limit of 8 connections to the system has not been exceeded. |
| You can establish a connection but cannot control the system, that is the Manual menu commands in the System Control are grey. | Check that no other user has a control mode connection. Check that you have sufficient access rights to control the system manually. |

Note: The **Method Wizard** can be used on a local system even if the network connection is not established.

The Connection field in System Control displays a "NO [X]"

The table below describes some connection problems and their solutions:

| Problem Description | Solution |
|---|---|
| The Connection field in the Run data pane in System Control says "NO [1]" or "NO [2]". | • Check that the UNICORN PC Control board is configured according to the settings made during the installation of the program. The same Control unit number, Address and IRQ must be set at the Control board, see the Administration and technical manual "Hardware installation". |
| | • The communication may also fail if there is a conflict between the UNICORN PC Control board configurations and other boards in the PC. If so, select a free Address and a free IRQ during UNICORN installation and at the Control Board, see the Administration and technical manual "Hardware installation". |

| Problem Description | Solution |
|--|---|
| The Connection field in the Run data pane in | • Choose Administration: System Setup in the UNICORN Manager. |
| System Control says "NO [3]". | - Select the system with problems in the dialog box and click the Edit button. |
| | - Check that the strategy, computer name and the control number are correct according to the installation at the local station which is physically connected to the system. See the Administration and technical manual "System definitions". |
| | If you connect remotely to a system |
| | - check that the local station which is physically connected to the system is turned on |
| | - check that the network is functioning at both the remote and the local station. |
| | • Check that the limit of eight connections to the system has not been exceeded. |

A.3 Methods and method runs

In this section

This section describes how to solve the following method and method run problems:

- Cannot perform Quit or Logoff
- Monitor signals do not appear in the Curves pane in **System Control**
- Error message "Couldn't create result file... Destination path could not be found"
- The Method-System Connection dialog box keeps appearing
- The Method Editor window does not fit on the screen
- There are red instructions in a method
- After Windows logout and login you cannot get a system connection
- The Print screen command does not send a copy of the screen to the printer

Cannot perform Quit or Logoff

The table below describes a problems and its solutions:

| Problem description | Solution |
|---|---|
| You are unable to perform Quit or Logoff from UNICORN for a connection. | You might be running a Scouting method or a MethodQueue . These functions require a control mode connection in order to start subsequent cycles correctly. Action: Stop the Scouting method or MethodQueue before you quit or log off. |

Monitor signals do not appear in the Curves panel in System Control

The table below describes a problem and its solution:

| Problem description | Solution |
|--|---|
| Monitor signals do not appear in the Curves pane in System Control . | Choose System:Settings in System Control |
| | Result: The System Instructions dialog box opens. |
| | • Choose the Curves group in the Instructions field. |
| | • Set the Store option to ON . |
| | Store C OFF |
| | Signals for which Store is set to ON can be selected from the View:Properties:Curves dialog box in System Control . |

Error message "Couldn't create result file... Destination path could not be found"

The table below describes a problem and its solution:

| Problem description | Solution |
|--|--|
| If you receive the error message "Couldn't create result file Destination path could not be found" at the end of a method, the local computer was unable to access the folder specified in the result file path. | This may happen if the specified folder is on the network server and network communication has been lost. The result file is saved in the Failed folder on the local station. |

The Method-System Connection dialog box keeps appearing

The table below describes a problem and its solution:

| Problem description | Solution |
|---|--|
| If the Method-System Connection dialog box keeps appearing you have some method(s) which is not connected to a system. | Connect the method(s) to the appropriate system. |
| Reason: Most likely you have imported some method(s) with the command File:Copy from External in the UNICORN Manager. | |

The Method Editor window does not fit on the screen

The Method Edit- The table below describes a problem and its solution:

| Problem description | Solution |
|---|---|
| The Method Editor window does not fit the screen and has scroll bars. Reason: The incorrect font size might be installed. | The display screen resolution may be set to "1024x768x65536" with "Large fonts". You need to install the "Small fonts". This requires that you have the Windows 2000 or Windows XP CD-ROM that was shipped with your Compaq computer. Insert the CD-ROM and follow the directions on the screen. |

Note: Always install the latest service pack after you have installed something from the Windows 2000/XP CD-ROM.

There are red instructions in a method

The table below describes some solutions to syntax error problems:

Problem description

Red instructions (instructions with a red dot) in a method are syntax errors and may be due to the following:

- The method was connected to the wrong system, that is the strategy of the system is incompatible with the method.
- The method instructions do not correspond to the components you have chosen for your system. Check your system components under Administration:System Setup in the **UNICORN** Manager.
- The **Copy** function was used instead of Copy from external when a method was imported from a diskette.
- The wrong system may have been selected in the Save As dialog box in the **Method Editor**.
- You may also have templates not intended for your system, which might be the case for custom designed systems.
- The systems strategy has been updated with a new strategy that differs in the instruction set.

Solution

There are several actions that you can

- Check that the method has been connected to the correct system in either of these ways:
 - in the System Method Connection dialog box when you use the Copy from external dialog box
 - in the Save As dialog box in Method Editor.
- If the system is custom designed, open the Method Editor, select the red instruction and either delete it or replace it with a corresponding instruction (if available) from the **Instruction box**. Repeat this for all red instructions before saving the method.

After Windows logout and login you cannot get a system connection

The table below describes a system connection problem. This applies only to local systems, not remote systems:

| Problem description | Solution |
|--|---|
| You have logged out of Windows 2000 and then logged in again, but you cannot get a system connection in UNICORN. | Restart the computer in order to obtain a system connection in UNICORN. |
| Reason: If you shut down Windows 2000 with the command Start:Shutdown:Close all programs and log in as a different user, you will not be able to obtain a System Control connection in UNICORN the next time you or another user logs on. This is because the described shutdown procedure automatically shuts down a number of processes, including those needed for system connection. The services are only started when the computer is booted up. | |

Print screen does not send a copy of the screen to the printer

The table below describes how to solve a printing problem:

| Problem description | Solution |
|--|---|
| The Print screen command only makes a copy of the screen to the clipboard and not to the default printer. | If you want to print the view on the screen, press the <print scrn=""></print> key and paste the image from the clipboard into an appropriate program, such as Microsoft® Paint, and then print out the image. |

Evaluation A.4

In this section

This section describes how to solve the following evaluation problems:

- Incorrect date and time in the result file
- Evaluation procedure aborts

Incorrect date and time in the result file

The table below describes a problem and its solution:

| Problem description | Solution |
|--|--|
| The result file shows incorrect date and time. | Check the system clock setting. The date and time recorded in the result file are taken from the PC system clock setting. |

Evaluation proced The table below describes a problem and its solution: **ure aborts**

| Problem description | Solution |
|----------------------------------|---|
| The evaluation procedure aborts. | Instructions in an evaluation procedure refer to curves by identification number irrespective of the curve names. Make sure that the curves processed when the procedure is executed are compatible with those processed when it was recorded. An evaluation procedure aborts if you try to store resulting curves at the position of an original raw data curve. |

A.5 ÄKTAdesign system specific problems

In this section

This section describes how to solve the following problems:

- Connected to a system but no system contact
- Flow scheme not displayed properly

Connected to a system but no system contact

The table below describes a problem and its solution:

| Problem description | Solution | |
|--|--|--|
| You are connected to a system but have no system contact. Indications: In the System Control, | Check that the system is turned on.Check that all the cable connections are intact. | |
| • the option Connection in the Run data pane says "Yes", | If the above actions do not help, try to restart both the computer | |
| • the option Instruments says "Scanning", | and the system. | |
| • there is no contact with the system after a period of waiting. | | |

Flow scheme not displayed properly

The table below describes a problem and its solution:

| Problem description | Solution |
|--|--|
| The flow scheme is not displayed properly. | Choose Settings:Control Panel: Display:Settings in the Windows Start menu to check that you have selected 65536 colors. |

Evaluation functions and instructions В

Introduction

This appendix describes the functions that are implemented in the **Evaluation** module.

In this appendix

This appendix contains these sections:

| Topic | See |
|------------------------------|-----|
| Smoothing algorithms | B.1 |
| Baseline calculation theory | B.2 |
| Peak table column components | B.3 |
| Procedure instructions | B.4 |

B.1 Smoothing algorithms

Introduction

This section describes how the smoothing functions are calculated. Choose **Operations:Smooth** in the **Evaluation** module to view and edit the options.

Moving Average

The table below describes the process when the **Moving Average** smoothing algorithm is used.

| Stage | Description |
|-------|---|
| 1 | For each data point in the source curve, the processed curve is calculated as the average of the data points within a window centered on the source data point. |
| | • The width of the window is determined by the parameter value, expressed as number of data points. |
| 2 | When the source point is less than half the window size from the beginning of the end of the curve, the average is calculated symmetrically round the source point over as many data points as possible. If you increase the window width, the smoothing effect is also increased. |

Note: The filter algorithm only accepts odd integer parameter values between 1 and 151. If an even number has been given, it is incremented by one (1).

Autoregressive

The table below describes the process when the **Autoregressive** smoothing algorithm is used:

| Stage | Description |
|-------|---|
| 1 | The first data point in the source curve is copied to the processed curve. |
| 2 | For each subsequent data point, the previous processed point is multiplied with the parameter value and added to the current source data point. |

| Stage | Description |
|-------|---|
| 3 | The result is then divided by the parameter value plus 1 according to the following formulae: |
| | $t_1 = S_1$ |
| | $t_{n} = \frac{(p * t_{n-1} + S_{n})}{(p+1)}$ |
| | Where: |
| | t_n = current processed point. |
| | t_{n-1} = previous processed point. |
| | S_n = current source point. |
| | p = smoothing parameter value. |
| | <i>Note</i> : If you increase the parameter value, the smoothing effect is also increased. |

Note: The filter algorithm only accepts integer parameter values between 1 and 25.

Median

The table below describes the process when the **Median** smoothing algorithm is used.

| Stage | Description |
|-------|---|
| 1 | For each data point in the source curve, the processed curve is calculated as the median of the data points within a window centered on the source data point. |
| | • The width of the window is determined by the parameter value, expressed as number of data points. |
| 2 | When the source point is less than half the window size from the beginning of the end of the curve, the median is calculated symmetrically round the source point over as many data points as possible. |
| | • If you increase the window width, the smoothing effect is also increased. |
| | To completely remove a noise spike, the window width should in effect be slightly more than twice the width of the spike. |

Note: The filter algorithm only accepts odd integer parameter values between 1 and 151. If an even number has been given, it is incremented by one.

Savitzky-Golay

The table below describes the process when the **Savitzky-Golay** smoothing algorithm is used.

| Stage | Description |
|-------|---|
| 1 | The algorithm is based on performing a least squares linear regression fit of a polynominal of degree k over at least k+1 data points around each point in the curve to smoothen the data. The derivate is the derivate of the fitted polynominal at each point. The calculation uses a convolution formalism to calculate 1st through 9th derivatives. |
| 2 | The calculation is performed with the data in low X to high X order. If the input trace goes from low to high, it is reversed for the calculation and is re-reversed afterwards. |

Note: See Gorry, Peter A, General Least-Squares Smoothing and Differentation by the Convolution (Savitsky-Golay) Method (Analytical Chemistry 1990, Volume 62, 570-573) for more information on the Savitzky-Golay algorithm.

B.2 Baseline calculation theory

Overall process

The table below describes the overall process of a baseline calculation.

| Stage | Description |
|-------|------------------------------------|
| 1 | The baseline segments are defined. |
| 2 | The baseline points are selected. |
| 3 | The baseline is drawn. |

Baseline segment definition

Baseline parameters are used to find the baseline segments. The default values for the parameters are determined from the source curve. The baseline segments are found by different parameters that are based on the type of algorithm that is

Note: The parameters can be displayed in the **Evaluation** module if you choose Integrate: Calculate baseline function. You can also click the Baseline settings button in the Integrate: Peak integrate dialog box.

Morphological algorithm

The **Morphological** algorithm searches for all parts of the source curve where:

- The curve parts come into contact at both ends of a horizontal line of the length defined in the **Structure width** parameter. The default value of this parameter is based on the widest detected peak in the curve. The horizontal line is moved along the curve up the peak until it reaches the contact points. The curve parts below the horizontal line and the line will now form a "curve" with a plateau. The center point in the plateau formed by the horizontal line will be the data point for the baseline.
- The data points fulfil the **Minimum distance between data points**. This parameter reduces the total number of data points that are created from a curve.

Classic algorithm

The **Classic** algorithm searches for all parts of the source curve where:

- The curve parts are longer than the **Shortest baseline segment**. This parameter determines the minimum length for a part of the source curve to be considered a possible baseline segment.
- The curve has no point outside the **Noise window**. The noise window is defined as a rectangular corridor parallel to the slope of the curve and centered on the first and last points within the currently inspected segment.
- The slope is less than the **Slope limit**. This limits the maximum slope of the baseline to differentiate baseline segments from peaks.
- The curve parts are lower than the **Max baseline level**. This parameter determines the highest acceptable signal level for the baseline.

