

Waters Micromass Quattro Premier Mass Spectrometer

Operator's Guide

Waters

34 Maple Street
Milford, MA 01757

71500073002 Revision B

NOTICE

The information in this document is subject to change without notice and should not be construed as a commitment by Waters Corporation. Waters Corporation assumes no responsibility for any errors that may appear in this document. This document is believed to be complete and accurate at the time of publication. In no event shall Waters Corporation be liable for incidental or consequential damages in connection with, or arising from, the use of this document.

© 2004 WATERS CORPORATION. PRINTED IN THE UNITED STATES OF AMERICA AND IRELAND. ALL RIGHTS RESERVED. THIS DOCUMENT OR PARTS THEREOF MAY NOT BE REPRODUCED IN ANY FORM WITHOUT THE WRITTEN PERMISSION OF THE PUBLISHER.

Micromass and Waters are registered trademarks, and ESCi, MassLynx, MaxEnt, MUX-technology, Quattro Premier, T-Wave, and ZSpray are trademarks of Waters Corporation.

PEEK is a trademark of Victrex Corporation.

Rheodyne is a registered trademark of Rheodyne, L.P.

Upchurch is a registered trademark of Scivex, Inc.

Windows is a registered trademark of Microsoft Corporation.

All other trademarks or registered trademarks are the sole property of their respective owners.



Safety Information

General

The Waters® Micromass® Quattro Premier™ Mass Spectrometer is designed solely for use as a mass spectrometer; any attempt to use it for any other purpose can damage the instrument, and will invalidate its warranty.

The Waters Micromass Quattro Premier Mass Spectrometer conforms to European standard EN61010-1:2001, Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements.

The instrument has been designed and tested according to recognized safety standards. If the instrument is used in a manner not specified by the manufacturer, the protection provided to the user may be impaired.

Whenever the safety protection of the instrument has been compromised, disconnect the instrument from all power sources, and secure the instrument against unintended operation.

The instrument must be installed so that the operator can easily isolate the instrument from the mains power supply at all times.

Biological Hazard

When you analyze physiological fluids, take all necessary precautions, and treat all specimens as potentially infectious. Precautions are outlined in “CDC Guidelines on Specimen Handling,” *CDC – NIH Manual*, 1984.

Suitable protection against biohazards must be taken during maintenance procedures and cleaning, as parts of the instrument are exposed to potentially infectious samples.

Chemical Hazard

Good Laboratory Practice should be adhered to when using potentially toxic, caustic, or flammable solvents and analytes.

Flammable Solvents Operation Hazard



Caution: *If the nitrogen supply pressure falls below 4 bar (58 psi), the instrument stops the nitrogen flow and admits air into the source. If flammable solvents are used, an ignition hazard exists under these conditions.*

If flammable solvents are used, ensure that the nitrogen supply pressure does not fall below 4 bar (58 psi) during the analysis. Also ensure that the Gas Fail connection (see [Section 1.6.3](#)) is connected to the HPLC system to ensure that the LC flow is stopped on nitrogen supply failure.

High Voltage Hazard



Warning: *Certain areas of the instrument may have high voltages present when the instrument is in Operate. To avoid non-lethal electric shock, make sure the instrument is in Standby before touching these areas.*

Certain areas of the instrument may have high voltages present when the instrument is in Operate. These areas are shown in [Figure 1](#), showing the instrument configured for ESI operation, and [Figure 2](#), showing the instrument configured for APcI operation.



Warning: *To avoid electric shock (non-lethal), any equipment connected to the ESI probe or optional MUX-technology interface should be earthed.*

Waters recommends that any equipment connected to the ESI probe or optional MUX-technology interface be earthed.

High Temperature Hazard



Warning: *To avoid burns, take care when working with the instrument as the source enclosure assembly may be at high temperature.*

The source enclosure assembly may be at high temperature, as shown in [Figure 3](#), showing the instrument configured for ESI operation, and [Figure 4](#), showing the instrument configured for APcI operation.

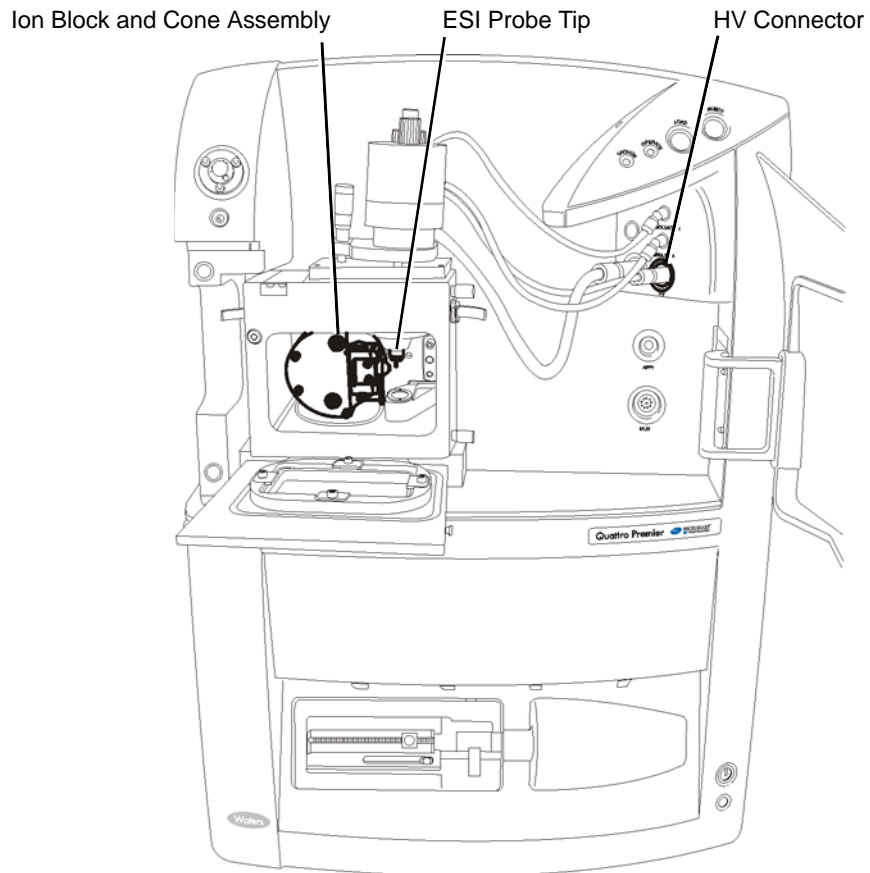


Figure 1 Quattro Premier High Voltage Hazards - ESI Operation

Ion Block and Cone Assembly

Corona Discharge Pin

HV Connector

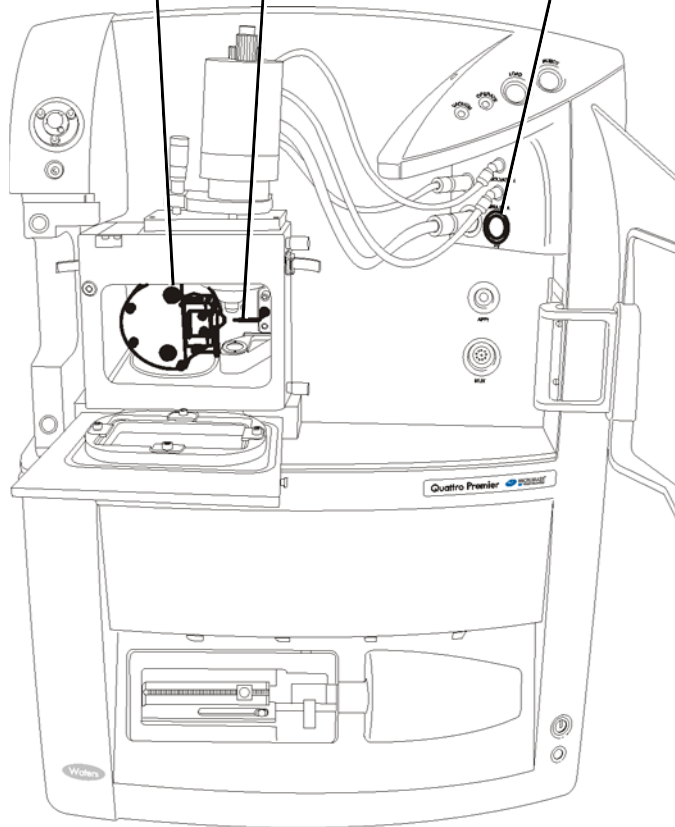


Figure 2 Quattro Premier High Voltage Hazards - APCl Operation

Source Enclosure Assembly

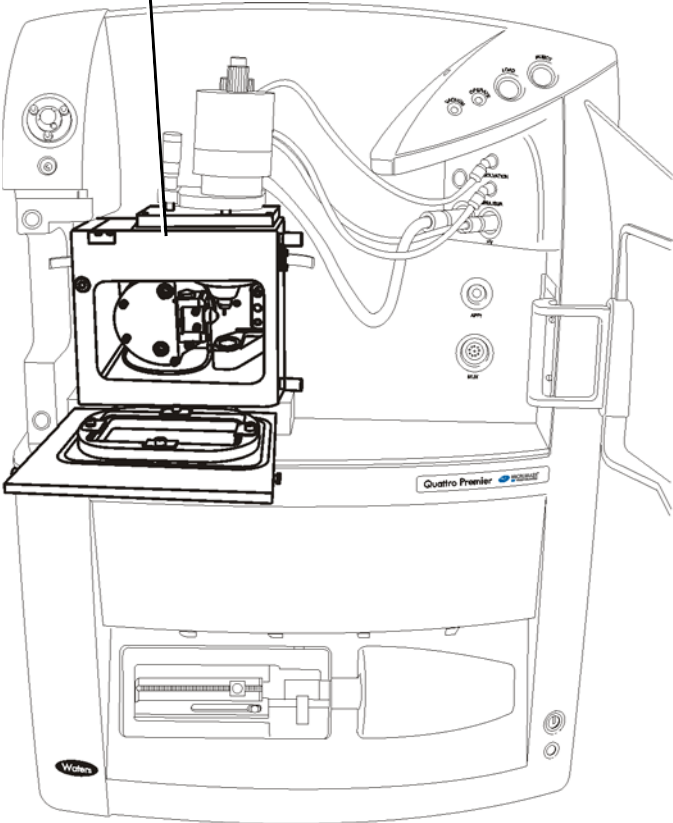


Figure 3 Quattro Premier High Temperature Hazard - ESI Operation

Source Enclosure Assembly

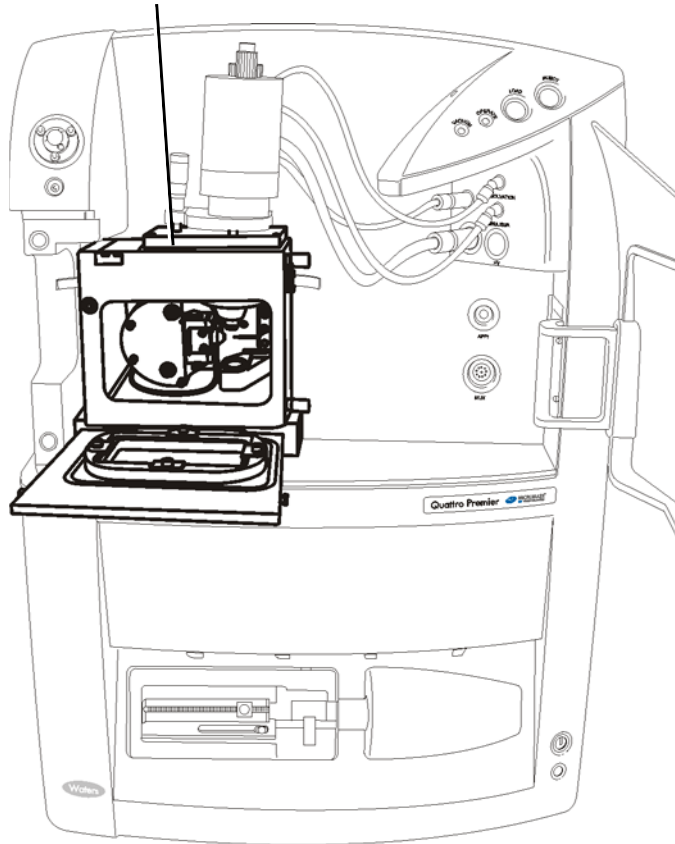


Figure 4 Quattro Premier High Temperature Hazard - AP/PCl Operation

Safety Symbols

Warnings in this guide or on the instrument must be observed during all phases of service, repair, installation, and operation of the instrument. Failure to comply with these precautions violates the safety standards of the design and intended use of the instrument.

Waters Corporation assumes no liability for the user's failure to comply with these requirements.

Safety symbols are used in this guide and on the instrument. A **Warning** is an instruction that draws the user's attention to the risk of injury or death; a **Caution** is an instruction that draws attention to the risk of damage to the instrument.

Consignes de sécurité

Généralités

Le Quattro Premier™ de Waters® Micromass® est destiné exclusivement à être utilisé comme spectromètre de masse. Tout usage détourné du Quattro Premier risquerait d'endommager l'instrument et invaliderait sa garantie.

Le spectromètre de masse Quattro Premier de Waters Micromass est conforme à la norme européenne EN61010-1:2001, Règles de sécurité pour appareils électriques de mesurage, de régulation et de laboratoire - Partie 1: Prescriptions générales.

Cet instrument a été conçu et testé conformément aux dispositions des normes de sécurité les plus reconnues. Toute utilisation de l'équipement non conforme aux instructions du fabricant risque de rendre déficiente la protection assurée par l'instrument.

Dans le cas où la sécurité de l'utilisateur se trouverait compromise, débranchez le cordon d'alimentation de l'instrument et assurez-vous qu'il ne pourra être mis en marche par mégarde.

L'instrument doit être installé de façon à faciliter l'accès de l'utilisateur au bloc d'alimentation électrique.

Risques biologiques

Lorsque vous analysez des fluides physiologiques, faites preuve d'une extrême prudence et considérez tous les spécimens comme potentiellement infectieux. Une liste des précautions à prendre figure dans le « CDC Guidelines on Specimen Handling », CDC - NIH Manual, 1984.

Portez des vêtements de protection adéquats lorsque vous procédez à une opération de maintenance ou de nettoyage, car les pièces de l'instrument peuvent comporter des traces des substances infectieuses auxquelles elles ont été exposées.

Risques chimiques

L'usage des solvants et analytes potentiellement toxiques, caustiques ou inflammables doit s'effectuer dans le respect des bonnes pratiques de laboratoire.

Risques liés à l'usage de solvants inflammables



Attention: Si la pression d'alimentation en azote tombe en dessous de 4 bars, l'instrument bloque automatiquement l'arrivée d'azote et déclenche une arrivée d'air dans la source. L'usage de solvants inflammables implique l'existence d'un risque d'ignition.

Lorsque vous utilisez des solvants inflammables, assurez-vous que la pression d'alimentation en azote ne tombe pas en dessous de 4 bars en cours d'analyse. De plus, assurez-vous que la connexion Gas Fail (voir [Section 1.6.3](#)) est correctement raccordée au système HPLC, de sorte que le débit LC soit interrompu en même temps que l'alimentation en azote.

Risques d'électrocution



Avertissement: Certaines parties de l'instrument peuvent être soumises à des tensions électriques élevées lorsque l'instrument est en mode de fonctionnement. Pour éviter toute électrocution accidentelle, veuillez placer l'instrument en mode veille.

Lorsque l'instrument est en mode de fonctionnement ou « Operate », certaines parties de l'instrument sont soumises à des tensions très élevées. Ces parties de l'instrument sont indiquées dans les figures 1 et 2. La [Figure 1](#) montre l'instrument configuré pour un usage en mode d'ionisation electrospray (ESI), et la [Figure 2](#) montre l'instrument configuré pour un usage en mode d'ionisation chimique à pression atmosphérique (APcI).



Avertissement: Pour éviter toute électrocution (non mortelle), tout l'équipement relié à la sonde ESI ou à l'interface MUX-technology, devrait être mis à la terre.

Waters recommande que tout l'équipement relié à la sonde ESI ou à l'interface MUX-technology soit mis à la terre.

Risques de brûlure



Avertissement: Pour éviter toute brûlure, faites attention lorsque vous travaillez à proximité de la source car elle peut atteindre des températures très élevées.

La source, représentée aux figures 3 et 4, peut atteindre des températures très élevées. La [Figure 3](#) montre l'instrument configuré pour un usage en mode ESI, et la [Figure 4](#) montre l'instrument configuré pour un usage en mode APcI.

Pictogrammes de sécurité

Les avertissements présents dans le manuel de l'utilisateur ou sur l'instrument-même doivent être scrupuleusement pris en considération, et ce à tout moment, que ce soit pendant l'entretien, la réparation, l'installation ou le fonctionnement de l'instrument. Tout défaut d'application de ces règles de sécurité serait considéré comme une violation des normes de sécurité relatives à la conception et à l'usage prévu de l'instrument.

Waters Corporation ne saurait voir sa responsabilité engagée en cas de manquement de l'utilisateur à respecter les consignes de sécurité.

Les pictogrammes qui suivent apparaissent dans le manuel de l'utilisateur ou sur l'instrument. L'**Avertissement** s'applique à toute instruction destinée à attirer l'attention de l'utilisateur sur l'existence d'un risque de blessure ou de mort. L'**Attention** s'applique à toute instruction destinée à informer l'utilisateur d'une situation qui peut endommager l'instrument.



Warning: General warning indicating a potential health or safety hazard. See the operator's guide for instructions.



Avertissement: Symbole d'avertissement indiquant qu'un produit ou composant pose un risque potentiel pour votre santé et sécurité. Consultez le manuel de l'utilisateur.



Warning: Hazardous voltages.



Avertissement: Tensions dangereuses.



Warning: Hot surfaces.



Avertissement: Surfaces chaudes.



Warning: Corrosive substances.



Avertissement: Substances corrosives.



Warning: Biological agents present that may constitute a serious health threat.



Avertissement: Présence d'agents biologiques susceptibles d'être nuisibles à la santé.



Warning: Toxic substances.



Avertissement: Substances toxiques.



Warning: Flammable substances.



Avertissement: Substances inflammables.



Warning: Laser radiation.



Avertissement: Rayonnements laser.



Warning: UV radiation.



Avertissement: Rayonnements UV.



Caution: Care must be taken to avoid the possibility of damaging the instrument, or affecting its operation.



Attention: Utilisez l'instrument en faisant preuve de beaucoup de précaution pour éviter de l'endommager et ainsi nuire à son fonctionnement.

Quattro Premier Mass Spectrometer Information

Intended Use

The Waters Micromass Quattro Premier Mass Spectrometer can be used as a research tool to deliver authenticated mass measurement in both MS and MS/MS modes.

The Quattro Premier is considered a General *in vitro* Diagnostic Device according to European Union Directive 98/79/EC.

Calibration

Follow acceptable methods of calibration with pure standards to calibrate methods. Use a minimum of five standards to generate a standard curve. The concentration range should cover the entire range of quality-control samples, typical specimens, and atypical specimens.

Quality Control

Routinely run three quality-control samples. Quality-control samples should represent subnormal, normal, and above-normal levels of a compound. Ensure that quality-control sample results are within an acceptable range, and evaluate precision from day to day, and run to run. Data collected when quality-control samples are out of range may not be valid. Do not report this data until you ensure that system performance is acceptable.

Table of Contents

Safety Information	iii
Consignes de sécurité	ix
Quattro Premier Mass Spectrometer Information	xiii
Chapter 1	
Instrument Description	1
1.1 Overview	1
1.2 Sample Inlet.....	3
1.3 Vacuum System	3
1.4 MassLynx Control System	3
1.5 Front Panel Controls, Indicators, and Connections.....	4
1.5.1 Power Switch	4
1.5.2 Embedded PC Reset Switch	4
1.5.3 Cone Gas, Desolvation Gas, and Nebulizer Gas Connections	6
1.5.4 Electrical Connections	7
1.5.5 Status Display	7
1.5.6 Divert/Injection Valve	8
1.6 Rear Panel Connections	10
1.6.1 Analog Channels	10
1.6.2 Contact Closure	11
1.6.3 Gas Fail	11
1.6.4 CE Int (Capillary Electrophoresis Interlock)	12
1.6.5 Comm For EPC	12
1.6.6 Com1	12
1.6.7 Aux O/P	12
1.6.8 Pump Relay	12
1.6.9 GAS IN	13

1.6.10	Collision Gas In	14
1.6.11	Exhaust	14
1.6.12	Waste	14
1.6.13	Mains Power Connection	14
1.6.14	Mains Power Fuses	14
1.6.15	Vacuum Connections	14
1.7	Top Panel.....	15
1.8	Mass Flow Controllers	15

Chapter 2

Setting Up and Tuning for ESI	17
2.1 Setting Up.....	17
2.1.1 Removing the APCI Probe	17
2.1.2 Removing the APCI Corona Discharge Pin	19
2.1.3 Installing the ESI (Electrospray) Probe	20
2.2 Tuning via Sample Infusion	22
2.2.1 Tuning for MS Operation	22
2.2.2 Tuning for MS/MS (Daughter Ion) Operation	29
2.2.3 Creating the MRM MS Method File	30
2.3 Preparing the Instrument for Data Acquisition	32
2.3.1 Reconfiguring the Connection to the Probe	32
2.3.2 Configuring the Inlet for LC Operation	33
2.3.3 Creating a Sample List and Starting Data Acquisition	33
2.3.4 Viewing and Printing the Tuning Parameters Associated with a Data File	34

Chapter 3

Setting Up and Tuning for APCI	35
3.1 Setting Up.....	35
3.1.1 Removing the ESI Probe	35
3.1.2 Installing the APCI Corona Discharge Pin	36

3.1.3	Installing the APCI Probe	38
3.2	Tuning via Sample Infusion	40
3.2.1	Tuning for MS Operation	40
3.2.2	Tuning for MS/MS (Daughter Ion) Operation	46
3.2.3	Creating the MRM MS Method File	48
3.3	Preparing the Instrument for Data Acquisition	50
3.3.1	Reconfiguring the Connection to the Probe	50
3.3.2	Configuring the Inlet for LC Operation	51
3.3.3	Creating a Sample List and Starting Data Acquisition	51
3.3.4	Viewing and Printing the Tuning Parameters Associated with a Data File	52
Chapter 4		
ESCI Multi-Mode Operation		53
4.1	Preparing the Instrument	53
4.2	Tuning the Instrument	54
4.3	Data Acquisition.....	55
Chapter 5		
Acquiring Data		57
5.1	Starting an Acquisition.....	57
5.1.1	Starting an Acquisition from the Tune Window	57
5.1.2	Starting Multiple-Sample Acquisition from the MassLynx Window	60
5.1.3	Automatic Quantification of the Sample List	63
5.2	Monitoring an Acquisition	65
5.2.1	Viewing the Status of an Acquisition	65
5.2.2	Viewing a Chromatogram in Real-Time	65
5.2.3	Viewing a Spectrum in Real-Time	65
5.3	Selecting Instrument Data Thresholds	66
5.3.1	Profile Data	67

5.3.2	Centroid Data	68
5.3.3	SIR Data	68
5.3.4	Ion Counting Threshold	68
5.3.5	Profile Data – Spike Removal	69
5.3.6	Analog Data	69
5.4	Checking the Status of the Communications Link.....	70
5.5	Stopping an Acquisition	70
5.6	Pausing an Acquisition.....	70
5.7	Setting-Up Scanning Functions.....	71
5.7.1	Creating a Function List	71
5.7.2	Setting-Up a Full Scan Function	77
5.7.3	Setting-Up an SIR Function	81
5.7.4	Setting-Up MS/MS Scanning Functions	84
5.7.5	Setting-Up an MRM Function	88
5.7.6	Setting-Up a Survey Function	88
5.7.7	Monitoring Acquisitions	111

Chapter 6

Setting-Up Mass Calibration	113
6.1 Overview	113
6.1.1 Types of Calibration	113
6.1.2 Calibration Process	114
6.2 Calibrating for Electrospray Operation.....	114
6.2.1 Preparing for Calibration	114
6.2.2 Selecting the Calibration Options	118
6.2.3 Setting the Calibration Parameters	119
6.2.4 Performing a Calibration	123
6.2.5 Calibration Failure	127
6.2.6 Incorrect Calibration	128
6.2.7 Manually Checking the Calibration	129

6.2.8	Manually Editing the Peak Matching	131
6.2.9	Saving the Calibration	132
6.2.10	Verifying the Calibration	132

Chapter 7

Maintenance Procedures 133

7.1	Maintenance Schedule.....	133
7.2	Safety and Handling.....	134
7.3	Gas-Ballasting the Rotary Pump (E2M28)	135
7.4	Checking the Rotary Pump Oil	137
7.5	Changing the Rotary Pump Oil	138
7.5.1	Required Materials	138
7.5.2	Procedure	138
7.6	Replacing the Oil Mist and Odor Filter Elements.....	139
7.6.1	Required Materials	139
7.6.2	Procedure	139
7.7	Cleaning the Source Components	142
7.7.1	Overview	142
7.7.2	Required Materials	143
7.7.3	Removing the Probe from the Source	143
7.7.4	Removing the Sample Cone	145
7.7.5	Removing the Gas Exhaust Port	150
7.7.6	Removing the Ion Source Enclosure and Ion Block	151
7.7.7	Removing the Source T-Wave Assembly from the Instrument	153
7.7.8	Disassembling the Source T-Wave Assembly	155
7.7.9	Disassembling the Source Ion Block	157
7.7.10	Cleaning the Sample Cone and Cone Gas Cone	165
7.7.11	Cleaning the Gas Exhaust Port	165
7.7.12	Cleaning the Source T-Wave Components	166
7.7.13	Cleaning the Ion Block and Extraction Cone	167

7.7.14	Cleaning the Isolation Valve Stem	168
7.7.15	Reassembling the Source Ion Block	168
7.7.16	Reassembling the Source T-Wave Assembly	169
7.7.17	Fitting the Source T-Wave Assembly to the Instrument	169
7.7.18	Fitting the Ion Block and Ion Source Enclosure	170
7.7.19	Fitting the Gas Exhaust Port	170
7.7.20	Fitting the Sample Cone	170
7.8	Cleaning or Replacing the ESI Probe Tip	171
7.8.1	Required Materials	171
7.8.2	Procedure	171
7.9	Cleaning or Replacing the Corona Discharge Pin.....	172
7.9.1	Required Materials	172
7.9.2	Procedure	172
7.10	Cleaning the APCI Probe Tip.....	173
7.11	Replacing the Ion Block Cartridge Heater	174
7.11.1	Required Materials	174
7.11.2	Procedure	174
7.12	Replacing the ESI Probe Sample Capillary.....	178
7.12.1	Required Materials	178
7.12.2	Removing the Existing Capillary	179
7.12.3	Installing the New Capillary	183
7.13	Replacing the APCI Probe Sample Capillary	187
7.13.1	Required Materials	187
7.13.2	Removing the Existing Capillary	188
7.13.3	Installing the New Capillary	192
7.14	Replacing the APCI Probe Heater.....	197
7.14.1	Required Materials	197
7.14.2	Procedure	198

Chapter 8

Troubleshooting	201
8.1 Spare Parts	201
8.2 Safety and Handling	201
8.3 System Troubleshooting	201
8.4 Component Hardware Troubleshooting	203
8.4.1 Power Switch Fails to Power-up the Instrument	203
8.4.2 No Peaks in the Tune Window (No Ion Beam)	203
8.4.3 Unsteady or Low Intensity Peaks (Ion Beam)	205
8.4.4 Unusually High LC Backpressure	206
8.4.5 Unusually Low LC Backpressure	207
8.4.6 Insufficient Vacuum	207
8.4.7 Leaking Nitrogen	208
8.4.8 Rotary Pump Oil Accumulated in the Exhaust Tubing	208
8.4.9 Ion Source Heater and Desolvation Heater are Not Working	209
8.4.10 APCI Probe Heater Not Working	209
8.4.11 Failure of the Fuse Supplying the Rotary Pump	209
8.4.12 Ion Mode Fault	210
8.4.13 Failure to Recognize a Particular Probe Type	210
8.4.14 Ripple	211
8.4.15 Loss of Communication with the Instrument	211
8.4.16 IEEE Communication Errors	212
8.5 High Noise Levels in MRM Analyses	212
8.5.1 Chemical Noise	213
8.5.2 Electronic Noise	213
8.6 Contacting Waters	214

Appendix A

Starting Up and Shutting Down the Instrument	215
A.1 Starting Up the Instrument	215

A.2	Shutting Down the Instrument	218
A.2.1	Emergency Shutdown	218
A.2.2	Overnight Instrument Shutdown	218
A.2.3	Complete Instrument Shutdown	218

Appendix B

Setting-Up the Syringe Pump	221
--	------------

Appendix C

The MassLynx Tune Window	223	
C.1	Opening the MassLynx Tune Window	223
C.2	Selecting the Ionization Mode.....	223
C.3	Controlling Gas Flows	225
C.3.1	Controlling the Nebulizer, Desolvation, and Cone Gas Flows	225
C.3.2	Controlling the Collision Gas Flow	225
C.4	Controlling the Syringe Pump.....	225
C.5	Selecting the Syringe Type.....	225
C.6	Selecting the Scan Time and Inter Scan Delay	226
C.7	Setting the Ramp Controls	226
C.7.1	Creating a Cone Voltage Ramp	226
C.7.2	Controlling the Cone Voltage Ramp	227
C.7.3	Creating a Collision Energy Ramp	227
C.7.4	Controlling the Collision Energy Ramp	228
C.8	Resetting the Zero Level	228
C.9	Controlling the Display of Readback Windows.....	228
C.10	Changing Tune Parameter Settings	229
C.11	Saving Instrument Tune Parameters.....	229
C.11.1	Creating a New Instrument Parameter File	230
C.11.2	Saving Tune Parameters in an Instrument Parameter File	230
C.11.3	Opening an Existing Instrument Parameter File	230
C.12	Printing Tune Information.....	230

C.13	Using the EasyTune Source Page.....	231
C.14	Changing Between the Peak and Vacuum Displays.....	231
C.15	Setting-Up Peaks for Tuning.....	231
	C.15.1 Selecting Peaks for Tuning	231
	C.15.2 Selecting the Operating Mode for a Peak	232
	C.15.3 Selecting the Tune Mass for a Peak	232
	C.15.4 Selecting the Span of a Displayed Peak	233
	C.15.5 Changing the Gain of a Displayed Peak	234
C.16	Customizing the Peak Display	234
	C.16.1 Opening the Peak Display Menu	234
	C.16.2 Customizing the Colors and Numbers of Displayed Traces	235
	C.16.3 Customizing the Peak Trace Line Appearance	236
	C.16.4 Customizing the Peak Intensity Display	236
	C.16.5 Customizing the Peak Display Grid	236
C.17	Selecting the Instrument Name	237
C.18	The Diagnostics Page.....	237
C.19	Manually Controlling the T-Wave Optics	237

Appendix D

Calibration Reference Information	241	
D.1 Overview	241	
D.2 Editing a Calibration Reference File	241	
D.3 Positive Ion Calibration Reference Files	242	
	D.3.1 Polyethylene Glycol	243
	D.3.2 Sodium Iodide and Cesium Iodide Mixture	243
	D.3.3 Sodium Iodide and Rubidium Iodide Mixture	244
D.4 Negative Ion Calibration Reference Files	244	
D.5 Preparing Reference Sample Solutions	245	
	D.5.1 Preparing the PEGNH ₄ Reference Sample Solution	245

D.5.2	Preparing the Sodium Iodide and Cesium Iodide Mixture Sample Solution	245
Appendix E		
Performance Specifications		247
E.1	Electrospray Positive Ion	247
E.2	Electrospray Negative Ion	247
E.3	MS Resolution	247
E.4	Mass Measurement Accuracy	248
E.5	APCI Positive Ion	248
Appendix F		
Theory and Principles of Operation		249
F.1	Ionization Techniques.....	249
F.1.1	Electrospray Ionization (ESI)	249
F.1.2	Atmospheric Pressure Chemical Ionization (APCI)	250
F.2	Ion Optics	250
F.3	MS Operating Modes	251
F.4	MS/MS Operating Modes	252
F.4.1	Daughter (Product) Ion Mode	252
F.4.2	Parent (Precursor) Ion Mode	254
F.4.3	Multiple Reaction Monitoring (MRM) Mode	255
F.4.4	Constant Neutral Loss Mode	256
F.4.5	Source and Collision Cell T-Wave Devices	257
Index		259

Chapter 1

Instrument Description

1.1 Overview

The Quattro Premier™ (Figure 1-1) is a high-performance tandem quadrupole mass spectrometer designed for routine LC/MS/MS operation.



Figure 1-1 Quattro Premier Mass Spectrometer

The Quattro Premier may be coupled to either of two types of inlet:

- An HPLC system, to provide molecular weight information from an LC run or perform target analysis and quantification.
- A syringe pump, for analysis of precious, low-concentration compounds.

The sample is ionized at atmospheric pressure in the source. The ions enter the vacuum system through a sampling cone, then pass through the source travelling-wave (T-Wave™) ion guide into the first quadrupole, where they are filtered according to their mass-to-charge ratio (m/z) (Figure 1-2). The mass-separated ions pass into the T-Wave collision cell where they either undergo collision-induced decomposition (CID) or pass to the second quadrupole. Any fragment ions are then mass-analyzed by the second quadrupole. The transmitted ions are detected by a conversion dynode, phosphor, and photomultiplier detection system. The output signal is then amplified, digitized, and passed to the control system.

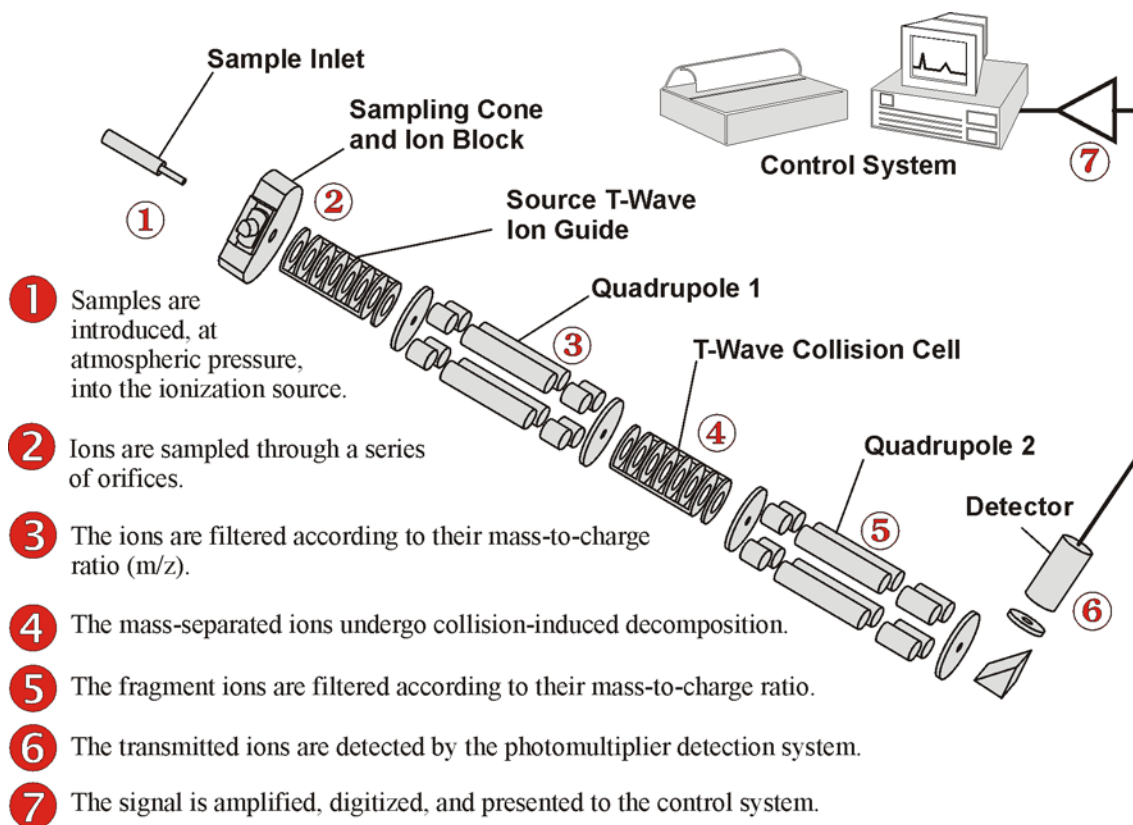


Figure 1-2 Quattro Premier Schematic

1.2 Sample Inlet

An HPLC system or infusion pump delivers sample to a ZSpray™ Ionization Source.

The ionization technique (see [Section F.1](#)) is selected by installing the appropriate probe. An ESI (electrospray ionization) probe is supplied as standard. An APCI (atmospheric pressure chemical ionization) is available as an option. Recognition pins on the probe identify the ionization method to the system. The source can also be operated as an ESCi™ multi-mode ionization source to combine ESI and APCI operation in a single run (see [Chapter 4](#)). In this case the ESI probe is used together with an APCI corona discharge pin, which is supplied as standard.

1.3 Vacuum System

An external backing pump and internal turbomolecular pumps generate vacuum in the instrument. The backing pump may be a rotary pump or, optionally, an oil-free scroll pump.

The control system monitors the turbomolecular pump speeds and continuously monitors the vacuum using an integral Pirani gauge. The Pirani gauge stops instrument operation if a vacuum loss is detected.

A vacuum isolation valve allows routine source maintenance to be performed without breaking the vacuum.

1.4 MassLynx Control System

The Quattro Premier is controlled by the PC-based MassLynx™ software. MassLynx also controls the HPLC system, autosampler, and divert/injector valve, if applicable. A second PC, embedded in the Quattro Premier, processes the acquired data. Communication between the MassLynx PC and the embedded PC is via a network link.

Analog inputs allow data acquisition from conventional LC detectors like an ultraviolet (UV) detector or evaporative light scattering detector (ELSD). Data from selected UV photo diode array detectors (for example, the Waters 996 PDA detector) can also be acquired.

The MassLynx software allows the following processes:

- Configuring the Quattro Premier.
- Creating inlet and MS methods that define operating parameters for a run.
- Tuning and calibrating the Quattro Premier.
- Running samples.
- Monitoring the run.
- Acquiring data.

See the *MassLynx User's Guide* and *MassLynx Help* for more information on installing and using the MassLynx software.

1.5 Front Panel Controls, Indicators, and Connections

Figure 1-3 shows a general view of the front of the instrument with the access door open.

1.5.1 Power Switch



Warning: *The power switch does not isolate the instrument from the mains power supply. To do this, disconnect the power supply cord (see [Section 1.6.13](#)) from the rear of the instrument.*

The instrument's power switch is located on the lower-right side corner of the front panel.

Note: *As the power switch does not isolate the instrument from the mains power supply, fans may be heard running, even when the instrument is off.*

1.5.2 Embedded PC Reset Switch

The embedded PC reset switch resets the embedded PC and instrument's electronics. This switch is accessed through a hole in the instrument's front panel and can be operated by means of a short length of PEEK™ tubing, or similar object.

If the instrument is in Operate, the embedded PC reset switch switches it into Standby in a controlled manner, even if communication with the MassLynx PC is lost. This may cause the backing pump to momentarily start or stop.

When the embedded PC reboots, it will take a short time to re-establish communication with the MassLynx PC.

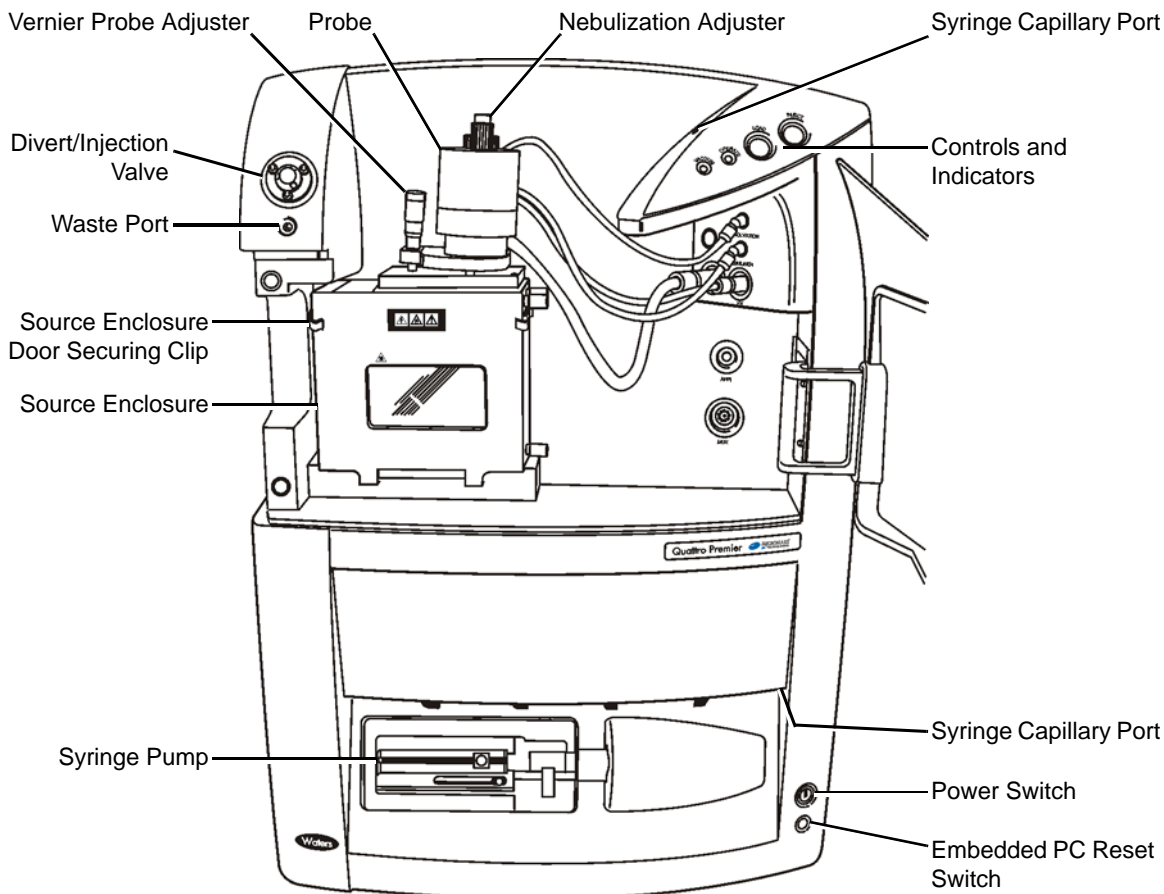


Figure 1-3 Front View with the Access Door Open

After the switch is operated, the high voltages will be switched off and the turbomolecular pumps will gradually slow down.

If the reset switch is operated when the MassLynx PC is offline, and the turbomolecular pump speed falls below 50% of full speed, the vent solenoid will open and admit air into the analyzer. The backing pump will be switched off a few seconds later.

If the instrument is vented fully, it must be evacuated again by selecting Options > Pump in the MassLynx Tune window (see [Appendix C](#)).

If the turbomolecular pump speed is above 50% of full speed when the embedded PC has booted-up, the embedded PC cancels the vent sequence, and the instrument evacuates again.

Note: On certain early instruments, the embedded PC will not cancel the vent sequence after it has booted-up. You must select Options > Pump on the Tune window to stop the instrument being vented.

1.5.3 Cone Gas, Desolvation Gas, and Nebulizer Gas Connections

The PTFE gas lines for the probe desolvation gas and nebulizer gas are connected to the front of the instrument by push-in fittings (Figure 1-4). The connection for the cone gas is inside the source; it also uses PTFE tubing.

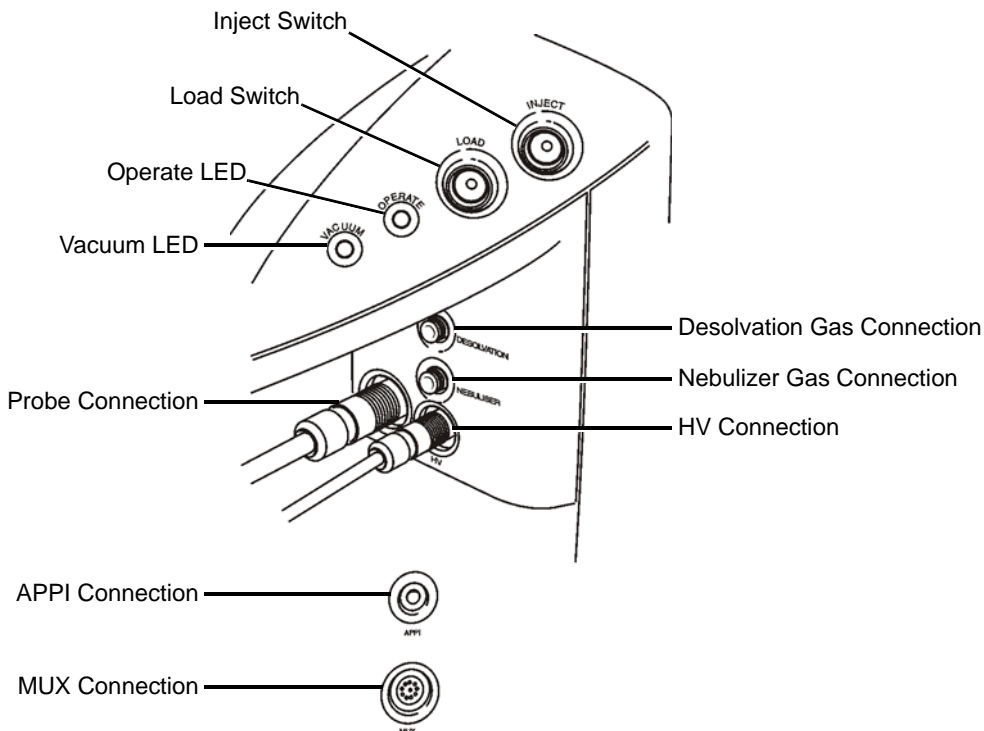


Figure 1-4 Front Panel Connections, Controls, and Indicators

1.5.4 Electrical Connections

The electrical connection for the ESI desolvation heater or APCI probe heater is via the Probe multi-way connector.

The high voltage connection for the ESI probe is via the front panel HV connection.

The APPI connection is used by the optional Combined APPI and APCI Source (see the *Waters Micromass Quattro Premier Combined APPI and APCI Source Operator's Guide* for details).

The MUX connection is used by the optional MUX-technology™ Interface (see the *Waters Micromass Quattro Premier MUX-technology Interface Operator's Guide* for details).

The high voltage connection for the corona discharge pin is inside the source.

1.5.5 Status Display

The Vacuum and Operate status light-emitting diodes (LEDs) are at the top-right corner of the instrument's front panel (see [Figure 1-4](#)).

The instrument's status is indicated as shown in [Tables 1-1 and 1-2](#).

Table 1-1 Vacuum LED Display

State	Vacuum LED Indication
Pumping	Flashing green
Pumped, below trip level	Steady green
Pumped, above trip level	Steady amber
Venting, if the turbomolecular pump speeds are above 50% of full speed	Flashing red
Venting, if the turbomolecular pump speeds are below 50% of full speed	Flashing amber, for approximately 10 seconds before the vented state is achieved
Vented	Steady red

Table 1-2 Operate LED Display

State	Operate LED Indication
Standby	No indication
Operate, above trip level	Steady amber
Operate, below trip level	Steady green
RF error	Flashing red
Tripped out of Operate due to low nitrogen gas supply pressure	Flashing amber
<p>Note: To return the instrument to Operate, you must:</p> <ol style="list-style-type: none"> 1. Re-establish the nitrogen supply. 2. In the Tune window, click Press for Standby. 3. In the Tune window, click Press for Operate. 	

Note: Any combination of LED indications not covered by [Tables 1-1 and 1-2](#) indicates an instrument fault. Contact Waters for advice (see [Section 8.6](#)).

1.5.6 Divert/Injection Valve



Warning: The liquids passing through the divert/injection valve may be biohazardous and/or toxic. Always wear nitrile gloves when working with the divert/injection valve.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when working with the divert/injection valve.

The divert/injection valve ([Figure 1-5](#)) is at the top-left corner of the instrument front panel (see [Figure 1-3](#)). It is an electrically driven Rheodyne[®] injector, which can be used in several ways, depending on the plumbing arrangement:

- As an injection valve (with the needle port and sample loop fitted).
- As a divert valve (to switch the flow of solvent during an LC run.)
- As a switching valve (for example, to switch between an LC system and a syringe pump containing calibrant).

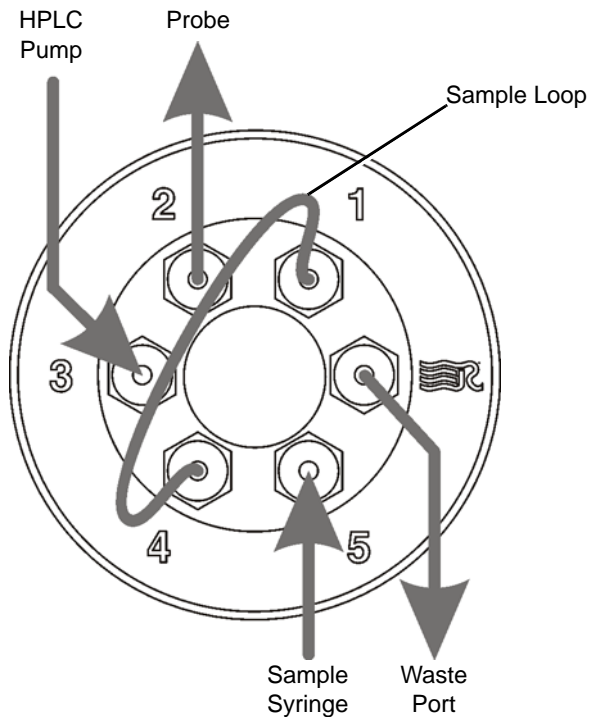


Figure 1-5 Divert/Injection Valve

The valve is controlled by MassLynx. The two switches, Load and Inject, at the top-right corner of the instrument front panel (see [Figure 1-4](#)), allow you to control the valve directly when making loop injections at the instrument.

For details of using the valve as a divert valve, see [“Setting Solvent Delays” on page 74](#).

1.6 Rear Panel Connections

The rear panel connections are shown in [Figure 1-6](#).

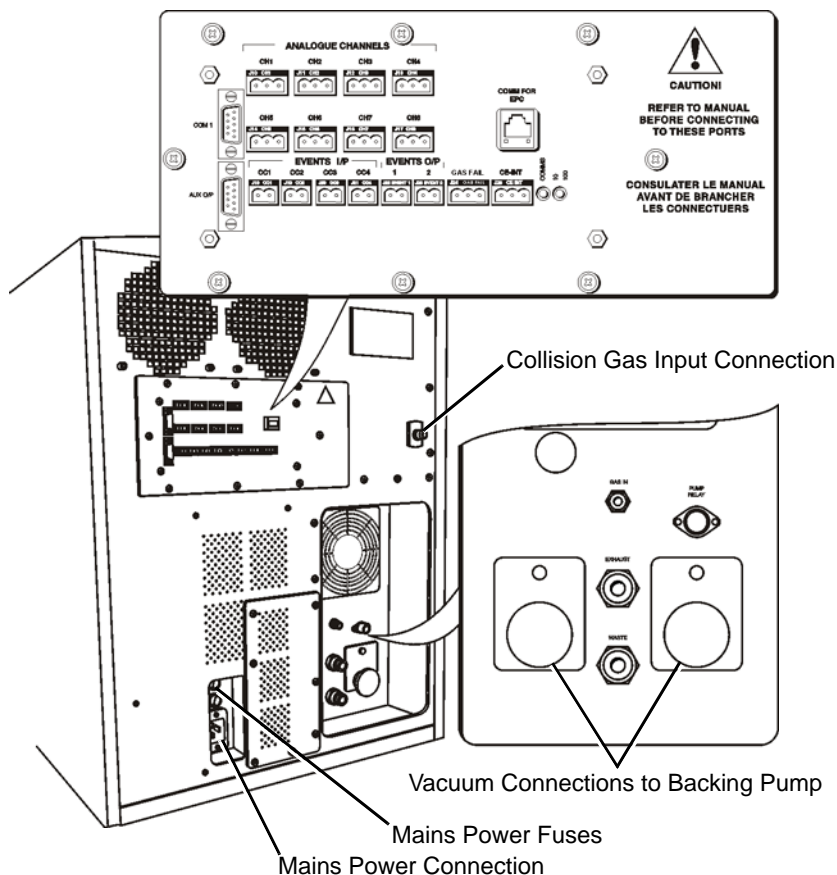


Figure 1-6 Rear Panel Connections

1.6.1 Analog Channels



Warning: To avoid electric shock and damage to the instrument, do not apply more than 16 V to any of the Analogue Channels connections.

Eight Analogue Channels inputs (CH1 to CH8) are available for acquiring simultaneous data such as a UV detector output (see [“Acquiring Analog Data”](#) on page 75 for further details).

Note: Although eight inputs are available, the MassLynx software can only control four of the channels.

Signals up to 2.5 V may be input. Analog data is processed by a 20-bit analog-to-digital converter. The maximum (overload) input is ± 16 V.

If the input cable is a two-wire assembly, the negative pole of each channel may need to be grounded (earthed).

1.6.2 Contact Closure



Warning: To avoid electric shock and damage to the instrument, do not apply more than 5 V to any of the Events I/P connections.



Warning: To avoid electric shock and damage to the instrument, do not apply more than 25 V to either of the Events O/P connections.

Two types of contact closure are available:

- **In** – Four inputs, Events I/P CC1 to CC4, allow external devices to start acquisition. Each event input signal can be transistor-transistor logic or contact closure. The maximum voltage is 5 V.
- **Out** – Two outputs, Events O/P 1 and 2, allow the mass spectrometer to trigger an external event. The maximum rating is 25 V, 0.5 A.

1.6.3 Gas Fail



Warning: To avoid electric shock and damage to the instrument, do not apply more than 25 V to the Gas Fail connection.

If the nitrogen supply pressure falls below 4 bar (58 psi), or the instrument's power supply fails, a contact closure signal is generated. This signal can be used to stop solvent flowing into the source by connecting this Gas Fail connection to the Stop Flow of the HPLC system. If the nitrogen supply fails, any solvent from the LC will be automatically drained from the source enclosure. The maximum rating is 25 V, 0.5 A.

1.6.4 CE Int (Capillary Electrophoresis Interlock)



Warning: To avoid electric shock and damage to the instrument, do not apply more than 25 V to the CE Int connection.

This connector interfaces with a capillary electrophoresis power supply so that the instrument is safely interlocked against high voltages. The maximum rating is 25 V, 0.5 A.

1.6.5 Comm For EPC

This RJ45 connector links the instrument's embedded PC to the MassLynx PC using the network cable supplied.

1.6.6 Com1

This connection can be used by a Waters field service engineer to communicate with the embedded PC.

1.6.7 Aux O/P



Caution: The Aux O/P connection must not be used, unless permitted by Waters.

This connection is used for connecting to auxiliary equipment.

1.6.8 Pump Relay

This connects to a backing pump interlock box, which allows the instrument to remotely control the backing pump ([Figure 1-7](#)).

Note: If the optional scroll pump is used, it may be supplied with a backing pump interlock box, or, alternatively, it may be supplied with a cable that directly connects the Quattro Premier to the scroll pump ([Figure 1-8](#)); this allows the Quattro Premier to control the scroll pump directly.

The scroll pump's on/off switch must be set to "on" if it is connected to a backing pump interlock box.

The scroll pump's on/off switch must be set to "off" if it is directly connected to, and controlled by, the Quattro Premier.



Warning: The backing pump is independently powered; hence, the pump interlock box can contain power even when the Quattro Premier is isolated from the mains power supply. To isolate the pump interlock box and backing pump, switch off the mains power to the pump interlock box.

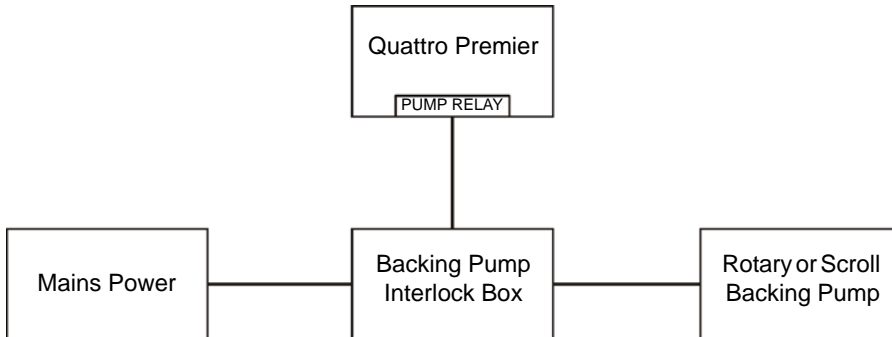


Figure 1-7 Rotary or Scroll Backing Pump Controlled Via a Backing Pump Interlock Box

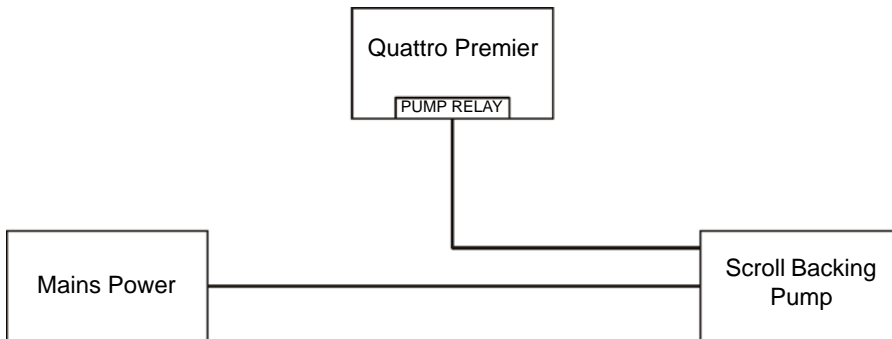


Figure 1-8 Scroll Backing Pump Controlled Directly by the Quattro Premier

1.6.9 GAS IN



Caution: If the nitrogen supply pressure falls below 4 bar (58 psi), the instrument stops the nitrogen flow and admits air into the source. If flammable solvents are used, an ignition hazard exists under these conditions.

The nitrogen gas supply is connected to this connection. The nitrogen must be dry and oil-free, with a purity of at least 95%. Regulate the supply at 6 to 7 bar (90 to 105 psi).

1.6.10 Collision Gas In

The collision gas supply is connected to this connection. The collision gas is argon; it must be dry and of high purity (99.9%). Regulate the supply at 0.5 bar (7 psi).

1.6.11 Exhaust



Warning: LC solvents and analytes may be carried in the nitrogen exhaust. Vent the nitrogen exhaust to atmosphere outside the laboratory.

This is the nitrogen gas exhaust connection. Vent the nitrogen to the atmosphere outside the laboratory.

1.6.12 Waste



Warning: The waste liquid from the source enclosure and the divert/injection valve comprises LC solvents and analytes. Always wear nitrile gloves while handling the container, and ensure that the waste liquid is correctly disposed of according to local environmental regulations.

Waste liquid from the top of the instrument, the source enclosure, and the divert/injection valve is drained from the instrument via this connection. The liquid must be passed into a suitable container, which you must empty at regular intervals.

1.6.13 Mains Power Connection

This is the mains power connection for the instrument.

1.6.14 Mains Power Fuses



Warning: Parts of the instrument may be electrically live even when a mains fuse has failed. To avoid electrical shock, isolate the instrument from the mains supply before replacing the mains fuses.

Two mains power fuses are located adjacent to the mains power connection. The fuses are rated at T10 AH 250 V.

If a one fuse fails, replace both fuses.

1.6.15 Vacuum Connections

Two vacuum connections are provided for connection to the backing pump.

1.7 Top Panel



Caution: To avoid accidental spillage damaging the instrument, the instrument's top panel must not be used for storing large volume solvent reservoirs.

The instrument's top panel may be used for storing small items, for example, small solvent bottles.

1.8 Mass Flow Controllers

The cone gas, desolvation gas, and collision gas flow rates are regulated by electronic mass flow controllers, which are controlled from the MassLynx Tune window. [Table 1-3](#) shows the flow rate ranges for these gasses.

If the nitrogen supply fails, solvent is prevented from accumulating in the source enclosure as described in [Section 1.6.3](#).

Table 1-3 Gas Flow Rate Ranges

Gas	Flow Rate Range
Cone (nitrogen)	0 to 300 L/h
Desolvation (nitrogen)	0 to 1200 L/h
Collision (argon)	0 to 1 mL/min

Chapter 2

Setting Up and Tuning for ESI

Note: For full details of the Tune window, see [Appendix C](#).

2.1 Setting Up

2.1.1 Removing the APCI Probe

You may need to remove the APCI probe and corona discharge pin from the instrument before fitting the ESI probe. [Figure 2-1](#) shows the APCI probe mounted on the instrument.




Warning: To avoid electric shock, ensure that the instrument is in Standby before starting this procedure.

1. In the MassLynx Tune window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.



Warning: The liquid passing through the HPLC pump, LC column, and APCI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.

2. Disconnect the LC system from the APCI probe.
3. Wait for three minutes to allow the desolvation gas flow to cool the probe and source.
4. In the MassLynx Tune window, click  to stop the nitrogen flow.



Warning: The probe and source may be hot. To avoid burns, take great care while working with the instrument's access door open.

5. Open the instrument's access door.
6. Disconnect the probe electrical connection on the instrument front panel.

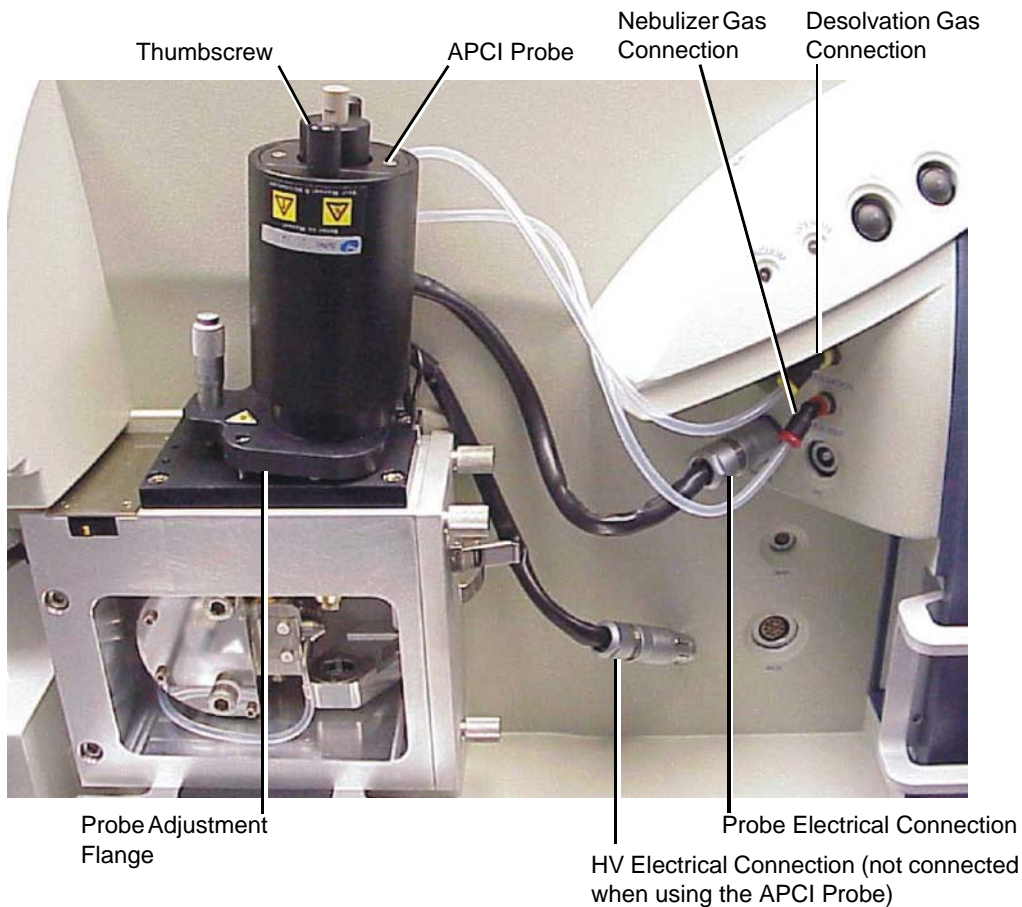


Figure 2-1 APCI Probe Mounted on the Source Enclosure

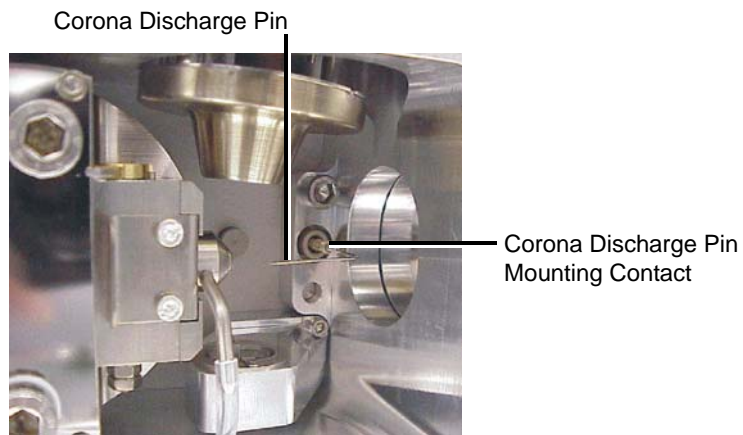
7. Disconnect the PTFE tubes at the nebulizer and desolvation gas connections on the front panel.
8. Undo the two thumbscrews securing the probe to the probe adjustment flange.
9. Carefully remove the probe from the probe adjustment flange.

2.1.2 Removing the APCI Corona Discharge Pin



Warning: To avoid electric shock, ensure that the instrument is in Standby when installing the corona discharge pin.

1. In the MassLynx Tune window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.
2. Unfasten the source enclosure door's securing clips and open the door.
3. Use needle-nose pliers to remove the corona discharge pin from its mounting contact. Store the corona discharge pin in a safe location.



4. Use the needle-nose pliers to fit the blanking plug to the corona discharge pin mounting contact.

Corona Discharge Pin Mounting Contact Blanking Plug



2.1.3 Installing the ESI (Electrospray) Probe



Warning: The probe and source may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while handling these components.



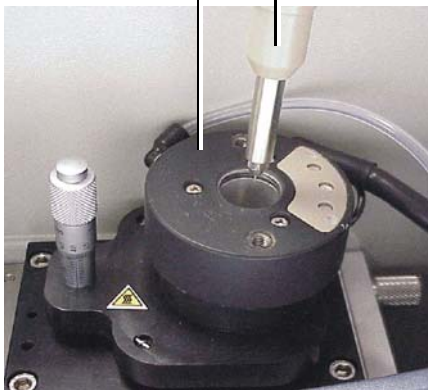
Warning: The probe and source may be hot. To avoid burns, take great care while working with the instrument's access door open.



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.

1. In the MassLynx Tune window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.
2. Open the instrument's access door.
3. Remove the protective sleeve, if fitted, from the electrospray probe tip.
4. Carefully slide the probe into the hole in the probe adjustment flange.

Probe Adjustment Flange Probe



5. Secure the probe by tightening the two thumbscrews ([Figure 2-2](#)).

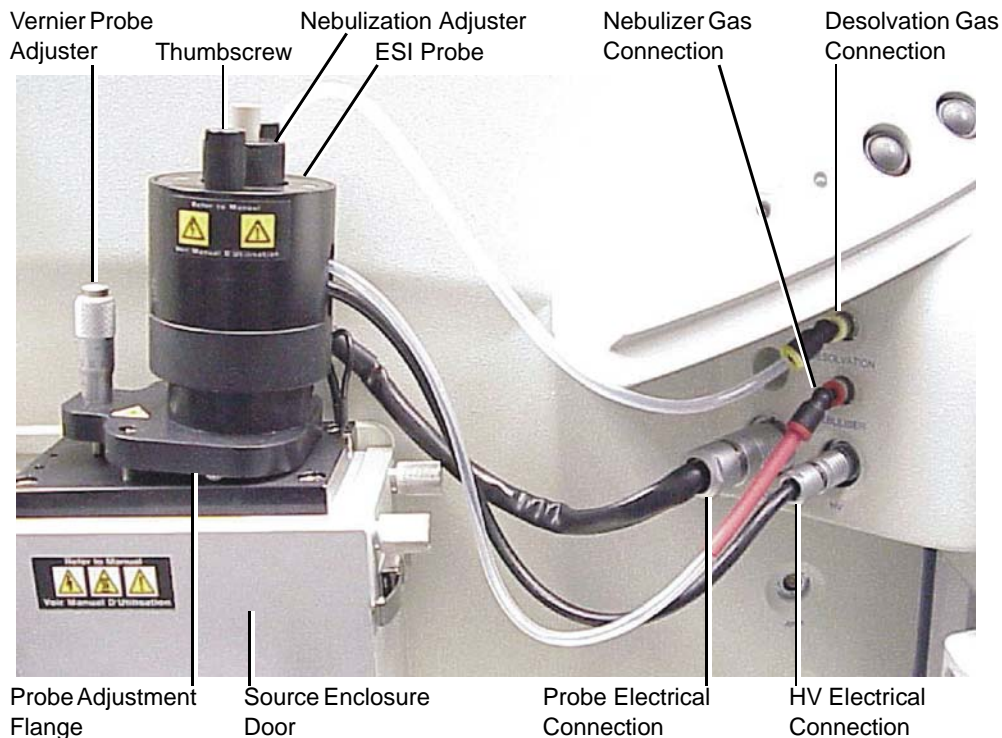


Figure 2-2 ESI Probe Mounted on the Source Enclosure, Showing the Connections to the Front Panel

6. Connect the probe adjustment flange electrical cable to the Probe connection.
7. Connect the probe adjustment flange PTFE tube to the Desolvation gas connection.
8. Connect the probe PTFE tube to the Nebuliser gas connection.
9. Connect the probe electrical lead to the HV connection.
10. Close the instrument's access door.

2.2 Tuning via Sample Infusion

Note: This example procedure specifies verapamil as the infused sample. Therefore, the choice of HPLC column and some parameter settings may be specific to that compound. You may tune the instrument using a different sample compound, however you may need to use a different type of column, and parameter settings may differ from the ones given here.

2.2.1 Tuning for MS Operation



Warning: The liquids passing through the HPLC pump, LC column, syringe pump, and ESI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when making the connections between the HPLC pump, LC column, syringe pump, and ESI probe.



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.

1. In the MassLynx Tune window, click Press for Standby, and confirm that the adjacent instrument status indicator shows red.
2. Complete the connections between the HPLC pump, syringe pump, and ESI probe as shown in [Figure 2-3](#).

The column used in this example is a Waters Symmetry[®] C₁₈ 2.1 × 100-mm, 3.5- μ m. The mobile phase used is 70:30 acetonitrile/water.

3. Look through the source's view port and confirm that the isolation valve lever is fully to the left (i.e., the valve is open) ([Figure 2-4](#)).



Warning: The source may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while handling this component.



Warning: The source may be hot. To avoid burns, take great care while working with the instrument's access door open.

If the isolation valve lever is in the wrong position:

- a. Open the instrument's access door.

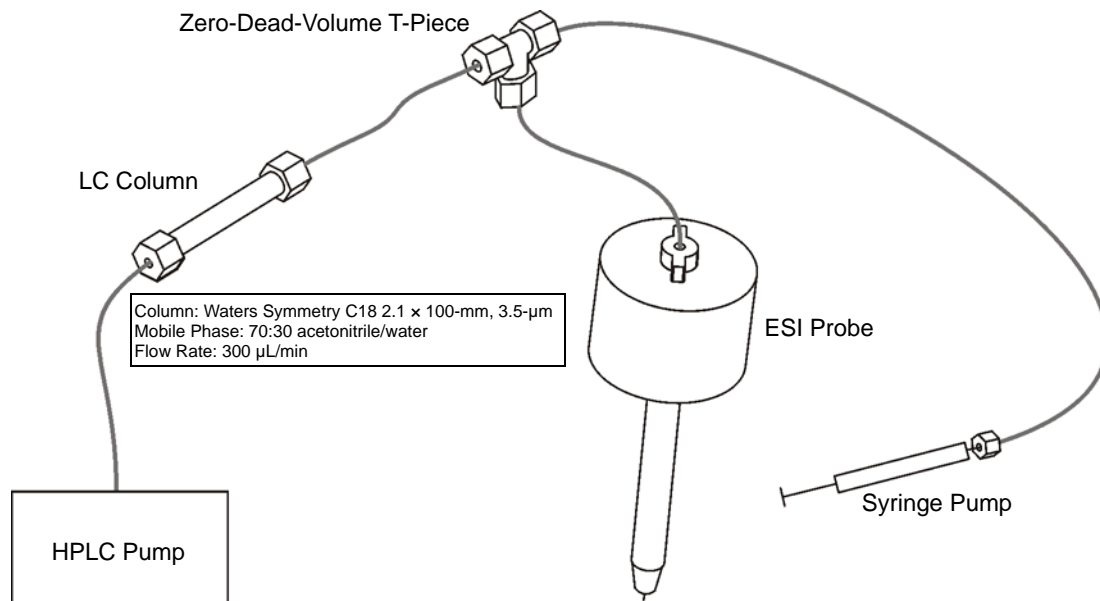


Figure 2-3 Syringe and Column Connections to the ESI Probe

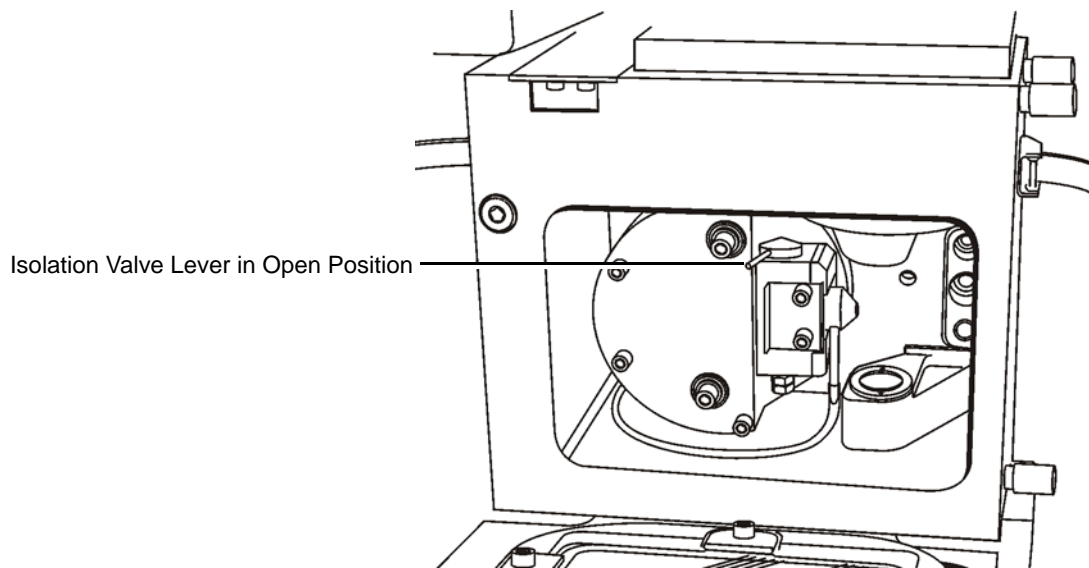


Figure 2-4 Isolation Valve in the Open Position (Source Enclosure Door Shown Open for Clarity)

- b. Unfasten the source enclosure door's securing clips, and open the door.
 - c. Move the lever to the open position.
 - d. Close the source enclosure door, and fasten the securing clips.
 - e. Close the instrument's access door.
4. In the MassLynx Tune window, select Ion Mode > Electrospray+ to display the ES+ Source page (Figure 2-5).

Note: The instrument can also be tuned via the EasyTune Source page (see Section C.13).

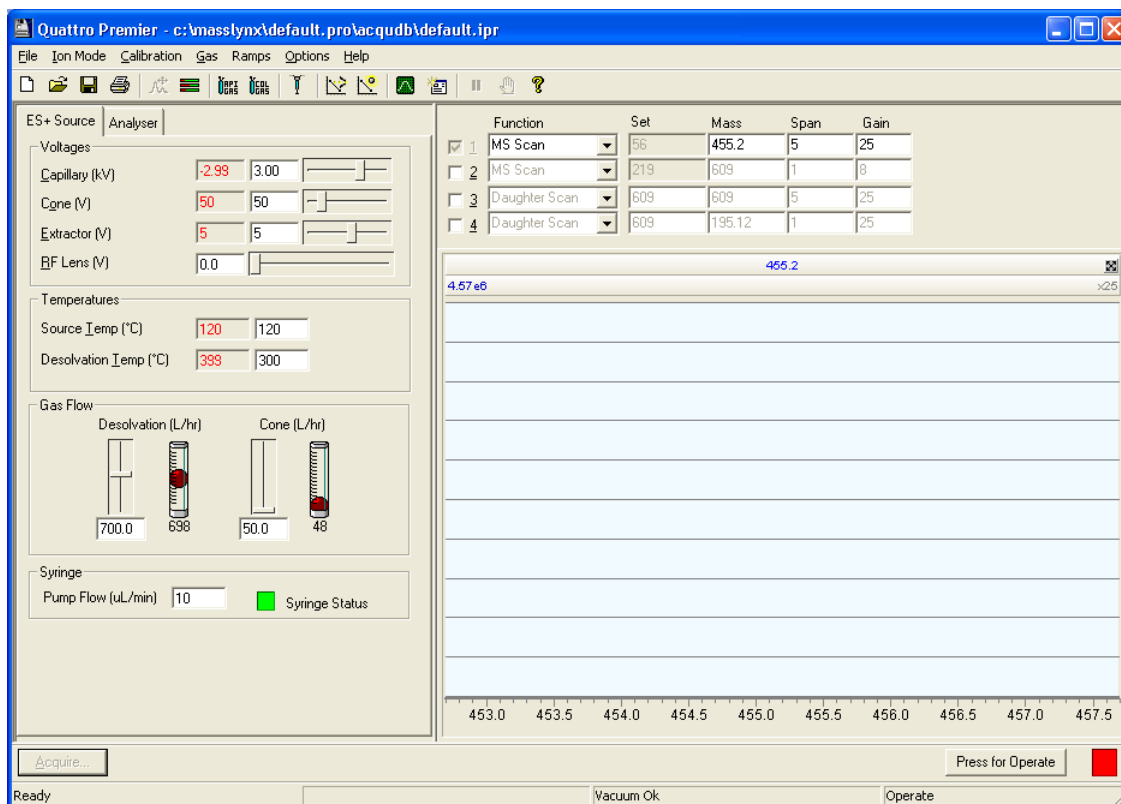


Figure 2-5 Tune Window ES+ Source Page

5. Set the parameters to the recommended values shown in Table 2-1.

Table 2-1 Recommended ES+ Source Page Parameter Values



Parameter	Recommended Value
Voltages	
Capillary (kV)	3.00
Cone (V)	50
Extractor (V)	5
RF Lens (V)	0.0
Temperatures	
Source Temp (°C)	120
Desolvation Temp (°C)	300
Gas Flow	
Desolvation (L/hr)	700.0
Cone (L/hr)	50.0

6. Click the Analyser tab.

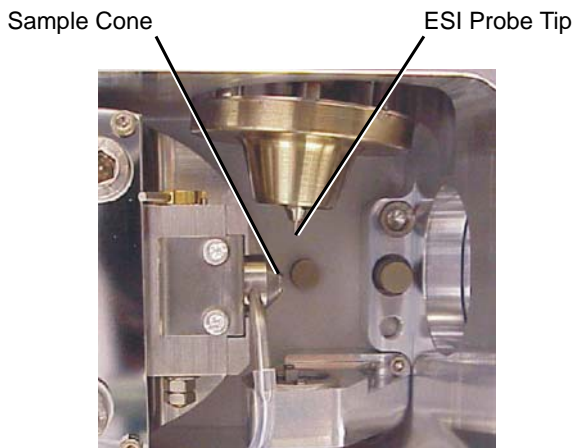
7. Set the parameters to the recommended values shown in [Table 2-2](#).

Table 2-2 Recommended Analyser Page Parameter Values

Parameter	Recommended Value
Analyser	
LM Resolution 1	15.0
HM Resolution 1	15.0
Ion Energy 1	0.5
Entrance	50
Collision	2
Exit	50.0
LM Resolution 2	15.0
HM Resolution 2	15.0
Ion Energy 2	3.0
Multiplier	650

8. Click  to start the nitrogen flow.
9. On the ES+ Source page, observe the Desolvation and Cone gas flows; confirm they are stabilized and correct.
10. Click Press for Operate, and confirm that the adjacent instrument status indicator shows green.
11. Turn on the LC system at a flow rate of 300 $\mu\text{L}/\text{min}$, and confirm that its pressure is stable.
12. Allow 15 minutes for the LC column to equilibrate.
13. Load the sample syringe with sample. In this example, verapamil (concentration 50 $\text{pg}/\mu\text{L}$, in 70:30 acetonitrile/water) is used.
14. Select the correct syringe type using this procedure:
 - a. Select Options > Syringe Type to open the Syringe Selection dialog box.
 - b. Choose the required syringe type from the drop-down list.
Note: *If the syringe type is not listed, click Edit, and add the relevant details.*
 - c. Click OK.
15. On the ES+ Source page, set Syringe, Pump Flow ($\mu\text{L}/\text{min}$) to 10.
16. Click  to start the syringe pump.
17. In the Tune window, select the Function 1 box.
18. Choose MS Scan from the adjacent drop-down list.
19. Enter Mass 455.2, Span 5, and Gain 25.
Note: *The above are recommended values and can vary from instrument to instrument.*
20. Observe the verapamil peak at m/z 455.2 ($M+H^+$) in the Tune window (see [Figure 2-6](#)).
21. Use the vernier probe adjuster, on the probe mounting flange (see [Figure 2-2](#)), to maximize the displayed peak intensity.

22. Use the vernier probe adjuster to move the probe tip as far away from the sample cone as possible without losing more than 20% of the maximum displayed peak intensity obtained in [step 21](#). This minimizes source contamination.



23. Use the nebulization adjuster, on the probe (see [Figure 2-2](#)), to give the best displayed peak intensity and stability.
24. In the Tune window ES+ Source page, adjust the Desolvation (L/hr) gas flow, in increments of 100; allow the pressure to stabilize after each adjustment. Set the gas flow to the value giving the highest displayed peak intensity.
25. Starting from a value of 0, increase the Cone (L/hr) gas flow in increments of 50; allow the pressure to stabilize after each adjustment. Set the gas flow to the highest value that does not significantly reduce the peak intensity. This minimizes solvent ion cluster formation.
26. Starting from a value of 2.6, increase the Capillary (kV) voltage in increments of 0.2. Set the capillary voltage to the value giving the highest displayed peak intensity.
27. Starting from a value of 15, increase the Cone (V) voltage in increments of 2. Set the cone voltage to the value giving the highest displayed peak intensity. Record this value.
28. Starting from a value of 0, increase the Extractor (V) voltage in increments of 1. Set the extractor voltage to the value giving the highest displayed peak intensity.
29. Adjust the RF Lens (V) voltage to the minimum value that maintains the highest peak intensity, without loss of resolution.

- Confirm that the displayed peaks have the correct resolution (the isotopes are resolved as shown in [Figure 2-6](#)). If necessary, adjust the Analyser page LM Resolution 1, HM Resolution 1, and Ion Energy 1 slider bars to achieve optimum resolution [typically 0.75 Da full width at half maximum (FWHM)].

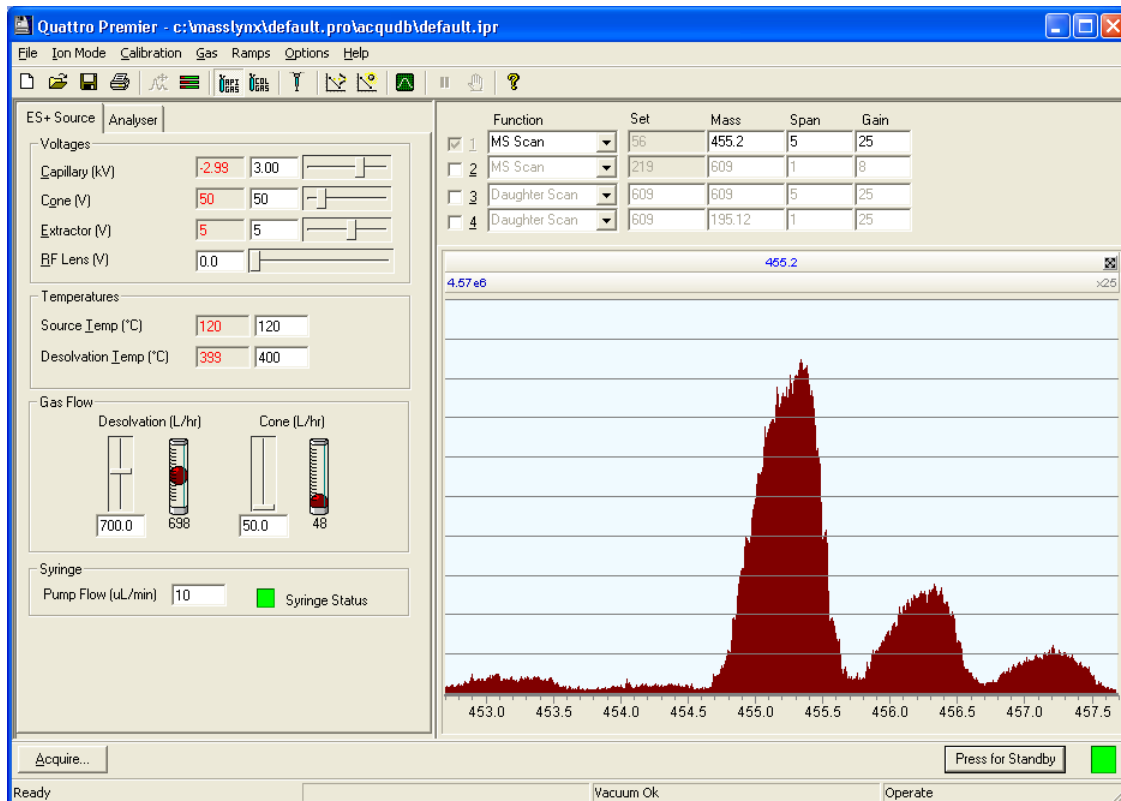


Figure 2-6 Tune Window with a Verapamil Peak

- In the Tune window, determine the m/z value of the peak's center, to one decimal place ([Figure 2-6](#)). Record this value.
- MS tuning is now complete. Perform the MS/MS tuning procedure ([Section 2.2.2](#)).

2.2.2 Tuning for MS/MS (Daughter Ion) Operation

1. In the Tune window, select the Function 2 box. The Function 1 box may now be cleared (Figure 2-7).

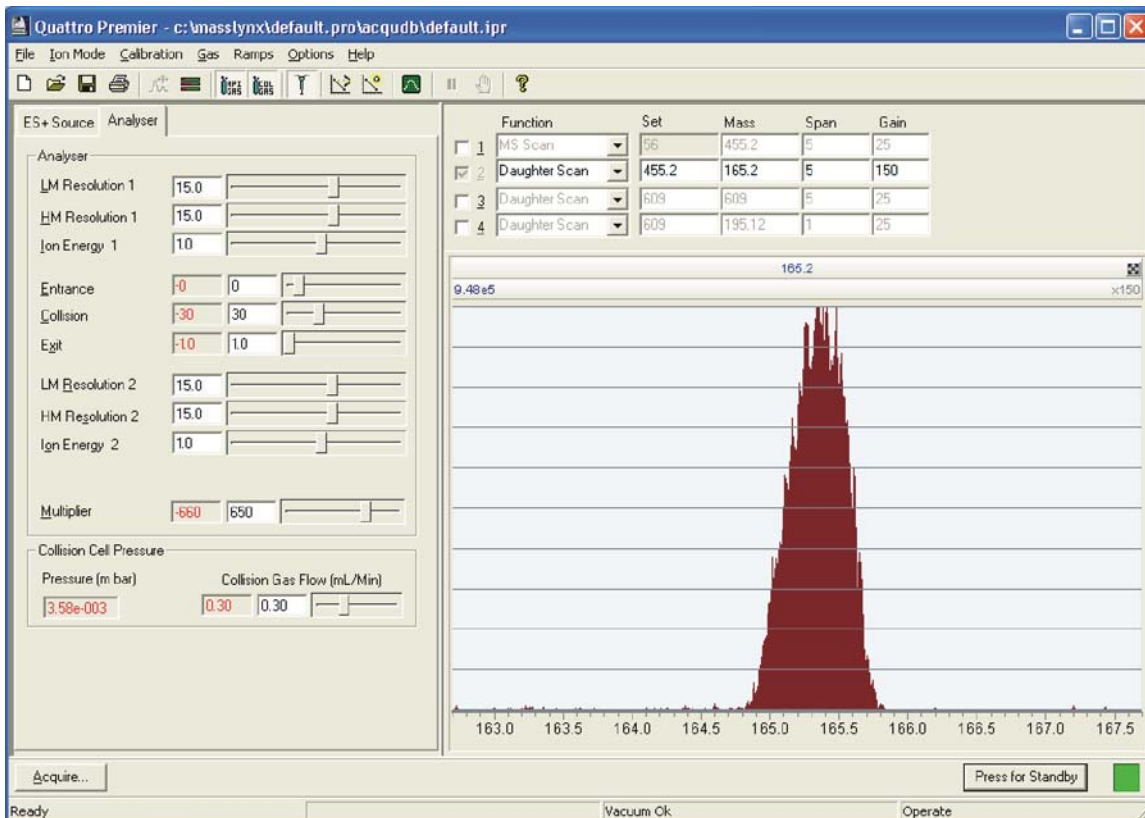



Figure 2-7 Tune Window with a Daughter Ion Peak

2. Select Daughter Scan from the adjacent drop-down list.
3. Enter Set 455.2 (i.e. the verapamil peak), Mass 165.2 (daughter ion), Span 5, and Gain 150.
4. Set Entrance to 0, Collision to 30, and Exit to 1.0.
5. On the Analyser page, set Ion Energy 2 to 1.0.

Note: Do not change the optimal Ion Energy 1 value obtained in [Section 2.2.1](#).

6. Click  to turn on the collision gas flow.

7. On the Analyser page, set the Collision Gas Flow (mL/min) to 0.3. This sets the Collision Cell Pressure to approximately 3 to 4×10^{-3} mbar.
8. Observe the daughter ion peak, at m/z 165.2, displayed in the Tune window.
9. If necessary, adjust the LM Resolution 2, HM Resolution 2, and Ion Energy 2 slider bars to achieve optimum resolution (typically <1 Da FWHM).
10. Starting at a value of 10, increase Collision (i.e., collision energy) in increments of 2. Set Collision to the value giving the highest displayed peak intensity. Record this value.
11. Optimize the Collision Gas Flow (mL/min) using increments of 0.05. Allow the Collision Cell Pressure readback to stabilize after each adjustment. Set the gas flow to the value giving the highest displayed peak intensity.
12. In the Tune window, determine the m/z value of the daughter ion peak's center to one decimal place (see [Figure 2-7](#)). Record this value.
13. MS/MS tuning is now complete. Create a Multiple Reaction Monitoring (MRM) MS method file (see [Section 2.2.3](#)).