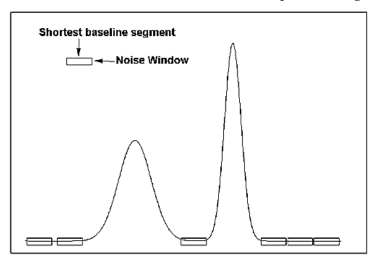
Baseline parameters

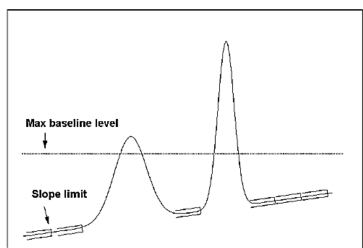
The baseline parameters can be illustrated as a rectangular box that the source curve has to fit into in order to be identified as a baseline segment, where:

- The length of the box corresponds to the Shortest baseline segment.
- The height of the box corresponds to the maximum level of noise on the baseline segments. This is referred to as the **Noise window**.
- The box is allowed to be tilted with a maximum slope corresponding to the **Slope limit**.
- The box is not allowed to move up above the **Max baseline level**.

Baseline parameters - illustration

The illustrations below shows the baseline parameters graphically.





Baseline segment identification

The table below describes the baseline segment identification process:

| Stage | Description |
|-------|---|
| 1 | The box is virtually moved along the source curve in steps of one third of the Shortest baseline segment length to look for baseline segments. |
| 2 | A baseline segment is found whenever the currently examined part of the source curve fits completely within the box. |
| 3 | The found baseline segments are joined by connecting adjacent segments, provided that the slope of the joining lines does not exceed the Slope limit . |

Baseline points (Classic algorithm)

When the baseline segments have been defined and joined, they are replaced by baseline points at the start and end of each segment. The line between these is also filled with points.

Note: The baseline points are shown as green squares in the **Integrate:Edit baseline** function of the **Evaluation** module.

Baseline drawing

The baseline points are used to create the baseline curve using a spline interpolation. The spline function ensures that the baseline curve is guided by the baseline points. However, the curve does not necessarily pass through the baseline points. The baseline will be a smoothly curved function passing close to or through the points.

To reduce the effect of noise at the peak integration, the created baseline is forced equal to the source curve in every position where the difference between the baseline and the source curve is small enough. Choose **Integrate:Calculate Baseline**. If the **Accept negative peaks** option is off, the baseline will be forced down to the level of the source curve whenever the created baseline goes above the source curve.

How to measure the baseline segment (Classic algorithm)

You can try to measure the **Shortest baseline segment** length directly on your chromatogram. The table below describes how to do this:

| Step | Action |
|------|---|
| 1 | Locate the shortest segment of the curve that you consider a part of the baseline. |
| 2 | Use the marker box on the chromatogram to measure the length of the segment. |
| 3 | Choose Integrate:Calculate Baseline and insert this value as the Shortest baseline segment value. |

How to measure noise level (Classic algorithm)

Curve coordinates can also be used to measure noise levels on the source curve. The table below describes how to do this:

| Step | Action |
|------|--|
| 1 | Use the Z00m function to focus on a part of the curve that is representative for the baseline noise. |
| 2 | Select an appropriate Y-axis scale. |
| 3 | Measure the Y-axis coordinates. |
| 4 | Calculate the noise range as the difference between the max. and min. values. Add an extra 20%. Choose Integrate:Calculate Baseline and insert this value as the Noise window value. |

How to measure the slope limit (Classic algorithm)

The table below describes how to measure the slope at any part of the curve.

| Stage | Description |
|-------|---|
| 1 | Select Operations:Differentiate in the Evaluation module. *Result: The Differentiate dialog box opens. |
| 2 | Select the desired source curve. Select the First order calculation option. Click OK. Result: The differentiated curve will appear in the active chromatogram. |
| 3 | Select an appropriate Y-axis scale, right-click and select Marker to measure the Y-axis values for the differentiated curve with the curve coordinates function. Result: The Y-axis value is interpreted as the UV curve slope at the selected retention point. |
| 4 | Determine the highest slope value of the baseline (non-peak) part of the curve. Add 10%. Select Integrate:Calculate Baseline and use this value as the Slope limit. |

Note: If the differentiated curve is very noisy, it can be filtered with a light **Moving** average filter in the **Operations:Smooth** function.

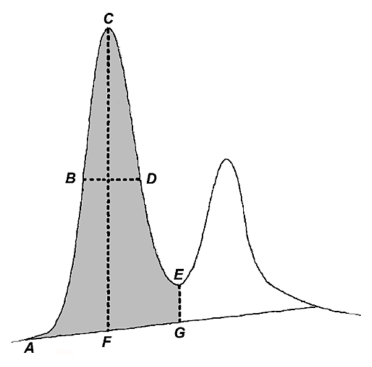
B.3 Peak table column components

Introduction

This section contains a list of peak parameters with explanations and calculation formulae when applicable.

Peak parameters - illustration

The diagram below illustrates the peak parameters. See the parameter list below for explanations.



Peak parameter descriptions

The list below contains descriptions of the peak parameters.

| Parameter | Description |
|-----------|--|
| Amount | Values calculated by the Analysis module. (Only available if the Quantitation module is installed.) |
| Area | Calculated as the area between the curve and baseline, between the peak start and peak end, time or volume base. (Gray area in the diagram above.) |
| Asymmetry | Peak asymmetry (indicator of column packing). See definition below this table. |

| Parameter | Description |
|-----------------------|---|
| Baseline height | Baseline amplitude at peak start, peak maximum and peak end. (A, F and G in the diagram above.) |
| Capacity factor | The capacity factor will only be calculated when the chromatogram is in volume base. The total liquid volume, Vt, must be entered in the Integrate dialog box for this parameter to be calculated. See definition below this table. |
| Concentration | Values calculated by the Analysis module. (Only available if the Quantitation module is installed.) |
| Fraction tube id | Fraction number at peak start, peak maximum and peak end. |
| Height | Maximum amplitude above the baseline. (C-F in the diagram above) |
| Kav | Gel phase distribution constant in gel filtration. Kav will only be calculated when a gel filtration column was used and when the chromatogram is in volume base. The void volume, V0, must be entered in the Integrate dialog box for this parameter to be calculated. See definition below this table. |
| Molecular size | Values calculated by the Analysis module. (Only available if the Quantitation module is installed.) |
| Plate height (HETP) | Height equivalent to theoretical plate and plates/meter. The column height must be entered in the Integrate dialog box for this parameter to be calculated. See definition below this table. |
| Peak endpoint heights | Amplitude above the baseline at left (A in the diagram above) and right peak limits (E-G in the diagram above). |

| Parameter | Description | |
|----------------------------|---|--|
| Peak endpoint retention | Retention value at peak start and peak end, time or volume base. (A and G in the diagram above.) | |
| Peak name | Name of the peak. | |
| Percent of total area | Peak area as a percent of the total area under the curve above the baseline. Time or volume base. Note: This value can differ in time and volume base if the flow rate is not constant throughout the method. | |
| Percent of total peak area | Peak area as a percent of the sum of all integrated peaks. Note: This value can differ in time and volume base if the flow rate is not constant throughout the method. | |
| Resolution | Peak resolution. See definition below this table. | |
| Retention | Retention at the peak maximum, time or volume base. (C in the diagram above.) | |
| Sigma | Standard deviation for a Gaussian-shaped peak. See definition below this table. | |
| Type of peak limits | Identifies the criteria for peak start and peak end as either the baseline intersection or dropline to the baseline or skim line. | |
| Width | Difference in retention between the peak end and peak start, time or volume base. (G-A in the diagram above.) | |
| Width at half height | Calculated by taking the maximum height of the peak above the baseline, then determining the peak width at half this value above the baseline. Time or volume base. (B-D in the diagram above, where BD bisects CF.) | |

Note: In the **Options** dialog box in the **UNICORN Manager** you can select if negative retentions should be displayed or not. The default selection is that negative retention is not displayed.

Sigma formula

The formula below is used to calculate **Sigma**.

Sigma =
$$\sqrt{\frac{\sum_{i=1}^{n} \left(y_i \left(x_i - x_{ymax}\right)^2\right)}{A_{peak}}}$$

Where:

- *n* is the number of data points.
- *x* is the volume or time value.
- x_{ymax} is the volume or time value at the maximum amplitude value.
- A_{peak} is the area of the peak.

Note: The peak width for a Gaussian peak is (4 x Sigma).

Peak resolution algorithms

The peak resolution is calculated with one of the following three algorithms:

- 1. $(V_{R2} V_{R1}) / ((W_{b2} + W_{b1}) / 2)$
- 2. $(V_{R2} V_{R1}) / ((Sigma_2 + Sigma_1) \times 2)$
- 3. $((V_{R2} V_{R1}) / (2 \times (W_{h2} + W_{h1}))) / 2.354$

Where:

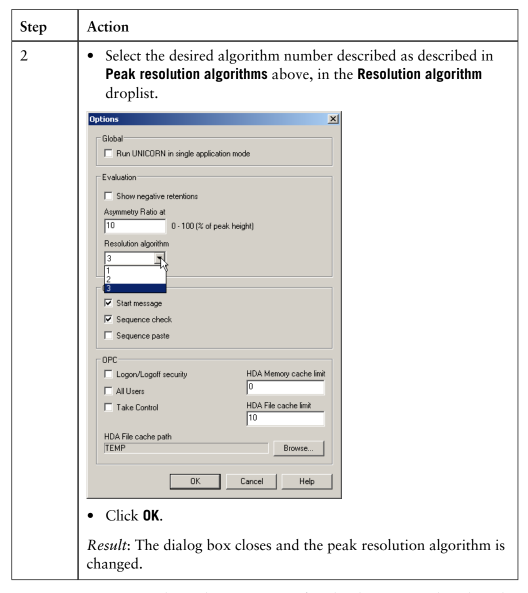
- V_{R1}, W_{b1}, Sigma₁ and W_{h1} are the retention, width, Sigma and width at half height of the previous peak.
- V_{R2}, W_{b2}, Sigma₂ and W_{h2} are the retention, width, Sigma and width at half height of the current peak.

Note: The **Resolution algorithm** variable in the **Options** dialog box in the **UNICORN Manager** determines which of the three algorithms is used. If this variable has the value 1, 2 or 3, then the algorithm with the corresponding number in the list above is used. The default value is 3.

How to change the peak resolution algorithm

The table below describes how to change the peak resolution algorithm in the **UNICORN Manager**.

| Step | Action |
|------|--|
| 1 | Choose the Administration:Options menu item. |
| | Result: The Options dialog box opens. |



Note: You must repeat the peak integrations after the change to update the values based on the new algorithm.

Capacity factor formula

The formula below is used to calculate the **Capacity factor**.

$$k^1 = \frac{V_R - V_t}{V_t}$$

Where:

- V_R = retention volume.
- V_t = total liquid volume.

Kav formula

The formula below is used to calculate **Kav**.

$$k_{av} = \frac{V_R - V_0}{V_C - V_0}$$

Where:

- V_R = retention volume.
- V_0 = void volume.
- V_C = column volume.

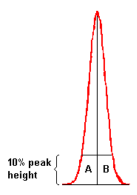
Asymmetry formula

The formula below is used to calculate the **Asymmetry**.

Asymmetry = B / A

Where:

- A is a partial peak width, measured at a percentage of the peak height, for the leading part of the peak.
- B is a partial peak width, measured at a percentage of the peak height, for the tailing part of the peak.



How to change the Asymmetry Ratio

The **Asymmetry Ratio** is selected in the **Options** dialog box in the **UNICORN Manager**. The table below describes how to select a value:

| Step | Action |
|------|---|
| 1 | Choose the Administration:Options menu item. |
| | Result: The Options dialog box opens. |
| 2 | Type a ratio value in the Asymmetry Ratio at text box. Click OK. |
| | Result: The ratio value is changed and the dialog box closes. |

Note: You must repeat the peak integrations after the change to update the values based on the new asymmetry ratio. The default ratio is 10%.

HETP formula

The formula below is used to calculate the **HETP** value.

HETP = L/N

 $N = 5.54 \times (V_R/w_h)^2$ assuming a Gaussian peak.

Where:

- N = no. of theoretical plates.
- L = bed height in cm.
- V_R = peak retention (elution) volume or time.
- w_h = peak width at half height expressed in the same units as V_R .

B.4 Procedure instructions

Introduction

This section contains lists of procedure instructions with descriptions. These instructions are used in the **Procedure Editor**. Choose **Procedures:Edit:New** in the **Evaluation** module to view the **Instruction** list.