2.2.3 Creating the MRM MS Method File

An MRM MS method file, containing the information obtained during the instrument tuning process, must now be created.

1. In the MassLynx window, click the MS Method icon to open the MS Method Editor ([Figure 2-8](#)).

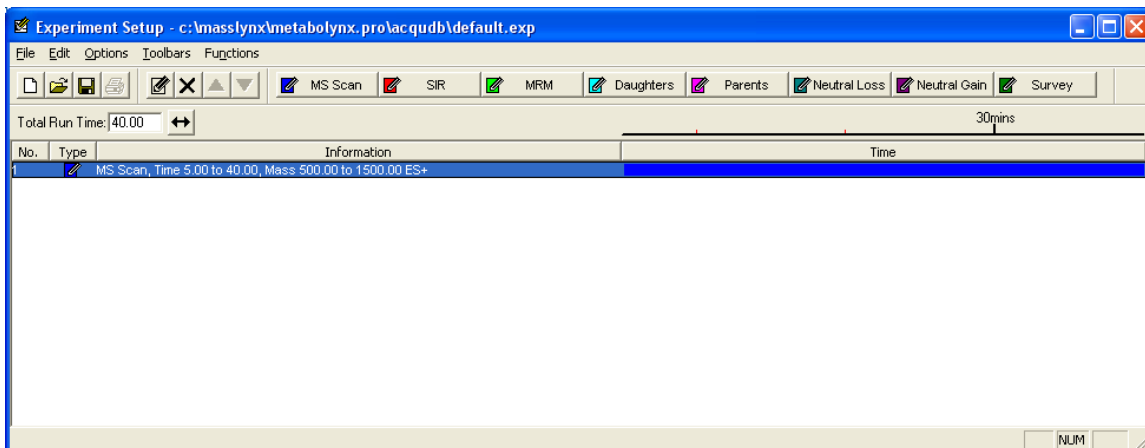




Figure 2-8 MS Method Editor

2. Click  to delete the current entry from the MS Method Editor.
3. Click  to open the MRM Function Editor.
4. Enter the value recorded in [Section 2.2.1, step 31 on page 28](#), in the Parent (m/z) box ([Figure 2-9](#)).

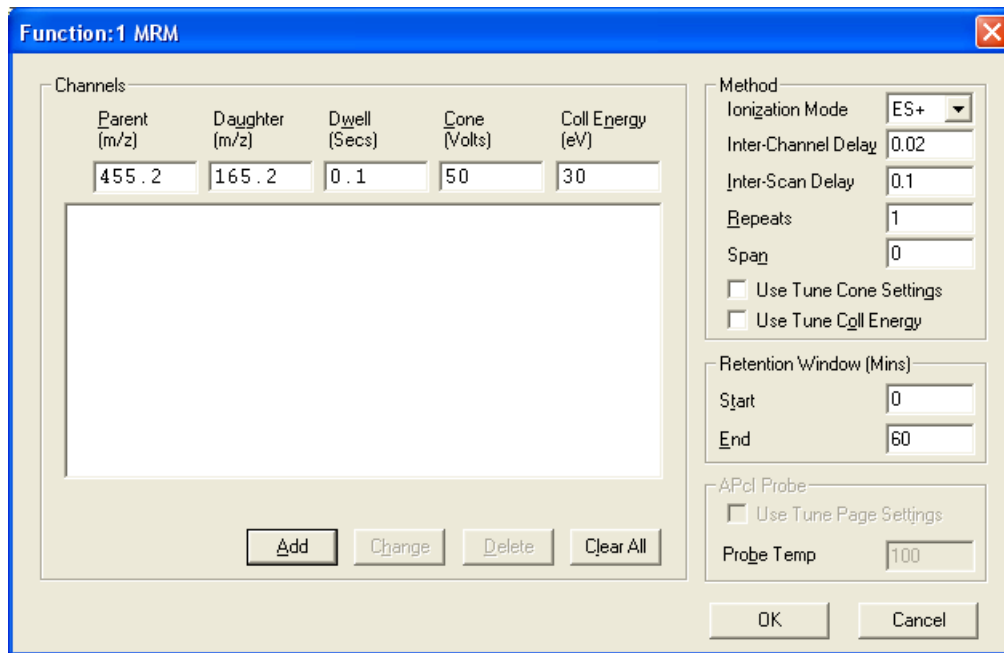



Figure 2-9 MRM Function Editor

5. Enter the value recorded in [Section 2.2.2, step 12 on page 30](#), in the Daughter (m/z) box.
6. Enter 0.1 in the Dwell (Secs) box.
7. Enter the value recorded in [Section 2.2.1, step 27 on page 27](#), in the Cone (Volts) box.
8. Enter the value recorded in [Section 2.2.2, step 10 on page 30](#), in the Coll Energy (eV) box.
9. Click Add; the values entered above are added to the Function List.
10. Select ES+ in the Ionization Mode drop-down list.
11. Set Span to 0.

12. Enter a correct LC run time in the Retention Window (Mins), End text box.
13. Click OK. The MRM Function Editor is closed and the values are included in the MS Method Editor.
14. Click  to open the Save As dialog box.
15. Save the experiment file as verapamil_1.exp.

The instrument is now ready for data acquisition in ESI mode (see [Section 2.3](#)).

2.3 Preparing the Instrument for Data Acquisition

2.3.1 Reconfiguring the Connection to the Probe




Warning: The liquid passing through the HPLC pump, LC column, and ESI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when making the connections between the HPLC pump, LC column, and ESI probe.



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.

1. Click  to stop the syringe pump.
2. Reconfigure the connections to the ESI probe, so that the LC column is connected directly to the probe; i.e., remove the zero-dead-volume T-piece and syringe pump connection ([Figure 2-10](#)).
3. In the Tune window, confirm that the daughter ion peak of interest has disappeared. If the peak does not disappear, consider two possible causes:
 - You tuned the instrument using a large background ion peak. Repeat the tuning procedure using a more suitable sample.
 - The source is contaminated with the sample compound. This may occur if the tuning standard sample concentration is too high. Clean the source (see [Section 7.7](#)), then repeat the tuning procedures, using a suitable sample concentration.

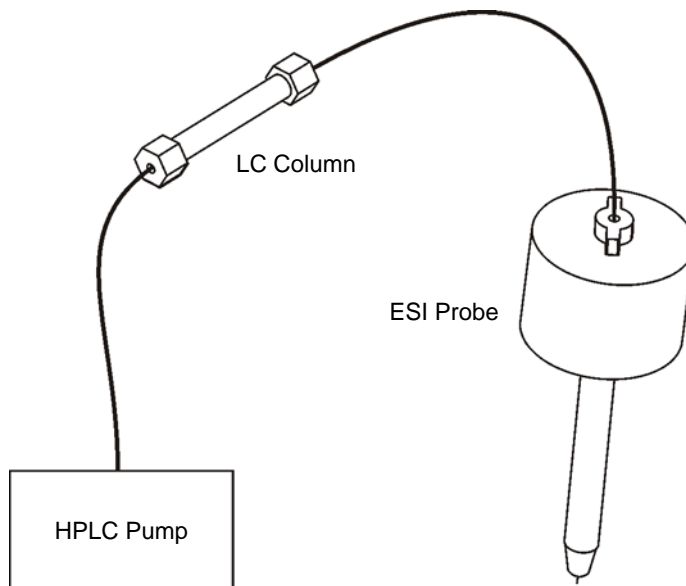


Figure 2-10 Column Connection to the ESI Probe


2.3.2 Configuring the Inlet for LC Operation

1. Click the MassLynx window Instrument shortcut bar Inlet Method icon to open the Inlet Method dialog box.
2. Select Tools > Instrument Configuration to open the Inlet Configuration dialog box.
3. Click Configure to open the Inlet Configuration Wizard.
4. Click Next.
5. Follow the on-screen instructions.
6. When the Inlet Configuration Wizard has finished, click Finish, Finish, and then close the Inlet Configuration dialog box.

2.3.3 Creating a Sample List and Starting Data Acquisition

Note: For comprehensive information on creating a MassLynx Sample List and starting data acquisition, see the MassLynx User's Guide.

1. In the MassLynx window, insert the required number of samples in the Sample List.
2. Enter the required file name(s) in the File Name column.
3. Enter the required text in the File Text column.

4. In the MS File column, select verapamil_1.exp.
5. Create a suitable inlet method file (see the *MassLynx NT Inlet Control Guide*).
6. Enter the inlet method file name in the Sample List Inlet File column.
7. Enter the bottle number(s) in the Bottle column.
8. Enter the injection volume(s) in the Inject Volume column.
9. Save the Sample List.
10. To start data acquisition, click  (see [Chapter 5](#) for details).

2.3.4 Viewing and Printing the Tuning Parameters Associated with a Data File

The tuning parameters associated with a data file are stored with the file as part of the experimental record. You can view or print these tuning parameters from the MassLynx Data Browser dialog box. See the *MassLynx User's Guide* for more information.

Note: *The readbacks incorporated in the experimental record are for indication purposes only. They are not true (calibrated) records of the actual voltages that were on the instrument during data acquisition.*

Chapter 3

Setting Up and Tuning for APCI

Note: For full details of the Tune window, see [Appendix C](#).

3.1 Setting Up

3.1.1 Removing the ESI Probe

You may need to remove the ESI probe from the instrument before fitting the APCI probe. [Figure 3-1](#) shows the ESI probe mounted on the instrument.




Warning: To avoid electric shock, ensure that the instrument is in Standby before starting this procedure.

1. In the MassLynx Tune window, click Press for Standby, and confirm that the adjacent instrument status indicator shows red.



Warning: The liquid passing through the HPLC pump, LC column, and ESI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.

2. Disconnect the LC system from the ESI probe.
3. Wait for three minutes to allow the desolvation gas flow to cool the probe and source.
4. In the MassLynx window, click  to turn off the nitrogen flow.



Warning: The probe and source may be hot. To avoid burns, take great care while working with the instrument's access door open.

5. Open the instrument's access door.

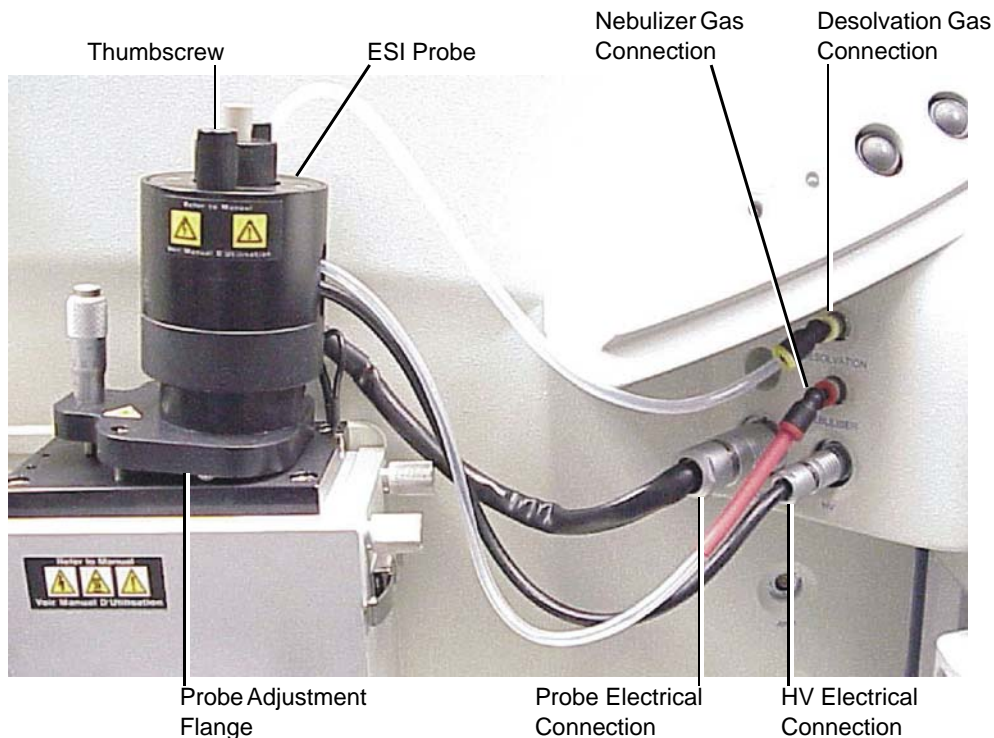


Figure 3-1 ESI Probe Mounted on the Source Enclosure

6. Disconnect the Probe and HV electrical connections on the instrument front panel.
7. Disconnect the PTFE tubes at the Nebuliser and Desolvation gas connections on the front panel.
8. Undo the two thumbscrews securing the probe to the probe adjustment flange.
9. Carefully remove the probe from the probe adjustment flange.

3.1.2 Installing the APCI Corona Discharge Pin



Warning: To avoid electric shock, ensure that the instrument is in Standby when installing the corona discharge pin.

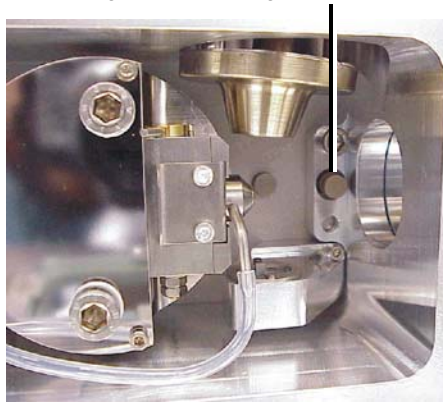
1. In the MassLynx Tune window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.



Warning: The source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

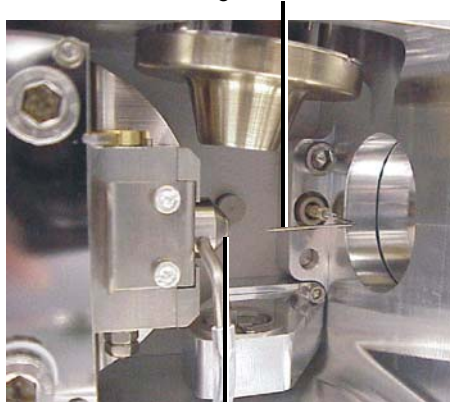
2. Unfasten the source enclosure door's securing clips and open the door.
3. Use needle-nose pliers to remove the blanking plug from the corona discharge pin mounting contact. Store the blanking plug in a safe location.

Corona Discharge Pin Mounting Contact Blanking Plug



4. Use the needle-nose pliers to fit the corona discharge pin to the mounting contact. Ensure that the corona discharge pin tip aligns with the sample cone tip.

Corona Discharge Pin



Sample Cone Tip

5. Close the source enclosure door and fasten the securing clips.

3.1.3 Installing the APCI Probe



Warning: To avoid electric shock, ensure that the instrument is in Standby when installing the APCI probe.

1. In the MassLynx Tune window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.



Warning: The probe may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

2. Carefully slide the APCI probe into the hole in the probe adjustment flange.
3. Secure the probe by tightening the two thumbscrews (Figure 3-2).
4. Connect the probe PTFE tubes to the nebulizer and desolvation gas connections on the front panel.
5. Connect the APCI probe electrical lead to the Probe connection on the front panel.

Note: The probe adjustment flange electrical lead is not connected when the APCI probe is fitted.

6. Close the instrument's access door.

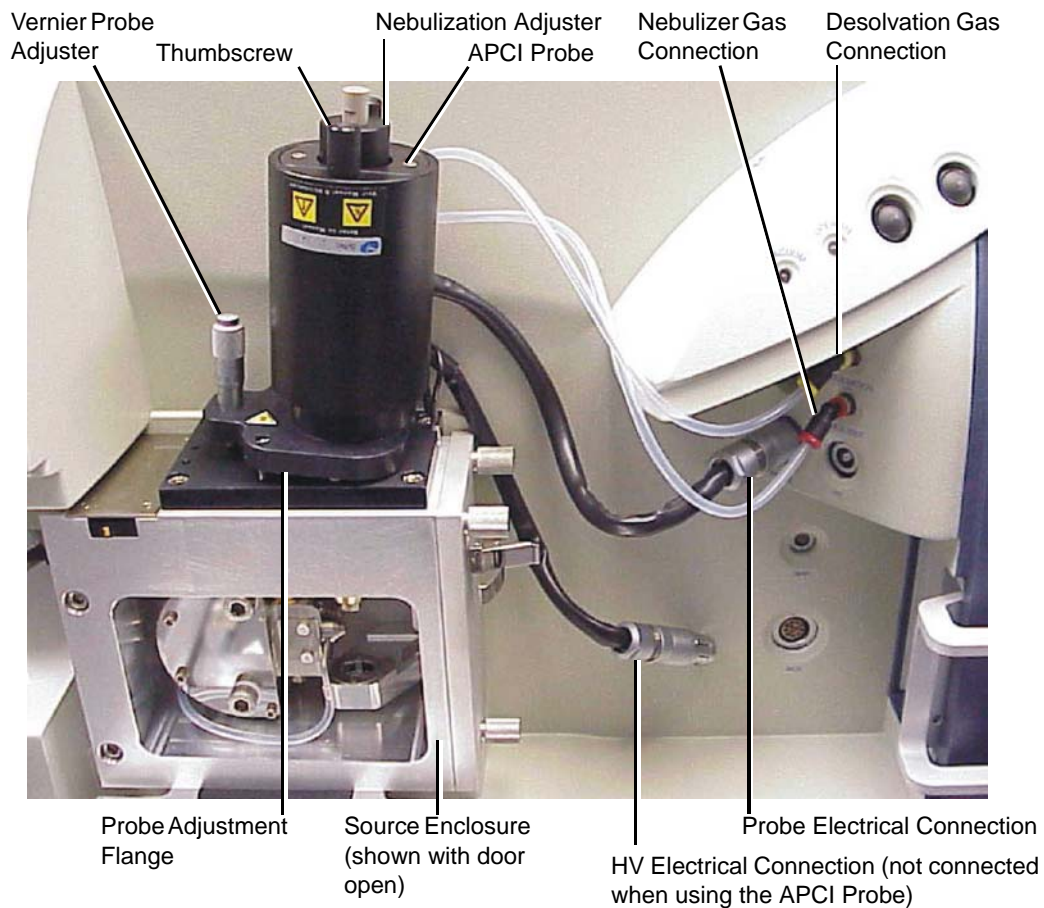


Figure 3-2 APCI Probe Mounted on the Source Enclosure

3.2 Tuning via Sample Infusion

Note: This example procedure specifies 17- α -hydroxyprogesterone as the infused sample. Therefore, the choice of HPLC column and some parameter settings may be specific to that compound. You may tune the instrument using a different sample compound, however you may need to use a different type of column, and parameter settings may differ from the ones given here.

3.2.1 Tuning for MS Operation



Warning: The liquid passing through the HPLC pump, LC column, syringe pump, and APCI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when making the connections between the HPLC pump, LC column, syringe pump, and APCI probe.



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.

1. In the MassLynx Tune window, click Press for Standby, and confirm that the adjacent instrument status indicator shows red.
2. Complete the connections between HPLC pump, LC column, syringe pump, and APCI probe as shown in [Figure 3-3](#).

The column used in this example is a Waters Symmetry C₁₈ 2.1 × 100 mm, 3.5 μ m. The mobile phase is 70:30 acetonitrile/water.

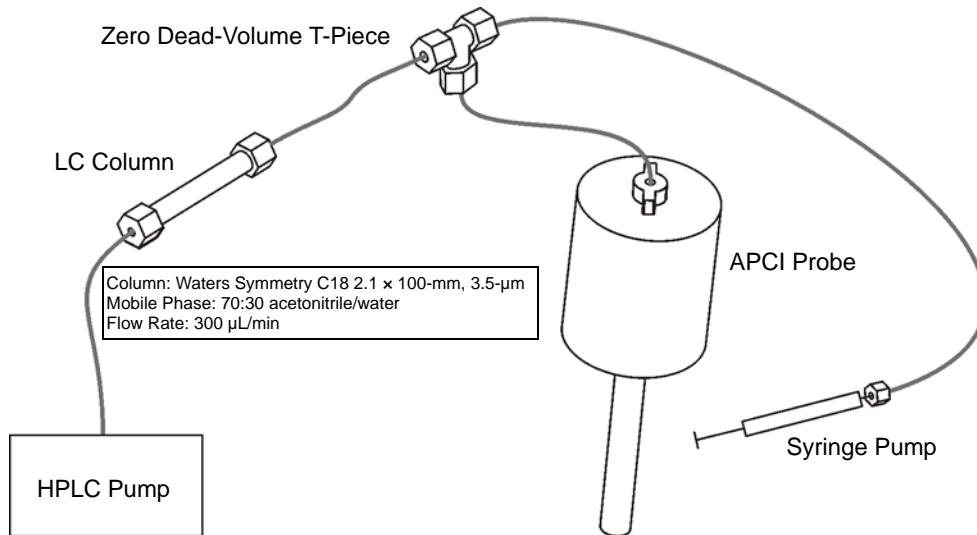


Figure 3-3 Pump and Column Connections to the APCI Probe

3. Look through the source's view port and confirm that the isolation valve lever is fully to the left (that is, the valve is open) (see [Figure 2-4 on page 23](#)).



Warning: The source may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while handling this component.



Warning: The source may be hot. To avoid burns, take great care while working with the instrument's access door open.

If the isolation valve lever is in the wrong position:

- a. Open the instrument's access door.
- b. Unfasten the source enclosure door's securing clips and open the door.
- c. Move the lever to the open position.
- d. Close the source enclosure door and fasten the securing clips.
- e. Close the instrument's access door.

- In the MassLynx Tune window, select Ion Mode > APCI+ to display the APCI+ Source page (Figure 3-4).

Note: The instrument can also be tuned using the EasyTune Source page (see Section C.13).

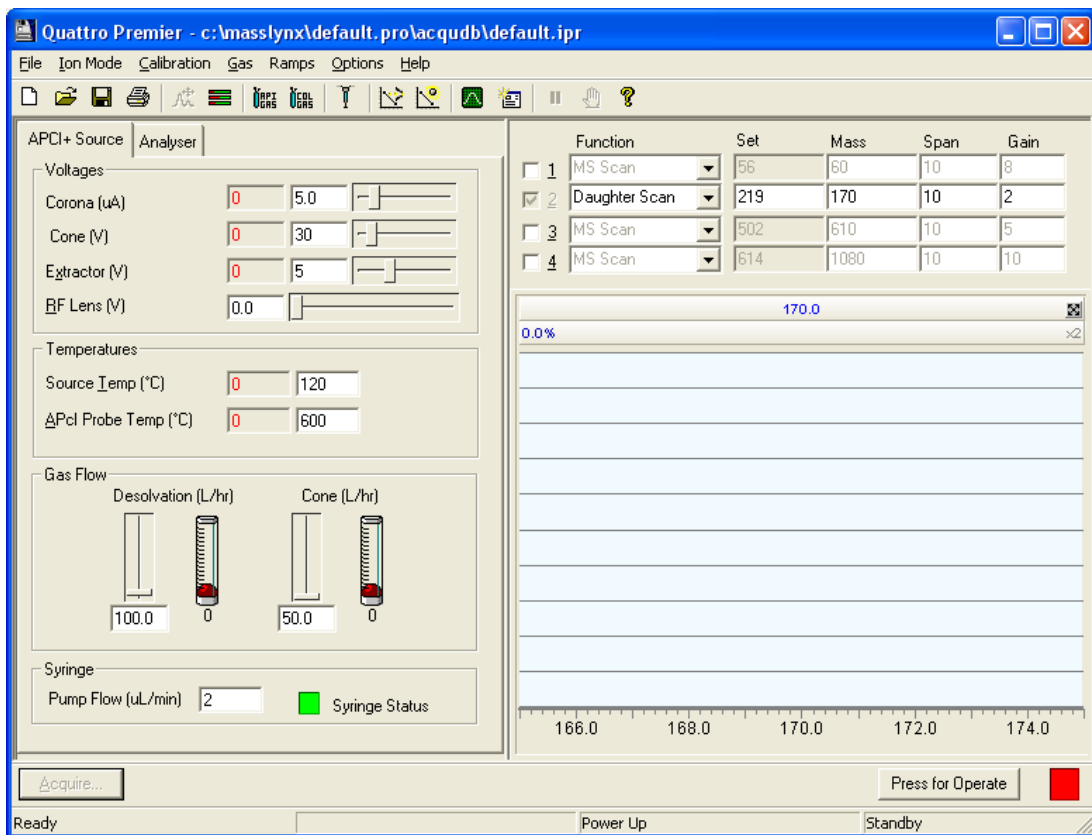


Figure 3-4 Tune Window APCI+ Source Page

- Set the parameters to the recommended values shown in Table 3-1.

Table 3-1 Recommended APCI+ Source Page Parameter Values


Parameter	Recommended Value
Voltages	
Corona (μA)	5.0
Cone (V)	30
Extractor (V)	5
RF Lens (V)	0.0
Temperatures	
Source Temp ($^{\circ}\text{C}$)	120
APCI Probe Temp ($^{\circ}\text{C}$)	600
Gas Flow	
Desolvation (L/hr)	100.0
Cone (L/hr)	50.0

6. Click the Analyser tab.

7. Set the parameters to the recommended values shown in [Table 3-2](#).

Table 3-2 Recommended Analyser Page Parameter Values

Parameter	Recommended Value
Analyser	
LM Resolution 1	15.0
HM Resolution 1	15.0
Ion Energy 1	0.5
Entrance	50
Collision	2
Exit	50.0
LM Resolution 2	15.0
HM Resolution 2	15.0
Ion Energy 2	1.0
Multiplier	650

8. Click  to start the nitrogen flow.
9. On the APCI+ Source page, observe the Desolvation and Cone gas flows; confirm they are stabilized and correct.
10. Confirm that the APcI Probe Temp (°C) readback shows approximately room temperature.
11. Click Press for Operate and confirm that the adjacent instrument status indicator shows green.
12. Confirm that the APcI Probe Temp (°C) readback reaches and stabilizes at the temperature set in [step 5 on page 42](#).
13. Turn on the LC system at a flow rate of 300 µL/min and confirm that its pressure is stable.
14. Allow 15 minutes for the LC column to equilibrate.
15. Load the syringe pump with sample. In this example, 17- α -hydroxyprogesterone (concentration 1 ng/µL, in 70:30 acetonitrile/water) is used.
16. Select the correct syringe type, using this procedure:
 - a. Select Options > Syringe Type to open the Syringe Selection dialog box.
 - b. Choose the required syringe type from the drop-down list.
Note: *If the syringe type is not listed, click Edit and add the relevant details.*
 - c. Click OK.
17. On the APCI+ Source page, set Syringe, Pump Flow (µL/min) to 10.
18. In the Tune window, select the Function 1 box.
19. Select MS Scan from the adjacent drop-down list.
20. Enter Mass 331.2, Span 3, and Gain 150.
Note: *The above are recommended values and can vary from instrument to instrument.*
21. Observe the 17- α -hydroxyprogesterone peak at m/z 331.2 (M+H⁺) in the Tune window (see [Figure 3-5](#)).
22. Starting with the probe tip midway between the cone and the end of the corona pin, use the vernier probe adjuster (see [Figure 3-2](#)) to maximize the displayed peak intensity.

23. Use the vernier probe adjuster to move the probe tip as far away from the cone as possible, without losing more than 20% of the maximum displayed peak intensity obtained in [step 22 on page 44](#). This minimizes source contamination.
24. Use the nebulization adjuster on the probe (see [Figure 3-2](#)) to give the best displayed peak intensity and stability.
25. On the Tune window's APCI+ Source page, optimize the Corona (μA) current to give the maximum displayed peak intensity. Do this by looking for the maximum peak intensity while performing the following steps:
 - a. Starting with a value of 5, decrease the value to 1, in decrements of 1.
 - b. Decrease the value to 0, in decrements of 0.1.
 - c. Reset the value to 5, and then increase the value in increments of 5.
 - d. Set the Corona (μA) current to the value giving the maximum displayed peak intensity.
26. Starting from a value of 0, increase the Cone (L/hr) gas flow in increments of 50. Allow the pressure to stabilize after each adjustment. Set the gas flow to the highest value that does not significantly reduce the peak intensity. This minimizes solvent ion cluster formation.
27. Starting from a value of 15, increase the Cone (V) voltage in increments of 5. Set the cone voltage to the value giving the highest displayed peak intensity. Record this value.
28. Starting from a value of 0, increase the Extractor (V) voltage in increments of 1. Set the extractor voltage to the value giving the highest displayed peak intensity.
29. Adjust the RF Lens (V) voltage to the minimum value that maintains the highest peak intensity, without loss of resolution.
30. Reduce the APCI Probe Temp ($^{\circ}\text{C}$) to 300, and then increase its value in increments of 50. Allow the temperature readback to stabilize after each adjustment. Set the temperature to the highest value that does not reduce the displayed peak intensity.

If the probe temperature is too low, the mobile phase may not be efficiently desolvated. This can result in chromatographic peak tailing.
31. Confirm that the displayed peaks have the correct resolution and the isotopes are resolved as shown in [Figure 3-5](#). If necessary, adjust the Analyser page LM Resolution 1, HM Resolution 1, and Ion Energy 1 slider bars to achieve optimum resolution.
32. In the Tune window, determine the m/z value of the peak's center to one decimal place ([Figure 3-5](#)). Record this value.

33. MS tuning is now complete, Perform the MS/MS tuning procedure (see [Section 3.2.2](#)).

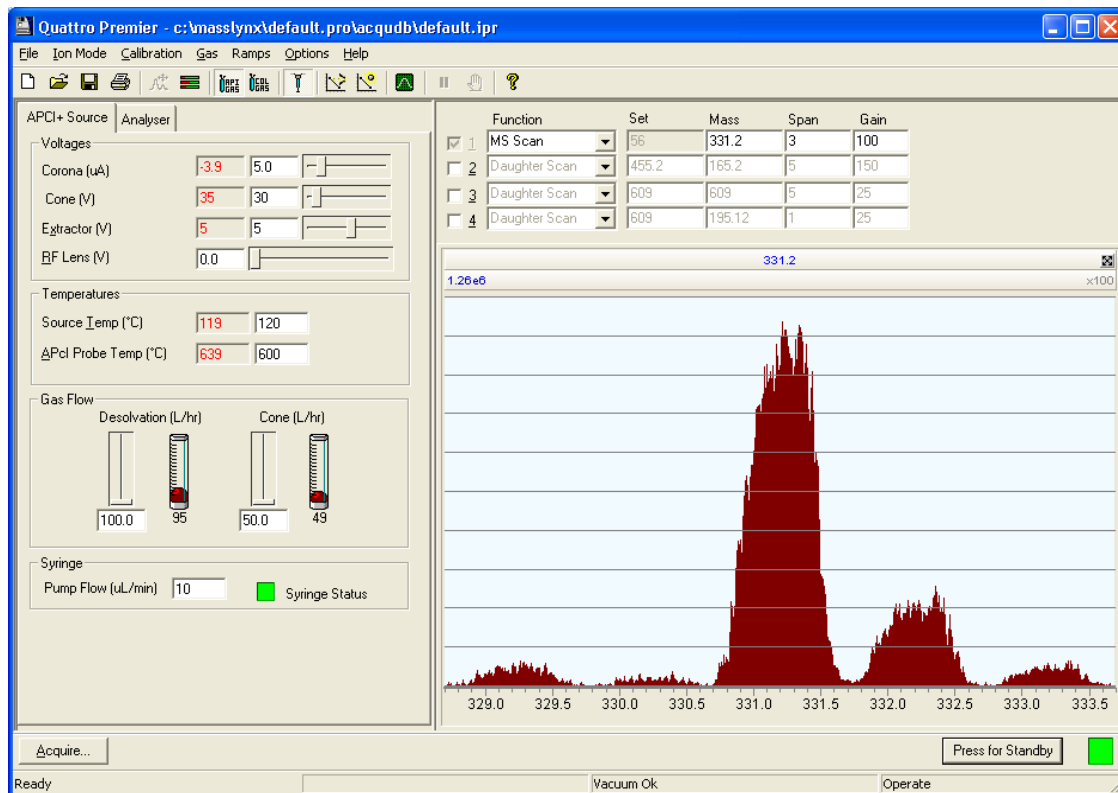



Figure 3-5 Tune Window with a 17- α -Hydroxyprogesterone Peak

3.2.2 Tuning for MS/MS (Daughter Ion) Operation

1. In the Tune window, select the Function 2 box. (The Function 1 box may be cleared, as shown in [Figure 3-6](#).)
2. Select Daughter Scan in the adjacent drop-down list box.
3. Enter Set 331.2 (i.e., the 17- α -hydroxyprogesterone peak), Mass 109.1 (daughter ion), Span 5, and Gain 600.
4. Set Entrance to 0, Collision to 30, and Exit to 1.0. Leave the other parameters at their current settings.
5. On the Analyser page, set Ion Energy 2 to 1.0.

Note: Do not change the optimal Ion Energy 1 value obtained in [Section 3.2.1](#).

6. Click  to start the collision gas flow.
7. On the Analyser page, set the Collision Gas Flow (mL/min) to 0.3. This sets the Collision Cell Pressure to approximately 3 to 4×10^{-3} mbar.
8. Observe the daughter ion peak, at m/z 109.1, displayed in the Tune window.
9. If necessary, adjust the LM Resolution 2, HM Resolution 2, and Ion Energy 2 slider bars to achieve optimum resolution (typically <1 Da FWHM).
10. Starting at a value of 10, increase the Collision (i.e., collision energy) in increments of 2. Set the collision energy to the value giving the highest displayed peak intensity. Record this value.
11. Optimize the Collision Gas Flow (mL/min), using increments of 0.05. Allow the Collision Cell Pressure readback to stabilize after each adjustment. Set the gas flow to the value giving the highest displayed peak intensity.
12. In the Tune window, determine the m/z value of the daughter ion peak's center to one decimal place (Figure 3-6). Record this value.

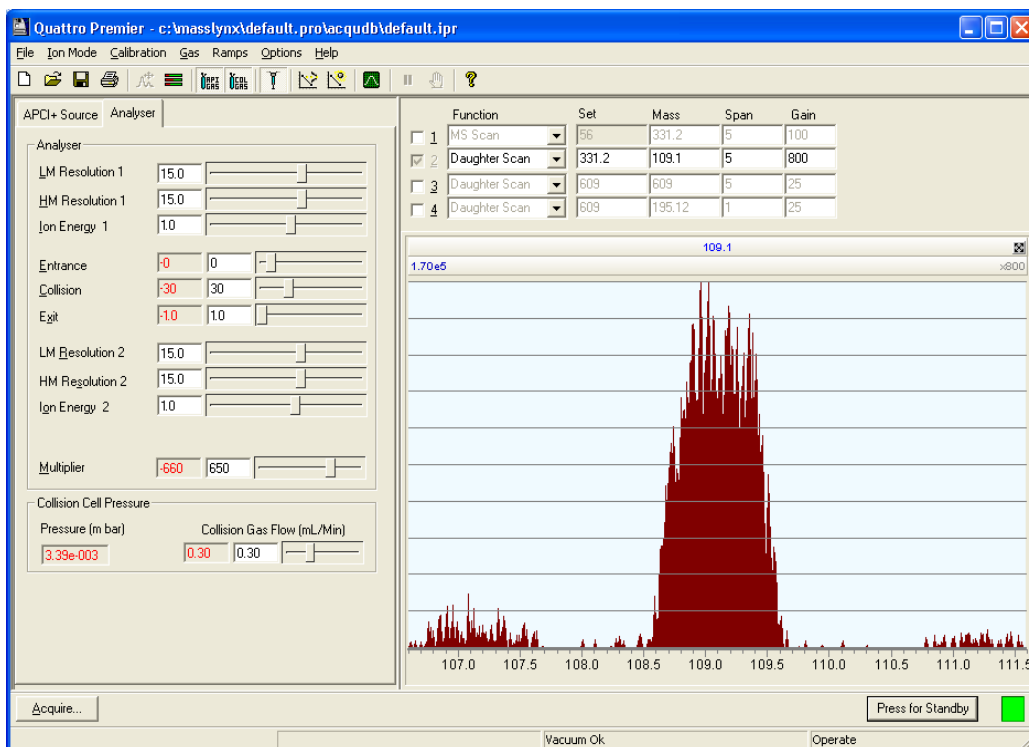


Figure 3-6 Tune Window with a Daughter Ion Peak

13. MS/MS tuning is now complete. Create an MRM MS method file (see [Section 3.2.3](#)).

3.2.3 Creating the MRM MS Method File

An MRM MS method file must now be created, containing the information obtained during the instrument tuning process.

1. In the MassLynx window, click the MS Method icon to open the MS Method Editor ([Figure 3-7](#)).

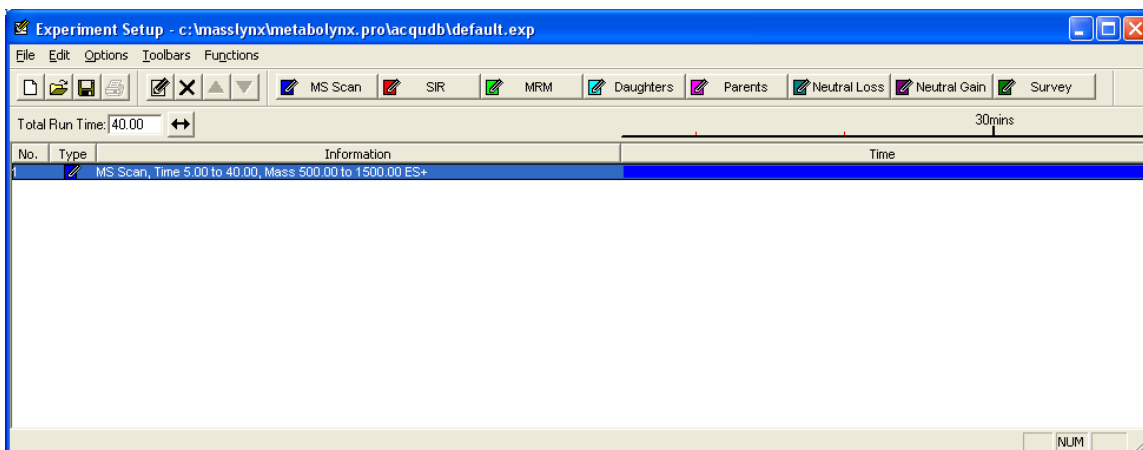




Figure 3-7 MS Method Editor

2. Click  to delete the current entry from the MS Method Editor.
3. Click  to open the MRM Function Editor ([Figure 3-8](#)).
4. Enter the value recorded in [Section 3.2.1, step 32 on page 45](#), in the Parent (m/z) box.
5. Enter the value recorded in [Section 3.2.2, step 12 on page 47](#), in the Daughter m/z) box.
6. Enter 0.1 in the Dwell (Secs) box.
7. Enter the value recorded in [Section 3.2.1, step 27 on page 45](#), in the Cone (Volts) box.
8. Enter the value recorded in [Section 3.2.2, step 10 on page 47](#), in the Coll Energy (eV) box.

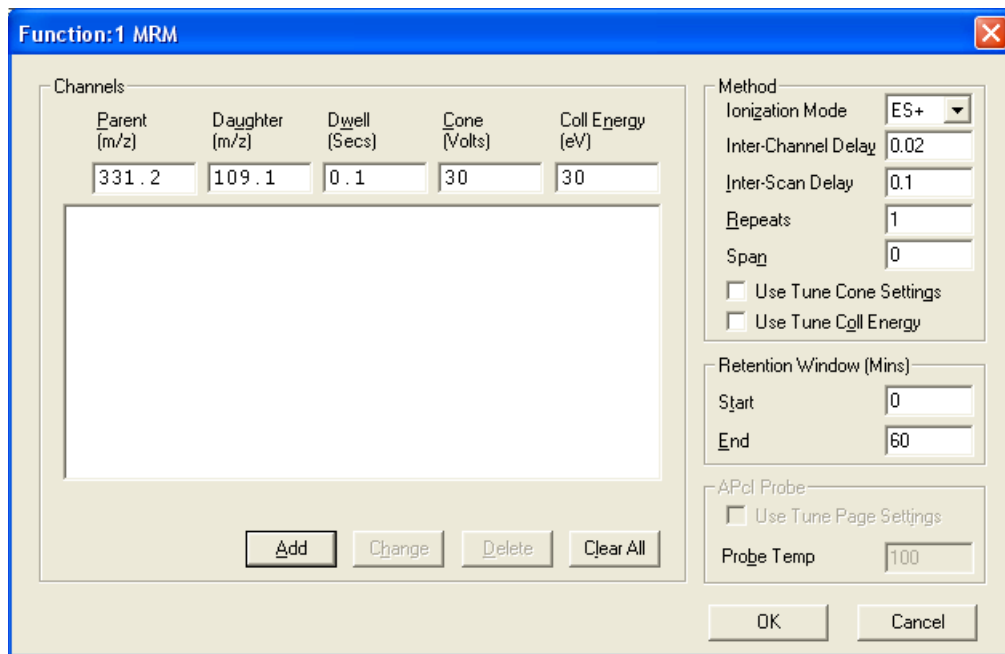



Figure 3-8 MRM Function Editor

9. Click Add. The values entered above are added to the Function List.
10. Select API+ from the Ionization Mode drop-down list.
11. Set Span to 0.
12. Enter a correct LC run time in the Retention Window (mins), End text box.
13. Click OK. The MRM Function Editor is closed, and the values are included in the MS Method Editor.
14. Click  to open the Save As dialog box.
15. Save the experiment file as hydroxyprogesterone peak_1.exp.

The instrument is now ready for data acquisition in APCI mode (see [Section 3.3](#)).

3.3 Preparing the Instrument for Data Acquisition

3.3.1 Reconfiguring the Connection to the Probe




Warning: The liquid passing through the HPLC pump, LC column, and APCI probe may be biohazardous and/or toxic. Always wear nitrile gloves when working with these items.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when making the connections between the HPLC pump, LC column, and APCI probe.



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.

1. Click  to stop the syringe pump.
2. Reconfigure the connections to the APCI probe, so that the LC column is connected directly to the probe; i.e., remove the zero-dead-volume T-piece and syringe pump connection (Figure 3-9).

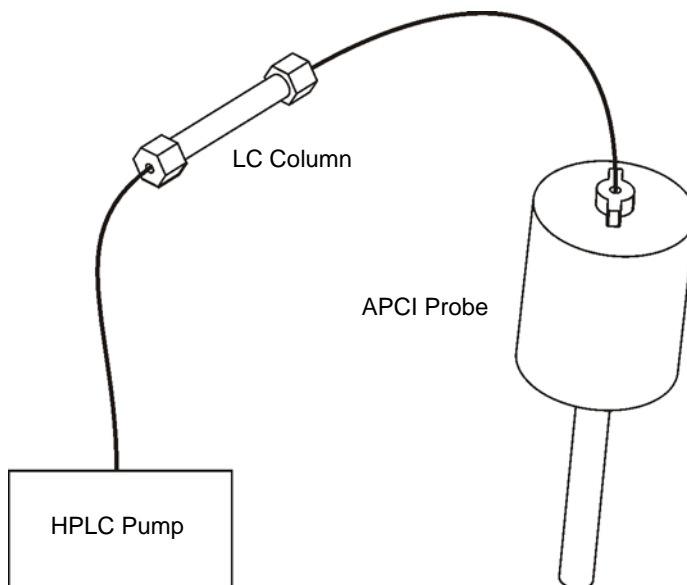


Figure 3-9 Column Connection to the APCI Probe


3. In the Tune window, confirm that the daughter ion peak of interest has disappeared. If the peak does not disappear, consider two possible causes:
 - You tuned the instrument using a large background ion peak. Repeat the tuning procedure using a more suitable sample.
 - The source is contaminated with the sample compound. This may occur if the tuning standard sample concentration was too high. Clean the source (see [Section 7.7](#)), then repeat the tuning procedures, using a suitable sample concentration.

3.3.2 Configuring the Inlet for LC Operation

1. Click the MassLynx window Instrument shortcut bar Inlet Method icon to open the Inlet Method dialog box.
2. Select Tools > Instrument Configuration to open the Inlet Configuration dialog box.
3. Click Configure to open the Inlet Configuration Wizard.
4. Click Next>.
5. Follow the on-screen instructions.
6. When the Inlet Configuration Wizard has finished, click Finish, Finish, and then close the Inlet Configuration dialog box.

3.3.3 Creating a Sample List and Starting Data Acquisition

Note: For comprehensive information on creating a MassLynx Sample List and starting data acquisition, see the *MassLynx User's Guide*.

1. In the MassLynx window, insert the required number of samples in the Sample List.
2. Enter the required file name(s) in the File Name column.
3. Enter the required text in the File Text column.
4. In the MS File column, select hydroxyprogesterone peak_1.exp.
5. Create a suitable inlet method file (see the *MassLynx NT Inlet Control Guide*).
6. Enter the inlet method file name in the Sample List Inlet File column.
7. Enter the bottle number(s) in the Bottle column.
8. Enter the injection volume(s) in the Inject Volume column.
9. Save the Sample List.
10. To start data acquisition, click  (see [Chapter 5](#) for further details).

3.3.4 Viewing and Printing the Tuning Parameters Associated with a Data File

The tuning parameters associated with a data file are stored with the file as part of the experimental record. You can view or print these tuning parameters from the MassLynx Data Browser dialog box (see the *MassLynx User's Guide* for more information).