Curve operation

The table below contains a list of instructions for curve operations.

| Instruction | Description |
|-------------|--|
| ADD | Adds two curves to produce a third curve, which is the sum of the two curves. The two source curves must have the same Y-axis unit and not be fraction or injection curves, or else a run time error will occur. |
| AMP_MUL | Multiplies the amplitude of the source curve by the multiplication factor and stores the result in the target curve position. |
| AMP_SHIFT | Shifts the amplitude of the source curve by the shift factor and stores the result in the target curve position. |
| CLEAR | Clears the specified curve from the working memory of the computer. |
| COPY | Copies the source curve to the target curve position. |
| CUT | Cuts out the part of the source curve between the Left and Right limits and stores the result in the target curve position. |
| DERIVATE | Differentiates the source curve (first or second order) and stores the result in the target curve position. The Y-axis of the target curve position will be a normalized scale without unit. |

| Instruction | Description |
|--------------------|---|
| DIV | Divides two curves to produce a third curve, which is the quotient of the two curves. The two source curves can have any Y-axis unit. The Y-axis of the target curve position will be a normalized scale without unit. |
| HISTOGRAM | Creates a histogram from any non-fraction curve (source curve 1) and a fraction curve (source curve 2_frac), and stores the result in the target curve position. If source curve 2 is not a fraction curve a run time error will occur. The Y-axis of the target curve position will be the same as that of the first source curve. |
| INTEGRATE | Performs a mathematical integration of the source curve and stores the result in a Result curve. This instruction is not the same as Peak integrate , which performs a real peak integration. |
| POOL_FRACTIONS | Pools fractions from the source curve and stores the result in the target curve position. The fractions are pooled from the first selected fraction to the last selected fraction. If the source curve is not a fraction curve, or First or Last is not an existing identification, a run time error will occur. |
| RET_MUL | Multiplies the retention of the source curve by the Multiplication factor and stores the result in the target curve position. |
| RET_SHIFT | Shifts the retention of the source curve by the Shift factor and stores the result in the target curve position. |
| SIMULATE_PEAK_FRAC | Simulates Peak Fractionation . |

| Instruction | Description |
|---------------|---|
| SMOOTH_AR | Smooths the source curve with an autoregressive filter and stores the result in the target curve position. The Filter parameter decides the strength of the filter. |
| SM00TH_MA | Smooths the source curve with a moving average filter and stores the result in the Resulting Curve . The Filter width parameter decides how many samples wide the filter is. |
| SMOOTH_MEDIAN | Smooths the source curve with a median filter and stores the result in target curve position. The Filter width parameter decides how many samples wide the filter is. |
| SMOOTH_SG | Smooths the curve with the Savitzky-Golay algorithm. |
| SUB | Subtracts two curves to produce a third curve, which is the difference of the two curves. The two source curves must have the same Y-axis unit and not be fraction or injection curves. |
| TDIV | Divides two curves to produce a third curve, which is the quotient of the two curves. The two source curves can have any Y-axis unit. The threshold values are used to avoid division of numbers close to zero. At those points where source curve 1 has an amplitude less than Threshold1 , or the source curve 2 has an amplitude less than Threshold2 , the result of the division is defined to be 1.0. |

Integration

The table below contains a list of instructions for integration.

| Instruction | Description |
|--------------------------|---|
| CALCULATE_BASELINE | Calculates a baseline from the source curve. The baseline is stored in the target curve position. DEFAULT can be selected in the Baseline parameters, which will then calculate default baseline parameters for each new curve. |
| CALCULATE_BASELINE_MORPH | Calculates a baseline from the curve crvSrc using a morphological method. DEFAULT can be selected in the Baseline parameters, which will then calculate default baseline parameters for each new curve. The baseline is stored in curve crvDst. |
| CLEAR_PEAKTABLE | Clears the peak table in Peak table source from the computer memory. |
| COPY_PEAKTABLE | Copies a peak table from Peak table source to Resulting peak table. |
| NEGATIVE_PEAKS | Controls the baseline behavior in subsequent baseline calculations. If ONOFF is ON then the baseline can be drawn above the curve and negative peaks can be detected by PEAK_INTEGRATE . If ONOFF is OFF then the baseline is never drawn above the curve. |
| PEAK_INTEGRATE | Performs a peak integration on the source curve and stores the resulting peak table in Resulting peak table . It is assumed that the baseline is subtracted. |
| PEAK_WINDOW | Specifies which part of the source curve that will be integrated. Peaks between retention Left limit and Right limit will be detected if the ONOFF parameter is set to ON . If ONOFF is set to OFF , the whole curve will be used for integration. |

| Instruction | Description |
|-----------------------|--|
| REJECT_PEAKS | Any combination of conditions is allowed. If all parameters are OFF then every detected peak is included in the peak table. |
| SET_COLUMN_HEIGHT | Sets the column height for the peak integration calculation of the HETP value. The Column height parameter is the height of the column in centimetres. If Column height is OFF then the HETP value is not calculated for the following integrations. |
| SET_COLUMN_VO | Sets void volume for Kav peak integration calculation. |
| SET_COLUMN_VT | Sets the total liquid volume for peak integration calculation of the capacity factor. |
| SET_SKIM_SIZE_RATIO | Sets the Skim size ratio to be used in the following peak integration(s). |
| WINDOW_PEAK_INTEGRATE | Integrates the curve within the peak window. All curve parts outside the peak window remain unchanged. |

File operation

The table below contains a list of instructions for file operations.

| Instruction | Description |
|--------------|---|
| CURVE_OPEN | Opens the curve specified in the Result file defined in File name and stores it in target curve position. If "*" is entered as File name the current result file will be used. The File name parameter may include a path from the users root folder. |
| IMPORT_CURVE | Imports a curve to the current chromatogram from another chromatogram (in the current file) and stores it in the target curve position. |

| Instruction | Description |
|------------------|---|
| IMPORT_PEAKTABLE | Imports a peak table to the current chromatogram from another chromatogram (in the current file) and stores it in the target curve position. |
| PEAKTABLE_OPEN | Opens the specified Peak table in the Result file defined in File name and stores it in the Resulting peak table. If "*" is entered as File name the current Result file will be used. The File name parameter may include a path from the current users root folder. |

Export

The table below contains a list of instructions for export operations.

| Instruction | Description |
|--------------------|---|
| EXPORT_CURVE_AIA | Exports the curve in AIA format. |
| EXPORT_CURVE_ASCII | Exports the Source curve to the file defined in Export to File in ASCII format . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. In the part of the source curve limited by the Left limit and Right limit every <n> sample is exported.</n> |
| EXPORT_CURVE_WKS | Exports the source curve to the file defined in Export to File in WKS format . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. In the part of the source curve limited by Left limit and Right limit every <n> sample is exported</n> |

| Instruction | Description |
|-----------------------|---|
| EXPORT_EVAL_LOG_ASCII | Exports an evaluation log in ASCII format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. |
| EXPORT_EVAL_LOG_WKS | Exports an evaluation log in WKS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. |
| EXPORT_EVAL_LOG_XLS | Exports an evaluation log in XLS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. |
| EXPORT_METHOD_ASCII | Exports a method to the file defined in Export to file in ASCII format . If "*" is entered as File name the current Result file will be used. If all parameters are OFF then no method is exported. If Main is ON then the main method is included and if Blocks is ON then all blocks are included in the exported file. |

| Instruction | Description |
|---------------------------|--|
| EXPORT_METHOD_WKS | Exports a method to the file defined in Export to file in WKS format. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. If all parameters are OFF then no method is exported. If Main is ON then the main method is included and if Blocks is ON then all blocks are included in the exported file. |
| EXPORT_METHOD_XLS | Exports a method to the file defined in Export to file in XLS format. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. If all parameters are OFF then no method is exported. If Main is ON then the main method is included and if Blocks is ON then all blocks are included in the exported file. |
| EXPORT_MULTI_CURVES_ASCII | Exports multiple curves (previously defined with EXPORT_SEL_CURVES instructions) in ASCII format to the file defined in Export to file. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |

| Instruction | Description |
|----------------------------|--|
| EXPORT_MULTI_CURVES_WKS | Exports multiple curves (previously defined with EXPORT_SEL_ CURVES instructions) in WKS format to the file defined in Export to file. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |
| EXPORT_MULTI_CURVES_XLS | Exports multiple curves (previously defined with EXPORT_SEL_ CURVES instructions) in XLS format to the file defined in Export to file. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |
| EXPORT_NORMALISE_RETENTION | Normalizes retention when exporting multiple curves. |
| EXPORT_PEAKTABLE_ASCII | Exports the peak table in Peak table source to the file defined in Export to file in ASCII format . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |
| EXPORT_PEAKTABLE_WKS | Exports the peak table in Peak table source to the file defined in Export to file in WKS format . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. |

| Instruction | Description |
|----------------------|---|
| EXPORT_PEAKTABLE_XLS | Exports the peak table in Peak table source to the file defined in Export to file in XLS format . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |
| EXPORT_PEAKTABLE_XML | Exports the peak table in Peak table source to the file defined in Export to file in XML format. If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name, a full search path must be entered in answer to the question. |
| EXPORT_SEL_CURVES | Selects a curve for subsequent export (using the EXPORT_MULTI-CURVES_* instruction). The curve is cut according to the right and left cut limit and the number of points to be exported may be set by the Export parameter (for example, every fifth point). |
| EXPORT_DOC_400_ASCII | Exports the documentation in the current result file in ASCII format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file. |

| Instruction | Description |
|--------------------|--|
| EXPORT_DOC_400_WKS | Exports the documentation in the current result file in WKS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file. |
| EXPORT_DOC_400_XLS | Exports the documentation in the current result file in MS Excel XLS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file. |
| EXPORT_DOC_WKS | Exports the documentation in the current result file in WKS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file. |

| Instruction | Description |
|------------------|---|
| EXPORT_DOC_XLS | Exports the documentation in the current result file in XLS format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file |
| EXPORT_DOC_ASCII | Exports the documentation in the current result file in ASCII format to the file defined in Export to file . If "*" is entered as File name the current Result file will be used. If "?" is entered followed by text, e.g. "Enter a file name", as File name , a full search path must be entered in answer to the question. If all parameters to this function are OFF then no documentation is exported. If at least one of them is ON then the documentation will be exported and the corresponding parts will be included in the exported file. |

Chromatogram functions

The table below contains a list of instructions for chromatogram functions.

| Instruction | Description |
|-------------|---|
| COPY_CHROM | Creates a copy of the specified chromatogram. If "*" is used as source then the current (default) chromatogram is used. If "*" is used as destination then a default name will be created for the copy. |

| Instruction | Description |
|----------------------------|--|
| CREATE_NEW_CHROM | Creates a new chromatogram with the given name. If "*" is used for the chromatogram name a default name will be generated and used. Note: It is a recommendation not to |
| | use only numbers as names for new chromatograms. |
| DELETE_CHROM | Deletes the named chromatogram. If trying to delete the current (default) chromatogram a run time error will be caused. |
| OPEN_CHROM | Opens the specified chromatogram from the specified file. |
| RENAME_CHROM | Renames the specified chromatogram. If "*" is used as From then the current (default) chromatogram is used. |
| RESTORE_DESTINATION_ CHROM | Resets the destination for the subsequent curve and peak table operations to the default chromatogram. Used in pair with the SET_DESTINATION_CHROM instruction. |
| SET_DESTINATION_CHROM | Opens the named chromatogram as destination for the subsequent curve and peak operations. Used in pair with the RESTORE_ DESTINATION_CHROM instruction. |

Other instructions The table below contains a list of instructions for other operations.

| Instruction | Description |
|-------------|--|
| BASE | Sets the X-axis base that the following calculations will be made in. If the value of the X-axis base is DEFAULT , then the default base is used (usually the base the method was run in). This instruction should be the first in the evaluation procedure, otherwise it will have no effect at all. |

| Instruction | Description |
|-------------|---|
| Comment | Inserts a comment below the marked instruction. |
| ENDLOOP | Marks the end of a LOOP statement. |
| LOOP | The instructions between this statement and the ENDLOOP statement are repeated n times. It is possible to have loops within loops as long as the number of LOOP statements matches the number of ENDLOOP statements. |
| MOLSIZE | Calculates the molecular sizes from a molecular size curve. |
| | A Mol. size column will be added to the Peak table. |
| QUANTITATE | Calculates the concentration and amounts in the sample from a quantitation table. |
| | Amount and Concentration columns will be added to the Peak table. |
| REPORT | Prints a report with the specified named report layout and title. If Title is "*" then the title in the report layout is used. If Report Layout is "*" then a default layout is used. |
| RUN_PROGRAM | Starts a program as a separate process. The Program name string contains the program name and parameters to start it with. |
| UPDATE | Updates a Quantitation table with new data from one standard concentration level. |
| | The default Limit(+/-) value of 12.5% will be used. |

Test instructions

The **Instruction** field also contains a group of test instructions. These instructions are only available for the UNICORN software development team.

| Instruction | Description |
|----------------------------|---|
| AUTOSAMPLER_PEAK_INTERVALS | Sets the area intervals for the AUTO-SAMPLER_PEAK_TEST. |
| AUTOSAMPLER_PEAK_TEST | Locates the first peak in the peak table. Compares the area of the peak in the peak table with the specified maximum and minimum areas. |
| GRADIENT_TEST_INTERVALS | Sets the level intervals for the GRADI-ENT_TEST . |
| GRADIENT_TEST | The theoretical straight line between the 0% and 100% levels are calculated. The deviation between the curve and the ideal straight line is compared in both directions from the center position (50%) until the deviation exceeds the defined maximum deviation. The calculated deviation points are checked against the defined limits. |
| STEP_RESPONSE_INTERVALS | Sets the level intervals for the STEP_RESPONSE_TEST. |
| STEP_RESPONSE_TEST | The relative amplitude is calculated at the specified retentions (The 0% and 100% amplitudes are used for reference). The calculated relative amplitudes are checked against the specified error margins. The 0% level amplitude is verified to be within the specified interval from the absolute 0 level. |
| UV_RESPONSE_INTERVALS | Sets the level intervals for the UV_RE-SPONSE_TEST. |

| Instruction | Description |
|------------------|--|
| UV_RESPONSE_TEST | The amplitudes for the 0% and 100% levels are calculated and the difference between the values are calculated. The results of (1) Curve2_Difference / Curve1_Difference and (2) Curve2_Difference / Curve3_Difference are calculated. The calculated points are checked if they are outside the defined limits from the 50% level. |

C Curve fit models and statistics

Introduction

The **Analysis** module (optional) is used to produce calibration curves and molecular size curves for analytical purposes. The quality of the curve fit model determines the accuracy of the curves. This appendix describes

- The available curve fit models.
- The statistical measurements in the **Analysis** module.