Note: *The readbacks incorporated in the experimental record are for indication purposes only. They are not true (calibrated) records of the actual voltages that were on the instrument during data acquisition.*

Chapter 4

ESCi Multi-Mode Operation

For ESCi operation, the instrument source is fitted with both the ESI probe and the APCI corona discharge pin. When acquiring data, the instrument alternates between ESI and APCI operation in a single run by switching between the electrospray capillary voltage and the APCI corona discharge pin current under the control of MassLynx.

4.1 Preparing the Instrument



Warning: To avoid electric shock, ensure that the instrument is in Standby before starting this procedure.

1. In the MassLynx Tune window, click Press for Standby, and confirm that the adjacent instrument status indicator shows red.
2. If the APCI probe is fitted to the probe adjustment flange, remove it (see [Section 2.1.1](#)), and fit the ESI probe (see [Section 2.1.3](#)).
3. If the APCI corona discharge pin is not fitted to the source, fit it (see [Section 3.1.2](#)).
4. If the ESCi multi-mode operation option has not been previously installed on the instrument, use the ESCi option key disk to upgrade MassLynx on the MassLynx PC; follow the on-screen instructions.

Note: If this software upgrade has not been installed, the ESCi option will not be available in the MassLynx Tune window.

4.2 Tuning the Instrument

1. In the MassLynx Tune window, select Options > Multiple ion modes > Dual ES/APCI, to display the ESCi Source page (Figure 4-1).

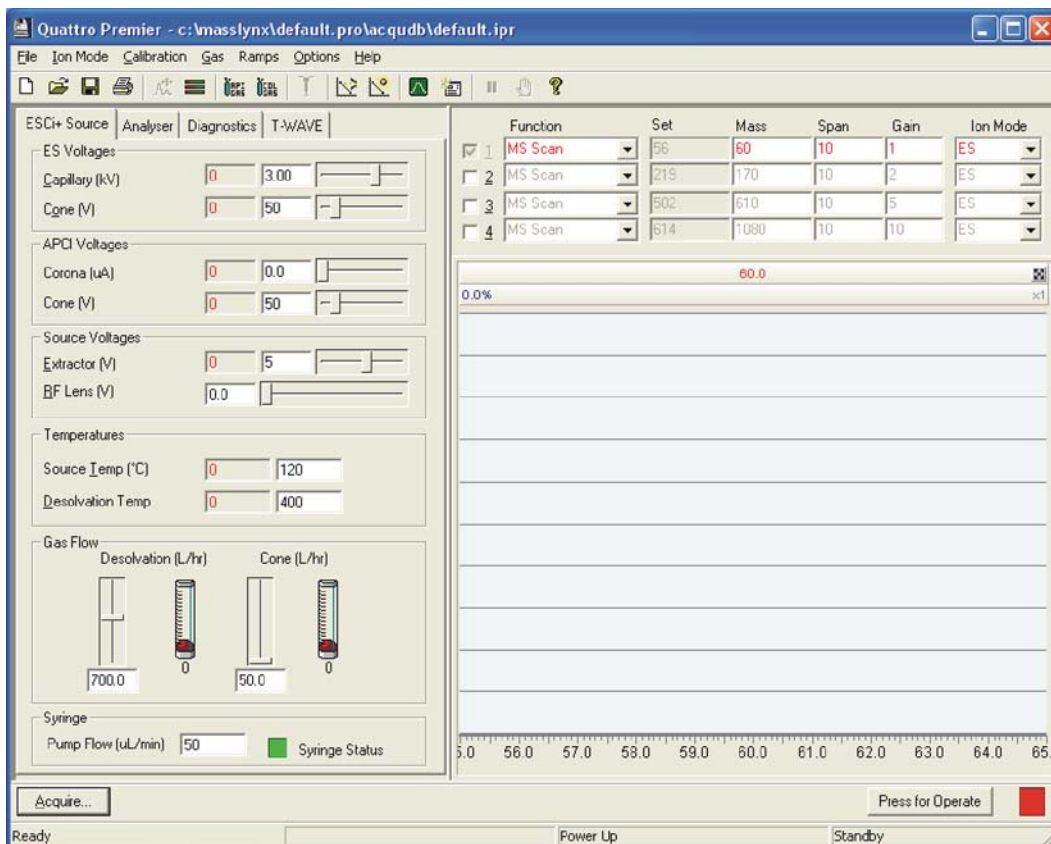


Figure 4-1 Tune Window ESCi Source Page


2. On the menu bar, select Ion Mode > ESCi+ to enable the positive ESCi ionization mode.
3. To tune the instrument for ESI mode:
 - a. Select ES in the Ion Mode drop-down list for Function 1.
 - b. Tune the instrument for ESI MS operation (see Section 2.2.1).

4. To tune the instrument for APCI mode:
 - a. Select APcI in the Ion Mode drop-down list for Function 1.
 - b. Set the APCI Voltages Corona (μA) parameter to 5.0.
 - c. Follow the procedure described in [Section 2.2.1](#) for ESI tuning, optimizing the APCI Voltages Corona (μA) parameter instead of the ES Voltages Capillary (kV) parameter.

4.3 Data Acquisition

1. In the MassLynx Tune window, select Options > Multiple ion modes > Dual ES/APcI, to display the ESCi Source page.
2. On the menu bar, select Ion Mode > ESCi+ or Ion Mode > ESCi- to enable the positive or negative ESCi ionization mode respectively.
3. Select the required mode (ES or APcI) for each Function from the Ion Mode drop-down list. Any combination of ES and APcI modes may be selected.

Note: The entries are color-coded; red denotes an ESI acquisition, blue an APCI acquisition. These colors are also initially applied to the resulting peaks displayed in the Tune window. You may modify the peak colors using the Customise Plot Appearance dialog box (see [Section C.16.2](#)).

4. In the MassLynx window create a Sample List (refer to the *MassLynx User's Guide* for comprehensive information on creating a Sample List).
5. To start data acquisition, click  (see [Chapter 5](#) for details).

Note: In the Tune Window, the ESI readbacks are valid only during ESI acquisition and, likewise, the APCI readbacks are valid only during APCI acquisition.

Chapter 5

Acquiring Data

5.1 Starting an Acquisition

There are two ways of starting an acquisition:

- A single sample acquisition from the Tune window ([Section 5.1.1](#)).
- A multiple sample acquisition from the MassLynx window ([Section 5.1.2](#)).

5.1.1 Starting an Acquisition from the Tune Window

The easiest way to acquire data is directly from the Tune window:

- Acquisitions can be started and stopped.
- Most of the scanning parameters can be controlled.

However, when using this method:

- Inlet programs cannot be used.
- SIR and MRM data cannot be acquired.
- Analog data cannot be acquired.
- Multiple sample sequences cannot be acquired.

To start a single sample acquisition:

1. In the Tune window, click **Acquire**, or select **Window > Acquire** to open the Acquisition Setup dialog box ([Figure 5-1](#)).
2. Enter the parameters required to accommodate the required mass range and scan times (see [Table 5-1](#) for details).
3. Click **Start**.

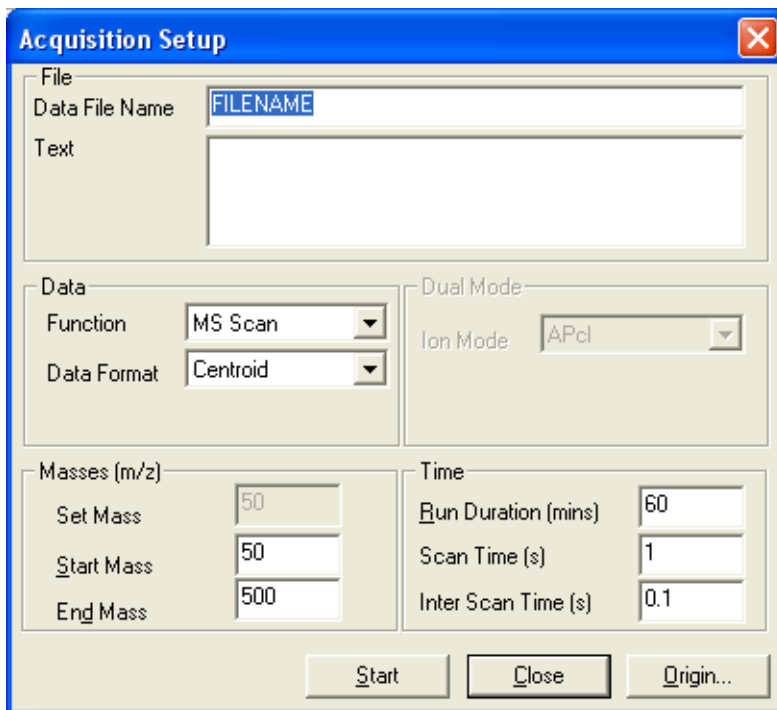


Figure 5-1 Acquisition Setup Dialog Box

Table 5-1 Acquisition Setup Dialog Box Parameters

Parameter	Description
File	
Data File Name	The Data File Name can include up to 128 characters. If the file already exists on disk, you are prompted to rename the file or overwrite the existing one. The file is written to the data directory of the current project.
Text	The sample description, which can include up to 74 characters. The description can be displayed on any output of the acquired data. To display text on multiple lines, press Ctrl+Return at the end of a line.

Table 5-1 Acquisition Setup Dialog Box Parameters (Continued)

Parameter	Description
Data	
Function	<p>The acquisition function used to collect the data can be any of:</p> <ul style="list-style-type: none"> • MS Scan • MS2 Scan • Daughter Scan • Parent Scan • Neutral Loss Scan • Neutral Gain Scan <p>See Section 5.7 for more information.</p>
Data Format	<p>The data format stored on disk can be any of:</p> <ul style="list-style-type: none"> • Centroid • Continuum • MCA <p>See “Types of Data Acquisition” on page 80 for more information.</p>
Masses (m/z)	
Set Mass	<p>Set Mass specifies the mass (Daughter Mass, Parent Mass, etc.) used for the particular function type. This control is disabled if the selected function does not require a set mass.</p>
Start Mass	<p>The mass (m/z) at which the scan starts.</p>
End Mass	<p>The mass (m/z) at which the scan stops.</p> <p>Note: <i>Start Mass must be lower than End Mass.</i></p>
Time	
Run Duration (mins)	<p>The length of the acquisition, in minutes.</p>
Scan Time (s)	<p>The duration of each scan, in seconds.</p>
Inter Scan Time (s)	<p>The time, in seconds, between a scan finishing and the next one starting. During this period no data are stored.</p>

Table 5-1 Acquisition Setup Dialog Box Parameters (Continued)

Parameter	Description
Origin	<p>Opens the Sample Origin dialog box. This allows you to specify additional information about the sample in the following fields:</p> <ul style="list-style-type: none"> • Submitter • Job • Task • Conditions


To change the project into which data are saved:

1. Cancel the acquisition.
2. In the MassLynx window, select File > Open Project to open an existing project, or File > Project Wizard to create a new project.


5.1.2 Starting Multiple-Sample Acquisition from the MassLynx Window

The MassLynx window contains a Sample List for defining multiple samples which may be used together to perform a quantitative analysis. The list of samples is created using a spreadsheet style editor, which you can tailor to suit individual requirements. See the *MassLynx User's Guide* for more detailed information about multiple-sample Sample List analysis.

Procedure

1. Create a Sample List in the MassLynx window.
2. Select Run > Start, or click  to open the Start Sample List Run dialog box (Figure 5-2).
3. Enter the appropriate parameters (see Table 5-2 for details).
4. Click OK.
5. Repeat steps 1 to 4 as required.

Sample Lists are added to a queue and run sequentially unless Priority or Night Time Process has been selected.

Note:  in the Sample List denotes the sample currently being acquired.

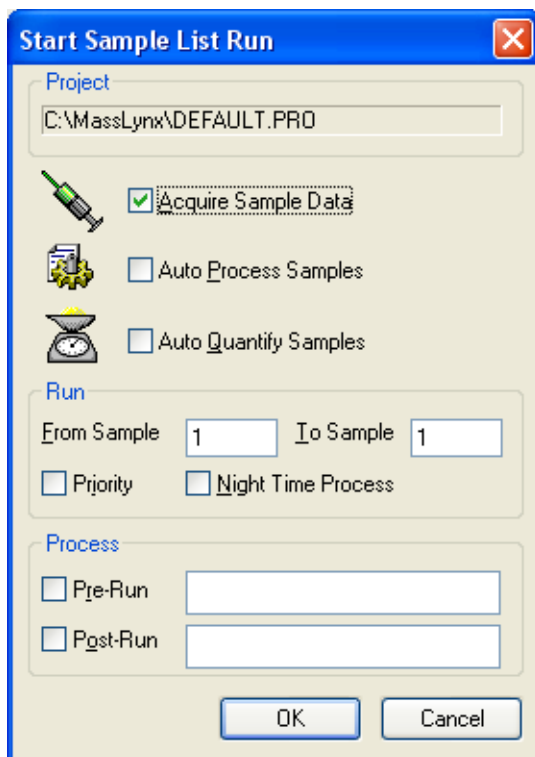


Figure 5-2 Start Sample List Run Dialog Box

Table 5-2 Start Sample List Run Dialog Box Parameters

Parameter	Description
Project	The name of the current project. To send acquired data to a different project, exit this dialog box, open the other project, and restart the acquisition.
Acquire Sample Data	Acquires data from all the samples in the list.
Auto Process Samples	Processes acquired data as specified in the Sample List Process column. This may be existing data, or data newly acquired when the Acquire Sample Data option has been selected (see the <i>MassLynx User's Guide</i> for further details).
Auto Quantify Samples	Quantifies the acquired data using the method specified in the Quantify Samples dialog box (see Section 5.1.3). The current method will be used if a method is not specified in the Quantify Samples dialog box. If this option is selected, the Quantify Samples dialog box will open when the Start Sample List Run dialog OK button is clicked (see the <i>MassLynx User's Guide</i> for further details).
Note: The above three actions can run together or independently; i.e., the User can acquire, process, and quantify data simultaneously, or acquire data in one run and process or quantify it later.	
Run	
From Sample	The sample number of the first sample to be acquired.
To Sample	The sample number of the last sample to be acquired.
Priority	Marks a job as a priority process; it will be placed at the top of the queue, to run after a currently running job.
Night Time Process	Marks this entry as a night-time process. This option is useful for time-consuming acquisitions that would interrupt work on smaller acquisitions during the day.
Process	
Pre-Run	Runs the external executable process specified in the adjacent text box, which performs pre-processing on the samples.
Post-Run	Runs the external executable process specified in the adjacent text box, which performs post-processing on the samples.

5.1.3 Automatic Quantification of the Sample List

The Quantify Samples dialog box (Figure 5-3) allows automatic processing of data files following acquisition. See the *MassLynx User's Guide* for more detailed information about using automated Sample List analysis.

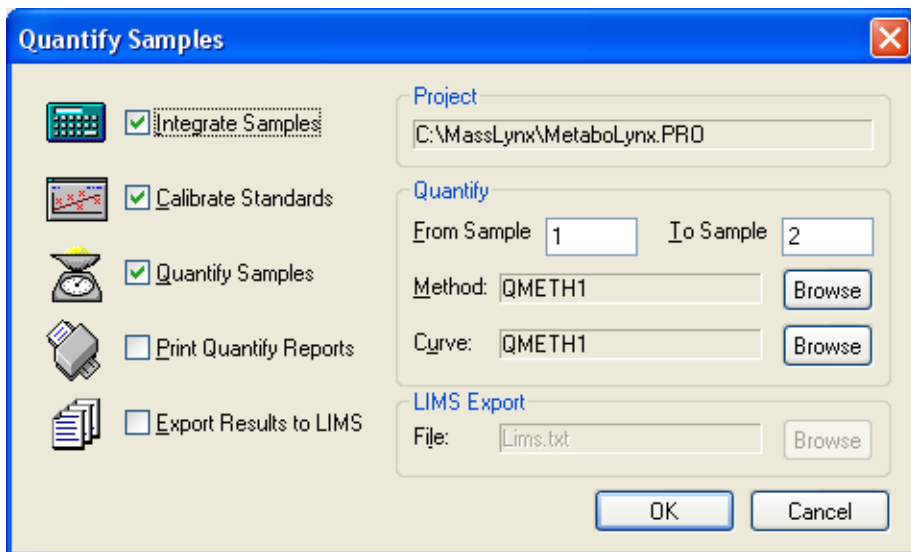


Figure 5-3 Quantify Samples Dialog Box

Procedure

1. Click Process Samples on the MassLynx Quantify shortcut bar, to open the Quantify Samples dialog box.
2. Enter the appropriate parameters (see Table 5-3 for details).
3. Click OK.

Table 5-3 Quantify Samples Dialog Box Parameters

Parameter	Description
Integrate Samples	Integrates all the sample data files named in the Sample List.
Calibrate Standards	Uses integration results to create and quantify calibration curves. Note: Do not select this option if an existing calibration is to be used; instead, use the Curve Browse button to select the required calibration file.
Quantify Samples	Uses integration results and quantify calibration curves to calculate compound concentrations.
Print Quantify Reports	Prints the results of integration and quantitation.
Export Results to LIMS	Produces a text file containing the quantitation results for use with a Laboratory Information Management System (LIMS). This option enables the LIMS Export pane File Browse button. Click Browse and select a file, or enter the name of a new one, then click Save.
Project	Displays the name of the current project.
Quantify	
From Sample:	The sample number of the first sample to be acquired.
To Sample:	The sample number of the last sample to be acquired.
Method	Displays the name of the current method file. To change the file, click the appropriate Browse button, and select a new file.
Curve	Displays the name of the current curve file. To change the file, click the appropriate Browse button and select a new file.


5.2 Monitoring an Acquisition

5.2.1 Viewing the Status of an Acquisition

Acquisition status is shown in the MassLynx window. The run time is shown on the MS Status shortcut bar. The scan status, sample number, and scan number are shown on the status bar at the bottom of the window.

To view a scan-by-scan statistical report of the progress of an acquisition, select Options > Acquisition Status in the Tune window. This opens the Scan Report window, which displays details of the scan currently being acquired.

5.2.2 Viewing a Chromatogram in Real-Time

1. In the MassLynx window Sample List menu bar, click Chromatogram to open the Chromatogram window.
2. Click  or select Display > Real-Time Update. The chromatogram display is updated as the acquisition proceeds.

5.2.3 Viewing a Spectrum in Real-Time

1. In the MassLynx window Sample List menu bar, click Spectrum to open the Spectrum window.
2. Select Display > Real-Time Update to open the Spectrum Real-Time Update dialog box (Figure 5-4).

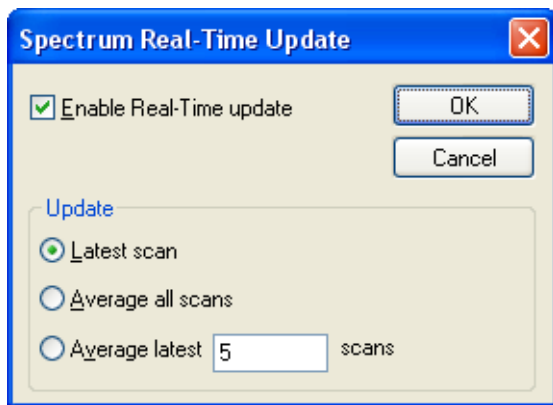



Figure 5-4 Spectrum Real-Time Update Dialog Box

3. Select Enable Real-Time update.
4. Select the required Update option (see the details below).
5. Click OK.

Note: Real-time updating can also be turned on and off by clicking .

When real-time update is enabled, the display is continually updated with spectra from the current acquisition. The actual information displayed is determined by clicking the appropriate update option:

- Latest scan displays the last acquired scan (the default option).
- Average all scans updates the display with spectra formed by averaging all the spectra acquired so far.
- Average latest scans updates the display with spectra formed by averaging the last n scans acquired, where n is specified in the associated text box.

5.3 Selecting Instrument Data Thresholds

Instrument data thresholding allows you to specify which data to acquire and store, and which to discard. Limiting the amount of data stored on disk can be useful when acquiring continuum data and performing long LC runs.

Instrument data threshold settings are specified in the Instrument Threshold Settings dialog box ([Figure 5-5](#)).

Changing Data Thresholding

1. In the Tune window, select Options > Set Instrument Threshold to open the Instrument Threshold Settings dialog box.
2. Make the required changes to the information. (The parameters are fully described in [Sections 5.3.1 to 5.3.6](#).)
3. Click OK.

The new parameters are downloaded at the start of the next acquisition scan.

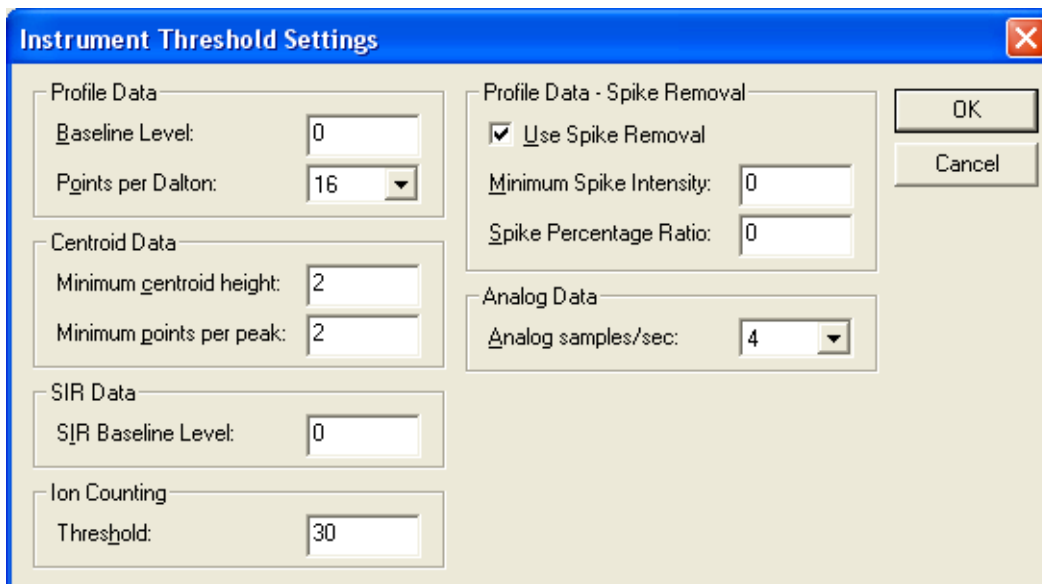


Figure 5-5 Instrument Threshold Settings Dialog Box

5.3.1 Profile Data

The Profile Data parameters control the amount of data collected during a continuum data acquisition.

Baseline Level changes the baseline position with respect to zero and is typically set to zero. A positive value increases the amount of noise seen. A negative value reduces the noise seen and acts as a form of thresholding applied to 1/16 Dalton-type samples.

The Baseline Level adjustment takes place after ion counting and, therefore, has a less significant effect than Ion Counting Threshold (see [Section 5.3.4](#)).

Points per Dalton can have one of three values: 4, 8, or 16:

- Acquiring data at 16 points per Dalton gives the greatest possible resolution.
- Selecting 8 points per Dalton instead of 16 produces data files approximately half as big.
- Acquiring data at 4 points per Dalton gives data with a smoothed appearance.

5.3.2 Centroid Data

Minimum centroid height sets a height below which detected peaks are ignored. This reduces the size of acquired data files and is useful when concentrating on larger peaks of interest. A suitable value can be deduced by evaluating spectral noise levels, and should be evaluated for each system.

The Minimum points per peak value specifies the minimum number of points that a continuum peak must comprise to initiate centroid processing. A typical value is 10.

5.3.3 SIR Data

The SIR Baseline Level value specifies the position of the SIR baseline above zero; it is typically set to zero. Increasing the value causes the baseline to appear higher in the spectrum.

5.3.4 Ion Counting Threshold

Ion Counting Threshold sets the intensity level below which a data point is ignored. This threshold is applied to all acquisitions, regardless of scanning mode. It is also the most significant data thresholding parameter, as it is the first to be applied to raw data.

When an acquisition is started, the instrument performs a prescan with the ion beam switched off. This allows measuring of the acquisition system electronic noise level and its standard deviation. Ion Counting Threshold affects only the system electronic noise level.

The specified Ion Counting Threshold level is multiplied by 1/10 of the standard deviation of the noise to determine the suitable intensity level. Hence a value of 10 equates to one standard deviation of the electronic noise level.

- Values can be set between 0 and 1000; the higher the number, the more data is discarded.
- If a value of 0 is entered, the intensity level is set so that it sits in the middle of the noise. This means that roughly half the noise data is acquired.
- A value of 10 places the threshold just above the noise, so almost all the noise data is acquired.
- If a value of 200 is specified, the threshold sits well above the noise level, so little noise data is acquired.
- A value of 30 is suitable for most data.

Note: Ion Counting Threshold should be set to 0 when acquiring data to be analyzed using MaxEnt™, as the MaxEnt algorithm needs to measure noise accurately within a data file (see the MassLynx User's Guide for full details).

Set Ion Counting Threshold so that background noise is removed without significantly reducing the intensity of the smallest peaks of interest.

5.3.5 Profile Data – Spike Removal

Spikes can be distinguished from real data because the peaks are very narrow and, when compared to their immediate neighbors, very intense. A data point determined to be a spike is removed by setting the value of the data point to the average of its immediate neighbors.

Spike removal involves additional processing during data acquisition and so reduces the maximum achievable acquisition rates by approximately 30%.

Performing Spike Removal During an Acquisition

1. In the Instrument Threshold Settings dialog box Profile Data - Spike Removal pane, select Use Spike Removal.
2. Examine the peak intensities displayed in the Tune window to deduce a suitable value for the intensity threshold below which spikes are ignored. Set Minimum Spike Intensity to this value.

A very low intensity signal may include single ion events that can be combined to produce significant peaks. For this type of data, set Minimum Spike Intensity to a suitable value such that these single ion events are not discarded as spikes.

3. Set a suitable value for Spike Percentage Ratio.

This ratio determines whether a data point is a spike by comparing the data point to its immediate neighbors. For example, with Spike Percentage Ratio set to 33%, a data point is regarded as a spike if its intensity is three times (or more) greater than both its immediate neighbors. A setting of 20% requires an intensity ratio of 5:1 to identify a spike.

4. Click OK to accept any changes.

5.3.6 Analog Data


Select the number of samples to acquire, per second, from the Analog samples/sec drop-down list.

5.4 Checking the Status of the Communications Link


To check the status of the communications link between the MassLynx PC and embedded PC, select Options > Communications Status, in the Tune window, to open the System Manager window. This displays details of the embedded system's status.

5.5 Stopping an Acquisition

You can stop an acquisition from either the Tune window or the MassLynx window. Data already acquired are saved.

- In the Tune window, click .



Note: Stopping an acquisition from the Tune window will stop only the current acquisition in the Sample List; data acquisition for the next entry in the Sample List will then start.

- In the MassLynx window, select Run > Stop, or click .

Note: Stopping an acquisition from the MassLynx window will pause the Sample List.

5.6 Pausing an Acquisition

You can pause an acquisition from either the Tune window or the MassLynx window:

- In the Tune window, click .
- In the MassLynx window, select Run > Pause or click .

Click the appropriate pause button again to restart the acquisition.

5.7 Setting-Up Scanning Functions

5.7.1 Creating a Function List

A function list contains the function(s) used by the mass spectrometer to scan the instrument during an acquisition; it is created using the MS Method Editor (Figure 5-6). A Function List can be a mixture of different scanning techniques that run either sequentially or concurrently during an acquisition.

Click the MS Method icon in the MassLynx window to open the MS Method Editor.

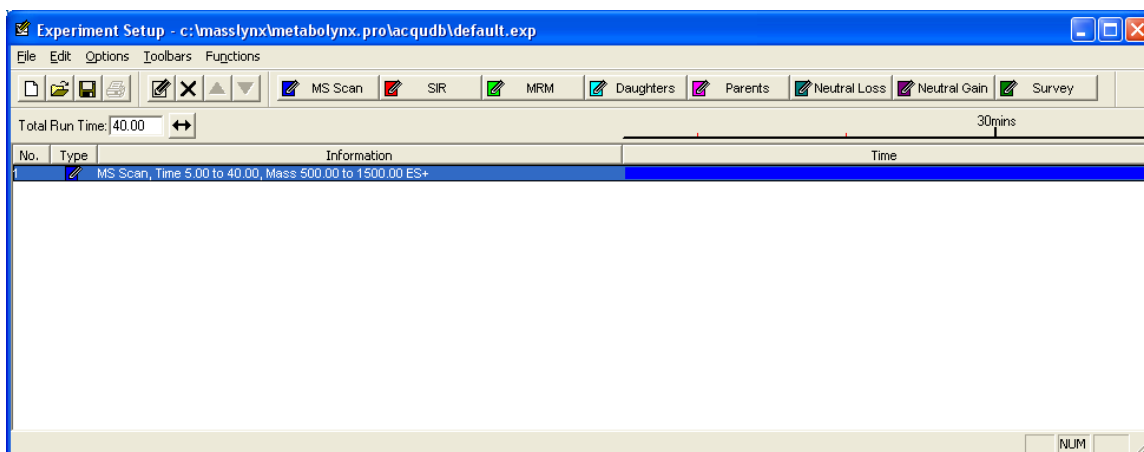


Figure 5-6 MS Method Editor

Mixed function acquisitions are typically used to acquire different SIR groups over different retention windows.

Once created, a Function List can be saved, and then reopened when an acquisition is started.

Figure 5-6 shows a simple Function List containing only one function: a full scan in centroid mode, between 500 and 1500 Da using ES+ ionization. Immediately above the function bar display is a time scale, which shows the elapsed time at which the function becomes active and for how long it runs. In this case, the function starts after 5 minutes and then runs for 35 minutes, terminating after a total elapsed time of 40 minutes.

A more complicated Function List, with four SIR functions, each running sequentially for 5 minutes, is shown in Figure 5-7.

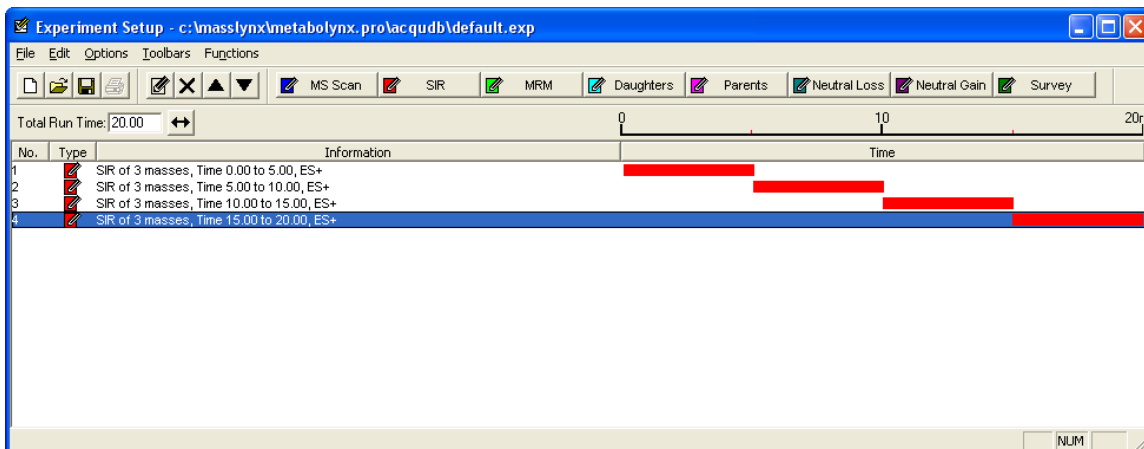


Figure 5-7 MS Method Editor with Four SIR Functions

The currently selected function is highlighted and enclosed in a rectangular frame. If the Function List contains several functions, a function can be selected by clicking on it or by using the keyboard arrow keys.

MS Method Editor Toolbar

The MS Method Editor toolbar functions are shown in [Table 5-4](#).

Table 5-4 MS Method Editor Toolbar Buttons









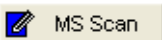
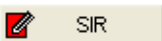

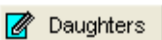
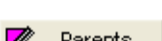


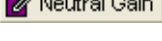
Toolbar Button	Function
	Create a new Function List.
	Open an existing Function List.
	Save the current Function List to disk.
	Print the current window in portrait format. Note: To print the current window in landscape format, select <i>File > Print</i> .
	Edit the selected function.
	Delete the selected function.

Table 5-4 MS Method Editor Toolbar Buttons (Continued)


Toolbar Button	Function
	Move the selected function up the list of functions.
	Move the selected function down the list of functions.
	Create a new Full Scan function (see Section 5.7.2).
	Create a new SIR function (see Section 5.7.3).
	Create a new MRM function (see Section 5.7.5).
	Create a new Daughter function (see “Setting-Up a Daughter Scan” on page 85).
	Create a new Parent function (see “Setting-Up a Parent Scan” on page 87).
	Create a new Neutral Loss function (see “Setting-Up a Neutral Loss Scan” on page 87).
	Create a new Neutral Gain function (see “Setting-Up a Neutral Gain Scan” on page 87).
	Create a new Survey function (see Section 5.7.6).

Adding a New Function to the List

1. Click one of the toolbar buttons, or choose the required function from the Functions menu to open the appropriate Function Editor, showing default values.
2. Make required changes to the parameters.
3. Click OK. The new function is added to the Function List.

Note: The Function Editors are described in [Sections 5.7.2 to 5.7.5](#).

Modifying an Existing Function in the List

1. Select the function.
2. Click  or double-click on the function to open the appropriate Function Editor, showing default values.
3. Make the required changes to the parameters.
4. Click OK. The Function List display is updated to show changes.

Note: The Function Editors are described in [Sections 5.7.2 to 5.7.5](#).

Copying an Existing Function in the List



1. Select the function.
2. Select Edit > Copy.
3. Select Edit > Paste. A copy of the function appears in the Function List.
4. Modify the parameters (see above).

Removing a Function from the List


1. Select the function.
2. Click , select Edit > Delete, or press the Del key.

Changing the Order of Functions in the List

Functions are displayed in ascending start and end time order; this order cannot be changed. For functions with identical start and end times, you can change the order in which they are performed:

1. Select the required function.
2. Repeatedly click  or  until the function is in the required position.

Setting the Maximum Retention Time

1. Enter the required value in the Total Run Time: text box.
2. Click . The ratio of the defined functions is maintained. For example, if two functions are defined, one from 0 to 5 minutes, and the other 5 to 10 minutes, then a Total Run Time of 10 minutes is displayed. If this value is changed to 20, the first function now runs from 0 to 10 minutes, and the second from 10 to 20 minutes.

Setting Solvent Delays

No data is stored during a solvent delay period, which means that solvent peaks that would normally be seen eluting on the Total Ion Current (TIC) chromatogram are not seen.

Select Options > Solvent Delay on the MS Method Editor to open the Solvent Delay dialog box ([Figure 5-8](#)). Up to four solvent delay periods can be entered in the dialog box.

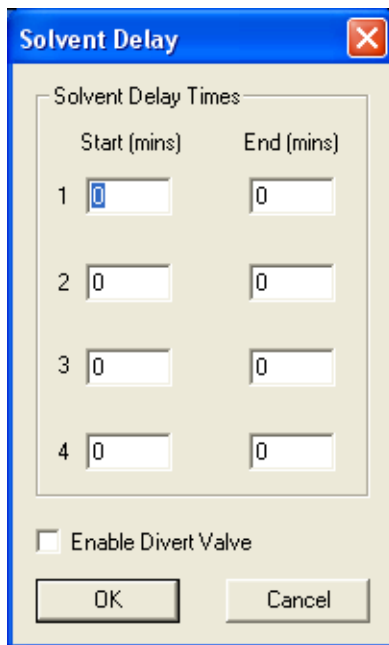


Figure 5-8 Solvent Delay Dialog Box

For APCI functions, the APCI probe temperature is set to the value specified on the Tune window APCI+ Source page, APCI Probe Temp field for each solvent delay period.

Select Enable Divert Valve to use the divert/injector valve in the divert mode. This diverts the flow of solvent during a solvent delay period either to, or away from, the source for the time period.

Acquiring Analog Data

Up to eight channels of analog data can be acquired. These are stored with the data acquired from the mass spectrometer.

Analog channels are typically used to collect data from external units such as UV detectors, which must be connected to the Analogue Channels connectors (see [Section 1.6](#)).

Note: Although eight channel inputs are available, the MassLynx software can only control four of them (see below).

A reading is made from the external channel at the end of each scan and stored with the data for that scan. The resolution of the chromatography for an analog channel therefore depends on the scan speed used to acquire the mass spectrometry data.

To store data for an analog channel:

1. On the MS Method Editor, select Options > Analog Data to open the Analog Data dialog box (Figure 5-9).

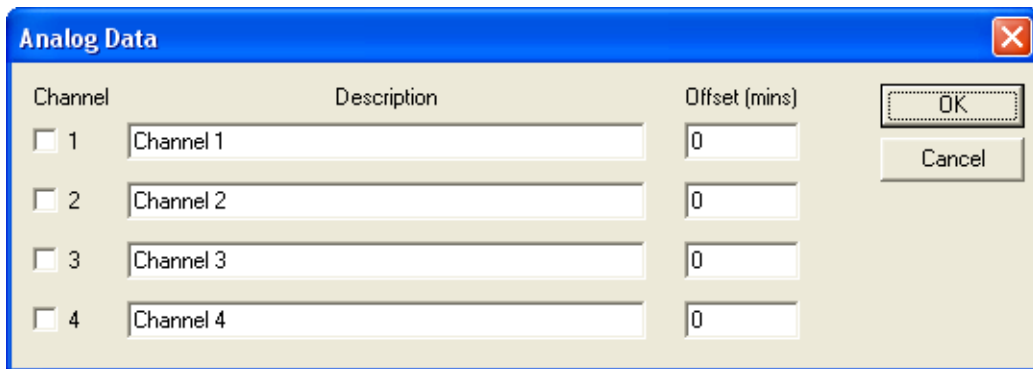


Figure 5-9 Analog Data Dialog Box

2. Select the box(es) for the required analog channel(s).
3. Enter a textual description for each selected channel.

This description is used on the analog Chromatogram dialog box as the channel description (see the *MassLynx User's Guide*).

4. Enter an Offset (mins) to align the external unit with the mass spectrometer.
5. Click OK.

Saving a Function List

1. On the MS Method Editor, select File > Save As to open a standard Save As dialog box.
2. Enter a new file name, or select an existing file from the displayed list.
3. Click Save.

Opening a Function List

1. On the MS Method Editor, select File > Open to open a standard Open dialog box.
2. Enter the required file name, or select an existing file from the displayed list.
3. Click Open.

5.7.2 Setting-Up a Full Scan Function

The full scan Function Editor (Figure 5-10) is used to set-up centroid, continuum, and MCA functions.

The screenshot shows a dialog box titled "Function: 2 MS Scan". It is divided into several sections for parameter configuration:

- Mass (m/z):** Start: 50, End: 600
- Method:** Ionization Mode: ES+, Data: Centroid
- Time (Mins):** Start: 0, End: 60
- Scan Duration (secs):** Scan Time: 1, Inter-Scan Delay: 0.1
- Cone Voltage:** Use Tune Page, Cone Voltage (V): 30, Use Cone Voltage Ramp, CV Ramp... button
- APCl Probe:** Use Tune Page Settings, Probe Temp: 100

Buttons for "OK" and "Cancel" are located at the bottom right of the dialog.

Figure 5-10 Full Scan Function Editor


1. On the MS Method Editor, click  MS Scan, or select Functions > MS Scan to open the full scan Function Editor.
2. Enter the required parameters (see Table 5-5 for details).

Table 5-5 Full Scan Function Editor Parameters

Parameter	Description
Mass (m/z)	
Start	The mass (m/z) at which the scan starts.
End	The mass (m/z) at which the scan stops. Note: <i>The Start mass must be less than the End mass.</i>
Time (Mins)	
Start	The retention time (in minutes) at which the scan starts.
End	The retention time (in minutes) at which the scan stops.
Cone Voltage	
Use Tune Page	When selected, the cone voltage set in the Tune window at the start of the acquisition is used. Note: <i>The cone voltage value cannot be altered during acquisition by typing new values into the Tune window, since the new values are not downloaded during acquisition.</i>
Cone Voltage (V)	The cone voltage.
Use Cone Voltage Ramp	When selected, applies a ramp to linearly increase the cone voltage (see “Applying a Linear Ramp to the Cone Voltage” on page 79).
CV Ramp	Opens the Cone Ramp dialog box (see “Applying a Linear Ramp to the Cone Voltage” on page 79). Note: <i>CV Ramp is available only when Use Cone Voltage Ramp is selected.</i>
Method	
Ionization Mode	Specifies the ionization mode and polarity applied during data acquisition.
Data	Specifies the type of data collected and stored on disk (see “Types of Data Acquisition” on page 80).
Scan Duration (secs)	
Scan Time	The duration of each scan in seconds.
Inter-Scan Delay	The time between a scan’s end and the next one’s start, in seconds. No data are stored during this period.

Table 5-5 Full Scan Function Editor Parameters (Continued)

Parameter	Description
APCI Probe	Note: The parameters in this pane are enabled when Ionization Mode is set to API.
Use Tune Page Settings	When selected, the APCI probe temperature set in the Tune window at the start of the acquisition is used. Note: The APCI probe temperature value cannot be altered by typing new values into the Tune window during the acquisition, since the new values are not downloaded during the acquisition.
Probe Temp	The probe temperature, in degrees centigrade.

Applying a Linear Ramp to the Cone Voltage

1. On the Full Scan Function Editor, select Use Cone Voltage Ramp.
2. Click CV Ramp to open the Cone Ramp dialog box (Figure 5-11).

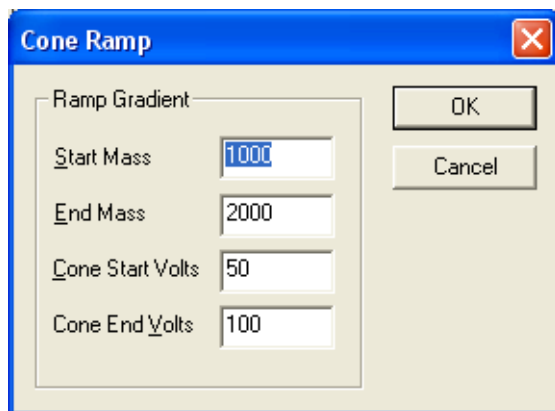


Figure 5-11 Cone Ramp Dialog Box

3. Enter the required parameters (see Table 5-6 for details).

Table 5-6 Cone Ramp Dialog Box Parameters

Parameter	Description
Ramp Gradient	
Start Mass	The mass (m/z) at which the lower cone voltage is specified.

Table 5-6 Cone Ramp Dialog Box Parameters (Continued)

Parameter	Description
End Mass	The mass (m/z) at which the upper cone voltage is specified.
Cone Start Volts	The lower cone voltage.
Cone End Volts	The upper cone voltage.

Note: The two values of cone voltage specify a cone voltage gradient that is then extrapolated to cover the full mass range.

Types of Data Acquisition

In the Full Scan Function Editor (see [Figure 5-10](#)), the Data parameter specifies the type of data to be collected and stored on disk:

- Centroid – Stores data as centroid, intensity, and mass-assigned peaks. Data are stored for every scan.
- Continuum – The signal received by the interface electronics is stored regularly to give an analog intensity picture of the data being acquired. Data are not processed into centroid peaks, but are stored for every scan.

As data are always acquired and stored, even when no peaks are acquired, data files tend to be significantly larger than those for centroid data. It is possible, however, to set a threshold below which the data are not stored. Depending on the nature of the data acquired, this can greatly reduce these effects. The threshold can be set so that data considered to be “noise” can be discarded, thus improving data acquisition speed and reducing data file sizes. For more information about setting data thresholds, see [Section 5.3](#).

- MCA (Multi Channel Analysis) – MCA data can be thought of as “summed continuum”, with only one intensity-accumulated scan stored for a given experiment. As each scan is acquired, its intensity data is added to the accumulated, summed data of previous scans.

An advantage of MCA is that random noise does not accumulate as rapidly as real data and therefore effectively averages out over a number of scans. This emphasizes the real data and improves the signal-to-noise ratio.

Another advantage of MCA is that scanning speeds can be increased and significantly less storage space is required, because data is stored only at the end of an experiment.

The disadvantage of MCA is that, because there is only one scan, it cannot be used for time-resolved data.

5.7.3 Setting-Up an SIR Function

The SIR (Selected Ion Recording) technique is typically used when only a few specific masses are to be monitored. Since most of the data acquisition time is spent on these masses, the technique is far more sensitive than full scanning.

The SIR Function Editor (Figure 5-12) is used to specify the masses to be monitored, along with their dwell times, spans, and inter-channel delay times.