In this appendix

This appendix contains these sections:

| Topic | See |
|------------------|-----|
| Curve fit models | C.1 |
| Statistics | C.2 |

C.1 Curve fit models

Calibration curve models

The **Analysis** module provides a comprehensive range of curve fit models. The following models are available for calibration curves:

- Linear.
- Linear through origin.
- Quadratic.
- Quadratic through origin.
- · Point to point.

Note: The average peak size for all points at a specific level is used to calculate the calibration curve.

Molecular size curve models

The following curve fit models are available for molecular size curves:

- Linear.
- Linear (log Mw).
- Quadratic.
- Quadratic (log Mw).
- Point to point.
- Point to point (log Mw).

Statistics

The **Analysis** module provides values for the appropriate constants that are used in each curve equation for all models, except for the point to point models. It also provides statistical data that you can use to assess the quality of fit of the curve to the data.

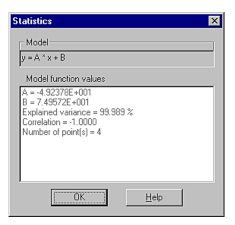
 Click the More... button in the Statistics field of the Quantitation table or Mol. **size table** dialog boxes to view the applied model statistics.

The Linear model The table below describes the features of the Linear curve fit model.

| Feature | Description |
|--|--|
| Equation. | y = Ax + B |
| Mathematical model. | The constants A and B are determined by linear least squares regression. |
| Minimum number of required points. | 2 (at least 4 points recommended) |
| Measuring range for the calibration curve. | Within the highest and lowest values for the points. |

Note: A variant of this model is available for the production of a molecular size curve. This uses the logarithm of the molecular size as the x value in the expression above.

The illustration below is an example of the statistical information for an applied **Linear** curve model:

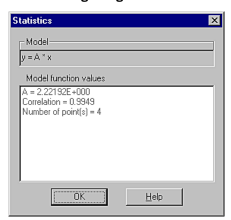


The Linear through origin model

The table below describes the features of the **Linear through origin** curve fit model:

| Feature | Description |
|--|--|
| Equation. | y = Ax |
| Mathematical model. | The constant A is determined by linear least squares regression. |
| Minimum number of required points. | 1 (at least 2 points recommended) |
| Measuring range for the calibration curve. | From the point with the highest value down to the origin. |

The illustration below is an example of the statistical information for an applied **Linear through origin** curve model:



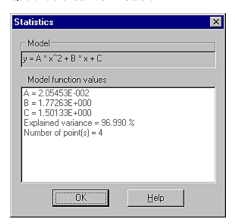
The Quadratic model

The table below describes the features of the **Quadratic** curve fit model:

| Feature | Description |
|--|---|
| Equation. | $y = Ax^2 + Bx + C$ |
| Mathematical model. | The constants A, B and C are determined by linear least squares regression. |
| Minimum number of required points. | 3 (at least 6 points recommended) |
| Measuring range for the calibration curve. | Within the highest and lowest values for the points. |

Note: A variant of this model is available for the production of a molecular size curve. This uses the logarithm of the molecular size as the x value in the expression above.

The illustration below is an example of the statistical information for an applied **Quadratic** curve model:

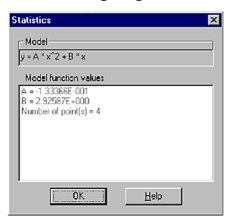


The Quadratic through origin model

The table below describes the features of the **Quadratic through origin** curve fit model:

| Feature | Description |
|--|--|
| Equation. | $y = Ax^2 + Bx$ |
| Mathematical model. | The constants A and B are determined by linear least squares regression. |
| Minimum number of required points. | 2 (at least 4 points recommended) |
| Measuring range for the calibration curve. | From the point with the highest value down to the origin. |

The illustration below is an example of the statistical information for an applied **Quadratic through origin** curve model:

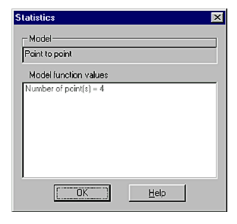


model

The Point to point The table below describes the features of the Point to point curve model.

| Feature | Description |
|--|--|
| Equation. | No single equation. |
| Mathematical model. | As these curves are not based on a single equation, no statistical data is available. The statistics table only contains information on the number of points in the curve. |
| Minimum number of required points. | 2 |
| Measuring range for the calibration curve. | Within the highest and lowest values for the points. |

The illustration below is an example of the statistical information for an applied Point to point curve model:



C.2 Statistics

Introduction

This section explains the correlation and explained variance calculations that are used by the **Analysis** module.

Correlation

The **Analysis** module calculates the correlation coefficient for linear models. This shows how well the data are linearly related. The correlation is displayed in the **Statistics** table.

If you are producing a calibration curve that relates peak area or height to amount or concentration, you aim to achieve a high positive correlation coefficient. A value of +1 indicates a perfect fit of all the data to the straight line. A molecular size curve has a negative slope, so the aim is towards a correlation coefficient of -1.

Too few data points

If you only have two data points for a **Linear** model, or only one point for a **Linear through origin** model, the fitted straight line will inevitably pass exactly through the points. By definition, this leads to a correlation of exactly +1, but this does not indicate a good fit, but instead indicates too few data points. In these cases the **Statistics** table will display a "---" symbol instead of the correlation value.

Correlation calculation

The correlation is derived as follows:

$$Correlation = \frac{\sum_{i} \left[(x_{i} - \overline{x}) (y_{i} - \overline{y}) \right]}{\sqrt{\left[\sum_{i} (x_{i} - \overline{x})^{2} \right] \left[\sum_{i} (y_{i} - \overline{y})^{2} \right]}}$$

Where:

- \bar{x} is the average of the x value.
- \overline{y} is the average of the y value.

For the molecular size model "Linear log(Mw)":

 $\bullet \bar{x}$ is the average of the logarithms of the molecular sizes.

Explained variance

Explained variance provides a measurement of how much of the variation in the data points (xy pairs) is due to the model. The remaining variation can be attributed to noise, i.e. random errors, or to the fact that an inappropriate model has been selected. This makes it possible to use the explained variance value for model selection, e.g. to decide if a quadratic model fits the data better than a linear model. This would be confirmed by a higher explained variance value.

Note: The explained variance is not calculated for curve models drawn through the origin.

Explained variance calculation

The explained variance is equal to R² adjusted for degrees of freedom. The illustration below shows the mathematical model:

Explained variance (%) = 100 x
$$\left[1 - \frac{SS_{residuals}/(n-k-1)}{SS_{total}/(n-1)} \right]$$

Where:

$$SS_{residuals} = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$
 (Residual Sum of Squares)

$$SS_{total} = \sum_{i=1}^{n} (y_i - \bar{y})^2$$
 (Total Sum of Squares)

- \bar{y} is the average of all y values. \hat{y}_i is a function value using the fitted model.

For example:
$$\hat{y}_i = Ax_i^2 + Bx_i + C$$

- n is the number of points (xy pairs).
- k is the number of x terms in the model.

For example, 1 for "Linear" and 2 for "Quadratic".

Undefined value for explained variance

You can only obtain a value for explained variance if you have sufficient data points on the curve. For instance, if you only have two points for a Linear model, or only three points for a **Quadratic** model, the fitted curve will pass exactly through the points. By definition, this leads to an undefined value for explained variance. In these cases the **Statistics** table will show a "---" symbol instead of an explained variance value.

The Column list D

Introduction

The Column List includes all available columns and their specific parameters. This appendix describes how to edit the Column List.

In this appendix

This appendix contains this section:

| Topic | See |
|------------------------------------|-----|
| How to edit the Column List | D.1 |

D.1 How to edit the Column List

Introduction

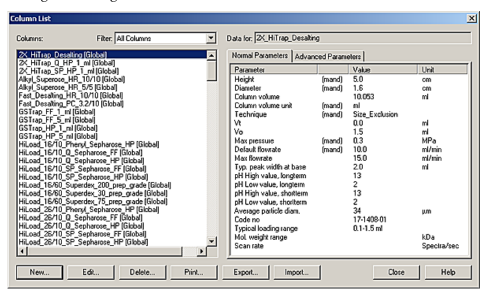
This section describes how to edit the list of available columns.

Available columns

When you create a new method and select a column, certain column-specific parameters are automatically copied into the method. The list of available columns is found in the **For column** field of the **New Method** dialog box. The **Column List** is not linked to a particular method, although the columns are edited within the **Method Editor**.

Columns are either globally available to all users, or only personally available. It is best not to edit the globally available columns, unless you save the changes under a new column name, since other users may not appreciate the changes.

Note: It is recommended that only a limited number of users are given access to the right to edit global columns. This is essential to avoid unintentional changes.



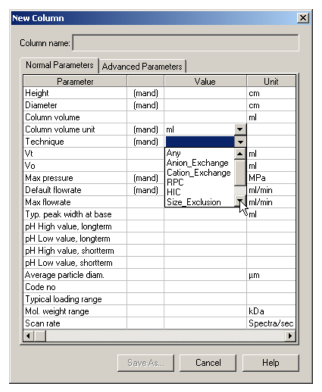
How to print the column list

The table below describes how to print the column list data.

| Step | Action |
|------|--|
| 1 | Click the print button. |
| | Result: The Print Column List dialog box opens. |
| 2 | Select to print a global or your personal column list. Click OK. |
| | Result: The column list is printed on the default Windows printer. |

dialog box

The New Column The illustration below shows the New Column dialog box:



column

How to add a new The table below describes how to add a new column to the Column List.

| Step | Action |
|------|--|
| 1 | Choose Edit:Column List in the Method Editor. |
| | Result: The Column List dialog box opens. |
| | Note: Select a column from the list to display the parameters in the field to the right. Most column parameters are displayed in the Normal Parameters tab. Additional parameters for special columns may be displayed in the Advanced Parameters tab. |
| 2 | Click the New button. |
| | Result: The New Column dialog box opens. |
| 3 | Select the appropriate parameter tab. |
| | Type the desired parameter values. |
| | Click the Save as button. |
| | <i>Note</i> : Mandatory parameters are labelled mand . The column cannot be saved unless all mandatory parameters are filled in. |
| | Result: The Save as dialog box opens. |

| Step | Action |
|------|---|
| 4 | Type the name of the new column. Click the Save as global checkbox if the column should be available to other users. |
| | <i>Note</i> : You must have Edit global lists authorization to save a column for global use. A global column cannot have the same name as a personal column. |
| | Click OK . Result: The new column is added to the Column List . |

Note: See column instruction to determine the back pressure over the system and the column.

The normal column parameters

The table below is a list of all the available normal column parameters:

| Parameter | Unit | Comment |
|--------------------|---------------------|--|
| Height | cm | Mandatory.Calculation of N/m. |
| Diameter | cm | Mandatory. |
| Column volume | nl, μl, ml or liter | Mandatory. Automatically calculated from Height and Diameter. User cannot set this parameter directly. |
| Column volume unit | nl, μl, ml or liter | Not mandatory. The column volume is calculated in the set unit. |
| Technique | | Mandatory. Decides which technique the column should be available for. |

| Parameter | Unit | Comment |
|-------------------------|--|--|
| Vt | nl, μl, ml or liter | Not mandatory. Total liquid volume. Used to calculate the capacity factor after an integration. |
| Vo | nl, µl, ml or liter | Not mandatory. Void volume. Used to calculate K_{av} after integration. |
| Max pressure | MPa | Mandatory. Used for setting pressure limit in a method automatically. |
| Default flowrate | nl/min, µl/min, ml/min or liter/min | Mandatory. Used to set the flowrate in a method automatically. |
| Max flowrate | nl/min, µl/min, ml/min or liter/min | Not mandatory. Used to give a warning if a higher flowrate is chosen when saving or starting a method. |
| Typ. peak width at base | nl, µl, ml or liter | Not mandatory. Used to set averaging time for UV detector. used to set peak fractionation parameters. |
| pH high value, longterm | | Not mandatory. Used to give a warning when saving or starting the method if the BufferPrep_pH value is higher than the set value. |

| Parameter | Unit | Comment |
|---------------------------|-------------|--|
| pH low value, longterm | | Not mandatory. Used to give a warning when saving or starting the method if the BufferPrep_pH value is lower than the set value. |
| pH high value, shortterm | | Not mandatory. Used to give a warning when saving or starting the method if the BufferPrep_pH value is higher than the set value. |
| pH low value, shortterm | | Not mandatory. Used to give a warning when saving or starting the method if the BufferPrep_pH value is lower than the set value. |
| Average particle diameter | μт | Not mandatory.Information only. |
| Code no | | Not mandatory.Information only. |
| Typical loading range | mg | Not mandatory.Information only. |
| Mol. weight range | kDa | Not mandatoryInformation only |
| Scan rate | spectra/sec | Not mandatory.Information only. |

Note: The values for the parameters **Max pressure**, **Default flowrate** and **Typical peak width at base** (used to set average time and peak fractionation parameter **MinWidth**) are only copied into the method if the corresponding instructions are available as variables.

How to edit column paramet-

The table below describes how to edit column parameters in the **Method Editor**:

| Step | Action |
|------|--|
| 1 | Choose Edit:Column List. |
| | Result: The Column List dialog box opens. |
| 2 | Select a column and click the Edit button. |
| | Result: The Edit Column dialog box opens. |
| 3 | Select the desired parameters and change the value settings. |
| 4 | Click the Save button. |
| | or |
| | Click the Save as button to save the column under a new name. |

Note: If a column has been selected and saved in a method, and the parameters for the column are changed later, the column in the method will not be updated automatically. When you open the method you will be asked if you want to update the parameters. The recommendation is that you answer Yes.

How to delete a column.

The table below describes how to delete a column:

| Step | Action |
|------|---|
| 1 | Choose Edit:Column List. Result: The Column List dialog box opens. |
| 2 | Select a column and click the Delete button. *Result: The Delete Column dialog box opens. |
| 3 | Click the checkbox for each column you want to delete. Click 0K. Result: The selected columns are deleted. |

How to export a column.

The column information for a system can be transferred to another by using the export and import functions in the column list. The table below describes how to export a column:

| Step | Action |
|------|---|
| 1 | Choose Edit:Column List. |
| | Result: The Column List dialog box opens. |

| Step | Action |
|------|---|
| 2 | Click the Export button. |
| | Result: The Export Column dialog box opens. |
| 3 | Click the checkbox for each column you want to export. |
| | Click Export. |
| | Result: The Export Column to file dialog box opens. |
| 4 | Select the desired folder in the navigation window. |
| | Type a new file name if neccessary. |
| | Choose the type of file to export (column file or Excel file) |
| | Click the Save button. |
| | Result: The column file is saved and the dialog box closes. |

Note: If a column is selected in the **Column List** when the **Export Column** dialog box is opened, this column will automatically be selected in the **Export Column** dialog box.

How to import a column.

The table below describes how to import a column:

| Step | Action |
|------|---|
| 1 | Choose Edit:Column List. |
| | Result: The Column List dialog box opens. |
| 2 | Click the Import button. |
| | Result: The Import Column dialog box opens. |
| 3 | Click the Browse button to locate the column file. |
| | Result: The Import Column from file dialog box opens. |
| 4 | Select a column file. |
| | Click Open. |
| | Result: The Import Columns dialog box opens. |
| 5 | Select the columns to import from the list. |
| | • Select Import as global to add the columns to the global column list if desired. |
| | Click Import. |
| | Result: The selected columns are imported and available in the column list. |

 $\it Note$: Select Import as global to import the columns to the global column list.

Ε

How to create and edit BufferPrep recipes

Introduction

The **BufferPrep** function is available for some ÄKTAdesign systems. This appendix describes how to create and how to edit the recipes for **BufferPrep**.

In this appendix

This appendix contains these sections:

| Topic | See |
|-----------------------------------|-----|
| How to create a BufferPrep recipe | |
| How to edit a BufferPrep recipe | |

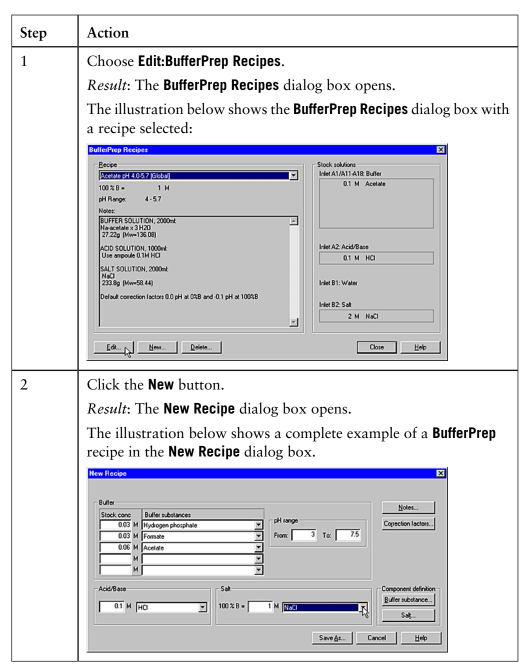
E.1 How to create a BufferPrep recipe

About BufferPrep recipes

New **BufferPrep** recipes are created in the **Method Editor**. The list of recipes is not linked to a specific method. Which recipe to use in a certain method is selected on the **BufferPrep** tab in the **Run Setup**.

How to create a recipe

The table below describes how to create a new **BufferPrep** recipe in the **Method Editor**:



| Step | Action |
|------|--|
| 3 | Select buffers from the Buffer substances droplists and type stock concentrations in the corresponding Stock conc box. |
| | See "How to define a new buffer substance" below if the desired substance is not available. |
| 4 | Select either HCI (acid) or Na0H (base) from the Acid/Base droplist and type the required stock concentration (typically 0.1 M) |
| 5 | Select a salt from the Salt droplist and type the maximum outlet concentration of the salt for 100%B (typically 1.0 M). |
| | See "How to define a new salt" below if the desired salt is not available. |
| 6 | Type the desired pH range minimum and maximum values in the From and To boxes. |
| | See "How to select the pH range" below this table. |
| 7 | Click the Notes button (optional). |
| | Type your notes about the recipe in the displayed dialog box. |
| | Click OK to return to the New Recipe dialog box. |
| 8 | Click Save as to save the recipe under a new name. |
| | <i>Note</i> : A warning message will appear if any of the recipe values are unfeasible. |
| 9 | Type a name in the dialog box. Click 0K. |
| | Result: The new recipe is added to the recipe list. |

Note: It is recommended that restricted access be given to the right to edit global recipes.

The recipes are either globally available to all users, or only personally available. It is best not to edit the globally available recipes, unless you save the changes under a new recipe name, since other users may not appreciate the changes.

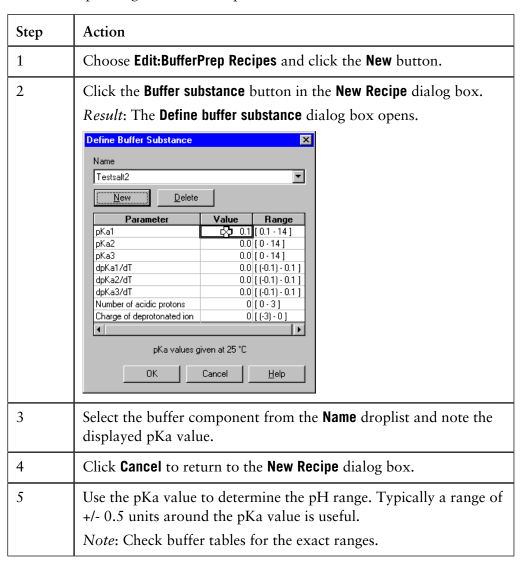
Buffer concentration

Use buffer concentrations that are 2-4 times higher than the concentration that is used in the normal preparation. When **BufferPrep** is used, the buffer will be diluted 2-10 times depending on the amount of acid/base that has to be used to reach the desired pH value.

Up to five different buffering components can be selected. To prevent a too high ionic strength, the sum of the concentrations for all selected buffers should be between 0.03 M and 0.2 M (typically 0.1 M).

How to select the pH range

The useful pH range depends on the pKa value. The table below describes how to determine a pH range based on the pKa value:



How to define a new buffer substance *Note*: Before you can define a new buffer substance you must ensure that all pKa values are available for the substance. The pKa values should be true (i.e. the pKa value at indefinite dilution) and not apparent pKa values (i.e. measured at a non-zero concentration). The pKa values should be given at 25° C.

The table below describes how to define a new buffer substance:

| Step | Action |
|------|---|
| 1 | Choose Edit:BufferPrep Recipes and click the New button. |
| 2 | Click the Buffer substance button in the New Recipe dialog box. |
| | Result: The Define buffer substance dialog box opens. |

| Step | Action |
|------|---|
| 3 | Click the New button. Result: The New component dialog box opens. |
| 4 | Type a name for the new component and click OK to return to the Define buffer substance dialog box. |
| 5 | Type appropriate values in the Value cells for each pKa and dpKa/dT parameter. Note: All values must fall within the stated Range limits. Up to three |
| | values can be entered for each buffering component. When the component has less than three pKa values, the other values should be set to zero. A dpKa/dT value of zero means that the pKa does not change with temperature. |
| 6 | • Type the number of acidic protons for the buffer substance in the form that it is actually weighed in. |
| | Example: The number is 2 for NaH₂PO₄, 1 for Na₂HPO₄ and 0 for Tris. Type the charge of the completely de-protonated ion. This will |
| | be a negative value for an acid and zero for a base. |
| | Example: The value is -3 for NaH₂PO₄ and 0 for Tris. Click 0K. |
| | Result: The new buffer substance is added to the list of available substances. |

How to define a new salt

Before you can define a new salt you must ensure that the new salt is inert, i.e. a salt with no buffering properties. The table below describes how to define a new salt:

| Step | Action |
|------|--|
| 1 | Choose Edit:BufferPrep Recipes and click the New button. |

| Step | Action |
|------|--|
| 2 | Click the Salt button in the New Recipe dialog box. |
| | Result: The Define salt dialog box opens. |
| | Name NaCi3 Parameter Value Range Charge of Anion [1·7] Charge of Cation [1·7] |
| 3 | Click the New button. Result: The New component dialog box opens. |
| 4 | Type a name for the new salt and click OK to return to the Define salt dialog box. |
| 5 | Type the appropriate charge of anion value in the corresponding Value cell. Example: The value for Cl⁻ is - 1. The value for SO₄²⁻ is -2. Type the appropriate charge of the cation value in the corresponding Value cell. Example: The value for Na⁺ is 1. The value for Mg²⁺ is 2. Click OK. Result: The new salt is added to the list of available salts. |

E.2 How to edit a BufferPrep recipe

Introduction

This section describes how to edit a **BufferPrep** recipe in the **Method Editor**.

How to edit a recipe

The table below describes how to edit a **BufferPrep** recipe:

| Step | Action |
|------|---|
| 1 | Choose Edit:BufferPrep Recipes. Result: The BufferPrep Recipes dialog box opens. |
| 2 | Select a recipe and click the Edit button. Result: The Edit Recipe dialog box opens. Edit Recipe Recipe name: Acetate pH 4.05.7 Buffer Stock conc O.1 M Acetate M Y |
| 3 | Change the substances and parameters as desired and click the Save button or the Save as button to save the new recipe. |

Changes to recipes in methods

If a recipe has been selected and saved in a method, and the recipe is later changed, the corresponding recipe in the method will not be updated automatically. When you open the method you will be asked if you want to update the parameters in the method recipe. The recommendation is that you answer **Yes**.

Note: The question will not appear if you only change the **Correction factors**. The **Correction factors** in the method recipe will not be updated.

How to determine if the Correction factors need to be changed

Correction factors can be set to fine-tune a recipe around a specific pH, to obtain high pH accuracy. The table below describes how to run the **BufferPrep** manually at 0% and 100% in the **System Control** module, to determine if the **Correction factors** need to be changed:

| Step | Action |
|------|---|
| 1 | Choose Manual:Other. |
| | Result: The System Other instructions dialog box opens. |

| Step | Action |
|------|---|
| 2 | Select the recipe from the Recipe Name droplist and click the Execute button. |
| | Result: The recipe instruction is added. |
| 3 | Click the Pump radio button and select BufferPrep_pH. Set the pH value in the pH parameter box and click the Execute button. |
| | Result: The BufferPrep pH value is added and the run starts. |
| 4 | Select Flow. |
| | • Set the flow rate in the FlowRate parameter box and click Execute . |
| | Result: The new flow rate is added. |
| 5 | Check the pH reading when it is stable in the BuffPre_pH meter in the Run Data pane. |
| | <i>Note</i> : At least 30 ml of eluent must pass through before the reading stabilizes. |
| | To display the BuffPre_pH meter, right-click and select Properties. Select BuffPre_pH on the Run Data Groups panel and click the OK button. |
| 6 | Select Gradient in the Instructions list. |
| | Type 100% in the Target parameter box, 0 in the Length parameter box and click Execute . |
| | Result: The gradient instruction is added. |
| 7 | Check the pH reading when it is stable at 100%. See "How to change the Correction factors " below. |

How to change the Correction factors If the readings described in the instruction above are acceptable at both 0% and 100%, the **Correction factors** do not need to be changed. If the **Correction factors** do not produce an acceptable result, they must be adjusted in the **Method Editor** module. The table below describes how to change the **Correction factors**:

| Step | Action |
|------|---|
| 1 | Choose Edit:BufferPrep Recipes. |
| | Result: The BufferPrep Recipes dialog box opens. |
| 2 | Select the recipe from the Recipe droplist and click the Edit button. |
| | Result: The Edit Recipe dialog box opens. |

| Step | Action |
|------|--|
| 3 | Click the Correction factors button. |
| | Result: The Correction Factors dialog box opens. |
| 4 | Type the deviation at 0% and 100%. |
| | <i>Example</i> : If the pH is set to 7.0 and the actual pH is 7.1, the Correction factor is 0.1. If the actual pH is 6.9, the Correction factor is -0.1. |
| | • Click 0K . |
| | • Click the Save button or the Save as button to save the recipe. |

Note: If there already are **Correction factors**, the measured pH deviation should be added to the old factors.