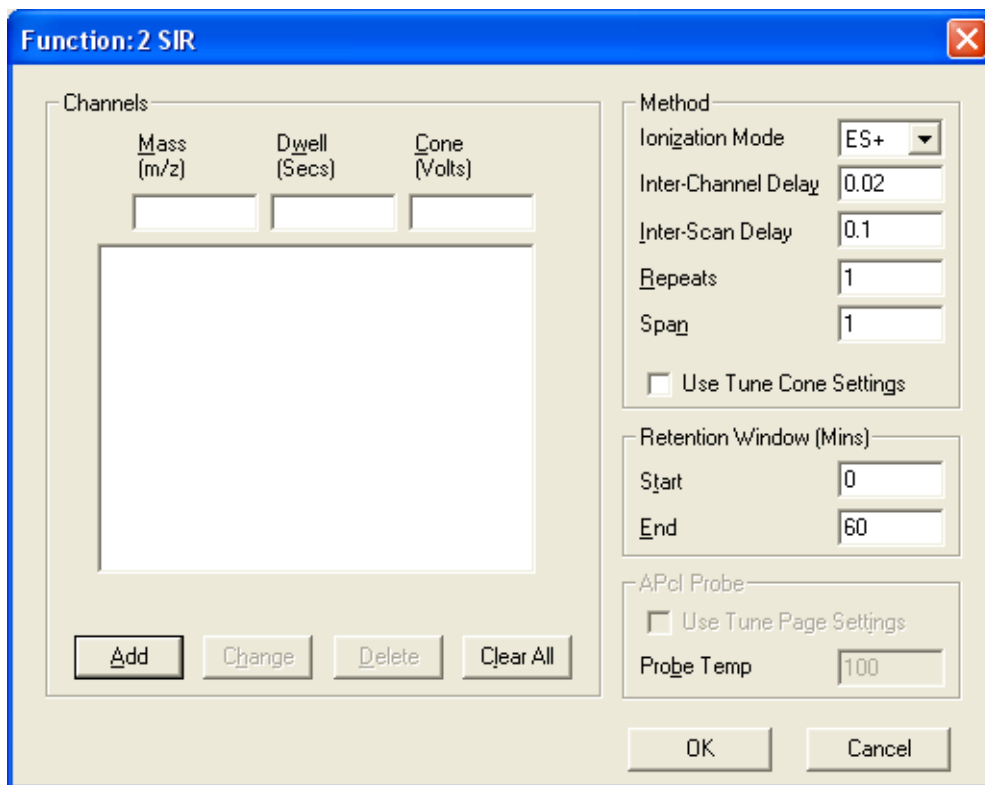


Figure 5-12 SIR Function Editor

Procedure

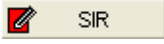
1. On the MS Method Editor, click , or select Functions > MS SIR to open the SIR Function Editor.
2. Enter the required parameters (see Table 5-7 for details).

Table 5-7 SIR Function Editor Parameters

Parameter	Description
Channels	
Mass (m/z)	The mass (m/z) to be scanned.
Dwell (Secs)	The length of time, in seconds, for which the highlighted mass is monitored.
Cone (Volts)	The cone voltage.
Add	Adds the current values in the Mass, Dwell, and Cone boxes to the Function List (see “Adding a Function to the SIR Function List” on page 83).
Change	Replaces the values in the selected function by the values currently in the Mass, Dwell, and Cone boxes (see “Modifying a Function in the SIR Function List” on page 83).
Delete	Deletes the selected function from the list (see “Deleting a Function from the SIR Function List” on page 83).
Clear All	Deletes all the functions from the list (see “Deleting All Functions from the SIR Function List” on page 83).
Method	
Ionization Mode	The ionization mode and polarity used during data acquisition.
Inter-Channel Delay	The time, in seconds, between finishing monitoring the current mass and starting monitoring the next mass in the Function List.
Inter-Scan Delay	The time, in seconds, between finishing one scan and starting the next scan.
Repeats	The number of repeats of the function. Repeats is only relevant for experiments having more than one function.
Span	Specifies a small mass window applied centrally about the current mass. During acquisition, this range is scanned over the specified Dwell time. A span of zero can be set to scan the specified mass without a mass window.
Use Tune Cone Settings	When selected, the cone voltage set in the Tune window at the start of the acquisition is used. Note: <i>The cone voltage value cannot be altered during acquisition by typing new values into the Tune window, since the new values are not downloaded during acquisition.</i>

Table 5-7 SIR Function Editor Parameters (Continued)

Parameter	Description
Retention Window (Mins)	
Start	The start of the retention time, in minutes, during which the current function is active.
End	The end of the retention time, in minutes, during which the current function is active.
APCI Probe	Note: The parameters in this frame are enabled when Ionization Mode is set to API.
Use Tune Page Settings	When selected, the APCI probe temperature set in the Tune window at the start of the acquisition is used. Note: The APCI probe temperature value cannot be altered by typing new values into the Tune window during the acquisition, since the new values are not downloaded during the acquisition.
Probe Temp	The probe temperature, in degrees centigrade.

Adding a Function to the SIR Function List

Up to 32 functions can be entered in the Function List.

1. Enter the required values into the Mass, Dwell, and Cone boxes.
2. Click Add.

Modifying a Function in the SIR Function List

1. Click on the function to select it.

The values for the selected function are displayed in the Mass, Dwell, and Cone boxes.

2. Change the values in the Mass, Dwell, or Cone boxes as required.
3. Click Change to update the values in the Function List.

Deleting a Function from the SIR Function List

1. Click on the function to select it.
2. Click Delete.

Deleting All Functions from the SIR Function List

To delete all the functions from the Function List, click Clear All.

5.7.4 Setting-Up MS/MS Scanning Functions

1. On the MS Method Editor (see [Figure 5-6](#)), click the appropriate toolbar function button to open the appropriate Function Editor. The MS/MS Scanning Function Editors are shown in [Figure 5-13](#).

Note: No toolbar button exists for the MS2 scan function. Select *Functions > MS2 Scan* to open the MS2 Function Editor.

2. Enter the required parameters (see the following sections).

Note: Many of the fields in the MS/MS editors are similar to those in the Full Scan Function Editor. Only fields that differ significantly are described.

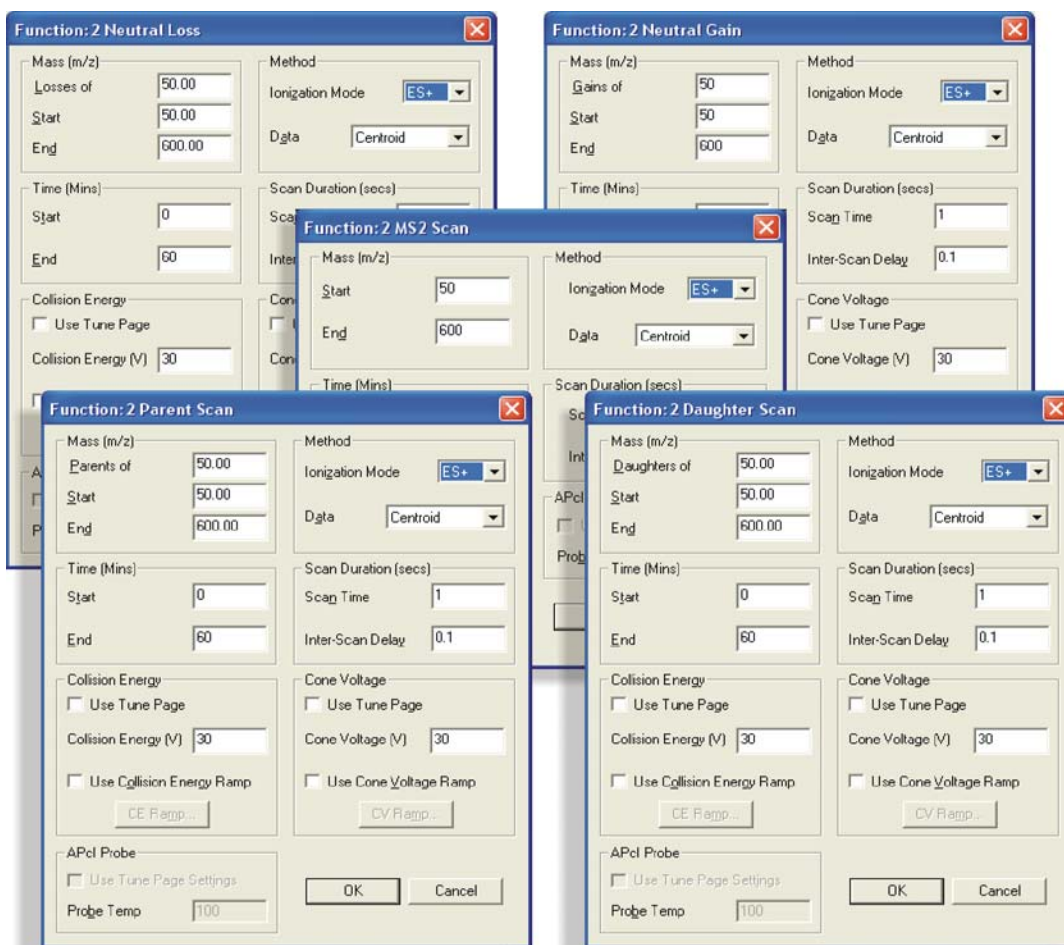


Figure 5-13 MS/MS Scanning Function Editors

Setting-Up a Daughter Scan

This is the most commonly used MS/MS mode; it is used to look at fragmentations of a particular ion. MS1 is set to the parent (precursor) mass and is not scanned.

The resolution of MS1 can be lowered until the peak width at the base is two masses wide, without the daughter spectrum containing any ions from the adjacent parent masses.

You may specify a daughter mass greater than the parent (precursor) mass. In this case, ions which have gained mass in the collision cell, or are of higher mass-to-charge ratio (for example, when a multiply-charged ion fragments and loses a charge) are detected.

[Table 5-8](#) describes the Daughter Scan Function Editor parameters.

Table 5-8 Daughter Scan Function Editor Parameters

Parameter	Description
Mass (m/z)	
Daughters of	The parent mass (m/z).
Start	The mass (m/z) at which the MS2 scan starts.
End	The mass (m/z) at which the MS2 scan ends.
Collision Energy	
Use Tune Page	When selected, the collision energy set in the Tune window at the start of the acquisition is used. Note: <i>The collision energy value cannot be altered during acquisition by typing new values into the Tune window, since the new values are not downloaded during acquisition.</i>
Collision Energy (V)	The collision energy, in electron volts, to be used for the collision cell during the scan.
Use Collision Energy Ramp	When selected, applies a linear ramp to the collision energy (see “Applying a Linear Ramp to the Collision Energy” on page 86).
CE Ramp	Opens the Collision Ramp dialog box (see “Applying a Linear Ramp to the Collision Energy” on page 86). Note: <i>CE Ramp is available only when Use Collision Energy Ramp is selected.</i>

Applying a Linear Ramp to the Collision Energy

1. On the Function Editor, select Use Collision Energy Ramp.
2. Click CE Ramp to open the Collision Ramp dialog box (Figure 5-14).

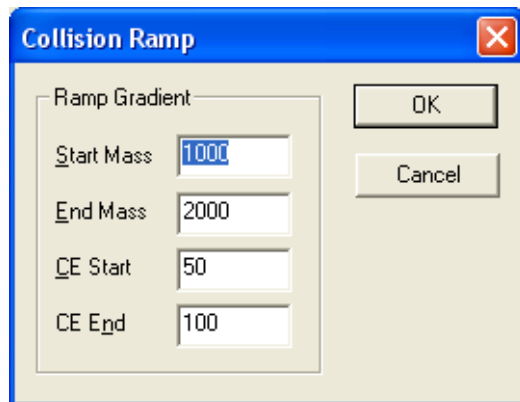


Figure 5-14 Collision Ramp Dialog Box

3. Enter the required parameters (see Table 5-9 for details).

Table 5-9 Collision Ramp Dialog Box Parameters

Parameter	Description
Ramp Gradient	
Start Mass	The mass (m/z) at which the lower collision energy is specified.
End Mass	The mass (m/z) at which the upper collision energy is specified.
CE Start	The lower collision energy.
CE End	The upper collision energy.

Note: The two values of collision energy specify a collision energy gradient that is then extrapolated to cover the full mass range.

Setting-Up a Parent Scan

This mode is used to look for the parent of a particular fragment.

MS2 is set to the mass of the fragment and is not scanned.

There are often several masses from which a daughter may come, so that any one fragment is derived from a number of different peaks.

[Table 5-10](#) describes the parameters unique to the Parent Scan Function Editor.

Table 5-10 Parent Scan Function Editor Parameters

Parameter	Description
Mass (m/z)	
Parents of	The daughter mass (m/z).
Start	The mass (m/z) at which the MS1 scan starts.
End	The mass (m/z) at which the MS1 scan ends.
	<i>Note: Start is normally set just below Parents of and End to a value above the highest expected parent mass.</i>

Setting-Up an MS2 Scan

In this mode, MS2 is resolving while MS1 transmits ions over a wide mass range. While this scanning mode can be used for acquiring data, it is mostly used in the Tune window for setting and optimizing the acquisition conditions.

Setting-Up a Neutral Loss Scan

In this mode, the peak in a spectrum that gives the neutral loss specified in Losses of is detected. The precursor mass is scanned in MS1 and MS2 is scanned at this mass less the neutral loss mass. Starting masses are therefore detected on the mass scale of MS1. Start (for MS1) should be greater than Losses of to give MS2 a valid start mass.

Setting-Up a Neutral Gain Scan

This is an infrequently used mode, since the mass selected by MS2 is seldom higher than that of MS1. It applies to studies where a precursor ion gains mass by ion molecule reaction or where multiply-charged ions fragment into particles with higher m/z values.

5.7.5 Setting-Up an MRM Function

Multiple reaction monitoring (MRM) functions are set-up in a similar way to SIR functions (see [Section 5.7.3](#)) but allow a number of MS/MS transitions (fragmentations) between MS1 and MS2 to be monitored.

All fields in the MRM Function Editor ([Figure 5-15](#)) are similar to those described in [Section 5.7.3](#).

Parent (m/z)	Daughter (m/z)	Dwell (Secs)	Cone (Volts)	Coll Energy (eV)
455.2	165.2	0.1	50	30

Figure 5-15 MRM Function Editor

5.7.6 Setting-Up a Survey Function

Survey scans are used to search for precursor ions.

1. In the MS Method Editor (see [Figure 5-6](#)), delete any existing functions from the Function List.
2. Click  Survey or select Functions > Survey Scan to open the Survey Scan Function Editor.

Note: The MS Method Editor does not add survey functions to the list if non-survey functions are present.

3. Select the required tab and specify the appropriate parameters (see below).

Setting-Up the Parameters for MS and MS/MS Scanning

The Survey and MSMS Template pages (Figure 5-16) allow you to specify the parameters for MS and MS/MS scanning during the survey; they are similar to the MS/MS scanning Function Editor pages (see Section 5.7.4). The MSMS Template page also allows you to select the required Instrument Parameter file (.ipr).

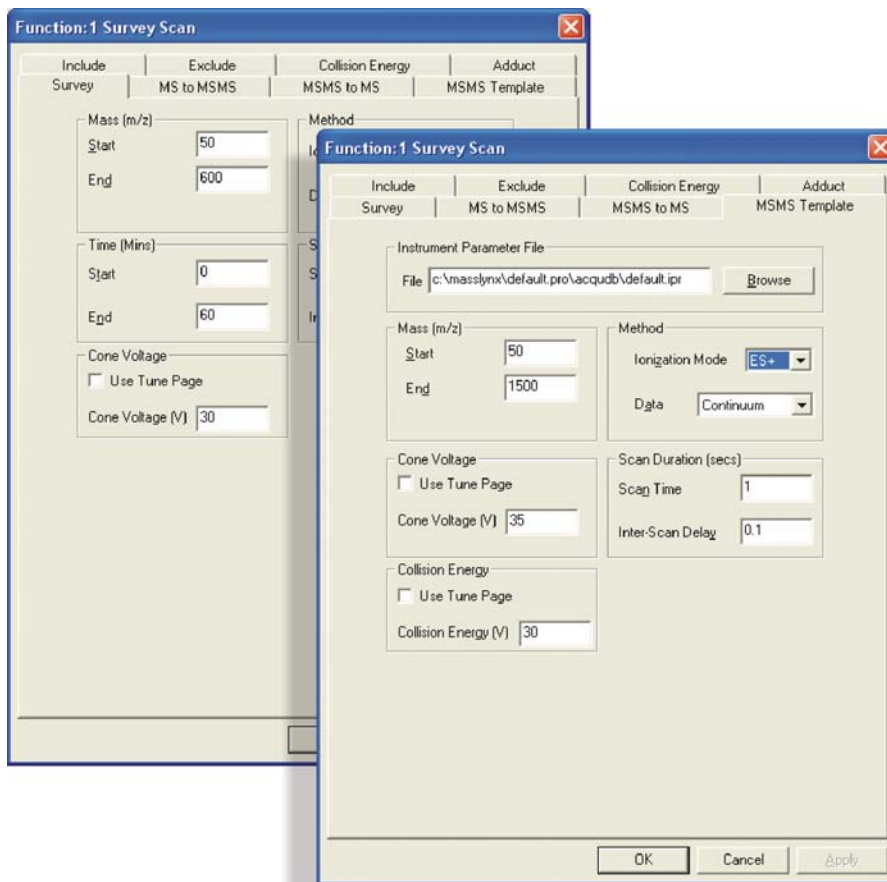


Figure 5-16 Survey Scan Function Editor: Survey and MSMS Template Pages

Setting-Up MS to MS/MS Switching

The MS to MSMS page (Figure 5-17) allows MS to MS/MS switching to be set-up.

Function:1 Survey Scan

Include Survey | Exclude MS to MSMS | Collision Energy MSMS to MS | Adduct MSMS Template

MS to MSMS Switch Criteria

TIC

Intensity

Threshold: 10

Detection Window (Da): 0.5

Number of Components: 1

Retention Time Window (s): 1

Charge State

Tolerance Window +/- (Da): 3

Extraction Window (Da): 2

Precursor Selection

Everything

Included Masses only

Included Masses Take Priority

Detected Precursor Inclusion

Auto Exclude

Always Include

Include After Time (s): 10

Data

Discard uninteresting survey scans

OK Cancel Apply

Figure 5-17 Survey Scan Function Editor: MS to MSMS Page

Table 5-11 describes the MS to MSMS page parameters.

Table 5-11 Survey Scan Function Editor: MS to MSMS Page Parameters

Parameter	Description
MS to MSMS Switch Criteria	
TIC	When selected, the switch from MS to MS/MS scanning occurs when the TIC of the spectrum rises above the specified Threshold value.
Intensity	When selected, the switch from MS to MS/MS scanning occurs when the intensity of the largest peak rises above the specified Threshold value.
Threshold	The value at which the switch from MS to MS/MS scanning occurs (TIC or Intensity).
Detection Window (Da)	When a peak is detected, no other peaks are sought within the mass range specified by plus or minus this value, centered about the detected peak.
Number of Components	Selects the maximum number of peaks of interest to detect from an MS scan. The software will attempt to find the Number of Components from an MS spectrum. Once the maximum number of components in the spectrum for the specified criteria is found, new MS/MS functions are generated automatically and a switch to MS/MS scanning performed.
Retention Time Window (s)	The Retention Time Window is used by the Include and Exclude lists. A mass will be included or excluded if it appears at the specified retention time plus or minus this value.
Charge State	
Tolerance Window +/- (Da)	If a peak is detected, but it is outside the Tolerance Window of where it should be, it is ignored by the Charge State Recognition routines.
Extraction Window (Da)	The window around the base peak for data to be extracted, in Da. For example, if the Extraction Window is 1 Da, the window will extend 0.5 Da either side of the base peak. The extracted data is then used to calculate the charge state.
Precursor Selection	
Everything	Monitors all the valid masses satisfying the selection criteria.

Table 5-11 Survey Scan Function Editor: MS to MSMS Page Parameters (Continued)

Parameter	Description
Included Masses only	Monitors only the masses in the Include List.
Included Masses Take Priority	Masses on the Include List are given priority. If no precursors are found, other valid masses are monitored. Note: A mass is valid if it is not on the Exclude List (see “Editing the Exclude List” on page 101) and it satisfies the precursor selection criteria.
Detected Precursor Inclusion	
Auto Exclude	Only switch on the masses once during an acquisition.
Always Include	Never exclude any masses from being switched on.
Include After Time (s)	Once a mass has been switched on, it will be excluded for the time specified in the adjacent text box. Once the retention time of a mass has exceeded this value, the mass can be considered as an peak of interest again.
Data	
Discard uninteresting survey scans	Stores only the survey scans that detect precursor ions. This saves on disk space because survey scans that contain no relevant data are rejected.

Setting-Up MS/MS to MS Switching

The MSMS to MS page (Figure 5-18) allows MS/MS to MS switching to be set-up.

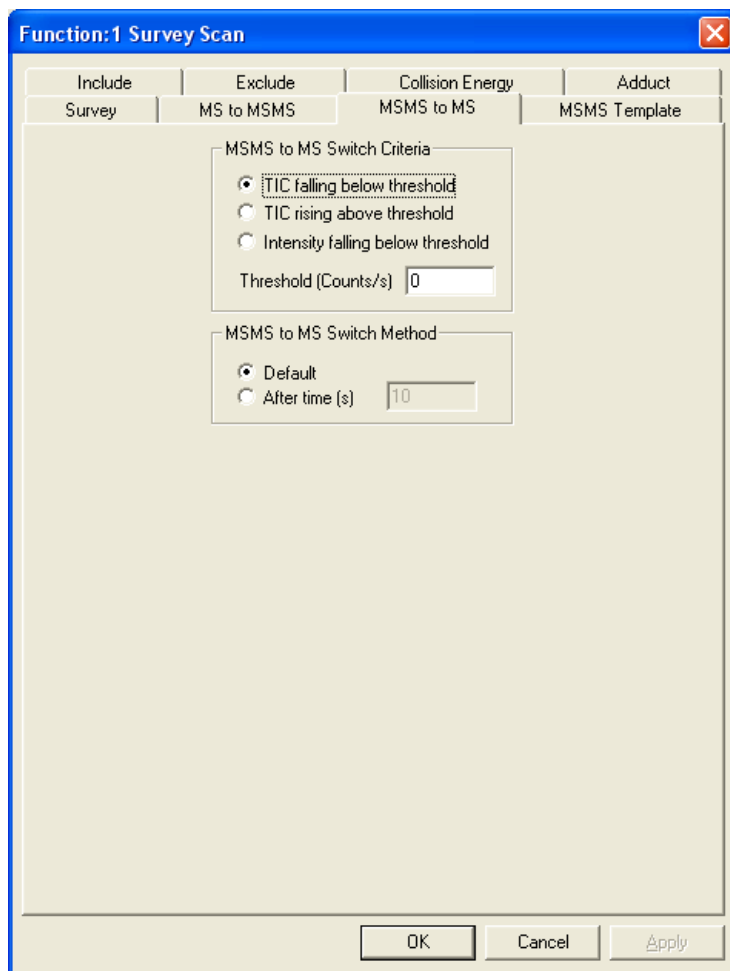


Figure 5-18 Survey Scan Function Editor: MSMS to MS Page

When MS/MS functions have been generated, they are carried out in parallel until the conditions for switching to MS are satisfied. When all MS/MS functions have stopped, the MS survey function is again carried out.

Table 5-12 describes the MSMS to MS page parameters.

Table 5-12 Survey Scan Function Editor: MSMS to MS Page Parameters

Parameter	Description
MSMS to MS Switch Criteria	
TIC falling below threshold	When selected, the switch from MS/MS to MS scanning occurs when the TIC of the spectrum falls below the specified Threshold value.
TIC rising above threshold	When selected, the switch from MS/MS to MS scanning occurs when the TIC of the spectrum rises above the specified Threshold value.
Intensity falling below threshold	When selected, the switch from MS/MS to MS scanning occurs when the intensity of the largest peak falls below the specified Threshold value.
Threshold (Counts/s)	The value at which the switch from MS/MS to MS scanning occurs.
MSMS to MS Switch Method	
Default	When selected, the MS/MS function stops when the MS/MS to MS switch criteria are met.
After time (s)	When selected, the MS/MS function stops when the MS/MS to MS switch criteria are met, or otherwise when the specified time has elapsed.

Including Masses

The Include page (Figure 5-19) allows masses (m/z) that are of interest, and prompt a switch to MS/MS operation, to be set-up.

The screenshot shows a dialog box titled "Function: 1 Survey Scan" with a close button (X) in the top right corner. The dialog is divided into several sections:

- Survey**: Includes tabs for "Include", "MS to MSMS", "MSMS to MS", and "MSMS Template".
- Include**: Contains sub-sections for "Exclude" and "Collision Energy".
- Include Mass**: A section with two checked options: "Range" and "File". Each has an adjacent text input field. A "Browse..." button is next to the "File" input.
- Table**: A table with columns: "Mass", "Retention Time", "Collision Energy", "Cone Voltage", "Charge State", and "Scan". The table is currently empty.
- Buttons**: A row of buttons: "New", "Add...", "Delete", "Save", and "Save As..".
- Include Window +/- (mDa)**: A text input field containing the value "100".
- Charge State**: A section with a checkbox labeled "Use Include By Charge State" (which is unchecked). Below it are two text input fields: "Charge State(s):" and "Number of Include Components:" (containing the value "10").
- Bottom Buttons**: "OK", "Cancel", and "Apply" buttons.

Figure 5-19 Survey Scan Function Editor: Include Page

These masses are listed in the Include List. It is normally used when the Survey Scan Function Editor MS to MSMS page Included Masses only option, or Included Masses Take Priority option is selected.

Table 5-13 describes the Include page parameters.

Table 5-13 Survey Scan Function Editor: Include Page Parameters

Parameter	Description
Include Mass	
Range	Specifies the required masses (m/z), or range of masses, to be used from those in the Include List Box. (Separate individual masses by commas; use an underscore to denote a range of masses, e.g., 510, 520, 550_600, 700.)
File	When selected, the name of the file in which the Include List is to be stored can be entered in the adjacent text box; alternatively, use the Browse button to select an existing file.
New	Resets all the options and clears the Include List.
Add	Opens the Include Masses dialog box, which allows the Include List to be edited (see “Editing the Include List” on page 97).
Delete	Deletes the selected entry from the list.
Save	Saves the Include List details to an existing file.
Save As	Saves the Include List details to a new file.
Include Window +/- (mDa)	Defines the Include Window size, in mDa. If an interesting mass is detected from the survey scan, and is close enough (within the Include Window) to the specified include mass, and all other criteria are met, the mass will be switched on. If the mass is outside the Include Window, it will not be switched on.
Charge State	
Use Include By Charge State	Select to use charge states.
Charge State(s)	Enter the required charge state(s).
Number of Include Components	Enter the number of components to be included.

Editing the Include List

1. On the Survey Scan Function Editor's Include page, click Add to open the Include Masses dialog box (Figure 5-20).

Note: To edit an existing entry in the Include List, double-click it.

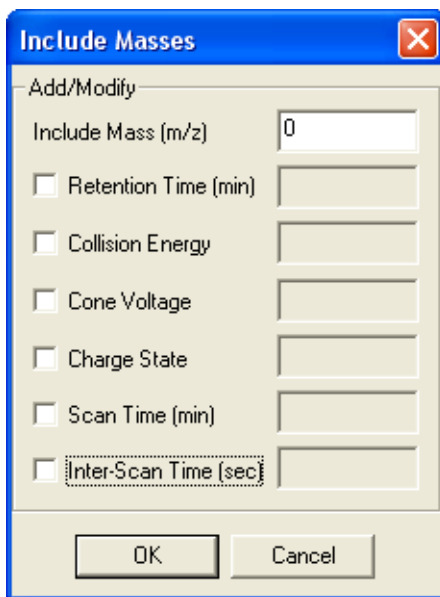


Figure 5-20 Include Masses Dialog Box

2. To enter a value, select the check box for the required option and enter the value in the adjacent text box.

Note: The text box becomes available only when you select an option.

Table 5-14 describes the Include Masses dialog box parameters.

Table 5-14 Include Masses Dialog Box Parameters

Parameter	Description
Add/Modify	
Include Mass (m/z)	The specified mass (m/z) to include in the Function Switching process.
Retention Time (min)	If the specified mass elutes at the specified Retention Time and is within the MS to MSMS page Retention Time Window (s) value, the mass is considered to be on the Include List. This allows masses eluting from a column to be included as a mass of interest.
Collision Energy	The collision energy used to split the detected mass for the MS/MS scans.
Cone Voltage	The Cone Voltage value applied during the MS/MS scans.
Charge State	Include the mass by its charge state.
Scan Time (min)	The Scan Time, in minutes, used during MS/MS scans for the detected mass.
Inter-Scan Time (sec)	The Inter-Scan Time used during MS/MS scans for the detected mass.

Excluding Masses

The Exclude page (Figure 5-19) is used to set-up masses that are to be ignored (i.e., not of interest) so that there is no switch to MS/MS operation. The Exclude List is a list of these masses.

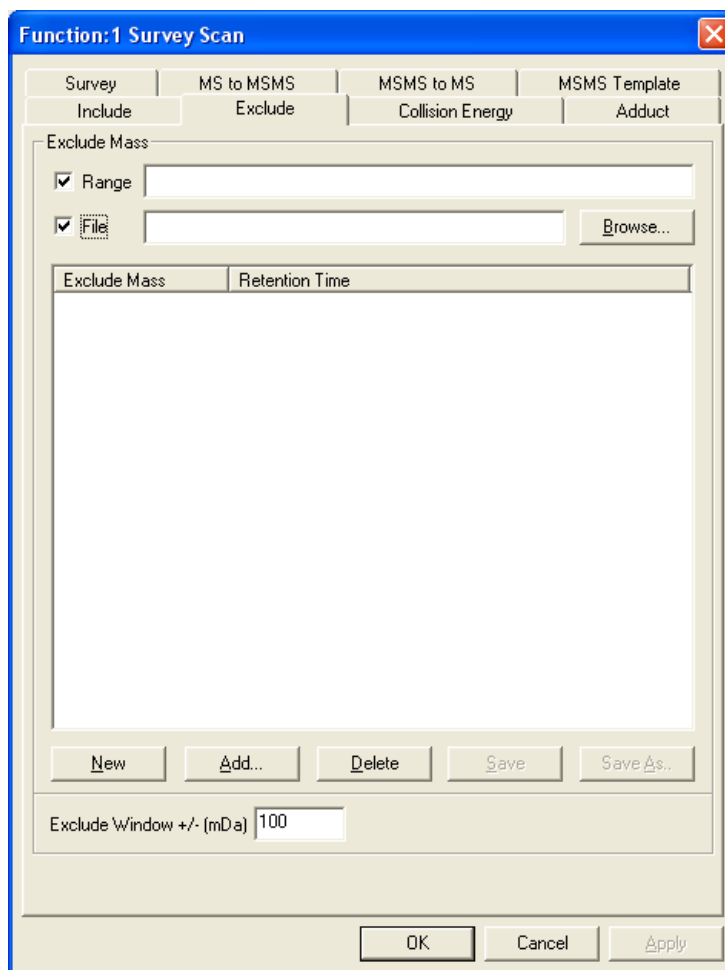


Figure 5-21 Survey Scan Function Editor: Exclude Page

Table 5-15 describes the Exclude page parameters.

Table 5-15 Survey Scan Function Editor: Exclude Page Parameters

Parameter	Description
Exclude Mass	
Range	Specifies the required masses (m/z), or range of masses, to be used from those in the Exclude List. (Separate individual masses by commas; use an underscore to denote a range of masses, e.g., 510, 520, 550_600, 700.)
File	When selected, the name of the file in which the Exclude List is to be stored can be entered in the adjacent text box; alternatively, click the Browse button to select an existing file.
New	Resets all the options and clears the Exclude List.
Add	Opens the Exclude Mass dialog box, which allows the Exclude List to be edited (see “Editing the Exclude List” on page 101).
Delete	Deletes the selected entry from the list.
Save	Saves the Exclude List details to an existing file.
Save As	Saves the Exclude List details to a new file.
Exclude Window +/- (mDa)	Defines the Exclude Window size, in mDa. If a mass is detected from the survey scan, it is close enough (within the Exclude Window) to the specified exclude mass, and all other criteria are met, the mass will be ignored.

Editing the Exclude List

1. On the Survey Scan Function Editor's Exclude page, click Add to open the Exclude Mass dialog box (Figure 5-22).

Note: To edit an existing entry in the Exclude List, double-click on it.

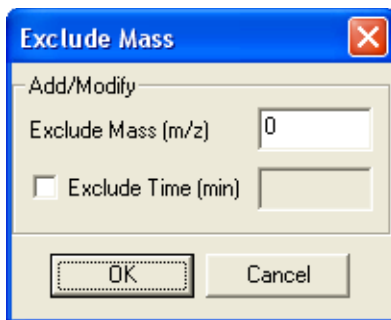


Figure 5-22 Exclude Mass Dialog Box

2. Enter the values as required.

Table 5-14 describes the parameters in the Exclude Mass dialog box.

Table 5-16 Exclude Mass Dialog Box Parameters

Parameter	Description
Add/Modify	
Exclude Mass (m/z)	The specified mass (m/z) to exclude from the Function Switching process.
Exclude Time (min)	If the Exclude Mass is detected at the Exclude Time, in minutes, and is within the MS to MSMS page Retention Time Window (s) value, it will be excluded from the Function Switching process.

Selecting Collision Energy Options

The Collision Energy page (see [Figure 5-19](#)) allows you to set-up or select one of three Collision Energy options.

Function: 1 Survey Scan

Survey | MS to MSMS | MSMS to MS | MSMS Template
Include | Exclude | Collision Energy | Adduct

Default Collision Energy

Use Default Collision Energy

Collision Energy Profile

Use Collision Energy Profile

File Browse...

Modify...

Charge State Recognition

Use Charge State Recognition

Maximum Number Of Charge States:

CS1 File Browse...

CS2 File Browse...

CS3 File Browse...

CS4 File Browse...

Modify...

OK Cancel Apply

Figure 5-23 Survey Scan Function Editor: Collision Energy Page

[Table 5-15](#) describes the Collision Energy page parameters.

Table 5-17 Survey Scan Function Editor: Collision Energy Page Parameters

Parameter	Description
Default Collision Energy	
Use Default Collision Energy	Uses a single default value, which is configured on the MSMS Template page (see Figure 5-16). This allows either the Tune window value or the value specified on the MSMS Template page to be used. If the Tune window value is to be used, the value can be changed during the scan, but it only becomes effective on the next switch from MS to MS/MS.
Collision Energy Profile	
Use Collision Energy Profile	Selecting this option enables the controls in the Collision Energy Profile frame. The Collision Energy Profile is a range of Collision Energy values, which are associated with a specified mass. You can enter up to five different Collision Energy values. When an interesting mass is detected in an MS scan, and the mass is on the Collision Energy Profile list, these Collision Energy values are used during the MS/MS scans for the mass.
File	The file name of the Collision Energy Profile file.
Browse	Allows you to browse for the required Collision Energy Profile file.
Modify	Opens the CE Profile dialog box, which allows a table of collision energy profiles to be created for masses that are being switched on (see “Creating a Table of Collision Energy Profiles” on page 105).

Table 5-17 Survey Scan Function Editor: Collision Energy Page Parameters (Continued)

Parameter	Description
Charge State Recognition	<p>Selecting this option enables the controls in the Charge State Recognition frame.</p> <p>When a mass of interest is detected in an MS scan, its charge state is calculated. Using the mass and its charge state, a Collision Energy value can be obtained from the entered charge state table. This value is then used during MS/MS scanning.</p> <p>The Include page's Include by Charge State and the Collision Energy page's Use Charge State Recognition functionalities can be used independently, or both can be configured to combine their individual functionalities (see “Charge State Recognition Functionality” on page 108, for further details).</p>
Maximum Number Of Charge States	<p>Enter the number of charge states to be considered (maximum four) in this text box; the following CSx File text boxes are enabled, as appropriate, by this action.</p>
CS1 File, etc.	<p>Enter the file name of the required charge state file, or use the adjacent Browse button to search for a file.</p>
Modify	<p>Opens the Modify Charge State dialog box; this is used to create, or modify charge state files (see “Creating or Modifying Charge State Files” on page 106, for details).</p>

Creating a Table of Collision Energy Profiles

To create a Collision Energy Profile Table for masses that are being switched on:

1. On the Survey Scan Function Editor's Collision Energy page, select Use Collision Energy Profile; the associated Modify button is enabled.
2. Click Modify to open the CE Profile dialog box (Figure 5-24).

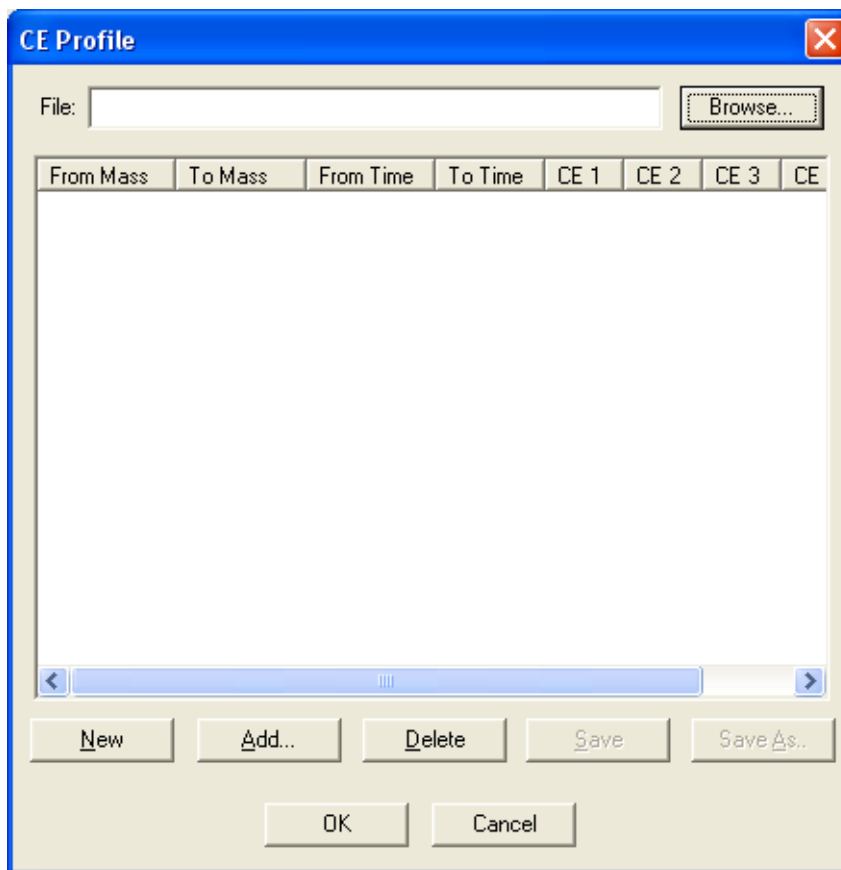


Figure 5-24 CE Profile Dialog Box

3. Click Add, or double-click an existing entry, to open the Collision Energy dialog box. This allows new entries to be added to the Collision Energy Profile Table.

[Table 5-18](#) describes the Collision Energy dialog box parameters.

Table 5-18 Collision Energy Dialog Box Parameters

Parameter	Description
Add/Modify	
From Mass (m/z)	Start mass (m/z) for the mass range in which the collision energy profile is to be used.
To Mass (m/z)	End mass (m/z) for the mass range in which the collision energy profile is to be used.
From Time (min)	Start time for the time range in which the collision energy profile is to be used.
To Time (min)	End time for the time range in which the collision energy profile is to be used.
CE1, etc.	The five allowed collision energy values, which will be applied to a relevant mass at a relevant time during MS/MS scanning. Selecting the check box for one value enables the check box for the following value.
OK	Closes the Collision Energy dialog box and enters the selected values in the CE Profile dialog box Collision Energy Profile Table.

Creating or Modifying Charge State Files

Use the Modify Charge State dialog box ([Figure 5-25](#)) to create or modify charge state files.

1. On the Survey Scan Function Editor's Collision Energy page (see [Figure 5-23](#)), select Use Charge State Recognition.
2. Enter the required Maximum Number of Charge States.
3. Click Modify to open the Modify Charge State dialog box.

Note: The number of Modify CS tabs displayed in this dialog box corresponds with the Maximum Number of Charge States selected in [step 2](#).

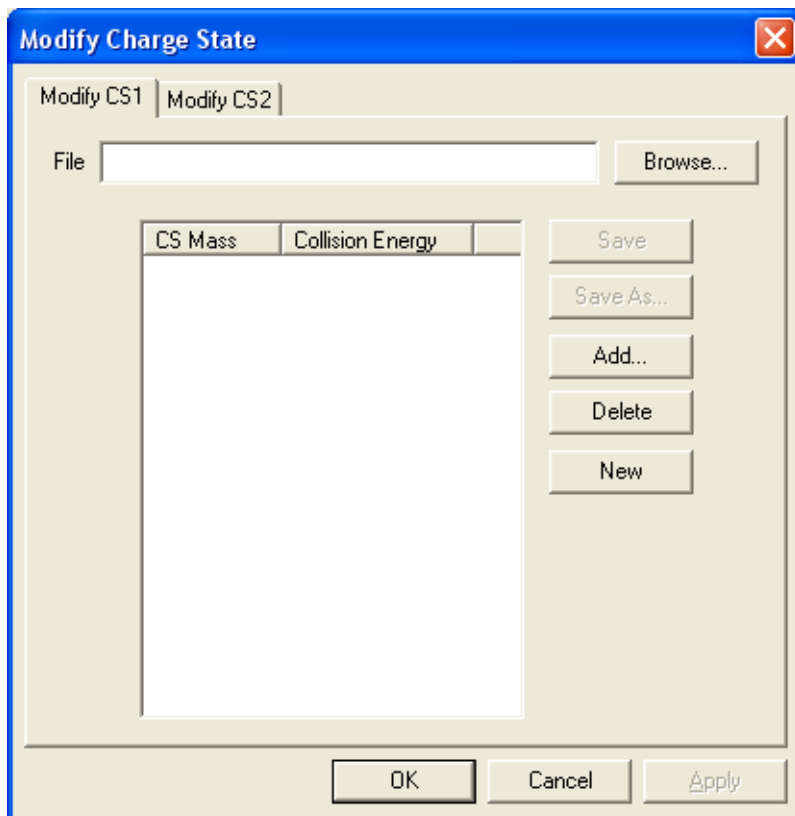


Figure 5-25 Modify Charge State Dialog Box

Adding New Entries to the Collision Energy Profile Table

1. Click the appropriate Modify CS tab.
2. Click Add or double-click an existing entry to open the Charge State Mass dialog box.
3. Enter the required values. [Table 5-19](#) describes the parameters in the Charge State Mass dialog box.

Table 5-19 Charge State Mass Dialog Box Parameters

Parameter	Description
Add/Modify	
CS Mass (m/z)	Specifies the mass (m/z) of interest.
Collision Energy	Sets the correct collision energy value to break up the mass of interest.
OK	Closes the Charge State Mass dialog box and enters the selected values in the Modify Charge State dialog box Mass List.

Saving a Collision Energy Profile Table

Click Save to save the collision energy profile table to a text file. If the collision energy profile table has not been saved previously, a standard Windows dialog box is opened, allowing you to specify the file name and path. Click Save As to save an existing list to a new filename.

Opening a Collision Energy Profile Table

Click Browse to navigate to the collision energy profile table text file, or enter the filename and path in the File text box.

Charge State Recognition Functionality

You can apply charge state recognition functionalities from the Survey Scan Function Editor Include (see [Figure 5-19](#)) and Collision Energy (see [Figure 5-23](#)) pages. The functionalities, Include by Charge State and Use Charge State Recognition, can operate independently or together.

Using Charge State Recognition with Include by Charge State Disabled

This allows the software to switch on any peak of interest, but a collision energy value will be calculated only for those masses with a charge state that matches those set up in the Charge State Recognition frame. For all other masses, the default collision energy will be applied; this is either the Tune window value, or the value specified in the Survey Scan dialog box MSMS Template page.

For example, if the Include by Charge State option is disabled, and the Charge State Recognition has been configured for Charge States 1 and 2, any detected mass of interest switches on in the normal way. Masses switched on with a charge state of 1 or 2 will have

a collision energy value calculated from the Charge State Recognition table. All other masses use the specified default collision energy value.

Using Charge State Recognition with Include by Charge State Enabled

This restricts the masses that are switched on to those with the correct charge state. A collision energy value will be calculated for those masses with a charge state that is configured in both Include by Charge State and Charge State Recognition. For masses with a charge state which is only on the Include by Charge State list, a default collision energy value will be used.

For example, if the Include by Charge State section is configured to allow charge states of 2 and 3, and the Charge State Recognition section is configured for Charge States 1 and 2, only masses with a charge state of 2 or 3 are switched on. Masses switched on with a charge state of 2 will have a collision energy value calculated from the Charge State Recognition table. Masses with a charge state of 3 will use the specified default collision energy value.

Creating an Adduct List

In MassLynx, an adduct is the mass of an ion added to an original mass.

The Adduct page ([Figure 5-26](#)) allows you to create a list of such masses.

If a mass of interest (target mass) on the Include List is detected, the mass spectrometer is switched to MS/MS operation and that mass is added to the Exclude List, so that it is ignored if detected again. Also, when the target mass is detected, any mass in the Adduct page adduct list is added to that target mass, and the resulting mass added to the Exclude List so that the adduct is also ignored if it is detected.

For example, for positive ion operation:

If the molecular target mass (M) is 200, the target mass (m/z) on the Include List is 201 ($M + H^+$). When detected, a switch to MS/MS operation is triggered and the mass is added to the Exclude List.

A typical adduct is sodium (Na, mass 23), for which the target adduct mass is 223 ($M + Na^+$). Hence the value to be added to the Adduct page adduct list is $(M + Na^+) - (M + H^+) = (223 - 201) = 22$.

Similarly, if the adduct is ammonium (NH_4 , mass 18), the target adduct mass is 218 ($M + NH_4^+$). Hence the value to be added to the Adduct page adduct list is $(M + NH_4^+) - (M + H^+) = (218 - 201) = 17$.