Method examples F

Introduction

This appendix contains practical method examples that can be applied in typical situations. The examples cover three different topic groups:

- Watch instructions
- Messages
- **Quality control**

Watch instructions allow the progress of a method run to be determined by the events during the method run, for example start collecting fractions when the first peak elutes, or equilibrate the column until the eluent conductivity has reached a given value. This is facilitated by the **Watch** instructions.

The system strategy includes **Watch** instructions for each monitor defined in the system. These instructions are used to survey method runs, and instruct the system to call a specified block or an instruction when a particular monitor signal meets a given condition. As long as the condition is not met, the block is not activated.

Messages can be used in a method to provide information to the operator but also for interaction between the system and the operator.

A Quality control procedure in a method can be used to ensure that the quality of the results remain consistent in a series of runs.

In this appendix

This appendix contains the following sections:

| Topic | See |
|--------------------------------------|-----|
| Simple equilibration | F.1 |
| Equilibration with simple safeguard | F.2 |
| Equilibration with extra safeguard | F.3 |
| Collection of absorbance peaks | F.4 |
| Collection of three absorbance peaks | F.5 |
| Messages | F.6 |
| Quality control procedure | F.7 |

F.1 Simple equilibration

Introduction

This section contains an example of how a **Watch** instruction for simple equilibration can be inserted into a method.

Example instruction

This is an example instruction as it would be presented in the **Text** pane.

0.00 Block EQUILIBRATE

(Equilibrate)

0.00 Base SameAsMain

0.00 Watch_Cond Less_than, 5 {mS/cm}, CONTINUE

0.00 Hold

0.10 Watch UV1 Less than, 100 {mAU}, CONTINUE

0.10 Hold

0.10 End Block

If you are not using ÄKTA instruments

If you are not using ÄKTA instruments, a delay should be added after the **Hold/Pause** instruction so that the following instruction will not be executed simultaneously with the **Hold/Pause** instruction.

This is what happens

The table below describes what happens in the above example:

| Stage | Description |
|-------|---|
| 1 | The Watch is started on the conductivity signal and the method is then put on Hold . |
| 2 | Continue is issued and Watch_cond is turned off automatically when the Watch_cond condition is fulfilled. |
| 3 | Method execution continues issuing a Watch_UV command. Again the method is put on Hold until the Watch condition is fulfilled. |

Note: Even though the line

Watch Cond Less than, 5 {mS/cm}, Continue

is in the method placed before **Hold**, the method is put on hold first and then continued only after the conductivity has reached a level less than 5 mS/cm. This is because **Hold** is an instruction that will be executed at its breakpoint, while **Continue** is not an instruction but rather an action for the **Watch** instruction.

Evaluation of the method

This method works satisfactorily although one drawback is that it might never end, and thus consume all of the buffer if the conditions for some reason are unfulfilled. See appendices F.2 Equilibration with simple safeguard on page 548 and F.3 Equilibration with extra safeguard on page 549.

F.2 Equilibration with simple safeguard

Introduction

This section contains an example of how a **Watch** instruction for simple safeguard can be inserted into a method.

Example instruction

This is an example instruction as it would be presented in the **Text** pane:

0.00 Block EQUILIBRATE

(Equilibrate)

0.00 Base SameAsMain

0.00 Watch_UV1 Less_than, 100 {mAU} END_BLOCK

5.00 Watch Off UV1

5.00 Message "The Condition was never reached", Screen, "No sound"

5.00 End Block

This is what happens

This is what happens in the above example:

The column is equilibrated until the UV has reached a level below 100 mAU or until the column has been equilibrated with five column volumes of buffer, whichever condition is met first. In this way, it is possible to equilibrate the column without the risk of running out of buffers.

F.3 Equilibration with extra safeguard

Introduction

This section contains an example of how a **Watch** instruction for extra safeguard can be inserted into a method.

Example instruction

This is an example instruction as it would be presented in the **Text** pane:

```
0.00 Block EQUILIBRATE
    (Equilibrate)
    0.00 Base SameAsMain
      0.00 Block COND LESS THAN
         (Cond less than)
        0.00 Base SameAsMain
        0.00 Watch Cond Less than, 5 {mS/cm} END BLOCK
        6.00 Message "Low conductivity not reached", Screen,
        "No sound"
        6.00 Pause INFINITE {Minutes}
        6.00 End Block
      0.00 Block COND STABLE
        (Cond stable)
        0.00 Base SameAsMain
        0.00 WatchPar Cond 0.500 {mS/cm}, 2 {mS/cm}
        0.00 Watch Cond Stable Baseline, 5 {Minutes},
        END BLOCK
        10.00 Message "Conductivity not stable", Screen,
        "No sound"
        10.00 Pause INFINITE {Minutes}
        10.00 End Block
```

Note: If you are not using ÄKTA instruments, a delay should be added after the **Hold/Pause** instruction so that the following instruction will not be executed simultaneously with the Hold/Pause instruction.

This is what happens

0.00 End Block

The table below describes what happens in the above example:

| Stage | Description |
|-------|--|
| 1 | The column is equilibrated until the conductivity is below 5 mS/cm. |
| 2 | If this value is not reached within 6 column volumes, the method is paused and a message is displayed. |

| Stage | Description |
|-------|---|
| 3 | Equilibration of the column is continued until the conductivity value is "stable" (allowed to vary by max. ±2 mS/cm) over a period of at least 5 minutes. |
| 4 | If this condition is not met within 10 column volumes, the method is again paused. |

Note: At each pause, the operator can decide whether to continue or abort the run.

F.4 Collection of absorbance peaks

Introduction

This section contains an example of how to collect absorbance peaks through outlets F3 and F4.

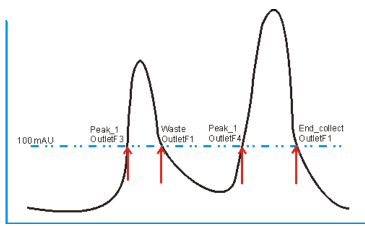
Example instruction

This is an example instruction as it would be presented in the **Text** window:

```
0.00 Block ELUTION
    (Elution)
    0.00 Base SameAsMain
    0.00 Gradient 100.0 {%}, 20.00 {base}
    0.00 Watch_UV1 Greater_Than, 100 {mAU}, Peak 1
      (Peak 1)
      0.00 Base SameAsMain
      0.00 OutletValve F3
      0.00 Watch_UV1 Less_Than_Or_Valley, 100 {mAU}, Waste
        (Waste)
        0.00 Base SameAsMain
        0.00 OutletValve WasteF1
        0.00 Watch_UV1 Greater_Than, 100 {mAU}, Peak_2
           (Peak 2)
          0.00 Base SameAsMain
          0.00 OutletValve F4
          0.00 Watch UV1 Less Than, 100 {mAU}, End collect
             (End collect)
             0.00 Base SameAsMain
             0.00 OutletValve WasteF1
             0.00 End Block
           0.00 End Block
        0.00 End block
      0.00 End Block
    20.00 End Block
```

Illustration

The illustration below shows peaks collected by the method in the example above.



This is what happens

In this example, one or two absorbance peaks are collected through outlets F3 and F4 respectively with waste fractions collected through outlet valve F1 (waste). Each called block (except **End_collect**) resets the **Watch** condition so that the method reacts correctly to subsequent changes in the UV absorbance.

Invalid Watch instructions

The design of a method of this kind (with several **Watch** instructions for the same monitor) is important. The construction in the following three lines appears simpler but is incorrect:

0.00 Watch_UV Greater_than, 100 {mAU}, Peak_2

0.00 Watch_UV Less_than, 100 {mAU}, End_collect

0.00 End_block

Here, the second **Watch** instruction will annul the first, since a signal can only be watched for one condition at a time.

F.5 Collection of three absorbance peaks

Introduction

This section contains an example of how to collect three absorbance peaks through outlets F3, F5 and F7 with waste fractions through outlets F4, F6 and F8.

The maximum number of peaks collected in this example is three due to the limited number of positions on the outlet valve.

Recommendations

Waste container needed

The waste fractions between the peaks are collected through the outlet valve positions F4, F6 and F8, so ensure that the tubing from these positions is lead to a suitably large container.

Condition for UV threshold

The UV threshold for collecting the waste fraction must be below the threshold for collecting the peak fraction so that the "waste" condition will not be fulfilled simultaneously or immediately after peak collection.

Example instruction

This is an example instruction as it would be presented in the **Text** window:

```
0.00 Block Eluate Fractionation
    (Eluate Fractionation)
    0.00 Base SameAsMain
    0.00 Watch UV1 Greater Than, 5 {mAU}, Peak
      0.00 Base SameAsMain
      0.00 OutletValve Feed
      0.00 Watch UV1 Less Than Or Valley, 4.75 {mAU}, Waste
        (Waste)
        0.00 Base SameAsMain
        0.00 OutletValve Feed
        0.00 Watch UV1 Greater Than, 5 {mAU}, Peak1
           (Peak1)
           0.00 Base SameAsMain
          0.00 OutletValve Feed
          0.00 Watch UV1 Less Than Or Valley, 4.75 {mAU},
          waste1
             (Wastel)
             0.00 Base SameAsMain
             0.00 OutletValve Feed
             0.00 Watch UV1 Greater Than, 5 {mAU}, Peak2
```

```
(Peak2)
           0.00 Base SameAsMain
           0.00 OutletValve Feed
           0.00 Watch_UV1 Less_Than_Or_Valley, 4.75
           {mAU}, Waste2
           0.00 End_block
        0.00 End_block
      0.00 End_block
    0.00 End_block
  0.00 End_block
0.00 End_block
```

pens

This is what hap- The table below describes what happens in the above example:

| Stage | Description |
|-------|--|
| 1 | When the UV reaches 5 mAU or more, the outlet valve is switched to the position for collecting the first peak. |
| 2 | When the UV reading goes down to 4.75 mAU, the outlet valve switches to the next position to separate the waste fraction from the collected peak fraction. |
| 3 | This process is repeated twice for the next two peaks so that when the UV reading rises above 5 mAU, the position switches to collect the peak fraction and the position switches again to collect the waste fraction when the UV reading falls again. |

F.6 Messages

When to use a message

Messages are used to inform the operator of the progress of the run. Messages can also be used for interaction between the operator and the system when necessary. A message can be for information in a screen only, or it can require a signature before the user can control the system. The messages are all added to the logbook text. This appendix describes how to add a message to a method. The appendix also gives two examples of how a message can be used.

How to add a Message instruction

The table below describes how to add a **Message** instruction to the method.

| Step | Action |
|------|--|
| 1 | Select Other in the Instructions field of the Instruction box. |
| | Select Message in the instructions list. |
| 2 | Type a message in the Message text box in the Parameters field. |
| 3 | Select one of the display options on the Mode menu: |
| | Screen, i.e. only a text message is displayed. |
| | • Noscreen, i.e. the message will not be displayed but only inserted into the logbook. |
| | • Authorize , i.e. the message will require a signature from the user before the user can interact with the system again. |
| 4 | Select a sound on the Sound menu if desired. Click the Insert button. |

Note: If the **Message** instruction is inserted in a conditional block it will only be displayed if the conditions of the block (for example a **Watch**) is fulfilled.

All messages are erased when the system reaches the **End** status. This also includes **Authorize** messages.

Protecting a method run with a message

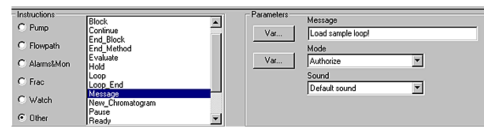
A message can be set up in the beginning of a method to protect the method run from unauthorized interference. Once the message is issued, the system is locked from interaction by any user unless the user provides an authorization signature. The only command that is available without authorization is **Pause**.

The illustration below shows the text instruction for the message described above:

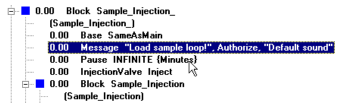
Pausing a method run for a manual sample injection

A message can be set up to pause the method until a sample has been injected manually. If a message requiring an authorization is followed by a **Pause** instruction the system will be paused until the message is acknowledged and signed. No other interaction with the system is available to the user. The operator will see a screen with a reminder to inject the sample before the method run proceeds.

The illustration below shows the selected message instruction in the **Instruction box** and the parameters for the message described above:



The illustration below shows the text instruction for the message described above:



Note: The message instruction must be followed immediately by the **Pause** instruction as shown above.

F.7 Quality control procedure

Introduction

When a series of runs is performed, irregularities in samples or in system or column performance can produce errors that will make the results inaccurate.

A quality control procedure can be added to a method to be used for a test run during the series of runs. The control procedure can ensure that the results remain within acceptable limits. If the result from the test run is unacceptable, the system can be paused so that the error is not repeated in subsequent runs.

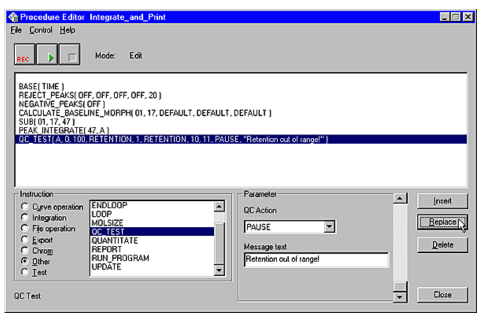
How to create the quality control procedure

The easiest way to create the quality control procedure is to edit an existing procedure that includes a peak integration. The table below describes how to do

| Step | Action |
|------|---|
| 1 | Choose Procedures:Edit:Open in the Evaluation module. |
| | Result: The Open Procedure dialog box opens. |
| 2 | Select the procedure (Global) Integrate_and_Print. Click the OK button. |
| | Result: The Procedure Editor opens with the procedure displayed. |
| 3 | Select the REPORT instruction in the procedure. Choose Other and QC_TEST in the Instruction field. |
| 4 | • Type appropriate values in the Parameter field. See "QC_TEST Parameter descriptions" below. |
| | Click the Replace button. |
| | <i>Result</i> : The REPORT instruction is replaced by the QC_TEST instruction. |
| 5 | Choose File:Save As. |
| | Result: The Save As dialog box opens. |
| | • Type a name for the procedure (for example QC_test). |
| | • Select the Global procedure check box if the procedure is to be available to all users. |
| | <i>Note</i> : If you select File:Save to save the procedure it will replace the (Global) Integrate_and_Print procedure. |

Illustration: The Procedure Editor

The illustration below shows the **Procedure Editor** with the **QC_TEST** instruction displayed:



QC_TEST parameter descriptions

The table below describes the parameters for the ${\tt QC_TEST}$ instruction.

The example values are used in the illustration above.

| Parameter | Description |
|-------------------|---|
| Peak table source | The peak table indicated in the PEAK_INTEGRATE instruction (Example: A). |
| Left limit | The retention value where the control instruction will begin (Example: 0). |
| Right limit | The retention value where the control instruction will end (Example: 100). |
| | <i>Note</i> : The control instruction will be applied to the run up to the sequence in the method where the control instruction is inserted: |
| | • The controlled part of the run will end at the Right limit if this retention value is reached before the control instruction is reached in the method. |
| | • If not, the controlled part of the run will end when the control instruction is reached in the method. |
| Peak selection on | The criteria for peak identification (Example: RETEN-TION). |
| Order number | The sequential order number of the peak (Example: 1). |

| Parameter | Description |
|----------------------|---|
| Peak table parameter | The peak table parameter that will be tested by the control instruction (Example: RETENTION). |
| Less than | Values less than the parameter value will be out of the acceptable range (Example 10). |
| Greater than | Values greater than the parameter value will be out of the acceptable range (Example 11). |
| QC Action | The action the system will take when the controlled value is out of the acceptable range (Example: PAUSE). |
| Message text | Free text message that is displayed when the controlled value is out of the acceptable range (Example: Retention out of range!) |

Note: All values must be included before the instruction can be inserted.

How to add the quality control procedure to a method

The table below describes how to add the quality control procedure to a method.

| Step | Action |
|------|---|
| 1 | Open the method in the Method Editor. Click the Run Setup icon. Result: The Run Setup for the method opens. |
| 2 | Select the Evaluation Procedures tab. Click the Import button. Result: The Import dialog box opens. |
| 3 | Select the quality control procedure you created and saved (Example: QC_test) in the Select field. Click the Import button. Result: The quality control procedure is added to the available evaluation procedures. Click the Close button. |
| 4 | • Click the check box to de-select the quality control procedure. Note: If the quality control procedure is selected it will initiate a new manual run at the end of the method run. |

| Step | Action |
|------|---|
| 5 | Click the Text Instructions icon. |
| | |
| | Select the last instruction in the method. |
| | Select Other:Evaluate in the Instructions field. |
| | Select the quality control procedure in the Procedure list. |
| | Click the Insert button. |
| 6 | Choose File:Save |
| | or |
| | • Click the Save icon. |
| | <i>Result</i> : When the method run is performed the quality control procedure will create a second chromatogram. If the controlled value is outside the acceptable range, the system will be paused. |

Α

Alarms

Description, 461

Alarms and warnings

Description, 215

Effects on the system, 215

Analysis

External standard quantitation, 397

Recovery calculation, description, 406

Quantitation reliability factors, 408

How to create a quantitation table, 411

Molecular size determination, overview, 449

How to create a molecular size table, 452

How to calculate molecular size, 455

Analysis module

How to install, 391

Automated quantitation

Basic conditions, 439

How to prepare the quantitation table, 439

How to set up the sample runs, 441

Automatic update with Replace, 442

Automatic update with Average, 443

How to perform updates in scouting runs, 444

Automated update with Average for scouting runs, 447

В

Baseline

Calculation options, 329

The Calculate function, 329

Reuse existing, 329

How to edit manually, 349

How to adjust the baseline graphically, 351

Definition of a segment, 493

Parameters, 494

Batch run

How to perform, 384

BatchID

Logbook illustration, 205

Blank curve

Calculate baseline based on, 329

BufferPrep

Description, 143

Stock solutions, 143

How to create a method, 144

How to adjust the correction factors, 145

How to create a recipe, 537

Buffer concentration, 538

How to select the pH range, 539

How to define a new buffer substance, 539

How to define a new salt, 540

How to edit a recipe, 542

How to verify correction factors, 542

How to change correction factors, 543

C

Calibration curve

How to update, 425

Chromatogram Layout

Curve tab, 236

Default curve names, 236

How to choose curve name appearance, 236

The Curve Style and Color tab, description, 238

Chromatogram window

How to display header information, 229

Shortcut menu, 231

How to optimize the workspace, 231

How to display a vertical marker, 232

How to display the Logbook overlay, 232

Chromatograms

Description, 227

Temporary chromatogram, 227

How to make layout changes, general, 235

How to change and fix the Y-axis, 240

How to add a second Y-axis, 240

How to change and fix the X-axis, 241

How to save a layout, 242

How to apply a layout, 242

How to cut a curve and store as new, 245

How to change the size of fraction marks, 246

How to change the size of injection marks, 246

How to change the size of logbook marks, 246

How to print active chromatograms, 247

How to add annotations, 278

How to edit annotation text, 278

How to rename, 286

How to open several to compare, 302

The command File:Open to compare, 302

How to open several with the File:Open command, 303

How to display several simultaneously, 304

Commands to import curves from result files, 305

How to import curves with File:Open to compare, 306

How to import curves with File:Open, 309

How to copy curves into a new, 310

How to set a reference point, 364

Classic algorithm

Definition, 340

Parameters, 340

How to set, 340

Shortest baseline segment, 341

Slope limits, 342

Noise window, 344

Missing peaks, 345

When to change the Max baseline level, 346

How to set Max baseline level, 346

Definition, 493

How to measure baseline segments, 495

How to measure noise level, 496

How to measure the slope limit, 496

Columns

Column prompt in manual instructions, 212

How to add a new, 529

Normal column parameters, 530

How to edit parameters, 533

How to delete, 533

How to export, 533

How to import, 534

Concentration levels

Levels in quantitation, definition, 413

Conditional call

Description, 100

Correlation

Explanation, 525

Curve fit models

Linear, 521

Linear through the origin, 522

Quadratic, 523

Quadratic through origin, 523

Point to point, 524

Curves

How to copy into the Temporary chromatogram, 227

Run curves default appearance, 231

How to choose the Y-axis scale, 231

Default curve names, 236

Peak labels, 238

Fraction text alignment options, 238

Logbook text alignment options, 238

How to change the color and style, 238

How to filter logbook information, 238

How to set a hatched background, 239

How to change and fix the Y-axis, 240

How to add a second Y-axis, 240

How to change and fix the X-axis, 241

How to save a layout, 242

How to apply a layout, 242

How to use the zoom function, 244

How to cut a curve and store as new, 245

How to reduce noise, 274

How to remove ghost peaks, 274

How to import a blank run curve, 276

How to subtract a blank curve, 276

How to add, 277

How to rename, 286

How to compare peaks in different curves, 288

Multifile Peak Compare Wizard, 288

Manual peak identification, 294

Commands to import curves into a chromatogram, 305

How to use the Open to compare command, 306

How to import using File:Open, 309

How to copy curves into one chromatogram, 310

How to align with Normalise, 312

How to move using the Shift function, 315

How to stretch or shrink using Multiply, 315

How to produce a mirror image, 316

How to shift a mirror image, 316

How to import, 319

Export options, 321

How to export, 321

How to export in AIA format, 323

How to delete unwanted curves, 326

How to divide, 366

How to reduce noise, 367

How to remove ghost peaks, 367

How to differentiate, 370

Simulated peak fractionation, 372

How to create, 373

Draw a straight curve between selected points, 376

How to create a fraction histogram, 377

Monitor signals stored as curves, 463

Curve settings, 463

Calibration curve models, 521

Molecular size curve models, 521

Curves pane in System Control

Description, 199

How to select curves to be monitored, 199

How to display a vertical marker, 199

How to set a reference point, 200

How to change curve colors and styles, 200

How to change scale of the Y-axis, 200

How to change scale of the X-axis, 201

How to zoom in regions of the pane, 202

Reduce scale of zoom, 202

How to select curve pressure units, 202

How to select text alignment, 202

How to display complete Logbook information, 203

D

Delete files and folders, 77

Delete Method blocks

How to use the Delete Block dialog box, 104

How to use the Block: Delete Block command, 105

How to delete unused blocks, 105

Documentation

How to view, 270 Documentation tabs, description, 270 Result information, 272 How to export, 324

Ε

Electronic signature

How to sign a result, 325

Evaluation

Chromatogram window views, 229

How to display chromatogram header information, 229

How to display peak table information, 230

Chromatogram window shortcut menu, 231

How to optimize the chromatogram workspace, 231

How to display a vertical marker, 232

How to set a reference point, 232

How to make chromatogram layout changes, general, 235

How to exit the module, 326

Evaluation logs

How to export, 324

Evaluation procedures

How to delete, 137

How to rename, 138

How to edit, 138

Explained variance

Definition, 525

External standard quantitation

How to perform, 397

F

File Navigator

How to open result files, 69

How to open, 222

How to locate files from the Files list, 222

How to use Find to locate a file, 223

How to open a recent run, 224

How to change preference settings for Recent Runs, 225

How to close, 225

Files and folders

How to copy, 75

How to move, 75

Copy to external, 75

How to copy from external, 76

Flow Scheme pane

Description, 204

Stretch to fit screen, 204

How to view manual instructions, 204

Folders

How to create a user-specific, 68

FPLCdirector

How to import data, 319

Frac-950

Defining the number of available tubes, 149

How to select the last tube, 150

Fraction Histogram

How to create a curve, 377

Fractions

How to set up the Frac unit, 149

How to view the contents of a fraction, 279

How to pool fractions, 279

How to create a pooled fraction curve, 280

Show only the pooled fraction curve, 281

How to calculate protein amounts and concentrations in pooled fractions, 282

How to determine the volume of a pooled fraction with a specific concentration, 282

How to use the Pool Fraction print list, 282

G

Generate Report Wizard

How to generate a report from the UNICORN Manager, 470

Gradient

Effects of Change and Replace on gradient length, 116

Gradients

Instruction parameters, 164

Step gradient instruction, 164

Gradient breakpoints, 164

Text instructions, 165

Define length as a variable, 165

ı

Instant Run
How to start, 191
Internal standard quantitation
Suitable components, 400
How to perform, 400
Reliability, 403

L

Linear flow rates

Description, 163

Log on and log off routines

How to start the program, 46

How to log on, 46

Log off alternatives, 47

Log off and set a password for a running process, 47

Unlock the system, 48

Quit UNICORN after log off, 49

Logbook

How to display an overlay in the Curves pane in System Control, 203 How to display an overlay in the chromatogram window, 232 How to filter the information, 238

Logbook pane

Description, 205
Autoscroll function, 205
How to filter the contents, 205
Search function, 206

M

Maintenance

How to view maintenance information, 465
How to set up a maintenance warning, 466
How to view warning parameters, 466
How to reset warning counters, 467
How to use the Generate Report Wizard from the UNICORN Manager, 470
How to use the Generate Report Wizard from the System Control, 473
Manual direct commands