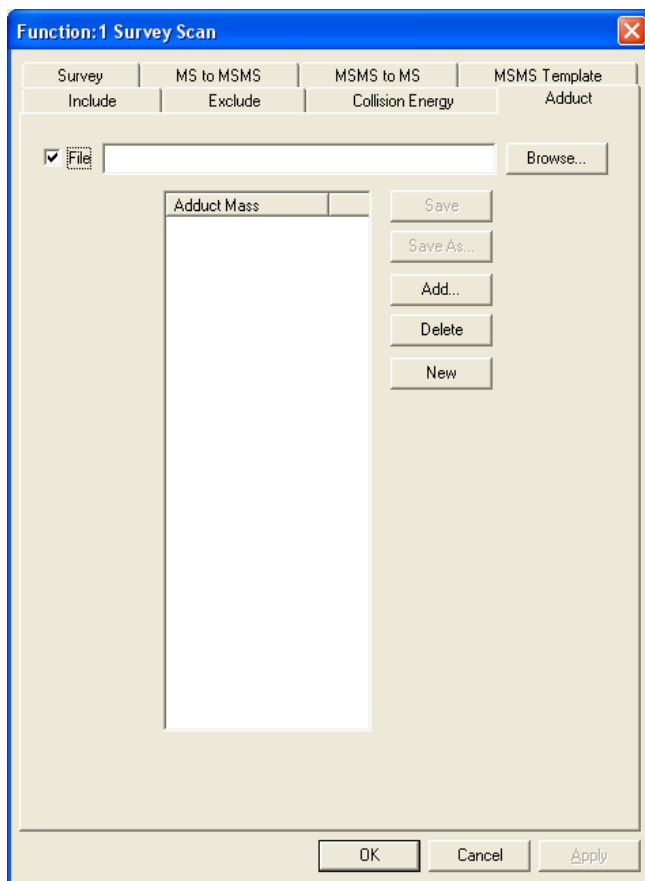


Figure 5-26 Survey Scan Function Editor: Adduct Page

Adding an Adduct Mass to the Adduct List

1. Click Add, or double-click an existing mass in the adduct list to open the Adduct Mass dialog box.
2. Enter the required value in the Add/Modify, Step Size (m/z) text box.
3. Click OK to add the value to the adduct list.

Saving an Adduct List

Click Save to save the adduct list to a text file. If the adduct list has not been saved previously, a standard Windows dialog box is opened, allowing you to specify the file name and path. Click Save As to save an existing list to a new filename.

Opening an Existing Adduct List

Click **Browse** to navigate to the adduct list text file, or enter the filename and path in the **File** text box.

5.7.7 Monitoring Acquisitions

When an acquisition is started, the **Function Switching Status** dialog box opens, showing the precursors currently running.

Chapter 6

Setting-Up Mass Calibration

6.1 Overview

MassLynx can perform a fully automated instrument mass calibration. This calibrates the instrument for static and scanning modes of acquisition over a variety of mass ranges and scanning speeds.

A mass spectrum of a reference sample (a calibration file) is acquired and is compared with the expected masses of the peaks in the sample (stored as a reference file). The mass differences between the reference peaks and calibration peaks are calculated; these are the calibration points. A calibration curve is then fitted through the calibration points.

Each calibration point's vertical distance from the curve is calculated. This represents the remaining (or residual) mass difference after calibration.

The standard deviation of the residual masses is also calculated. This value is the best single indication of the calibration's accuracy.

6.1.1 Types of Calibration

Each quadrupole analyzer requires up to three calibration curves:

- A static calibration is used to “park” the analyzer accurately on a specific mass of interest (for example, in tuning, SIR and MRM).
- A scanning calibration enables peaks acquired in a scanning acquisition to be mass measured accurately.
- A scan speed compensation calibration compensates for lag time in the system when the instrument is scanned rapidly.

A separate mass spectrum of the reference sample is acquired for each selected calibration type (see the *MassLynx User's Guide* for details).

6.1.2 Calibration Process

The calibration process comprises the following steps:

1. Tuning the instrument.
2. Selecting the appropriate reference file for the reference sample to be used.
3. Starting an automatic calibration.
4. Checking the resulting calibration report.

6.2 Calibrating for Electrospray Operation

When a calibration ends, the instrument may be used to acquire data over any mass range and scan speed within the calibrated range.

Calibration over the instrument's full mass range can be achieved by using a mixture of sodium iodide and rubidium iodide, however the following example uses ammoniated PEG to calibrate over a more limited mass range such as that typically used by most operators.

6.2.1 Preparing for Calibration

Introducing the Reference Sample

This calibration example describes an automatic calibration, which requires reference sample to be present for several minutes. The reference sample is best introduced using the instrument's syringe pump (see [Appendix B](#)).

1. Fill the syringe with the reference sample, PEG-NH₄⁺ (see [Appendix D](#)).
2. Couple the syringe to the electrospray probe, using fused silica tubing.
3. Fit the syringe to the syringe pump.
4. In the MassLynx Tune window, select the type of syringe used (see [Section C.5](#)).
5. On the ES+ Source page, set Pump Flow (μL/min) to 10.

Tuning

1. If the MassLynx Tune window (Figure 6-1) is not already open, click the MassLynx window Instrument shortcut bar MS Tune icon.
2. If the ES+ Source page is not displayed, select Ion Mode > Electrospray+.
3. Set the parameters to the recommended values shown in Table 6-1.

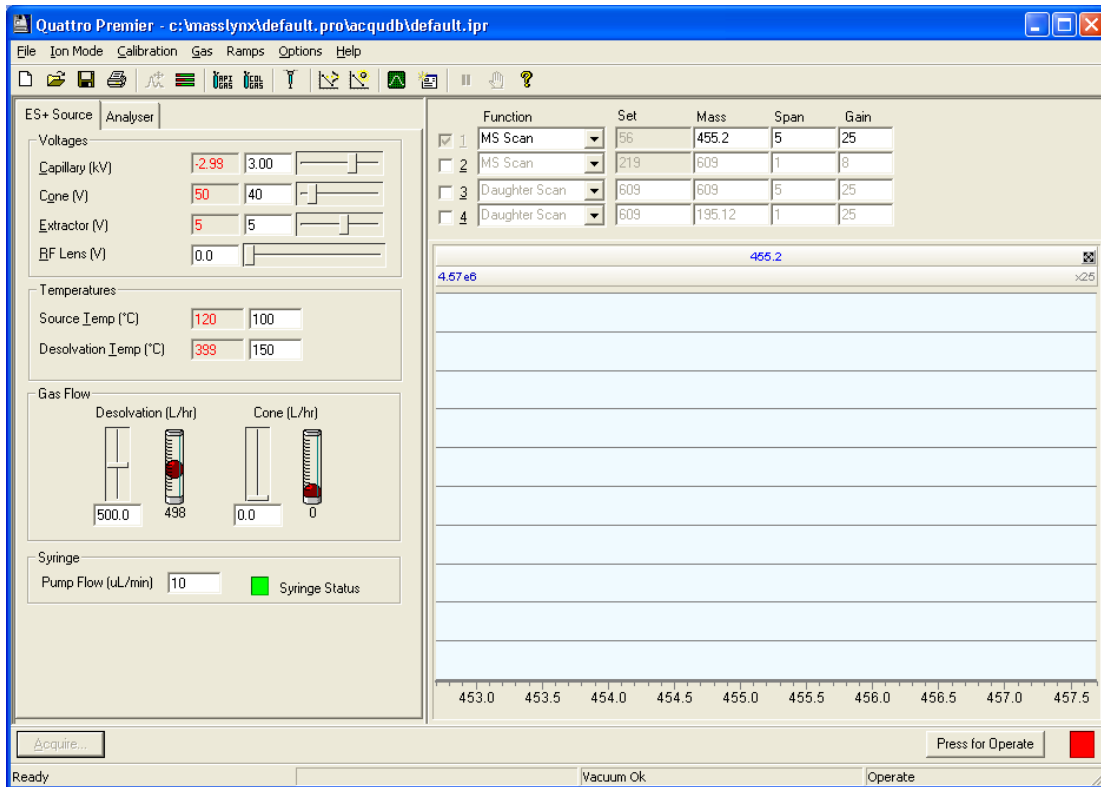


Figure 6-1 Initial Tune Window



Table 6-1 Recommended ES+ Source Page Parameter Values

Parameter	Recommended Value
Voltages	
Capillary (kV)	3.00
Cone (V)	40
Extractor (V)	5
RF Lens (V)	0.0
Temperatures	
Source Temp (°C)	100
Desolvation Temp (°C)	150
Gas Flow	
Desolvation (L/hr)	500.0
Cone (L/hr)	0

4. Click the Analyser tab.
5. Set the Analyser page parameters to the recommended values shown in [Table 6-2](#).

Table 6-2 Recommended Analyser Page Parameter Values

Parameter	Recommended Value
Analyser	
LM Resolution 1	15.0
HM Resolution 1	15.0
Ion Energy 1	1.0
Entrance	50
Collision	2
Exit	50.0
LM Resolution 2	15.0
HM Resolution 2	15.0
Ion Energy 2	1.0
Multiplier	650

6. Click  to start the nitrogen flow.
7. On the ES+ Source page, observe the Desolvation and Cone gas flows; confirm they are stable and are correct.
8. Click Press for Operate, and confirm that the adjacent instrument status indicator shows green.
9. Click  to start the syringe pump.
10. In the Tune window, observe the displayed PEG masses (Figure 6-2).

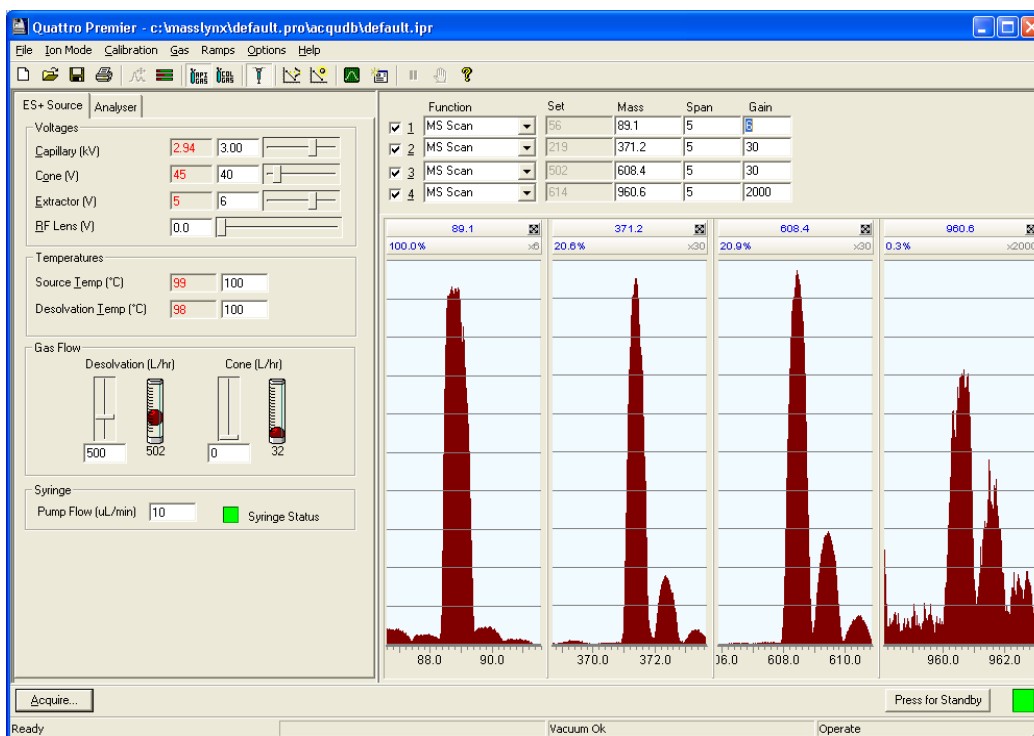


Figure 6-2 Tune Window with PEG Peaks

11. On the Analyser page, set LM Resolution 1, HM Resolution 1 and Ion Energy 1 to give unit mass resolution for all the displayed peaks (the isotopes should be resolved).
12. Confirm that no ions are saturated. (If a peak is at full screen and at a gain of 1, it is saturated; adjust the value of Multiplier, or dilute the sample.)

13. Select MS2 Scan in each of the Function drop-down lists.
14. Observe the displayed PEG masses, and confirm that the resolution is correct. Adjust LM Resolution 2, HM Resolution 2, and Ion Energy 2 if necessary.

6.2.2 Selecting the Calibration Options

In the Tune window, select Calibration > Calibrate Instrument to open the Calibration Options window (Figure 6-3).

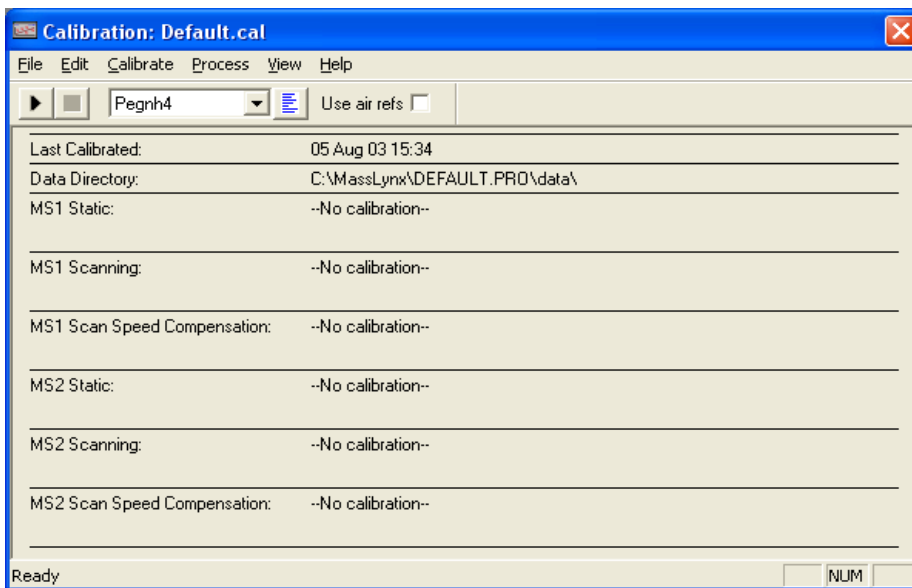


Figure 6-3 Calibration Options Window

Selecting the Reference File

For the PEG reference sample, select Pegnh4 from the reference file drop-down list.

Removing the Current Calibrations

1. Select Calibrate > Default.
2. When prompted, click Yes to save the changes to the default.cal file.

This ensures that a file with no calibration is currently active on the instrument and prevents any previously saved calibrations from being overwritten or modified.

6.2.3 Setting the Calibration Parameters

When the software is initially loaded, default calibration parameters are set. These usually give a suitable calibration, but under some conditions they may need adjusting.

Setting the Automatic Calibration Check Parameters

Automatic calibration check parameters define limits that the calibration must attain before the instrument is successfully calibrated. These parameters are set in the Automatic Calibration Check dialog box (Figure 6-4), which is opened by selecting Edit > AutoCal Check Parameters in the Calibration Options window. Table 6-3 gives details of the Automatic Calibration Check dialog box parameters.

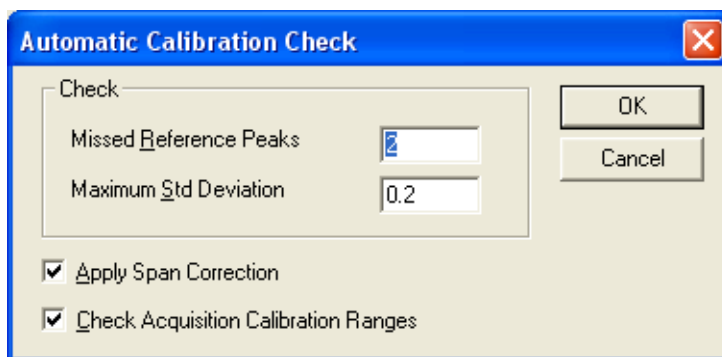


Figure 6-4 Automatic Calibration Check Dialog Box

Table 6-3 Automatic Calibration Check Dialog Box Parameters

Parameter	Description
Check	
Missed Reference Peaks	Specifies the maximum number of consecutive peaks that may not be matched when comparing the reference spectrum and the acquired calibration spectrum. The calibration fails if this number is exceeded. The default value for this parameter, 2, is suitable in most cases.
Maximum Std Deviation	This parameter is set to a default value of 0.2. During calibration, the difference between the measured mass in the acquired calibration file and the true mass in the reference file is determined for each pair of matched peaks. If the standard deviation of the resulting set of mass differences exceeds the specified value, the calibration fails. Reducing the value of Maximum Std Deviation gives a more stringent limit. Increasing the value of Maximum Std Deviation means that the requirement is easier to meet but may allow incorrect peak matching. Values greater than 0.2 should not be used except under unusual conditions.
Apply Span Correction	This option should always be selected; it allows different mass ranges to be scanned, within the calibrated range, without affecting mass assignment
Check Acquisition Calibration Ranges	This option allows warning messages to be displayed if an attempt is made to acquire data from outside the calibrated range for mass and scan speed. You should leave this option selected.

Setting General Calibration Parameters

Set the calibration parameters in the Calibration Parameters dialog box ([Figure 6-5](#)), which is opened by selecting Edit > Calibration Parameters in the Calibration Options window.

The Peak Match parameters determine the limits within which the acquired data must lie for the software to recognize the calibration masses and result in a successful calibration. The default values are shown in [Figure 6-5](#).

Note: For low scan speeds (up to 1000 Da/s), set Initial error (Da) to 2. For higher scan speeds (up to 4000 Da/s), increase this value to 4.

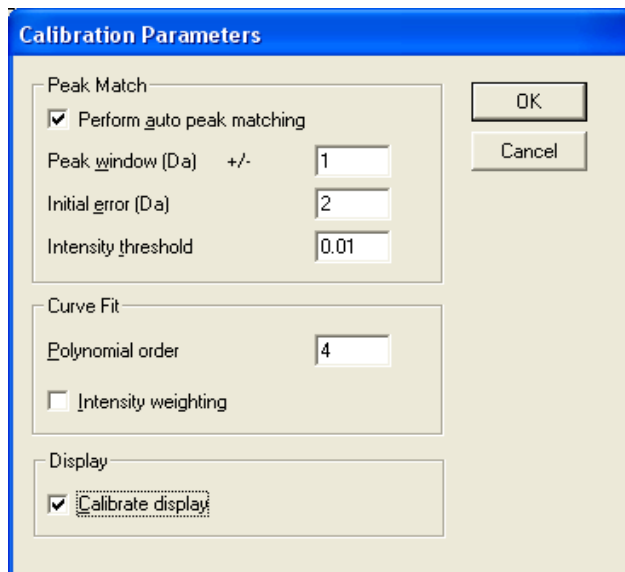


Figure 6-5 Calibration Parameters Dialog Box

Increasing the Peak window and Initial error values gives a greater chance of incorrect peak matching. All peaks in the acquired spectrum below the Intensity threshold value (measured as a percentage of the most intense peak in the spectrum) are not used in the calibration procedure.

The Polynomial order of the curve has values from 1 to 5 as the available options. You should only use values 2 and 4:

- An order of 2 is suitable for wide mass ranges (e.g., 2 to 3000 Da), or for calibrating with widely spaced reference peaks. Sodium iodide, in particular, has widely spaced peaks (150 Da apart), and horse heart myoglobin is used to calibrate higher up the mass scale, hence this is the recommended polynomial order for these calibrations.
- An order of 4 is typically used for calibrations which include the lower end of the mass scale, and have closely-spaced reference peaks. This is suitable for calibrations using PEG (e.g., 50 to 1050 Da), however it is not suitable for calibrations with widely-spaced reference peaks.

6.2.3.1 Selecting Mass Measure Parameters

Specify the mass measure parameters and parameter values in the Mass Measure Parameters dialog box (Figure 6-6), which is opened by selecting Edit > Quad Mass Measure Parameters in the Calibration Options window.

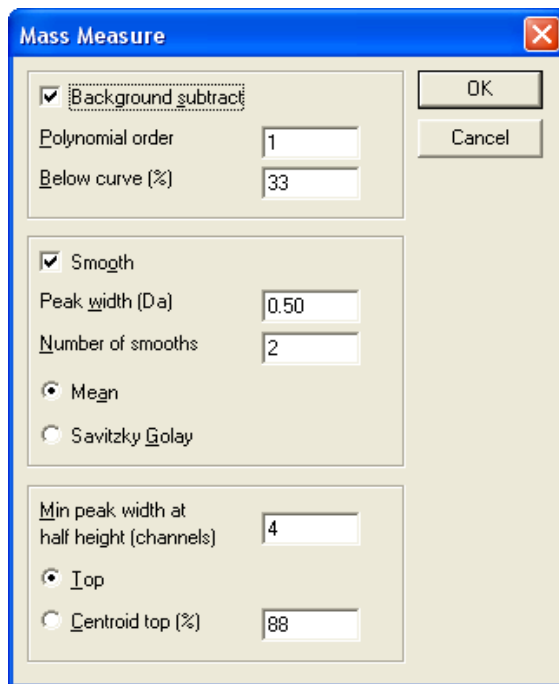


Figure 6-6 Mass Measure Dialog Box

If continuum or MCA data are acquired for calibration, these parameters must be set before the calibration is performed. If centroided data are used for calibration, the mass measure parameters are not used. The data type (continuum, MCA, or centroided) is selected in the Calibration Acquisition Setup dialog box (see “[Selecting the Acquisition Parameters](#)” on page 124).

With electrospray calibrations, particularly with sodium iodide which has some low intensity peaks at higher mass, it is recommended that continuum or MCA data are acquired.

Note: *It is important that the data are smoothed correctly, and that the peak width at half height (PWHH) is entered in the smoothing parameters, as shown in [Figure 6-6](#).*

At high scan speeds, instrument resolution may decrease. Ensure that the centroiding parameters are set to use the top of the peak so that mass assignment of peaks is accurate.

6.2.4 Performing a Calibration

Three types of calibration are available with MassLynx:


- Static
- Scanning
- Scan speed compensation

It is recommended that all three types of calibration are performed, so that mass ranges and scan speeds can be changed while maintaining correct mass assignment. However, it is possible to have any combination of these calibrations:

- If only a static calibration is present, the instrument is calibrated for acquisitions where the quadrupoles are held at a single mass, as in SIR or MRM.
- If only a scanning calibration is present, the instrument is only correctly calibrated for scanning acquisitions over the same mass range, and at the same scan speed, as those used for the calibration.
- For the scan speed compensation to be used correctly, a scanning calibration should also be performed.
- If static and scanning calibrations are both present, the instrument is calibrated for acquisitions where the quadrupole is held at a single mass, and for scanning acquisitions with a mass range which lies within the mass range of the scanning calibration, providing that the same scan speed is used.

For example, if the instrument is calibrated from m/z 100 to 900 with a 2 s scan (400 Da/s), data can be acquired from 100 to 500 Da with a 1 s scan time (also 400 Da/s) whilst maintaining correct mass assignment. In this case, the static calibration would be used to determine the start mass of the acquisition, and the scanning calibration would be used for mass assignment and scan range.

- If scanning calibration and scan speed compensation are present, the instrument is only calibrated for scanning acquisitions over the same mass range as that used for the calibration, but the scan speed can be changed, provided that it remains within the scan speeds used for the two calibrations. The mass range should not be changed, as there is no static calibration to locate the start mass.

Select the calibration types in the Automatic Calibration dialog box (Figure 6-7), which is opened by clicking , or by selecting Calibrate > Start from the Calibration Options window.

Note: *Data acquisition will not start at this point.*

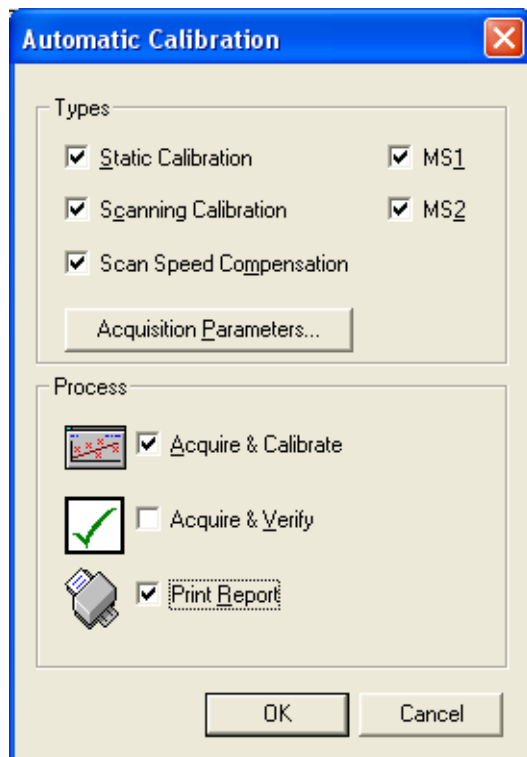


Figure 6-7 Automatic Calibration Dialog Box

If a complete calibration is required:

1. Select Static Calibration, Scanning Calibration, and Scan Speed Compensation.
2. Select MS1 and MS2.
3. Select the Acquire & Calibrate and Print Report check boxes.

Selecting the Acquisition Parameters

1. Click Acquisition Parameters in the Automatic Calibration dialog box to open the Calibration Acquisition Setup dialog box (Figure 6-8), and set the mass range, scan speeds, and acquisition mode.

The parameters in this dialog box define the limits of scan range and speed for the instrument and calibration parameters. When the instrument is fully calibrated, any mass range or scan speed is allowed within the upper and lower limits dictated by the parameters (50 to 1050 Da and 200 to 2000 Da/s, respectively, in the example shown in Figure 6-8).

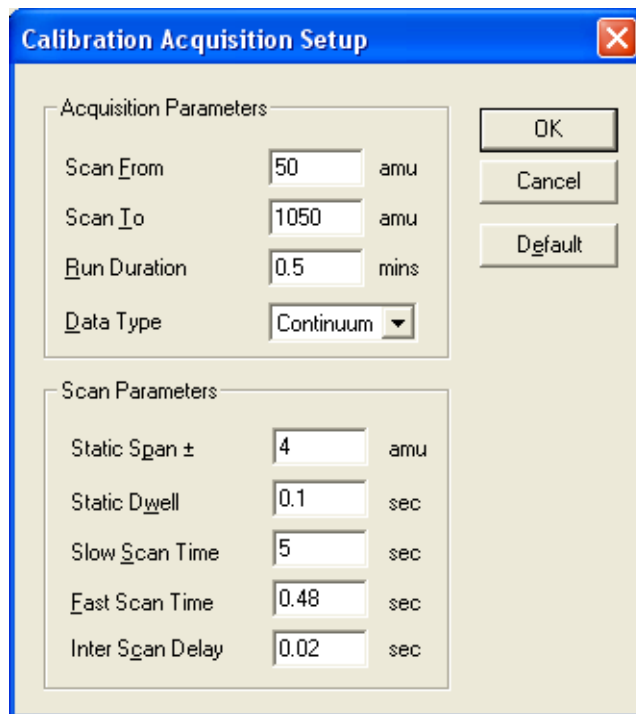



Figure 6-8 Calibration Acquisition Setup Dialog Box

2. When the required parameters have been set, click OK to return to the Automatic Calibration dialog box.

Starting the Calibration Process

1. Ensure that the syringe contains sufficient reference sample to complete the calibration (enough for 3 minutes in this example, i.e., 30 μ L minimum).
2. In the Tune window, click  to start the syringe pump.
3. Click OK on the Automatic Calibration dialog box (see [Figure 6-7](#)).

The instrument acquires all of the calibration files in the order shown in [Table 6-4](#).

Once all of the data have been acquired, the functions in each data file are combined to give a single spectrum which is then compared against the reference spectrum to form a calibration. This process takes place in the same order as shown in [Table 6-4](#). If the Calibration Options dialog box is open, a constantly updated status message for the calibration is displayed.

Table 6-4 Calibration Order and Data Files

Calibration	Data File
MS1 static	STATMS1
MS1 scanning	SCNMS1
MS1 scan speed compensation	FASTMS1
MS2 static	STATMS2
MS2 scanning	SCNMS2
MS2 scan speed compensation	FASTMS2

If the calibration statistics meet the requirements specified by the selected calibration parameters when the process is completed, a message appears informing you that the calibration has been successful and the Calibration Options window is updated to reflect the calibration results (Figure 6-9). A calibration report is then printed showing a calibration curve for each of the calibration processes.

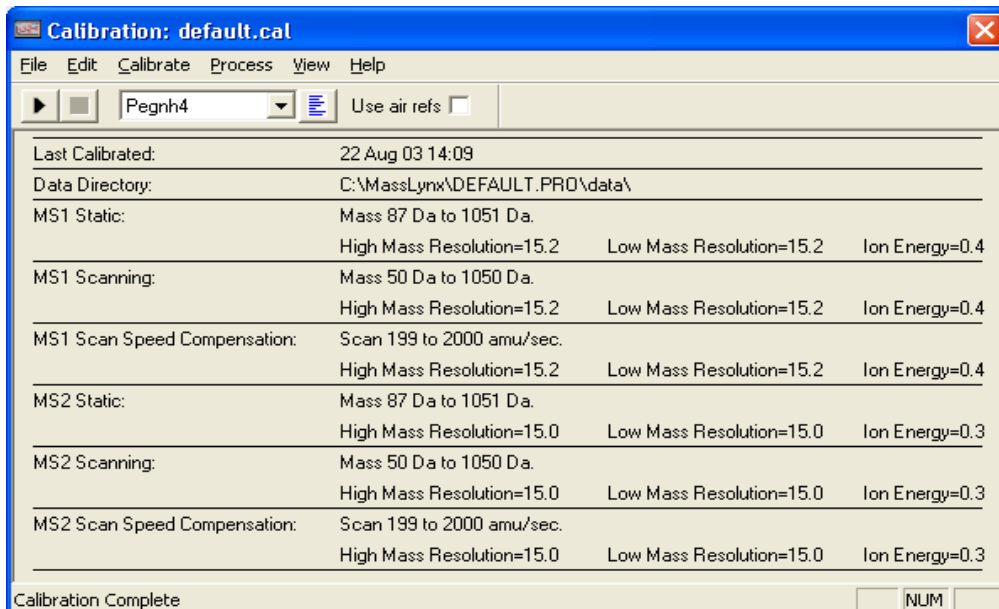


Figure 6-9 Calibration Options Window with Completed Calibration Details

Save the acquisition under a suitable file name for it to be effective (see Section 6.2.9).

6.2.5 Calibration Failure

If the calibration statistics do not meet the requirements, a message describes at what point, and why, the calibration failed. This message also states where the attempted calibration's data can be viewed so that the exact cause of failure can be determined.

A calibration fails under these conditions:

- No peaks are detected.
- Too many consecutive peaks are missed.

The Calibration Fails Because No Peaks Are Detected

The calibration fails if the acquired calibration data file contains no peaks. This may be due to:

- Lack of reference sample.
- Wrong scans or wrong data file being used for the calibration.
- No flow of solvent into the source.
- Multiplier (on the Tune window Analyser page) set too low.

The Calibration Fails Because Too Many Consecutive Peaks Are Missed

The calibration fails if the number of consecutive peaks not found exceeds the value in the Missed Reference Peaks parameter, which you specify in the Automatic Calibration Check dialog box (see [Figure 6-4](#)). Peaks may be missed for the following reasons:

- The reference sample supply is nearly depleted, so the less intense peaks are not detected.
- The Multiplier value, specified on the Tune window Analyser page, is too small, so the less intense peaks are not detected.
- An incorrect ionization mode is selected. Ensure that the data have been acquired with Ion Mode set to ES+ in the Tune window.

Note: *It is possible to calibrate in negative ion mode electrospray, using the `naineg.ref` reference file with a suitable reference sample.*

- The Intensity threshold value, specified in the Calibration Parameters dialog box (see [Figure 6-5](#)), is too high. Peaks appear in the acquired calibration file but are ignored because they fall below this threshold level.

- The value for the Initial error (Da), or Peak window (Da) parameter, specified in the Calibration Parameters dialog box (see [Figure 6-5](#)), is too small. Hence, the calibration peaks lie outside the limits set by these parameters.
- The Maximum Std Deviation, specified in the Automatic Calibration Check dialog box (see [Figure 6-4](#)), has been exceeded.
- The wrong reference file has been selected. Ensure that the correct file (nairb.ref in this case) is selected in the Calibration Options window (see [Figure 6-3](#)).

If too many consecutive peaks have been missed, view the data in the on-screen calibration report to see whether the missed peaks are present in the acquired calibration file.

Correcting the Calibration Failure

1. Determine the failure's cause, as explained above, and rectify it.
2. Repeat the calibration.
3. If the Intensity threshold, Initial error (Da) and Peak window (Da) have been adjusted to obtain a successful calibration, view the on-screen calibration report to confirm that the correct peaks have been matched.

Note: *With a low threshold, and wide ranges set for the Initial error (Da) and Peak window (Da), it may be possible to select the wrong peaks and still get a "successful" calibration. This is particularly relevant for calibrations using PEG, where there may be peaks from PEG+H⁺, PEG+NH₄⁺, PEG+Na⁺, and doubly-charged species.*

4. Click OK in the calibration report window to accept the new calibration, or select Cancel to retain the previous calibration.

6.2.6 Incorrect Calibration

If the suggested calibration parameters have been used and, providing that good calibration data have been acquired, then the instrument should be calibrated correctly, that is, the acquired spectrum looks like the reference spectrum and all of the expected peaks are highlighted.

However, in some circumstances it is possible to meet the calibration criteria without matching the correct peaks. This situation is unusual, but the on-screen calibration report should always be checked to verify that the correct peaks have been matched.

Peak matching errors may occur under the following conditions:

- In the Calibration Parameters dialog box (see [Figure 6-5](#)):

- The Peak window (Da) value is too high (>1.5).
- The Initial error (Da) value is too high (>2.0).
- The Intensity threshold is set to 0.
- In the Automatic Calibration Check dialog box (see [Figure 6-4](#)):
 - The Maximum Std Deviation value is too high (>0.2).

An alternative cause of incorrect calibration is from contamination or background peaks. If a contamination or background peak lies within one of the peak matching windows and is more intense than the reference peak in that window, then the wrong peak is selected. Under some conditions this may happen with PEG. There are two ways to counter this:

1. If the reference peak is closer to the center of the peak window, the peak window can be narrowed until the contamination peak is excluded. Take care to ensure that no other reference peak is excluded.
2. If the reference peak is not closer to the centre of the peak window, or if by reducing the window other reference peaks are excluded, then the calibration can be edited manually (see [Section 6.2.8](#)).

6.2.7 Manually Checking the Calibration

1. View the calibration (whether successful or failed) in more detail by selecting Calibrate > From File in the Calibration Options window (see [Figure 6-3](#)) to open the Display Calibration Graphs dialog box ([Figure 6-10](#)). With the required calibration selected, the correct calibration file is automatically opened.

The Browse button allows you to select a file for use by the calibration. The selected file must be from the appropriate project.

2. Click OK to repeat the calibration procedure for that particular file and display a calibration report (see [Figure 6-11](#)).

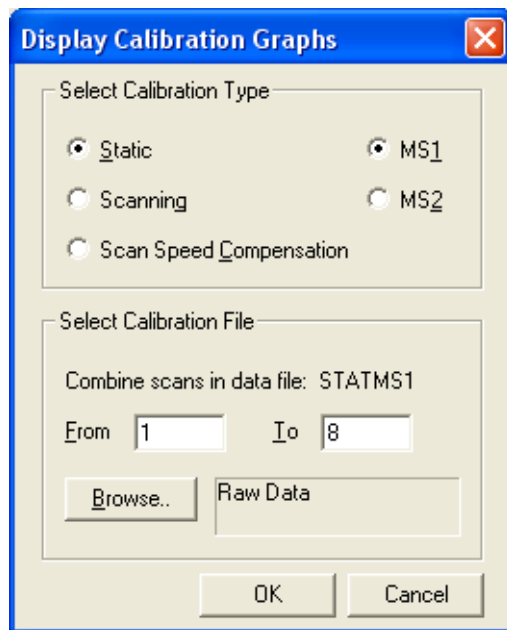


Figure 6-10 Display Calibration Graphs Dialog Box

The calibration report contains four displays:

- The acquired spectrum
- The reference spectrum
- A plot of mass difference against mass (the calibration curve)
- A plot of residual against mass

Display an expanded region by clicking and dragging with the mouse button. This allows the less intense peaks in the spectrum to be examined to verify that the correct peaks have been matched. The peaks in the acquired spectrum that have been matched with a peak in the reference spectrum are highlighted in a different color.

Calibration has been successful if the acquired spectrum looks like the reference spectrum and all of the expected peaks are highlighted.

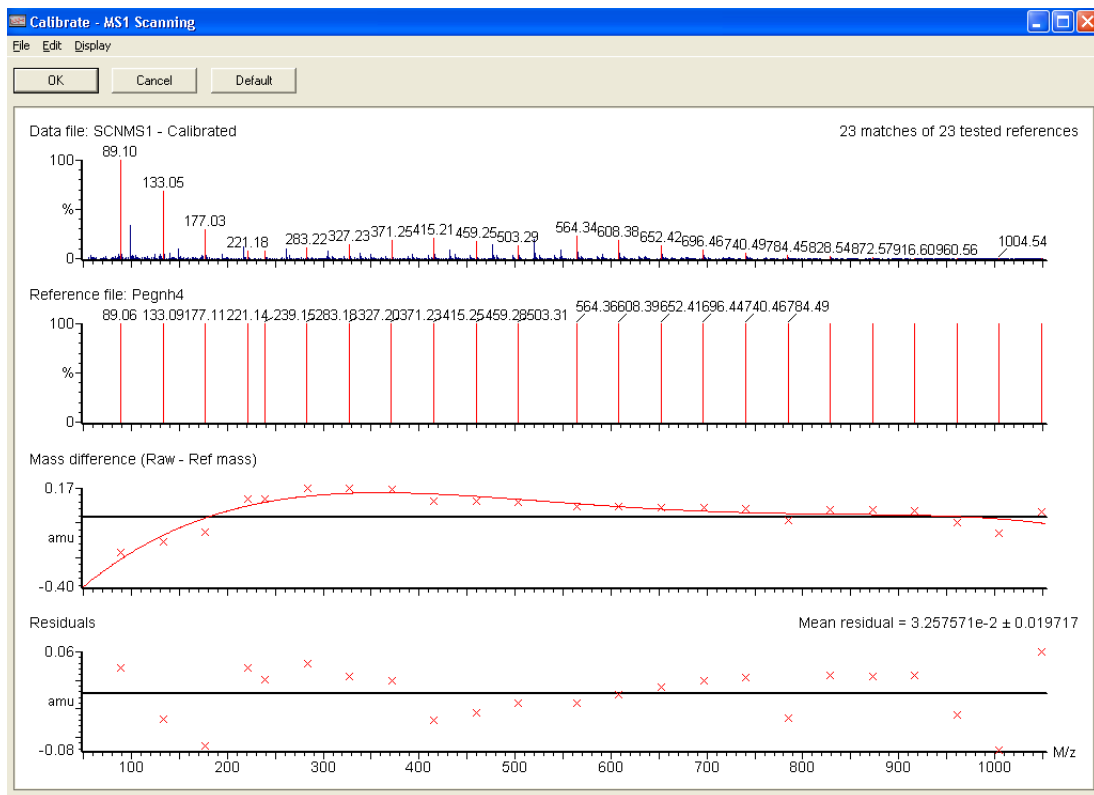


Figure 6-11 Calibration Report

6.2.8 Manually Editing the Peak Matching

If an incorrect peak has been matched in the calibration process, it can be excluded manually from the on-screen calibration report as follows:

1. Right-click the peak in the displayed reference spectrum.
2. Right-click the peak in the displayed acquired spectrum.

The peak is excluded and is no longer highlighted.

If the true reference sample peak is present, but has not been matched, it can be included in the calibration by right-clicking on it to match it with the closest peak in the reference spectrum.

Note: Manually editing a peak does not affect the other matched peaks in the calibration.


6.2.9 Saving the Calibration

1. If the displayed calibration is acceptable, click OK in the Calibration Report (see [Figure 6-11](#)). The Calibration Options window (see [Figure 6-3](#)) is reopened and updated to display the current calibration.
2. Select File > Save As to save this calibration file using the Save As dialog box.

When recalled, the calibration has the same constraints of mass range and scan speed. The ion energy and resolution settings used for the calibration acquisition are also recorded, as these can affect the mass assignment.

6.2.10 Verifying the Calibration

Once a full instrument calibration is in place, it is not always necessary to repeat the full calibration procedure when the instrument is next used. Instead a calibration verification can be performed. (There is no benefit in verifying each calibration individually, recalibration is just as quick.)

1. In the Tune window, select Calibration > Calibrate Instrument to open the Calibration Options window (see [Figure 6-3](#)). Confirm that the current calibration is correctly displayed.
2. Open the Automatic Calibration dialog box by clicking  or selecting Calibrate > Start.
3. Clear the Acquire & Calibrate check box, and select Acquire & Verify.
4. Ensure that the syringe contains sufficient reference sample to complete the calibration (enough for 3 minutes in this example, i.e., 30 μ L minimum).
5. Click OK to start the verification procedure.

When the acquisition ends, the data are combined into single spectrum which is compared against the reference file. A calibration curve is drawn and a report printed in a similar way to when the original calibration was performed.

Unlike the original calibration procedure, the instrument calibration is not changed and the a verification report is printed.

Chapter 7

Maintenance Procedures

7.1 Maintenance Schedule

[Table 7-1](#) lists periodic maintenance schedules that ensure optimum instrument performance.

The maintenance frequencies shown apply to instruments that normally receive moderate use.

Table 7-1 Maintenance Schedule

Maintenance Procedure	Frequency
Gas-ballast the rotary pump (see Section 7.3).	Weekly (ESI). Daily (APCI).
Inspect and adjust the rotary pump oil level (see Section 7.4).	Weekly.
Change the rotary pump oil (see Section 7.5).	Every 3000 hours of pump operation.
Replace the rotary pump oil mist filter element (see Section 7.6).	Six-monthly. Note: For applications that contaminate the rotary pump oil, this period will be reduced and must be determined from experience.
Replace the rotary pump odor filter element (see Section 7.6).	Monthly, or whenever the rotary pump emits an oily odor.
Replace the scroll pump seals.	Annually (see the Edwards document <i>XDS 35i Instruction Manual A730-01-880</i> , supplied with the instrument).
Clean the source assembly (see Section 7.7).	When sensitivity decreases to unacceptable levels.
Clean the ESI probe tip (see Section 7.8).	When sensitivity decreases to unacceptable levels.

Table 7-1 Maintenance Schedule (Continued)

Maintenance Procedure	Frequency
Clean the APCI probe tip (see Section 7.10).	When sensitivity decreases to unacceptable levels.
Clean the corona discharge pin (APCI mode) (see Section 7.9).	When sensitivity decreases to unacceptable levels.
Clean the ion block assembly (see Section 7.7).	When it is visibly fouled. When background or high peak contaminants are unacceptably high.
Clean all source components (see Section 7.7).	When sensitivity decreases to unacceptable levels. When cleaning the cone gas cone, sample cone, and gas exhaust port fails to improve analytical results.
Replace the ESI probe capillary (see Section 7.12).	When sensitivity decreases to unacceptable levels.
Perform ZSpray APCI probe maintenance (see Section 7.13).	When sensitivity decreases to unacceptable levels.

7.2 Safety and Handling

Bear in mind the following safety considerations when performing maintenance procedures.



Warning: *The instrument components may be contaminated with biologically hazardous materials. Always wear nitrile gloves while handling the components.*



Warning: *To prevent injury, always observe good laboratory practices when handling solvents, changing tubing, or operating the instrument. Know the physical and chemical properties of the solvents used (see the Material Safety Data Sheets for the solvents in use).*



Warning: *To avoid electric shock, do not remove the instrument's panels. There are no user-serviceable items inside the instrument.*



Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing any maintenance.



Warning: The probe and source may be hot. To avoid burns, take great care while working with these components.

7.3 Gas-Ballasting the Rotary Pump (E2M28)

When the rotary pump draws large quantities of solvent vapors, the vapors tend to condense in the pump oil, reducing pump efficiency. Gas-ballasting purges condensed contaminants from the oil and returns any oil to the pump from the oil mist filter.

Gas-ballast the rotary pump when any of the following conditions apply:

- With ESI operation, once a week.
- With frequent APCI operation, once a day.
- If the pump oil appears cloudy.
- If the vacuum pressure is higher than normal.
- If condensate forms in the rotary pump exhaust line.
- When changing the rotary pump oil.
- If the level of accumulated oil in the oil mist filter is high.



Warning: To avoid burns, take great care while working with the rotary pump, as it may be hot.



Caution: Failure to routinely gas-ballast the rotary pump shortens oil life and consequently pump life.



Caution: Do not vent the instrument when the rotary pump is gas-ballasting.



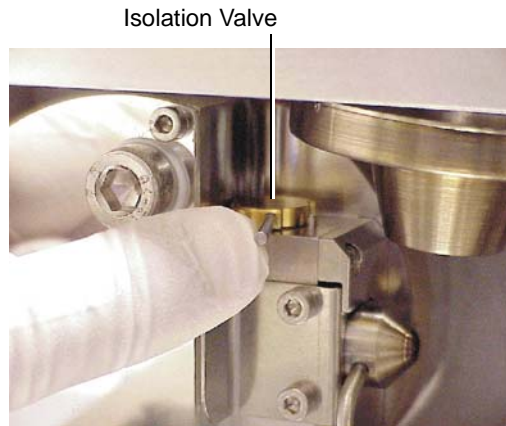
Caution: Do not gas-ballast the rotary pump while the instrument is in Operate.



Caution: Never gas-ballast the rotary pump for more than 2 hours.