Buttons in System Control, 208

Manual instructions in System Control

During a method run, 212

Column prompt, 212

Functions of buttons, 213

How to save results manually, 214

Measurements

How to make direct, 363

Messages

Usage, 160

How to issue, 160

How to add a message instruction to a method, 555

Method blocks

Description, 98

Blocks in the Text pane, 98

How to show or hide instructions, 98

Blocks in the Block pane, 99

Blocks in the Gradient pane, 99

Calls, 100

Unconditional calls, 100

Conditional calls, 100

Use the Instruction box to add, 101

Use the New Block dialog box to add, 102

Fields of the New Block dialog box, description, 102

Use the Delete Block dialog box, 104

How to use the Delete Block command, 105

How to delete unused blocks, 105

How to rename blocks, 106

How to find text strings, 107

How to copy a block, 107

How to move a block, 108

How to import, general information, 109

How to import, 109

Base instructions, descriptions, 155

Block length, 158

Method Editor

Modes, 28

Text instructions display panes, 29

The Block pane, 29

The Flow Scheme pane, 30

The Gradient pane, 30

The Text pane, 30 The Instruction box, 31 Icon descriptions, 94 Log Format, 159

Method files

How to open in the UNICORN Manager, 69 How to connect a method to a system, 77

Method instructions

Instruction markings, 112

How to add an instruction, 113

Pause, Hold and Hold_until instructions, 113

How to delete instructions, 114

Undo delete, 114

How to change, 115

Difference in function between Change and Replace command, 115

Move an instruction within the same breakpoint, 117

Move an instruction to another breakpoint, 117

Instructions at the same breakpoints, 157

Method runs

Start from the UNICORN Manager, 190

Start from System Control, 190

How to define methods as menu commands, 190

How to start an Instant Run, 191

How to use the Start Protocol, 191

How to start a method run when the system is busy, 191

Run Data pane, description, 196

Curves pane in System Control module, 199

Flow Scheme pane, description, 204

Logbook pane, description, 205

Scouting runs, 216

How to perform a MethodQueue run, 217

If network communication fails, 219

Method templates

How to create a method, 85

Template information, 86

Save a new method, 86

How to create a template from a method, 171

How to delete a template, 171

Method Wizard

How to create a method, 81

How to save a new method, 83

MethodQueue

How to create a new, 183

How to use several systems in a queue, 184

Relative timing of steps, 185

Unattended execution, 185

Temporary hold when system is busy, 186

Folder handling, 187

File handling, 187

How to edit a MethodQueue, 187

How to perform a MethodQueue run, 217

Unattended operation, 217

Start when the system is busy, 217

How to display and edit pending and running MethodQueues, 218

Methods

How to create using a wizard, 81

How to sign, 91

Different method editing operations, 96

Method variables, general description, 118

How to select a column, 156

Hold instruction, description, 162

Pause instruction, description, 162

Hold_until instruction, description, 162

Linear flow rates, 163

Gradient instruction parameters, 164

How to print, 172

How to export, 174

How to export, 324

Monitor signal limits, 460

Molecular size calculation

Overview, 450

Preparations before curve creation, 451

How to create a size table, 452

How to open an existing table, 454

How to rename a molecular size table, 454

How to delete a molecular size table, 455

How to calculate the size, 455

Error signs, 457

Procedure instruction, 457

Molecular size curve

Overview, 449

Description, 453

Morphological algorithm

Description, 336

How to set, 336

Structure width, 337

Incorrect structure width, 338

Noise window, 338

Minimum distance between points, 339

Definition, 493

Multifile Peak Compare Wizard

How to select the operation, 288

How to select data to compare, 290

How to select the peaks, 292

Manual peak identification, 294

How to select the Peak Data, 295

The Data View dialog box, 296

How to use the 2D data view, 297

How to use the 3D data view, 299

How to save the settings, 301

How to open saved settings, 301

P

Peak integration

How to perform, 330

Differences between to filter peaks and to reject peaks, 333

How to display peak labels, 334

How to select part of a curve for peak integration, 358

Peak skim

Compared to drop-lines, 360

How to select a ratio, 360

Peak table

How to display information, 230

How to rename, 286

How to export, 323

How to select contents, 364

Peaks

How to filter from view, 333

Labels, 334

How to display peak labels, 334

How to open the peak table, 351

How to delete a peak, 353

How to add a fill color and pattern, 354

Drop-lines, description, 355

How to split a peak, 355

How to join peaks, 356

How to add peak names, 356

How to exclude before integration, 359

Include negative peaks in integration, 359

How to select a skim ratio, 360

Edit integration for part of a curve, 361

Peak purity, 366

Peak identification through the absorbance ratio, 366

Peak parameters, 497

How to change the peak resolution algorithm, 500

How to change the Assymetry Ratio value, 502

Pool Fraction print list

How to use, 282

PrimeView

How to import data, 320

Problem reports

How to use the Generate Report Wizard from the UNICORN Manager, 470

How to use the Generate Report Wizard from the System Control, 473

Procedure instructions

Curve operations, 504

Integration, 507

File operations, 508

Export functions, 509

Chromatogram functions, 515

Miscellaneous, 516

Test instructions, 518

Procedures

How to record, 379

Global procedures, 380

How to build a procedure with instructions, 380

How to edit, 382

How to add instructions, 383

How to run a single procedure, 384

How to batch run, 384

How to rename, 387

How to delete, 387

Protein activity

Match to UV curve, 285

Protein amounts

Calculation formula, 282

Protein concentrations

Calculation formula, 282

Q

Quality control

How to create a control procedure, 557

How to add a control procedure to a method, 559

Quantitation

General description, 394

Process steps, 394

Procedure instructions, 396

How to use an external standard, 397

External standard reliability, 399

Internal standard, description, 400

How to use an internal standard, 400

Standard addition, description, 404

How to use standard addition, 404

Standard addition reliability, 405

Recovery calculation, description, 406

How to use recovery calculation, 406

Recovery calculation, reliability, 407

General reliability factors, 408

Preparations before, 410

Standard concentration levels, 410

Reject peaks, 410

How to create a quantitation table, 411

How to select table components, 414

Peak identification, 415

Relative retention, 416

How to adjust peak identification settings, 416

Peak identification criteria, 417

How to create a calibration curve, 418

How to enter standard data, 418

How to create a calibration curve, 418

How to enter standard data, 418

How to select an Internal Standard, 420

Statistics, 421

How to open a table for editing, 422

How to update a calibration curve, 423

How to rename the table, 427

How to delete a quantitation table, 427

How to prepare for, 429

How to calculate the amount and concentration, 430

How to view the results, 431

Calculation error signs, 431

Standard addition stages, 432

How to prepare for standard addition, 432

How to select the standard addition component, 433

How to prepare for recovery calculation, 435

How to calculate the recovery, 435

Automated sample runs, 441

Automatic update with Replace, 442

Automatic update with Average, 443

Automated in scouting runs, 444

Automated update with Average for scouting runs, 447

Quick View

How to preview result files, 70

R

Recovery calculation

How to perform, 406

Reliability, 407

Recovery factor calculation

How to prepare for recovery calculation, 435

How to calculate the recovery, 435

Error signs, 436

Reference curves

In Run Setup, 140

Rename files and folders, 77

Reports

How to create a blank customized report, 250

Edit mode toolbar buttons, 251

How to add or delete pages, 252

How to change the page setup, 252

How to add objects to a report, 253

How to add free text, 254

How to add picture objects, 255

How to include chromatograms, 256

How to include a peak table, 256

How to include a pool table, 256

How to include Method objects, 257

How to add documentation, 258

How to add the Evaluation log, 258

How to include Quantitate and Mol. Size data, 259

How to include Frac-950 data, 259

Toolbar icons in Report Edit Mode, 260

How to print, 261

How to save the report in PDF format, 262

How to save the report format, 263

How to create a Standard report, 264

How to print a standard report, 265

How to edit a standard report, 267

How to edit a customized report, 268

Result file name

Name options, 147

Serial numbers, 148

Unique identifier, 148

Result files

How to open in the UNICORN Manager, 69

How to open in the File Navigator, 69

Automated printing of, 136

Specify folder for storing, 148

How to open in UNICORN Manager, 221

How to open in UNICORN Manager, 221

Electronic signature, 325

How to save, 326

Run Data pane

Description, 196

How to change the appearance, 196

How to change text color or background, 197

How to set pressure units, 197

How to view and edit manual instructions, 197

Run Setup

Description, 28

Run Setup from the Method Wizard, 82

Tabs, description, 123

Variable tab view options, 125

How to change variable values, 125

Blue variable values, 125

How to delete variables, 126

How to rename variables, 126

How to change a variable into a detail variable, 126

How to change a detail variable into a regular variable, 127

Scouting tab, 128

Questions tab, description, 129

Question status, alternatives, 129

Mandatory questions, 129

Authorized questions, 129

Chromatogram questions, 129

Questions, answer types, 129

How to insert a question, 130

How to edit questions, 131

Gradient tab, description, 133

Gradient zoom function, 133

Gradient marker line, 134

Gradient, change X-axis base, 134

Gradient hatch marks, 134

Notes, description, 135

How to write method notes, 135

How to search for text in the method notes, 135

Evaluation Procedures tab, 136

How to import evaluation procedures in the method, 137

How to edit method procedures, 138

Reference curves, 140

How to add reference curves, 140

Columns tab, 142

BufferPrep, description, 143

How to create a BufferPrep method, 144

BufferPrep correction factors, 145

Method Information tab, description, 146

Result Name tab, description, 147

Result file serial numbers, 148

Result file unique identifier, 148

Batch ID, 148

Start Protocol contents, 151

How to export values, 153

Scouting tab buttons, 176

S

Scouting

Specify folder for storing results, 148

Start Protocol settings, 152 Usage, 176 Changing variables, 176 Change variable values, 176 How to set up a Scouting Scheme, 177 How to edit a Scouting Scheme, 177 How to rename variables, 179 How to delete a variable, 179 How to change a variable into a detail variable, 179 How to copy variable content, 180 How to define columns, 181 How to perform a Scouting run, 216 Result files, 216 Scouting runs Change of variables during a run, 216 Searches General functions, 37 Security Backup, 78 Set_Mark Usage, 160 How to issue, 161 Slope values Usage, 369 How to measure, 370 SMART Manager How to import data, 319 Smoothing algorithms Moving average, 490 Autoregressive, 490 Median, 491 Savitzky-Golay, 492 Snapshots How to view, 41 How to add a text instruction, 90 Standard addition quantitation How to perform, 404 Reliability, 405 How to identify sample peaks, 433 How to select the component, 433

Start Protocol

How to use, 191

Statistics

Correlation, 525

Explained variance, 525

Stock solutions

Description, 143

Strategy

How to display the strategy instructions, 90

System Control module

Description, 32

Overview, 194

How to select the displayed panes, 194

How to customize the panes, 195

Toolbar buttons, 208

Manual instructions during a method run, 212

Manual instructions, 212

How to save manual results, 214

Alarms and warnings, 215

System Control status bar

Description, 210

Watch status, 211

System Control Toolbar

Manual command buttons, availability, 208

Manual command buttons, functions, 208

Windows buttons, 209

System Access buttons, 210

System data

How to backup, 63

How to restore backup data, 64

System operation

Automated workstation lock or logoff, 48

Unlock the system, 48

Unlock system locked by other users, 48

How to connect to a system, 58

Connection modes, 59

How to leave control of the system, 60

How to disconnect a system, 60

System settings

System summary, 61

How to change default settings, 459

How to assign a new value, 460

T

Temporary chromatogram

Description, 227

Text instructions

Message instruction, 160

Set_Mark instruction, 161

Hold instructions, 162

Pause instruction, 162

Hold_until instruction, 162

Linear flow rates, 163

How to form a step gradient instruction, 164

Gradient breakpoints, 164

Gradient instructions, 165

Watch instructions, 166

How to insert a Watch instruction, 166

Watch parameter options, 167

Text Instructions Editor

When to use, 88

How to edit instructions, 88

Save a new method, 89

How to select panes, 95

Toolbar icons

In the System Control module, 33

Troubleshooting

Logon problems, 477

Strategy file error, 478

Access problems, 479

Connection problems, 479

Method problems, 482

Incorrect time and date, 487

Evaluation procedure aborts, 487

ÄKTAdesign system problems, 488

U

Unconditional call

Description, 100

Unconditional method instructions

Base instruction, 155

UNICORN Manager

Limited access to, 27

٧

Variables

General description, 118
Identification in text instructions, 118
How to change method variable values, 119
Breakpoints or gradient lengths, 119
How to define new method variables, 119
Variable names, 120
How to rename a method variable, 120
How to remove a method variable, 121

W

Warnings

Description, 461

Watch instructions

Description, 100

Standard Watch conditions, 166

How to insert an instruction, 166

Parameter options, 167

Air sensors, 167

Permanent settings, 168

Temporary settings, 168

Delta_Peak settings, 168

Delta_Base settings, 169

Watch Stable_baseline, 170

Wizard

How to create a method, 81

Υ

Y-axis

How to choose the Y-axis scale, 231

Z

Zero baseline

Index

Definition, 329

Zoom function

How to enlarge parts of a curve, 244