Procedure

1. Close the isolation valve by moving its lever fully to the right.



2. Operate the gas-ballast control ([Figure 7-1](#)).
3. When the oil is clear and has drained back to the rotary pump, return the gas-ballast control to its normal position.
4. Open the isolation valve.

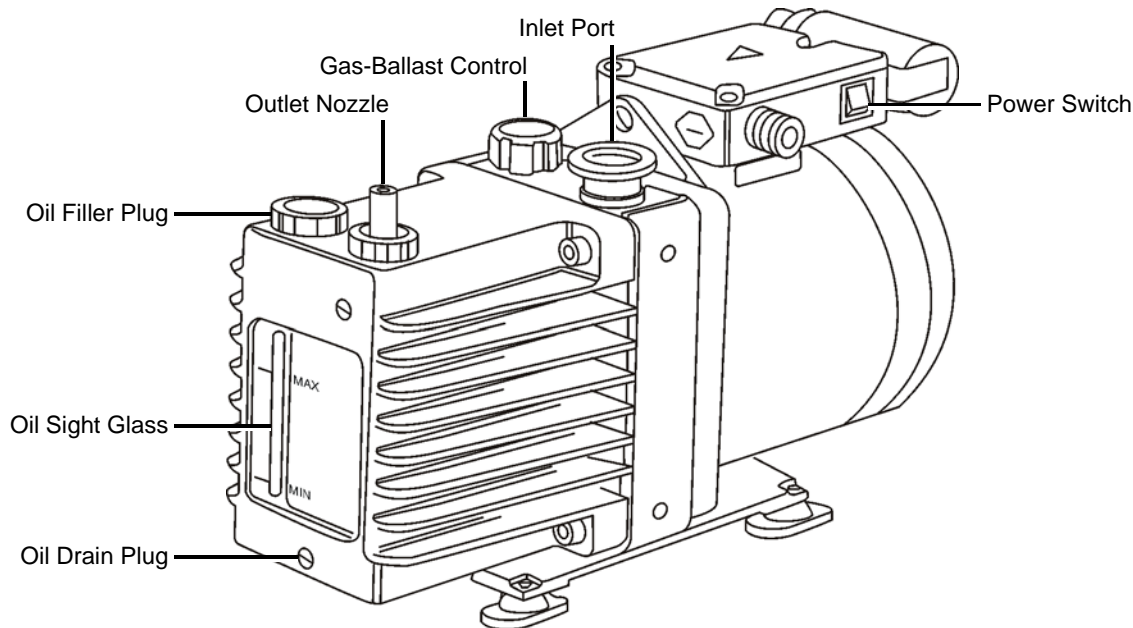


Figure 7-1 Rotary Pump

7.4 Checking the Rotary Pump Oil



Warning: To avoid burns, take great care while working with the rotary pump, as it may be hot.

The oil level can be checked while the pump is operating. However, the instrument must be vented and shut down before adding oil.

The rotary pump oil level appears in the oil level sight glass on the pump (see [Figure 7-1](#)).

Check the oil level at weekly intervals; at all times it should be at, or near, the MAX level indicated by the markings beside the sight glass. If oil must be added, vent and shut down the instrument before removing the oil filler plug (see [Section A.2.3](#)).

Examine the oil each time the its level is checked. It should be colorless and free of visible contaminants. If the oil is discolored, change it (see [Section 7.5](#)).

7.5 Changing the Rotary Pump Oil

Change the rotary pump oil every 3 to 4 months, or whenever it becomes noticeably discolored.

7.5.1 Required Materials

- Nitrile gloves
- Flat-blade screwdriver
- Container for used oil
- Funnel
- Vacuum oil – use only Ultragrade 19 or Inland Q45 (Edwards 45) vacuum pump oil

7.5.2 Procedure



Warning: *The rotary pump oil may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while handling the oil and ensure that the waste oil is correctly disposed of according to local environmental regulations.*

1. Operate the pump to warm the oil before draining it.
2. Gas-ballast the rotary pump (see [Section 7.3](#)).
3. Vent and shut down the instrument (see [Section A.2.3](#)).
4. Power-off the power supply to the backing pump interlock box.



Warning: *To avoid burns, take great care while working with the rotary pump and pump oil, as they may be hot.*

5. Raise the pump 6 to 8 inches (150 to 200 mm) above the floor, if necessary.
6. Place an object under the motor to tilt the pump toward the side on which the oil drain plug is located (see [Figure 7-1](#)).
7. Remove the oil filler plug.
8. Use the flat-blade screwdriver to remove the oil drain plug.
9. Let the oil drain completely.
10. Fit the oil drain plug.

11. Remove the object used to tilt the pump.
12. Fill the pump until the oil in the sight glass reaches the MAX level.
13. Allow a few minutes for the oil to drain into the pump.
14. Recheck the oil level and, if necessary, add more oil.
15. Fit the oil filler plug and, if applicable, lower the pump to the floor.
16. Switch on the pump, by operating the power switch.
17. Switch on the power supply to the backing pump interlock box.

7.6 Replacing the Oil Mist and Odor Filter Elements

7.6.1 Required Materials

- Nitrile gloves
- 4-mm hex (Allen) key
- Lint-free cloth

7.6.2 Procedure



Warning: The rotary pump oil may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while handling the oil mist filter assembly.



Warning: To avoid burns, take great care while working with the rotary pump and oil mist filter assembly, as they may be hot.

1. Gas-ballast the rotary pump to drain the oil from the oil mist filter (see [Section 7.3](#)).
2. Turn off the power supply to the rotary pump interlock box.
3. Wipe clean the outside of the oil mist filter body.

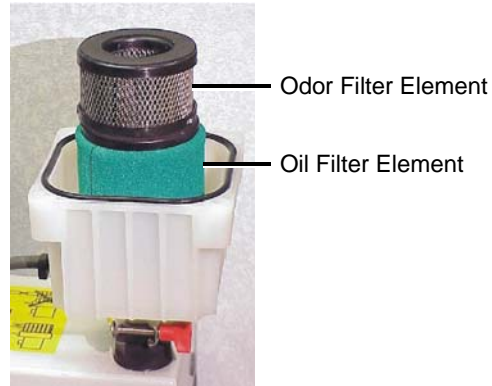
4. Use the hex (Allen) key to remove the four screws that secure the filter upper body to the lower body.



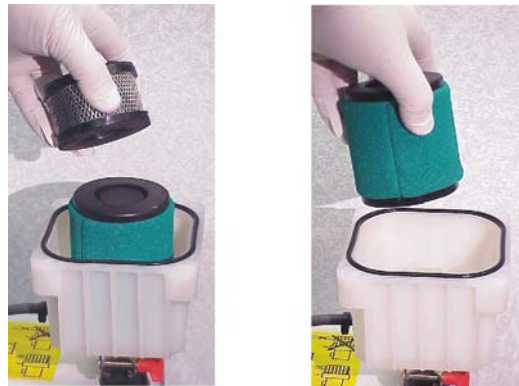
5. Remove the upper body from the lower body.



This exposes the oil filter element and odor filter element.

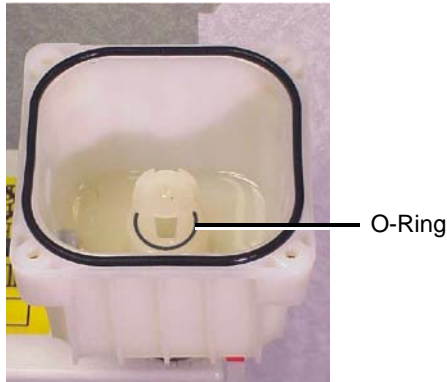


6. Lift out the element(s) that are to be replaced.



7. Wipe clean the insides of the upper and lower body assemblies.

8. Ensure that the oil filter element O-ring is in position on the lower body assembly.



9. Fit the new filter elements to the lower body assembly, ensuring that the foam sealing rings are correctly seated on the top and bottom of the elements.
10. Fit the upper body assembly to the lower body assembly.
11. Fit and tighten the four screws that secure the upper body to the lower body.

7.7 Cleaning the Source Components

7.7.1 Overview

Clean the source components (sample cone, cone gas cone, and gas exhaust port) when:

- They are visibly fouled.
- LC and sample-related causes for decreased signal intensity have been dismissed.

If cleaning these parts fails to increase signal sensitivity, also clean the extraction cone, source T-Wave, and ion block.

Procedure


1. Disassemble the source components and source T-Wave assembly (see [Sections 7.7.3 to 7.7.8](#)).
2. Clean the source components and source T-Wave assembly (see [Sections 7.7.10 to 7.7.14](#)).

3. Reassemble the source T-Wave assembly and the source components (see [Sections 7.7.15 to 7.7.20](#)).

7.7.2 Required Materials

- Nitrile gloves
- Needle-nose pliers
- Two wrenches
- Hex (Allen) keys
- Jeweller's screwdriver
- Large, flat-blade screwdriver
- Glass-fiber pen
- Appropriately sized glass vessels in which to completely immerse components when cleaning. Use only glassware not previously cleaned with surfactants.
- HPLC-grade methanol
- HPLC-grade water
- Formic acid
- Ultrasonic bath
- Source of oil-free, inert gas (nitrogen or helium) for drying (air-drying optional).
- Lint-free paper towels

7.7.3 Removing the Probe from the Source

1. Disconnect the LC system from the probe.
2. In the MassLynx window, click Press for Standby and confirm that the adjacent instrument status indicator shows red.
3. Wait for 3 minutes to allow the desolvation gas flow to cool the probe and source.
4. Click  to turn off the nitrogen flow.



Warning: *The probe and source may be hot. To avoid burns, take great care while working with the instrument's access door open.*

5. Open the instrument's access door.
6. Disconnect the electrical connection(s) at the instrument front panel ([Figure 7-2](#)).

Note: The probe electrical connection will not be connected if the APCI probe is being used.

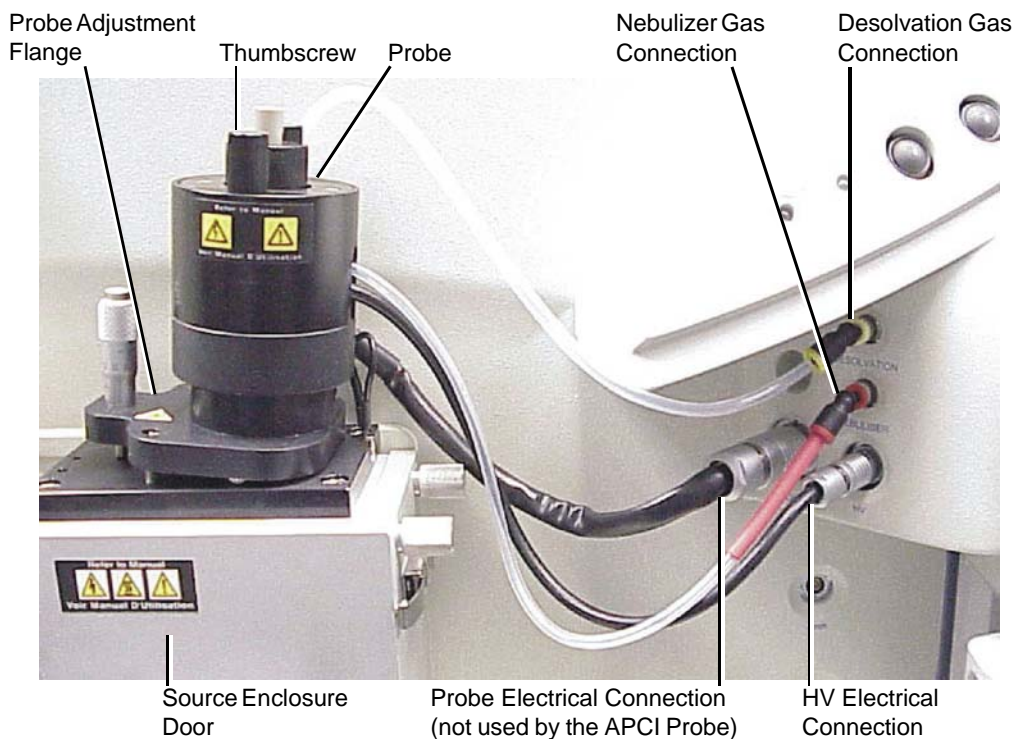


Figure 7-2 Probe Connections (ESI Probe Shown as an Example)

7. Disconnect the PTFE tubing at the Nebulizer gas connection on the front panel.
8. Undo the two thumbscrews securing the probe to the probe adjustment flange.



Warning: The probe may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



Caution: To avoid damaging the probe seals when removing an APCI probe, confirm that the probe temperature is less than 150 °C, as displayed on the Tune window APCI Probe Temp (°C) readback, before removing the probe.

9. Carefully remove the probe from the probe adjustment flange.

7.7.4 Removing the Sample Cone



Warning: The source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



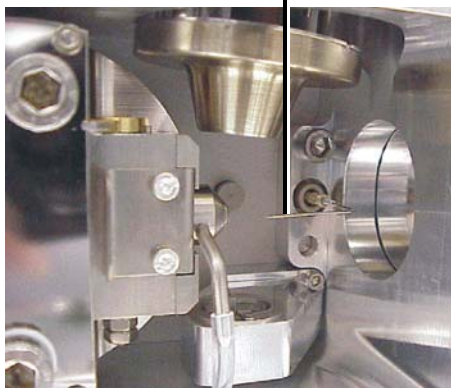
Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.



Warning: The source may be hot. To avoid burns, take great care while performing this procedure.

1. Unfasten the source enclosure door's securing clips and open the door (see [Figure 7-2](#)).
2. If using ESCi mode or an APCI probe, carefully remove the corona discharge pin.

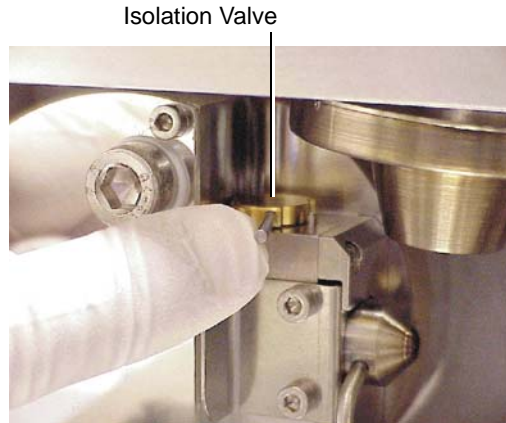
Corona Discharge Pin





Caution: Failure to close the isolation valve before removing the sample cone may damage the instrument.

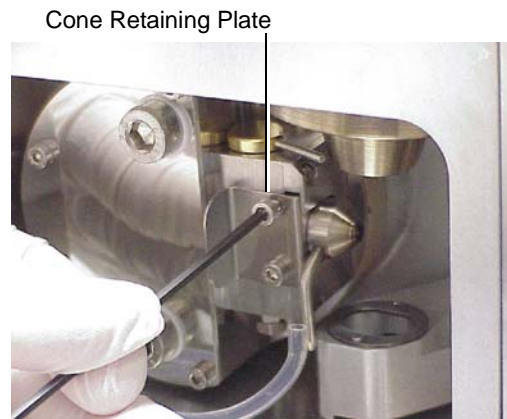
3. Close the isolation valve by moving its lever fully to the right.



4. Disconnect the PTFE tube from the cone gas cone.



5. Use a hex (Allen) key to remove the two cone retaining plate securing screws.



6. Remove the cone retaining plate.



7. Carefully remove the sample cone/cone gas cone assembly from the isolation valve body.

Sample Cone/Cone Gas Cone Assembly



8. Use a jeweller's screwdriver to carefully remove the O-ring from the sample cone/cone gas cone assembly.



Caution: The sample cone is fragile. To avoid damaging it, never place the sample cone on its tip; always place it on its flanged base.

9. Separate the sample cone from the cone gas cone.

Figure 7-3 shows the sample cone, cone gas cone, and O-ring.

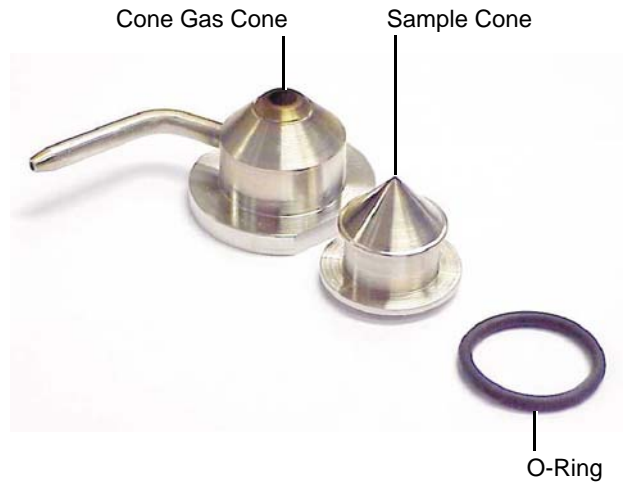


Figure 7-3 Sample Cone/Cone Gas Cone Assembly Components

10. Clean the sample cone and cone gas cone (see [Section 7.7.10](#)).

7.7.5 Removing the Gas Exhaust Port



Warning: The source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



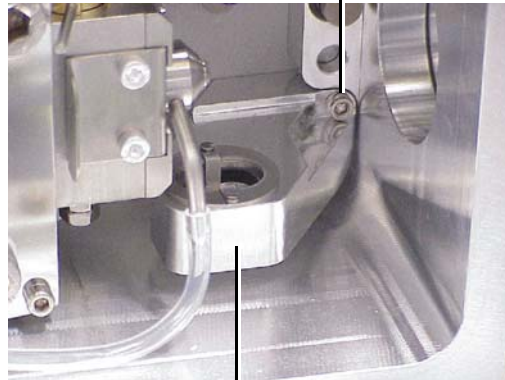
Warning: To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.



Warning: The source may be hot. To avoid burns, take great care while performing this procedure.

1. Use a hex (Allen) key to remove the gas exhaust port securing screw.

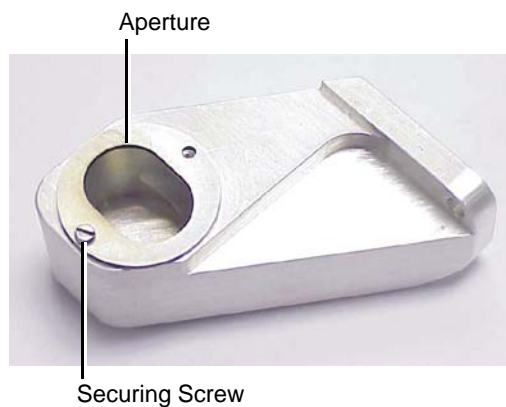
Gas Exhaust Port Securing Screw



Gas Exhaust Port

2. Remove the gas exhaust port from the source enclosure.

3. Remove the two screws securing the aperture to the gas exhaust port.



4. Remove the aperture from the gas exhaust port.
5. Clean the gas exhaust port (see [Section 7.7.11](#)).

7.7.6 Removing the Ion Source Enclosure and Ion Block

1. Shut down the instrument (see [Section A.2](#)).



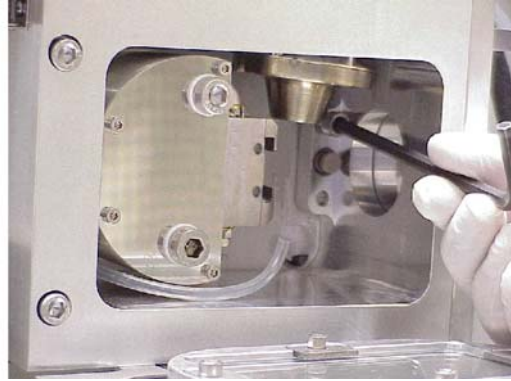
Warning: The source may be hot; to avoid burns, allow it to cool down for at least 30 minutes before proceeding.

2. Disconnect the Probe electrical connection at the instrument front panel (see [Figure 7-2](#)).
3. Disconnect the PTFE tubing at the Desolvation gas connection on the front panel (see [Figure 7-2](#)).

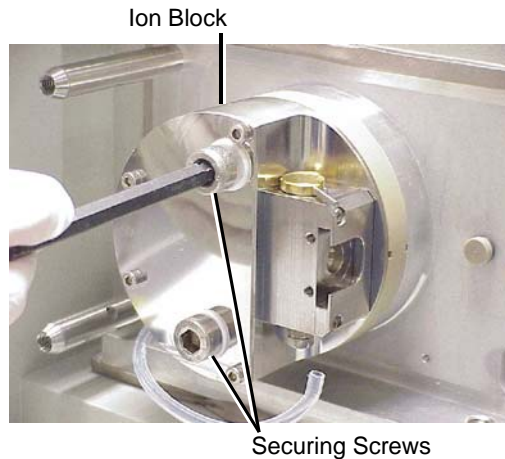


Warning: The source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

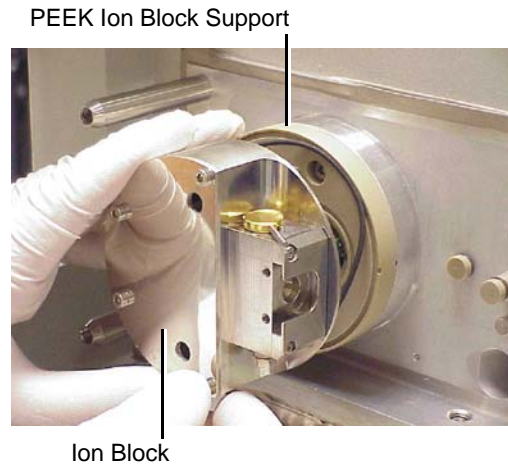
4. Use a hex (Allen) key to loosen the three captive source enclosure securing screws.



5. Remove the source enclosure from the instrument.
6. Use the 6-mm hex (Allen) key to remove the two ion block securing screws.



7. Remove the ion block from the PEEK ion block support.

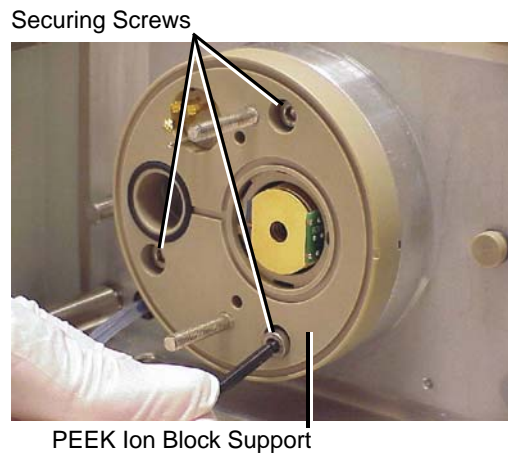


7.7.7 Removing the Source T-Wave Assembly from the Instrument

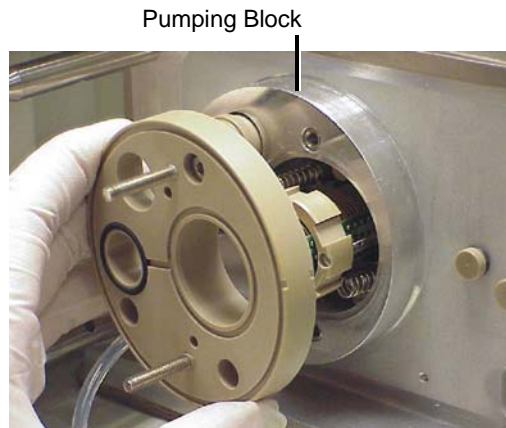


Warning: The source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

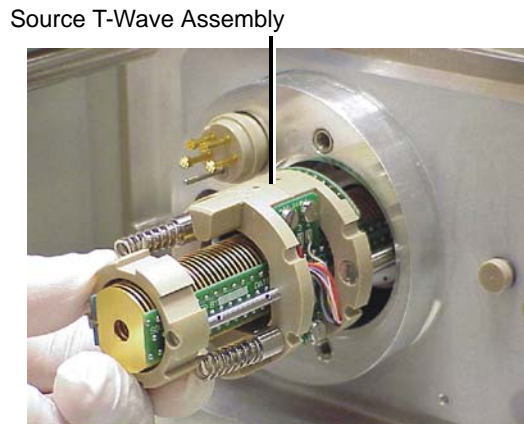
1. Use a hex (Allen) key to remove the three PEEK ion block support securing screws.



2. Remove the PEEK ion block support from the pumping block.



3. Carefully remove the source T-Wave assembly (see [Figure 7-4](#)) from the pumping block.



Note: Later versions of the source T-Wave assembly are fitted with PEEK covers over the electrical wiring.

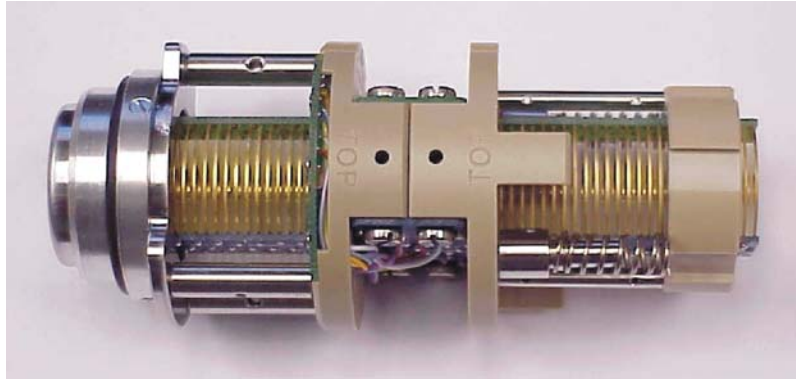


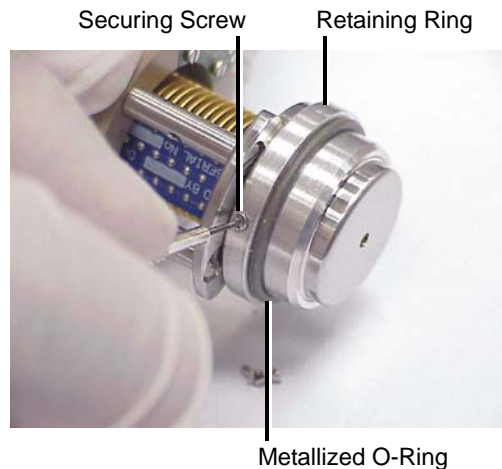
Figure 7-4 Source T-Wave Assembly

7.7.8 Disassembling the Source T-Wave Assembly

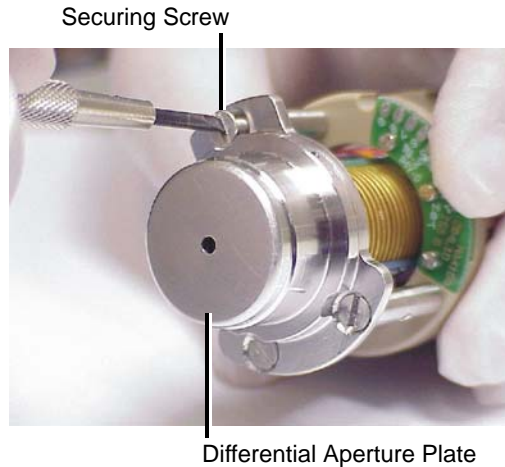


Warning: The source T-Wave components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. On later versions of the source T-Wave assembly only, remove the PEEK covers after removing the two screws securing the covers to the assembly.
2. Use the jeweller's screwdriver to remove the three screws securing the retaining ring to the source T-Wave assembly.

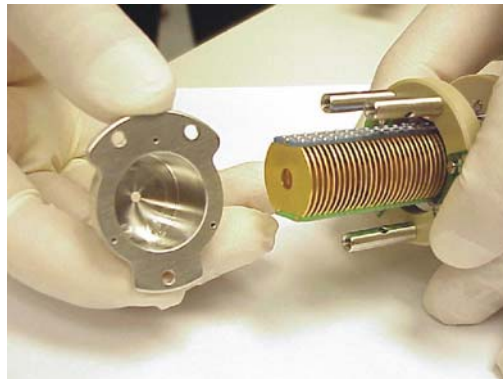


3. Remove the metallized O-ring and retaining ring from the assembly.
4. Remove the three screws securing the differential aperture plate to the assembly.



5. Remove the differential aperture plate from the assembly. The source T-Wave assembly components are shown in [Figure 7-5](#).

Note: No further disassembly of the source T-Wave assembly is required.



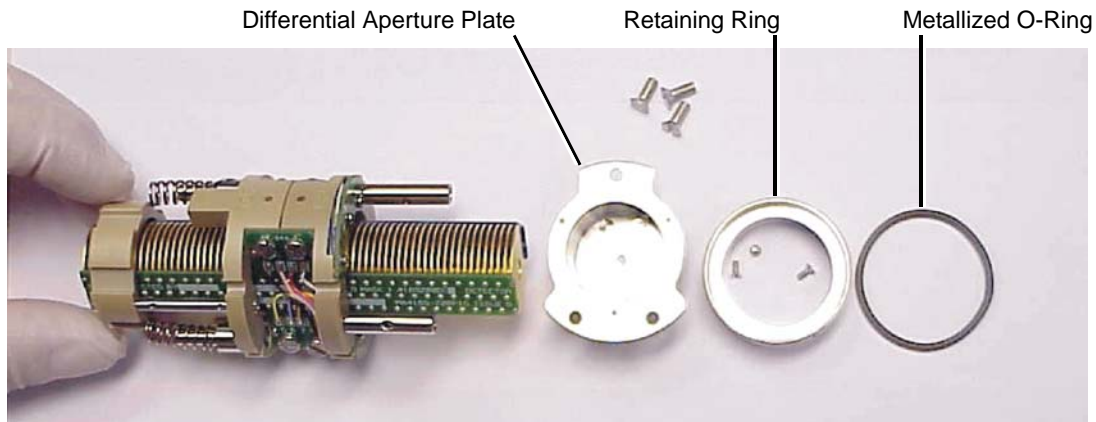


Figure 7-5 Source T-Wave Assembly Components

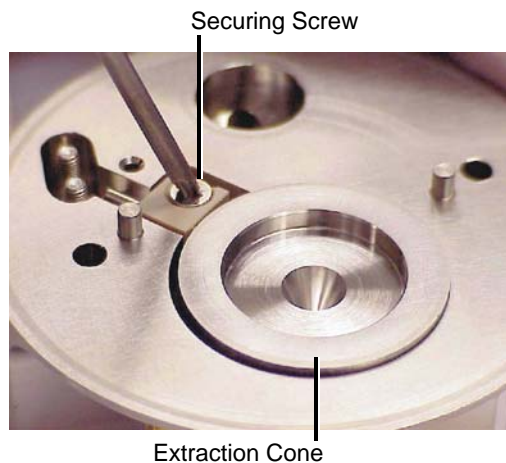
6. Clean the source T-Wave assembly and differential aperture plate (see [Section 7.7.12](#)).

7.7.9 Disassembling the Source Ion Block



Warning: The source ion block components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. Remove the extraction cone securing screw.





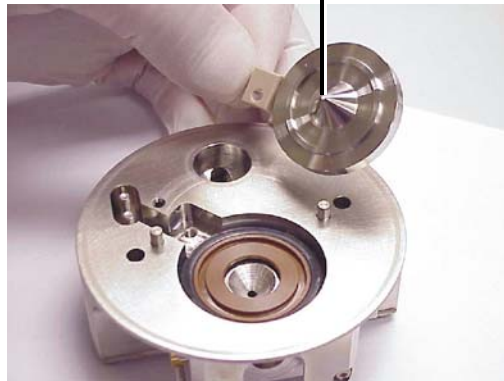
Caution: Take great care not to damage the extraction cone aperture when removing the extraction cone from the ion block.



Caution: The extraction cone is fragile. To avoid damaging it, never place the extraction cone on its tip, always place it on its flanged base.

2. Remove the extraction cone from the ion block.

Extraction Cone Aperture

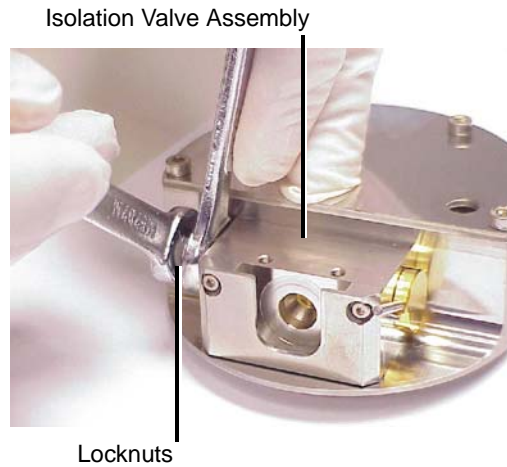


3. Clean the extraction cone (see [Section 7.7.13](#)).
4. Remove the extraction cone seal from the ion block.

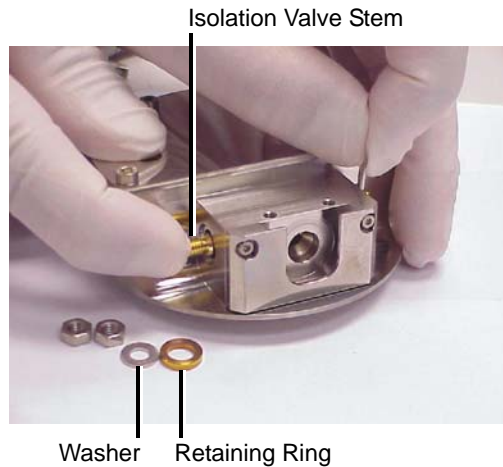
Extraction Cone Seal



5. Remove the two locknuts from the bottom of the isolation valve body.



6. Remove the washer and retaining ring from the isolation valve stem.



7. Remove the isolation valve stem from the isolation valve body (gently push the stem while repeatedly opening and closing the isolation valve).

The isolation valve stem components are shown in [Figure 7-6](#).



Figure 7-6 Isolation Valve Stem Components

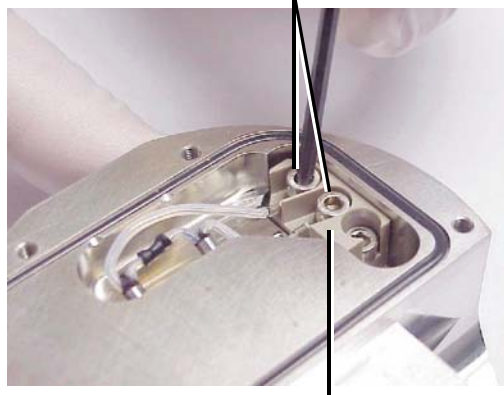
8. Clean the isolation valve stem as described in [Section 7.7.14](#).
9. Use a hex (Allen) key to remove the four ion block cover plate securing screws.



10. Remove the ion block cover plate.

11. Use a hex (Allen) key to remove the two screws securing the heater cartridge wires to the PEEK terminal block.

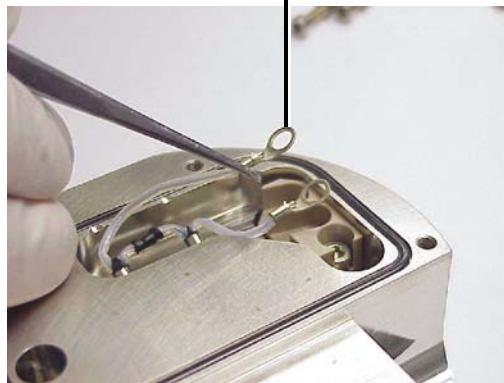
Cartridge Heater Wire
Securing Screws



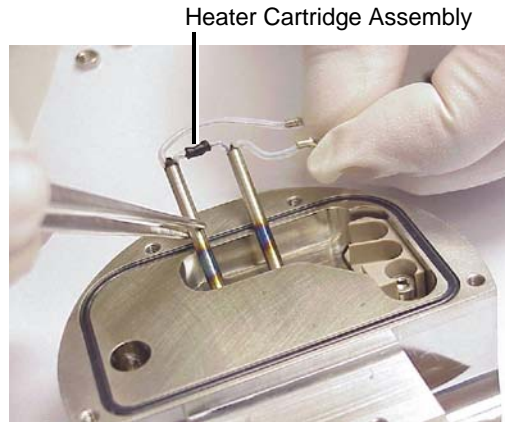
PEEK Terminal Block

12. Use the needle-nose pliers to carefully swing the ring terminal tags out of the terminal block.

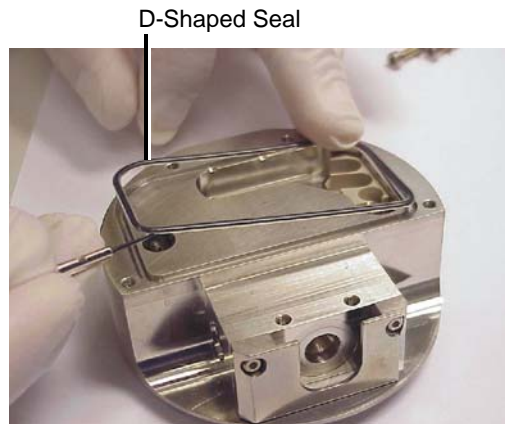
Ring Terminal Tag



13. Use the needle-nose pliers to gently slide the heater cartridge assembly out of the ion block.

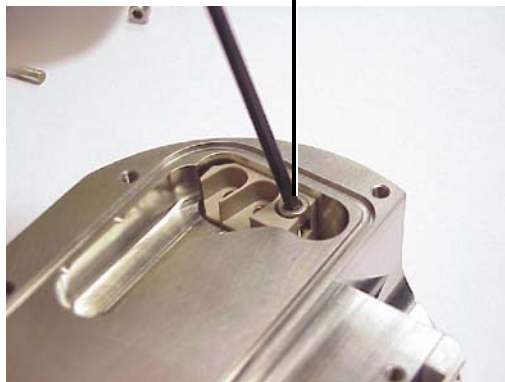


14. Use the jeweller's screwdriver to carefully remove the D-shaped seal from the ion block.



15. Use a hex (Allen) key to remove the PEEK terminal block securing screw.

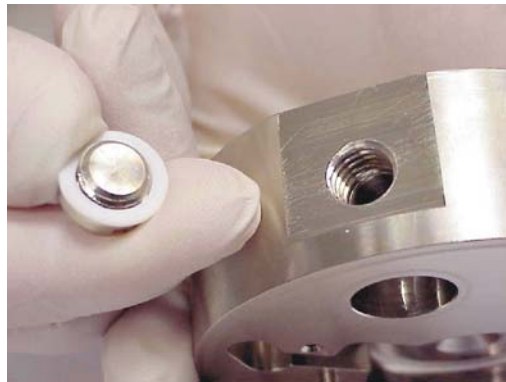
PEEK Terminal Block Securing Screw



16. Use the needle-nose pliers to remove the PEEK terminal block from the ion block.



17. Use the screwdriver to remove the ion block blanking plug.



18. Clean the ion block as described in [Section 7.7.13](#).

19. Reassemble the ion block as described in [Section 7.7.15](#).

7.7.10 Cleaning the Sample Cone and Cone Gas Cone



Warning: The sample cone and cone gas cone may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



Caution: The sample cone is fragile. To avoid damaging it, never place the sample cone on its tip; always place it on its flanged base.

1. Use a glass-fiber pen to remove gross contamination from the cone gas cone surface by gentle abrasion.



Warning: Use extreme care when working with formic acid. Use a fume hood and appropriate protective equipment.

2. If the sample cone contains debris, place a drop of formic acid on its orifice.
3. Immerse the sample cone and cone gas cone separately in glass vessels containing 1:1 methanol/water.

Note: If the parts are obviously contaminated, use 45:45:10 methanol/water/formic acid.

4. Place the vessels in the ultrasonic bath for 30 minutes.
5. If formic acid was used in the cleaning solution:
 - a. Rinse the parts by immersing them separately in glass vessels containing water, and placing the vessels in the ultrasonic bath for 20 minutes.
 - b. Displace the water by immersing the parts in separate glass vessels containing methanol, and placing the vessels in the ultrasonic bath for 10 minutes.
6. Carefully remove the parts from the vessels, and blow-dry them with inert, oil-free gas. Alternatively, place the components on lint-free towels, and allow them to air dry. Wipe off any water spots with a lint-free cloth.

7.7.11 Cleaning the Gas Exhaust Port



Warning: The gas exhaust port may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. Remove gross contamination from the gas exhaust port and aperture by rinsing them in water.

Note: Gentle abrasion, using a glass-fiber pen, may be required to remove solid deposits.

2. Immerse the gas exhaust port and aperture in a glass vessel containing 1:1 methanol/water.
3. Place the vessel in the ultrasonic bath for 30 minutes.
4. Carefully remove the components from the vessel, and blow-dry them using inert, oil-free gas. Alternatively, place the components on lint-free towels, and allow them to air dry. Wipe off any water spots with a lint-free cloth.

7.7.12 Cleaning the Source T-Wave Components



Warning: The source T-Wave components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

Cleaning the Source T-Wave Differential Aperture Plate

1. Use a glass-fiber pen to gently remove ion burn marks, paying particular attention to the inner surfaces of the differential aperture plate.
2. Immerse the differential aperture plate in a glass vessel containing 1:1 methanol/water.
3. Place the vessel in the ultrasonic bath for 30 minutes.
4. Carefully remove the differential aperture plate from the vessel, and blow-dry it using inert, oil-free gas. Alternatively, place the differential aperture plate on lint-free towels, and allow it to air dry. Wipe off any water spots with a lint-free cloth.

Cleaning the Source T-Wave Assembly



Caution: Use of acetone, chlorinated solvents, or acid as solvents when cleaning the source T-Wave assembly will damage the assembly. Use only methanol or water.



Caution: Take great care not to damage the T-Wave assembly's plates when using the wire brush for cleaning.

1. Gently insert the wire brush (provided) into the T-Wave assembly's aperture, and use a rotary motion to clean the plates along the length of the device.

2. Use a glass-fiber pen to remove any burn marks on the assembly's entrance and exit plates.
3. Flush-out the assembly, using methanol from a wash-bottle.
4. Immerse the assembly in a glass vessel containing 1:1 methanol/water.
5. Place the vessel in the ultrasonic bath for 30 minutes.



Caution: Do not dry the source T-Wave assembly by any method other than blow drying, otherwise reintroduced contamination may lead to difficulty in pumping down the instrument.

6. Carefully remove the assembly from the vessel, and blow-dry it using inert, oil-free gas.
7. Visually inspect the assembly to confirm that no fibers from the wire brush are lodged in it. If fibers are present, repeat the procedure from [step 3](#) onwards.

7.7.13 Cleaning the Ion Block and Extraction Cone



Warning: The ion block and extraction cone may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



Warning: Use extreme care when working with formic acid. Use a fume hood and appropriate protective equipment.



Caution: The extraction cone is fragile. To avoid damaging it, never place the extraction cone on its tip.

1. Immerse the ion block and extraction cone separately in glass vessels containing 1:1 methanol/water.

Note: If the components are obviously contaminated, use 45:45:10 methanol/water/formic acid.

2. Place the vessels in the ultrasonic bath for 30 minutes.
3. If formic acid was used in the cleaning solution:
 - a. Rinse the ion block and extraction cone by immersing them separately in glass vessels containing water, and placing the vessels in the ultrasonic bath for 20 minutes.

- b. Displace the water by immersing the ion block and extraction cone separately in glass vessels containing methanol, and placing the vessels in the ultrasonic bath for 10 minutes.
4. Carefully remove the ion block and extraction cone from the vessels, and blow-dry them using inert, oil-free gas. Alternatively, place the components on lint-free towels, and allow them to air dry. Wipe off any water spots with a lint-free cloth.

7.7.14 Cleaning the Isolation Valve Stem



Warning: The isolation valve components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. Use a glass-fiber pen to remove carbon deposits by gentle abrasion.
2. Rinse the valve stem by placing it in a vessel containing 1:1 methanol/water, and placing the vessel in an ultrasonic bath for 20 minutes.

7.7.15 Reassembling the Source Ion Block

Note: Use clean nitrile gloves when reassembling the source components to avoid re-contaminating them.

1. Examine the condition of all O-rings and seals; replace any that are damaged.
2. Fit the ion block blanking plug to the ion block and tighten.
3. Fit the PEEK terminal block to the ion block.
4. Fit and tighten the PEEK terminal block securing screw.
5. Fit the D-shaped seal to the ion block, ensuring that it is correctly seated.
6. Use the needle-nose pliers to slide the new heater cartridges into the ion block.
7. Position the two heater cartridge ring tags onto the PEEK block terminals.
8. Use a hex (Allen) key to fit and tighten the two screws securing the heater cartridge wires to the PEEK terminal block.
9. Fit the ion block cover plate.
10. Fit and tighten the four ion block cover plate securing screws.
11. Fit the isolation valve stem to the isolation valve body.
12. Fit the washer and retaining ring to the bottom of the isolation valve stem.
13. Fit and tighten the two lock nuts to the bottom of the isolation valve stem.
14. Fit the extraction cone seal to the ion block.

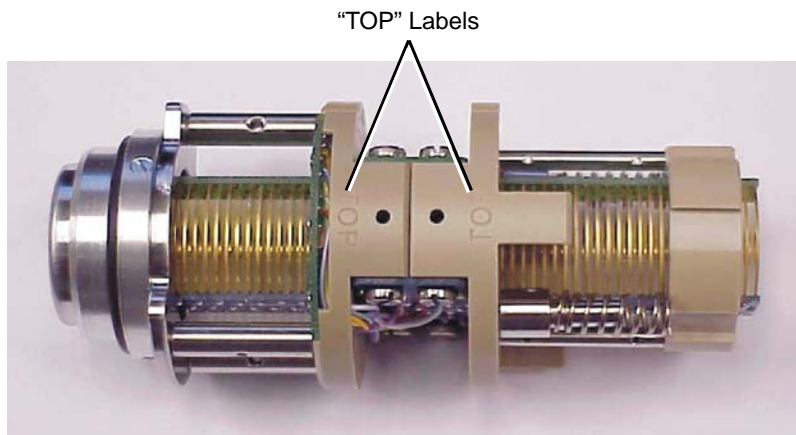
15. Fit the extraction cone to the ion block.
16. Fit and tighten the extraction cone securing screw.

7.7.16 Reassembling the Source T-Wave Assembly

1. Fit the differential aperture plate to the source T-Wave assembly.
2. Fit and tighten the three screws that secure the differential aperture plate to the source T-Wave assembly.
3. Fit the retaining ring to the assembly.
4. Check the condition of the metallized O-ring; if it is damaged, replace it.
5. Fit the metallized O-ring to the assembly.
6. Fit and tighten the three screws that secure the retaining ring to the source T-Wave assembly.
7. On later versions of the source T-Wave assembly, fit the PEEK covers to the assembly, then fit and tighten the two screws securing the covers to the assembly.

7.7.17 Fitting the Source T-Wave Assembly to the Instrument

1. Ensuring that the “TOP” labels (stamped on the assembly’s PEEK supports) are uppermost, carefully slide the source T-Wave assembly into the pumping block.



2. Fit the PEEK ion block support to the pumping block.
3. Fit and tighten the two PEEK ion block support securing screws.

7.7.18 Fitting the Ion Block and Ion Source Enclosure

1. Check the condition of all O-rings; replace any that are damaged.
2. Ensure that all the O-rings are in position on the ion block.
3. Fit the ion block to the PEEK ion block support.
4. Fit and tighten the two ion block securing screws.
5. Fit the source enclosure to the pumping block.
6. Fit and tighten the three source enclosure securing bolts.
7. Connect the Probe electrical connection at the instrument's front panel (see [Figure 7-2](#)).
8. Connect the PTFE tubing to the Desolvation gas connection at the instrument's front panel (see [Figure 7-2](#)).
9. If required, the instrument can be evacuated as described in [Section A.1](#).

7.7.19 Fitting the Gas Exhaust Port

1. Fit the aperture to the gas exhaust port.
2. Fit and tighten the two aperture retaining screws.
3. Fit the gas exhaust port into the source enclosure.
4. Fit and tighten the gas exhaust port securing screw.

7.7.20 Fitting the Sample Cone



Caution: *The sample cone is very fragile. To avoid damage, never place the sample cone on its tip; always place it on its flanged base.*

1. Fit the sample cone into the cone gas cone.
2. Check the condition of the sample cone/cone gas cone assembly O-ring; replace it, if it is damaged.
3. Fit the O-ring to the sample cone/cone gas cone assembly.
4. Fit the sample cone/cone gas cone assembly to the side of the isolation valve body.
5. Fit the cone retaining plate.
6. Fit and tighten the two cone retaining plate securing screws.
7. Connect the PTFE tube to the cone gas cone.

8. Open the isolation valve by moving its lever fully to the left.
9. If an APCI probe is to be used, fit the corona discharge pin.
10. Close the source enclosure door and fasten the clips.

7.8 Cleaning or Replacing the ESI Probe Tip

Clean the ESI probe tip if a blockage occurs in the internal metal sheathing through which the stainless steel capillary passes.

Replace the ESI probe tip if the threads are damaged.

Replace the O-ring if gas leaks from the O-ring.

7.8.1 Required Materials

- Nitrile gloves.
- ¼-inch (6-mm) wrench.
- Appropriately-sized glass vessels, in which to completely immerse components when cleaning. Use only glassware not previously cleaned with surfactants.
- HPLC-grade methanol.
- HPLC-grade water.
- Ultrasonic bath.
- Source of oil-free, inert gas (nitrogen or helium) for drying (air-drying optional).

7.8.2 Procedure



Warning: The probe and source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. Remove the probe from the source (see [Section 7.7.3](#)).
2. Use the ¼-inch (6-mm) wrench to unscrew and remove the probe tip.
3. If the probe tip is damaged, replace it; alternatively, clean the probe tip as follows:
 - a. Immerse the probe tip in a glass vessel containing 1:1 methanol/water.
 - b. Place the vessel in the ultrasonic bath for 20 minutes.

- c. Carefully remove the probe tip from the vessel, and blow-dry it using inert, oil-free gas.
4. If necessary, remove the O-ring and fit a new one.
5. Fit and tighten the probe tip to the probe.
6. Adjust the probe tip so that the fully extended capillary (when the probe nebulizer adjuster knob is fully screwed down) protrudes by approximately 1 to 1.5 mm.
7. Fit the probe to the source.

7.9 Cleaning or Replacing the Corona Discharge Pin

Clean the corona discharge pin if it appears to be corroded or black, or when the signal intensity weakens.

7.9.1 Required Materials

- Nitrile gloves
- Needle-nose pliers
- Lapping film
- HPLC-grade methanol
- Lint-free tissue

7.9.2 Procedure



Warning: *The probe and source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.*



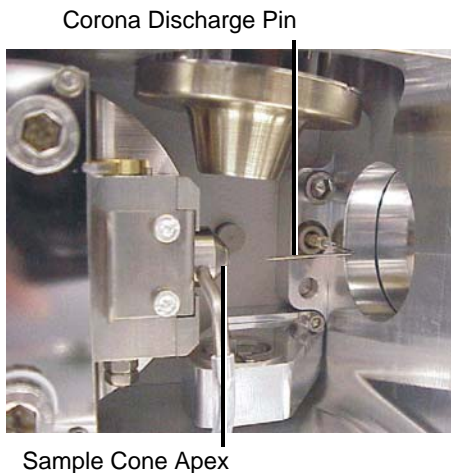
Warning: *To avoid electric shock, ensure that the instrument is in Standby before commencing this procedure.*



Warning: *The probe and source may be hot. To avoid burns, take great care while performing this procedure.*

1. Remove the probe from the source, as shown in [Section 7.7.3](#).
2. Unfasten the source enclosure door's securing clips, and open the door (see [Figure 7-2](#)).


3. Use the needle-nose pliers to remove the corona discharge pin from the source, pulling it straight out.
4. Clean and sharpen the tip of the pin with the lapping film, then wipe it clean with a methanol-saturated tissue. Replace the pin if it is deformed or otherwise damaged.
5. Reinstall the pin with the tip pointing toward the sample cone apex.



6. Close the source enclosure door, and fasten the clips.
7. Fit the probe.
8. Reconnect the front panel gas and electrical connections.

7.10 Cleaning the APCI Probe Tip

Clean the APCI probe tip when a buffer build-up is detected on the probe tip, or when the signal intensity weakens.

1. Stop the liquid flow.
2. In the Tune window, click  to start the nitrogen flow.
3. Set Desolvation (L/hr) to approximately 650.
4. Set APCI Probe Temp (°C) to 650.

5. Click Press for Operate, and wait 10 minutes with the APCI probe heater temperature at 650 °C. This will remove any chemical contamination from the probe tip.

7.11 Replacing the Ion Block Cartridge Heater

Replace the cartridge heater if it fails to heat.

7.11.1 Required Materials

- Nitrile gloves
- Flat-blade screwdriver
- Needle-nose pliers
- Hex (Allen) keys

7.11.2 Procedure



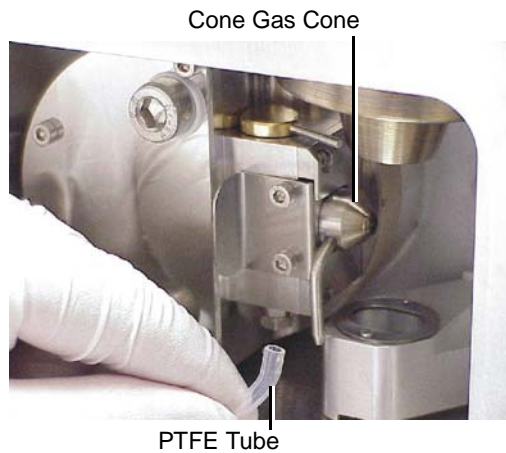
Warning: The probe and source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



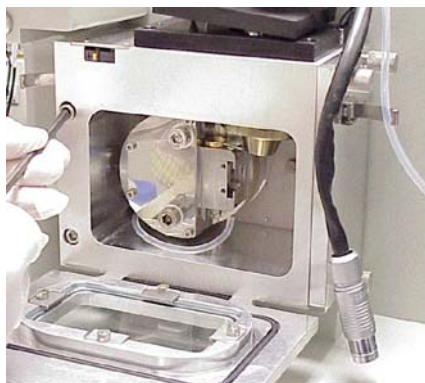
Warning: The probe and source may be hot. To avoid burns, take great care while performing this procedure.

1. Follow the procedure for venting the instrument (see [Section A.2.3](#)).
2. Remove the probe from the source (see [Section 7.7.3](#)).
3. If using an APCI probe, carefully remove the corona discharge pin.

4. Disconnect the PTFE tube from the cone gas cone.

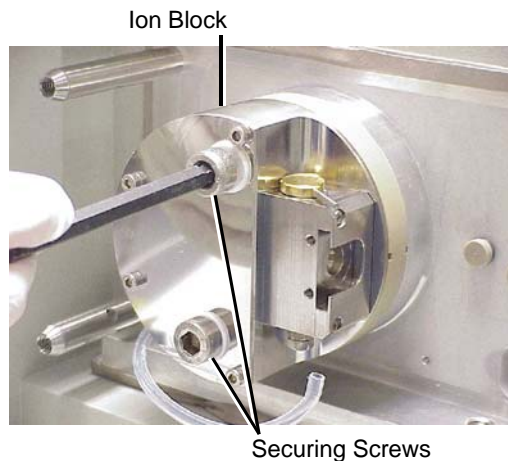


5. Disconnect the Probe electrical connection from the instrument's front panel (see [Figure 7-2](#)).
6. Disconnect the PTFE tubing at the Desolvation gas connection from the instrument's front panel (see [Figure 7-2](#)).
7. Use a hex (Allen) key to remove the three source enclosure securing bolts.



8. Remove the source enclosure from the instrument.

9. Use the 6-mm hex (Allen) key to remove the two ion block securing screws.



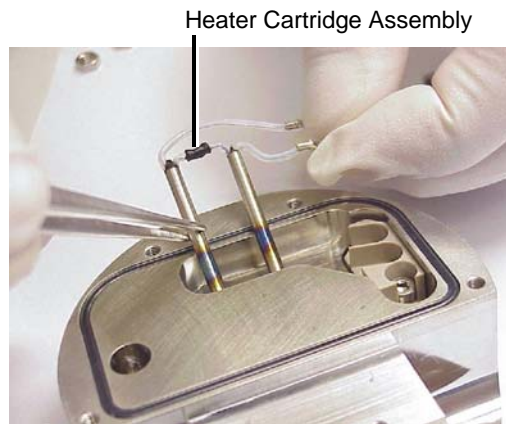
10. Remove the ion block from the PEEK ion block support.
11. Use a hex (Allen) key to remove the four ion block cover plate securing screws.
12. Remove the ion block cover plate.
13. Use a hex (Allen) key to remove the two screws securing the heater cartridge wires to the PEEK terminal block.



14. Use the needle-nose pliers to carefully swing the ring terminal tags out of the terminal block.



15. Use the needle-nose pliers to gently slide the heater cartridge assembly out of the ion block.



16. Use the needle-nose pliers to slide the new heater cartridges into the ion block.
17. Position the two heater cartridge ring tags onto the PEEK block terminals.
18. Use a hex (Allen) key to fit and tighten the two screws securing the heater cartridge wires to the PEEK terminal block.
19. Fit the ion block cover plate.

20. Fit and tighten the four ion block cover plate securing screws.
21. Check the condition of all O-rings; if any are damaged, replace them.
22. Ensure that all the O-rings are in position on the ion block.
23. Fit the ion block to the PEEK ion block support.
24. Fit and tighten the two ion block securing screws.
25. Fit the source enclosure to the pumping block.
26. Fit and tighten the three source enclosure securing bolts.
27. Connect the PTFE tube to the cone gas cone.
28. If using an APCI probe, carefully fit the corona discharge pin.
29. Connect the Probe electrical connection on the instrument's front panel (see [Figure 7-2](#)).
30. Connect the PTFE tubing to the Desolvation gas connection on the instrument's front panel.
31. Fit the probe on the source.

7.12 Replacing the ESI Probe Sample Capillary

The stainless steel sample capillary in the ESI probe must be replaced if it becomes blocked and cannot be cleared, or if it becomes contaminated or damaged.

7.12.1 Required Materials

- 7-mm wrench
- ¼-inch (6-mm) wrench
- 5/16-inch wrench
- Needle-nose pliers
- Hex (Allen) key

7.12.2 Removing the Existing Capillary



Warning: The probe and source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

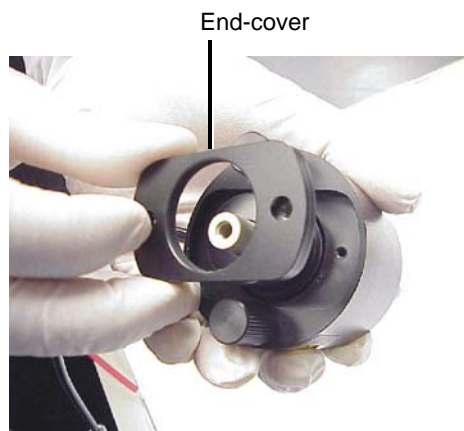


Warning: The probe and source may be hot. To avoid burns, take great care while performing this procedure.

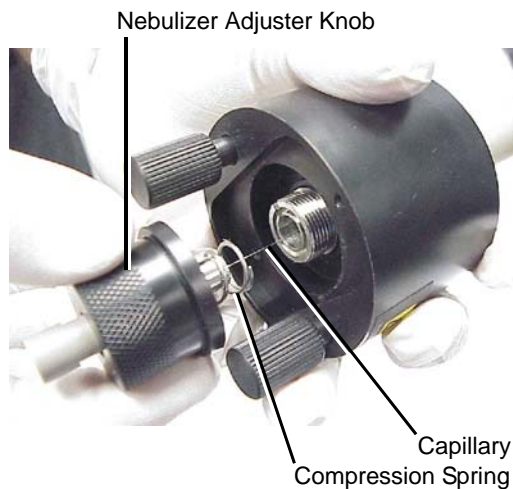
1. Follow the procedure in [Section 7.7.3](#) to remove the probe from the source.
2. Use the hex (Allen) key to remove the two probe end-cover retaining screws.



3. Remove the end-cover.



4. Unscrew and remove the nebulizer adjuster knob to reveal a PEEK union/UNF coupling assembly, compression spring, and the capillary.

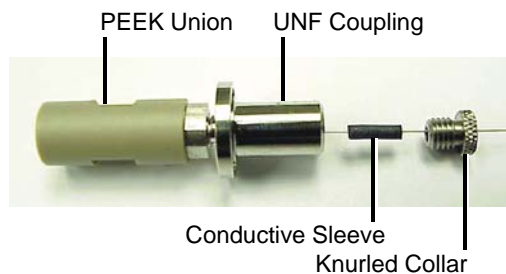


5. Remove the nebulizer adjuster knob, PEEK union/UNF coupling assembly, compression spring, and capillary from the probe.
6. Remove the PEEK union/UNF coupling assembly, compression spring, and capillary from the nebulizer adjuster knob.

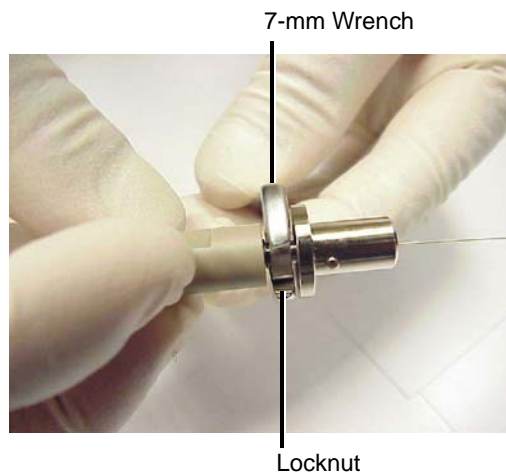


7. Remove the compression spring from the PEEK union/UNF coupling assembly and capillary.

8. Unscrew and remove the knurled collar from the UNF coupling to reveal a conductive sleeve on the capillary.



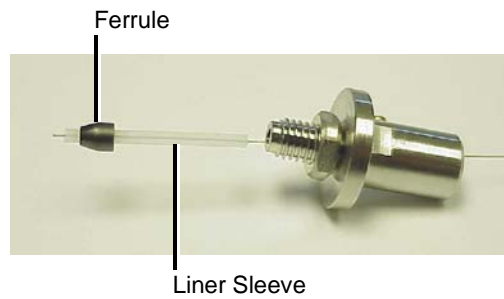
9. Remove the knurled collar and conductive sleeve from the capillary.
10. A locknut is used to secure the PEEK union and UNF coupling. Use the 7-mm wrench to loosen the locknut.



11. Unscrew the PEEK union from the UNF coupling (this connection is finger-tight only).



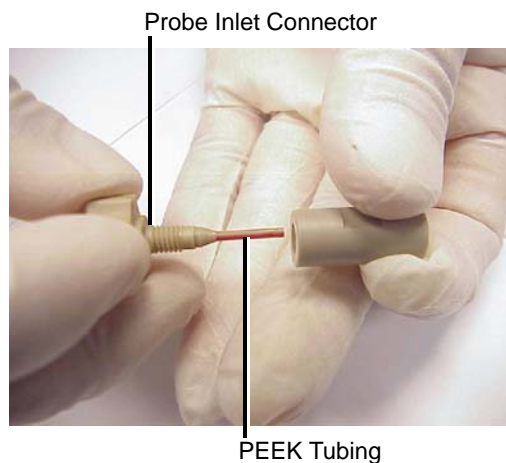
This reveals a ferrule and liner sleeve:



12. Remove the ferrule and liner sleeve from the capillary.
13. Remove the capillary from the UNF coupling.

7.12.3 Installing the New Capillary

1. Insert a square-cut length of red PEEK tubing in the probe inlet connector, and screw the connector, finger-tight, into the PEEK union. This ensures a minimum dead volume when fitting the capillary.



2. Fit the UNF coupling to the new capillary.
3. Use the needle-nose pliers to slide a new liner sleeve and ferrule onto the capillary.
4. Insert the capillary in the PEEK union, and ensure that it is fully seated.
5. Screw the UNF coupling into the PEEK union, finger-tight only.
6. Pull on the capillary gently, testing to ensure that it stays in place.
7. Use the 7-mm wrench to tighten the locknut against the PEEK union until the union can no longer be twisted.
8. Slide a new conductive sleeve and the knurled collar over the capillary.
9. Tighten the knurled collar to the UNF coupling.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when performing the leak test.

10. Check for leaks in the assembly by attaching the free end of the PEEK tubing to an LC pump and pumping 50:50 acetonitrile/water through it, at 1 mL/min.



If leakage occurs, disassemble and remake the connection, and repeat the leak test.

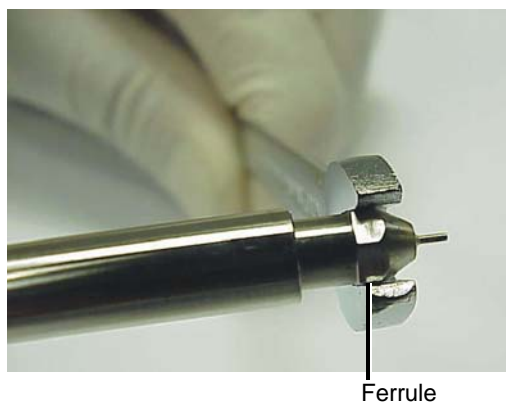


Leaking Liquid

11. When performing the leak test, check the backpressure on the LC pump, which will be high if the capillary is blocked. If this is the case, replace the capillary.
12. When the leak test has been performed successfully, disconnect the PEEK tubing from the LC pump.
13. Remove the probe inlet connector and PEEK tubing from the PEEK union.
14. Fit the PEEK union/UNF coupling assembly to the nebulizer adjuster knob.
15. Fit the compression spring to the capillary and PEEK union/UNF coupling assembly.



16. Use the 1/4-inch (6-mm) wrench to remove the probe tip from the probe.



17. Carefully thread the capillary through the probe assembly.

18. Depress the PEEK union so that the locating pin on the UNF coupling is fully engaged in the locating slot at the head of the probe assembly. When the union is fully depressed, tighten the nebulizer adjuster knob. Do not tighten the knob fully.

Locating Pin
(compression spring removed for clarity)



Locating Slot

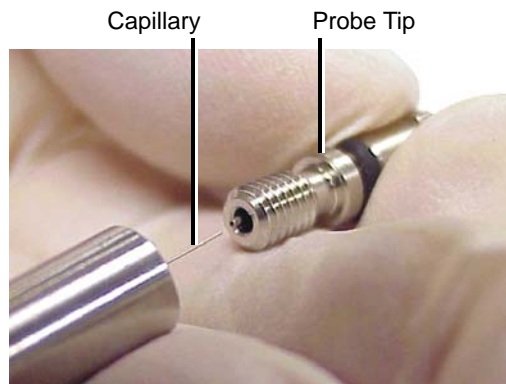



Locating Pin Locating Slot



19. Fit the probe end-cover to the probe assembly.
20. Fit and tighten the two end-cover securing screws.

21. Fit the probe tip over the capillary, and screw the tip onto the probe assembly.



22. If necessary, adjust the probe tip so that the fully extended capillary (when the nebulizer adjuster knob is fully screwed down) protrudes by approximately 1 to 1.5 mm.
23. Use the nebulizer adjuster knob to adjust the capillary so that the capillary protrudes by approximately 0.5 mm from the end of the probe.
24. Attach the nebulizer gas connection, and start the nitrogen flow by clicking  in the MassLynx Tune window.
25. Check the probe tip for nitrogen leaks. If a leak is found, replace the probe tip assembly and its O-ring (see [Section 7.8](#)).
26. Fit the probe to the instrument.

7.13 Replacing the APCI Probe Sample Capillary

Replace the stainless steel sample capillary in the APCI probe if it becomes blocked and cannot be cleared, or if it becomes contaminated or damaged.

7.13.1 Required Materials

- 7-mm wrench
- ¼-inch (6-mm) wrench
- 5/16-inch wrench
- Needle-nose pliers

- Jeweller's screwdriver
- Hex (Allen) key

7.13.2 Removing the Existing Capillary



Warning: The probe and source components may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.



Warning: The probe and source may be hot. To avoid burns, take great care while performing this procedure.

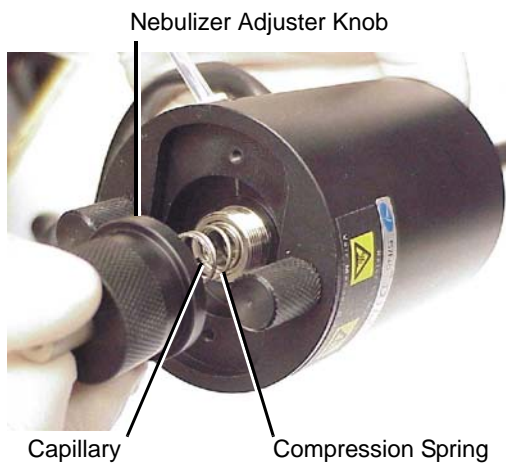
1. Follow the procedure detailed in [Section 7.7.3](#), to remove the probe from the source.
2. Use the hex (Allen) key to remove the two probe end-cover retaining screws.



3. Remove the end-cover.



4. Unscrew and remove the nebulizer adjuster knob to reveal a PEEK union/UNF coupling assembly, compression spring, and the capillary.



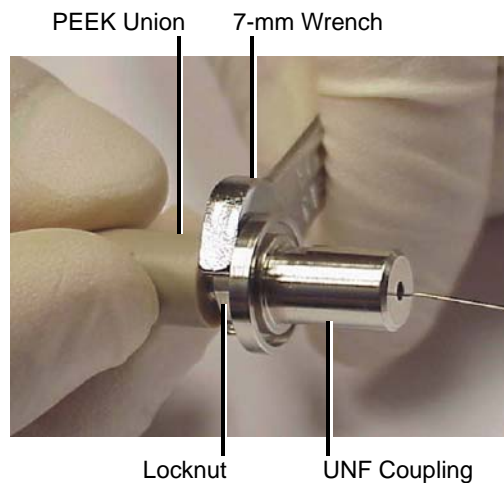
5. Remove the nebulizer adjuster knob, PEEK union/UNF coupling assembly, compression spring, and capillary from the probe.

6. Remove the PEEK union/UNF coupling assembly, compression spring, and capillary from the nebulizer adjuster knob.

PEEK Union/UNF Coupling Assembly



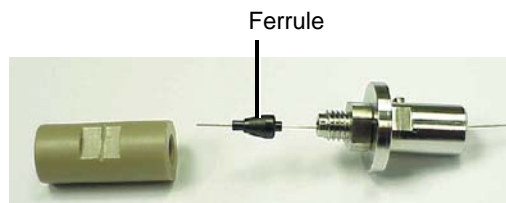
7. Remove the compression spring from the PEEK union/UNF coupling assembly and capillary.
8. A locknut is used to secure the PEEK union and UNF coupling. Use the 7-mm wrench to loosen the locknut.



9. Unscrew the PEEK union from the UNF coupling (this connection is finger-tight only).



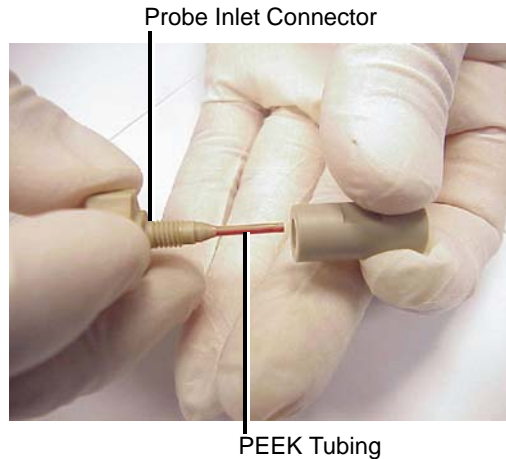
This reveals a ferrule.



10. Remove the ferrule from the capillary.
11. Remove the capillary from the UNF coupling.

7.13.3 Installing the New Capillary

1. Insert a square-cut length of red PEEK tubing in the probe inlet connector, and screw the connector, finger-tight, into the PEEK union. This ensures a minimum dead volume when fitting the capillary.



2. Fit the UNF coupling to the new capillary.
3. Use the needle-nose pliers to slide a new ferrule onto the capillary.
4. Insert the capillary in the PEEK union, and ensure that it is fully seated.
5. Screw the UNF coupling into the PEEK union, finger-tight only.
6. Pull on the capillary gently, testing to ensure that it stays in place.
7. Use the 7-mm wrench to tighten the locknut against the PEEK union.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when performing the leak test.

8. Check for leaks in the assembly by attaching the free end of the PEEK tubing to an LC pump and pumping 50:50 acetonitrile/water through, at 1 mL/min.

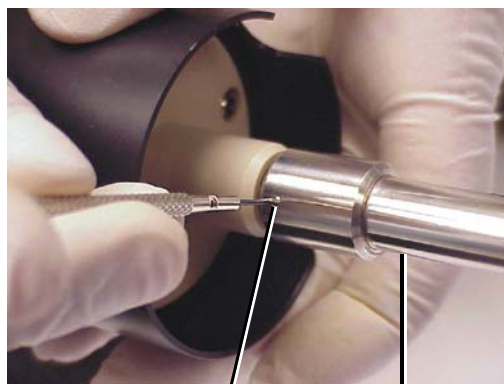


If leakage occurs, disassemble and remake the connection, and repeat the leak test.



Leaking Liquid

9. When performing the leak test, check the LC pump's backpressure, which will be high if the capillary is blocked. If this is the case, replace the capillary.
10. When the leak test has been performed successfully, disconnect the PEEK tubing from the LC pump.
11. Remove the probe inlet connector and PEEK tubing from the PEEK union.
12. Use the jeweller's screwdriver to loosen the two set screws securing the probe heater cover to the probe.



Set Screw

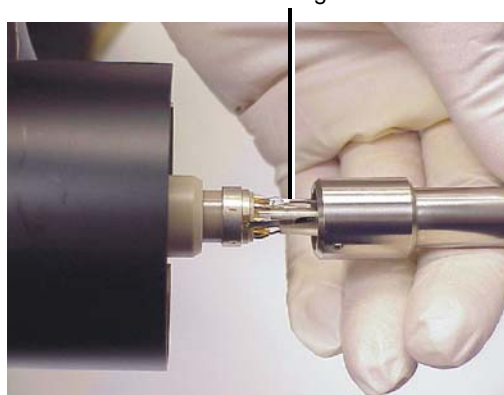
Probe Heater Cover



Caution: Take great care not to damage the probe heater's electrical wiring when removing the probe heater cover, or while the probe heater is exposed.

13. Carefully pull the probe heater cover off the probe, revealing the probe heater.

Probe Heater Electrical Wiring





Caution: When handling the probe heater, take great care to grip the heater so as not to damage its electrical wiring.



Caution: To avoid damaging the electrical connections to the probe heater, do not twist the heater when removing it from the probe assembly.

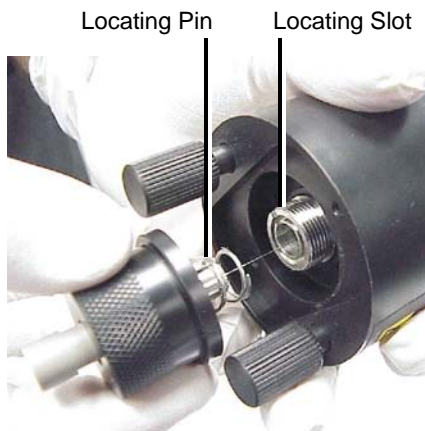
14. Gripping the probe heater as shown, carefully pull it off the probe assembly.

Probe Heater



15. Fit the PEEK union/UNF coupling assembly to the nebulizer adjuster knob.
16. Fit the compression spring to the capillary and PEEK union/UNF coupling assembly.
17. Carefully thread the capillary through the probe assembly.

18. Depress the PEEK union so that the locating pin on the UNF coupling is fully engaged in the locating slot at the head of the probe assembly. When the union is fully depressed, tighten the nebulizer adjuster knob. Do not tighten the knob fully.



19. Fit the probe end-cover to the probe assembly.
20. Fit and tighten the two end-cover securing screws.

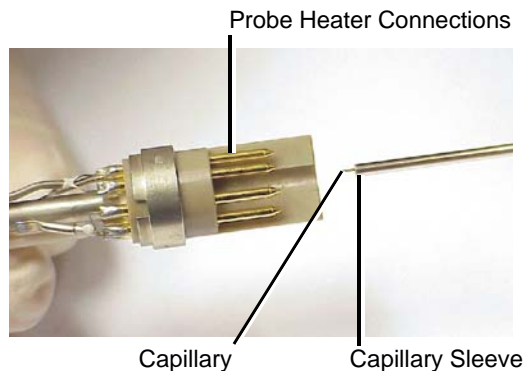


Caution: When handling the probe heater, take great care to grip the heater so as not to damage its electrical wiring.



Caution: Take great care not to damage the probe heater's electrical connections, capillary sleeve, or capillary when fitting the heater over the capillary sleeve.

21. Carefully slide the probe heater over the capillary sleeve on the probe assembly.





Caution: To avoid damaging the electrical connections to the probe heater, do not twist the heater when fitting it to the probe assembly.

22. Fit the probe heater to the probe assembly, ensuring that the heater is fully seated on the probe assembly.



23. Fit the probe heater cover to the probe assembly.
24. Use the jeweller's screwdriver to tighten the two set screws securing the probe heater cover to the probe.
25. Use the nebulizer adjuster knob to adjust the capillary so that the capillary protrudes by approximately 0.5 mm from the end of the probe.
26. Fit the probe to the instrument.

7.14 Replacing the APCI Probe Heater

Replace the APCI probe heater if it fails to heat.

7.14.1 Required Materials

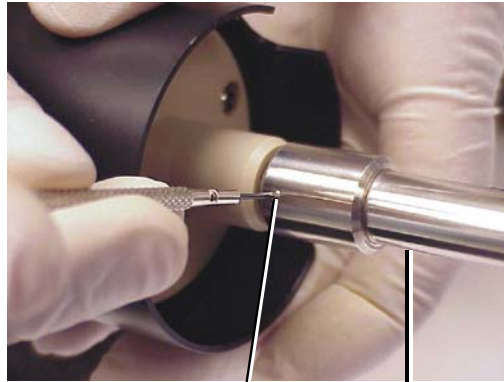
Jeweller's screwdriver

7.14.2 Procedure



Warning: The probe may be contaminated with biohazardous and/or toxic materials. Always wear nitrile gloves while performing this procedure.

1. Use the jeweller's screwdriver to loosen the two set screws securing the probe heater cover to the probe.



Set Screw

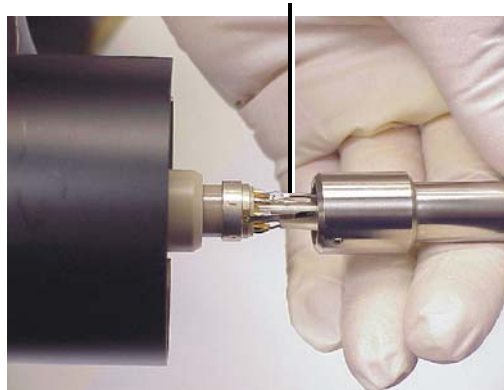
Probe Heater Cover



Caution: Take great care not to damage the probe heater's electrical wiring when removing the probe heater cover, or while the probe heater is exposed.

2. Carefully pull the probe heater cover off the probe, revealing the probe heater.

Probe Heater Electrical Wiring





Caution: When handling the probe heater, take great care to grip the heater so as not to damage its electrical wiring.



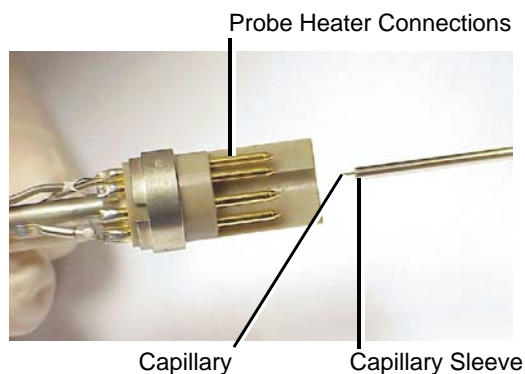
Caution: To avoid damaging the electrical connections to the probe heater, do not twist the heater when removing it from the probe assembly.

3. Gripping the heater as shown, carefully pull the probe heater off the probe assembly.



Caution: Take great care not to damage the probe heater's electrical connections, capillary sleeve, or capillary when fitting the heater over the capillary sleeve.

4. Carefully slide the new probe heater over the capillary sleeve on the probe assembly.





Caution: To avoid damaging the electrical connections to the probe heater, do not twist the heater when fitting it to the probe assembly.

5. Fit the probe heater to the probe assembly, ensuring that the heater is fully seated on the probe assembly.



6. Fit the probe heater cover to the probe assembly.
7. Use the jeweller's screwdriver to tighten the two set screws securing the probe heater cover to the probe.
8. Use the nebulizer adjuster knob to adjust the capillary so that the capillary protrudes by approximately 0.5 mm from the end of the probe.
9. Fit the probe to the instrument.

Chapter 8

Troubleshooting

This chapter describes how to troubleshoot the Quattro Premier Mass Spectrometer with the help of recommended troubleshooting procedures. This chapter covers:

- Safety and handling
- General troubleshooting
- Component hardware troubleshooting

8.1 Spare Parts

Waters recommends that the customer only replace parts mentioned in this document.

8.2 Safety and Handling

When troubleshooting the Quattro Premier, keep the following safety considerations in mind.



Warning: To avoid electric shock, do not remove the instrument's panels. There are no user-serviceable items inside the instrument.



Warning: To prevent injury, always observe good laboratory practices when handling solvents, changing tubing, or operating the Quattro Premier. Know the physical and chemical properties of the solvents used (see the Material Safety Data Sheets for the solvents in use).

8.3 System Troubleshooting

Examine the system, checking the simple things first. Is something obvious causing the problem, for example, are the instrument and its cables improperly connected, or is there any leakage of fluid, vacuum, or gas?

Compare current system operation with the way the system operated before the problem appeared. To help identify normal operating conditions:

- Make a note of the LC system (tubing and electrical connections).
- Keep a daily log.
- Run test samples regularly. Check the instrument performance with known samples, preferably the ones used for instrument qualification.

This illustrates the importance of keeping track of system parameters and performance during normal operation. Troubleshooting is easier when typical operating conditions are known. For example, are the system tuning parameters values similar to those when a test sample was run previously? Are the lens settings required for optimum sensitivity higher than those obtained previously? If extreme values have to be used to achieve good results, some part of the system is likely to require attention.

Methodically check and eliminate possible causes to identify the system parameter that is atypical.

See the troubleshooting information in the following sections to identify possible causes of symptoms and suggest corrective actions.

If you determine that a problem relates to another system component (for example, HPLC, autosampler, UV detector), see the appropriate operator's guide.

8.4 Component Hardware Troubleshooting

The following sections provide suggestions for resolving hardware problems.

8.4.1 Power Switch Fails to Power-up the Instrument



Warning: Parts of the instrument may be electrically live even when a mains power fuse has failed. To avoid electric shock, isolate the instrument from the mains supply before replacing the mains power fuses.

Possible Cause	Corrective Action
Mains power fuse failure.	Replace both mains power fuses (see Section 1.6.14).

8.4.2 No Peaks in the Tune Window (No Ion Beam)

Possible Cause	Corrective Action
The Tune window parameters are improperly set.	Optimize the parameters (see Sections 2.2.1 and 3.2.1). Once a beam has been obtained, ensure that all lenses affect the beam as expected.
Cables are not properly connected.	Ensure that all the cables have been correctly attached to the source and probe.
The instrument is not in Operate.	Put the instrument into Operate by clicking Press for Operate in the Tune window. (When in Operate, the adjacent icon and the Operate LED on the instrument's front panel are both green.)
Communication failure.	Reinitialize the instrument by selecting Options > Reinitialize in the Tune window. Reboot the embedded PC using the embedded PC reset switch (see Section 1.5.2).
No sample is present.	Ensure that sample is loaded correctly in the autosampler or syringe pump syringe.

Possible Cause	Corrective Action
The isolation valve is closed.	Open the isolation valve (see Figure 2-4).
The source components are dirty.	Clean the source components (see Section 7.7).
Insufficient nitrogen flow.	Ensure that the nitrogen pressure is 6 to 7 bar (90 to 105 psi) and the gas flow rate in the Tune window is >100 L/h.
No LC flow.	Look for solvent flow from the autosampler or syringe pump.
Fluid leak in the HPLC system.	Replace the APCI capillary (see Section 7.13).
The source components have been incorrectly assembled.	Ensure that the source and probe voltage readbacks vary with the Tune window settings. If any voltage is absent, disassemble and correctly reassemble the source and T-Wave assemblies (see Section 7.7).
The ESI or APCI sample capillary is blocked.	Replace the capillary (see Section 7.12 or Section 7.13).

8.4.3 Unsteady or Low Intensity Peaks (Ion Beam)

Possible Cause	Corrective Action
Poor nebulization.	<p>In the Tune window, ensure that the source and desolvation temperature, and gas flow settings are suitable for the flow rate.</p> <p>Liquid inside the source enclosure indicates that the source temperature is too low; in the Tune window Source page, increase the Source Temp (°C) parameter value.</p> <p>Ensure that the nitrogen pressure is 6 to 7 bar (90 to 100 psi).</p> <p>Check the stability of the nitrogen flow (use a good-quality two-stage regulator).</p> <p>In the Tune window Source page, ensure that the Desolvation Gas Flow (L/h) parameter is greater than 100.</p>
Problem with the sample delivery (autosampler, syringe pump, or HPLC system).	<p>Troubleshoot the autosampler.</p> <p>Inspect the syringe in the syringe pump for leaks and ensure that it is correctly grounded (earthed).</p> <p>Ensure that there is sufficient sample in the vials.</p> <p>Look for pressure variation on injecting the sample.</p>
Fluid leak in the HPLC system.	<p>Look for leaks in the HPLC system, and rectify them.</p>
The source components require cleaning.	<p>Clean the source components (see Section 7.7).</p>
The lens settings are wrong or atypical.	<p>Ensure that all the settings are correct.</p> <p>Ensure that the Tune window readbacks have reasonable values.</p> <p>Ensure that all the lens parameters affect the beam.</p>

Possible Cause	Corrective Action
The cone or collision cell voltage ramp is on.	Turn off the voltage ramp (see Section C.7).
The ESI or APCI sample capillary is not properly installed.	Ensure that the probe position is correct. Ensure that the ESI or APCI probe sample capillary protrudes 0.5 mm (see Sections 7.12 and 7.13).
The ESI probe tip subassembly O-ring is damaged.	Replace the O-ring (see Section 7.8).
The corona discharge pin is not correctly aligned.	Ensure that the corona discharge pin is correctly aligned (see Section 7.9).
The CID gas pressure is incorrect.	Infuse sample and optimize the gas pressure. Check that the CID gas regulator is set to 0.5 bar, and is not leaking.
The collision cell parameter values are incorrect.	In the Tune window, confirm that the Entrance, Exit, and Collision parameters are optimized, and have reasonable readbacks.
The analyzer and multiplier parameters incorrect.	In the Tune window, ensure that Multiplier is set to 650. Ensure that the Ion Energy and Resolution parameters are set correctly for the acquisition.

8.4.4 Unusually High LC Backpressure



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when inspecting the sample capillary, injection loop, or LC system tubing.

Possible Cause	Corrective Action
There is a blockage in the sample capillary or injection loop due to particulate matter from the sample.	Remove the probe from the source and increase the solvent flow to 500 $\mu\text{L}/\text{min}$ to clear the blockage.

Possible Cause	Corrective Action
The tubing from LC system is blocked.	Remove the finger-tight nut and tubing from the back of the probe. If the backpressure remains high, replace the tubing.
The ESI probe sample capillary is blocked.	Replace the sample capillary (see Section 7.12).
The ESI or APCI probe sample capillary is not fully seated in the LC union.	Remove and disassemble the probe, and reseal the sample capillary correctly in the union (see Section 7.12 or Section 7.13).

8.4.5 Unusually Low LC Backpressure

Possible Cause	Corrective Action
Leaking connector.	Inspect all the fittings and tighten them if necessary.
Problem with the LC solvent delivery.	Troubleshoot the LC system.

8.4.6 Insufficient Vacuum

There is insufficient vacuum when the Pirani gauge is reading greater than 5×10^{-4} mbar, when the CID gas is off.

Possible Cause	Corrective Action
The ion block O-rings are leaking.	Disassemble the source and check the condition of the ion block O-rings (see Section 7.7).

Possible Cause	Corrective Action
The backing pump is not operating correctly.	If using a rotary pump, gas ballast the pump to return accumulated oil from the oil mist filter (see Section 7.3). Check the rotary pump oil (see Section 7.4). If the oil is dirty, change the oil (see Section 7.5). Repeat if necessary. If using a scroll pump, replace the scroll pump seals (see the Edwards document <i>XDS 35i Instruction Manual A730-01-880</i> , supplied with the instrument).
Leak in the vacuum backing line.	Inspect the vacuum hose for cracks or vacuum leaks.
Restriction in the vacuum pump exhaust tubing.	Inspect the exhaust line for restrictions.
Turbo pump not operating properly.	Contact Waters for advice (see Section 8.6).

8.4.7 Leaking Nitrogen

A hissing sound or solvent smell can indicate a nitrogen leak.

Possible Cause	Corrective Action
Poor seal around the source enclosure.	Visually inspect the source enclosure sealing surfaces for imperfections or nicks. Examine the condition of the encapsulated O-rings.

8.4.8 Rotary Pump Oil Accumulated in the Exhaust Tubing

Possible Cause	Corrective Action
The oil mist filter needs replacement.	Replace the oil mist and odor filter elements (see Section 7.6).

8.4.9 Ion Source Heater and Desolvation Heater are Not Working

Possible Cause	Corrective Action
The ion source heater has failed.	Check the Source Temp (°C) readback on the Tune window Source page. Replace the heater if necessary (see Section 7.11).
The main system printed circuit board fuse has failed.	Check the Desolvation Temp (°C) readback on the Tune window Source page. Contact Waters if the readback is incorrect (see Section 8.6).

8.4.10 APCI Probe Heater Not Working

Possible Cause	Corrective Action
If the desolvation heater is working in ESI mode, the APCI probe heater may need replacing.	Replace the APCI probe heater (see Section 7.14).

8.4.11 Failure of the Fuse Supplying the Rotary Pump

Note: This fuse is not supplied by Waters – it is in the user's power supply.

Possible Cause	Corrective Action
The oil mist filter element is saturated. Vacuum oil may also be accumulating in the exhaust tubing.	Replace the oil mist and odor filter elements (see Section 7.6).
The system needs to be ballasted.	Ballast the pump for 20 to 30 minutes (see Section 7.3).
The mains supply voltage is less than 208 V a.c.	The mains supply voltage to the instrument must be measured by a qualified electrician.
The rotary pump oil is very dirty.	Change the rotary pump oil (see Section 7.5).

8.4.12 Ion Mode Fault

The Tune window drop-down menu options are unavailable, or the instrument spontaneously switches probe type.

Possible Cause	Corrective Action
One, or both, of the probe contact pins are jammed inside the probe and are not making contact with probe support plate.	Remove the probe cover, free the contact pin, and ensure that both the pins and their associated springs move freely within the bushing.

8.4.13 Failure to Recognize a Particular Probe Type

Possible Cause	Corrective Action
A problem with the probe.	Remove the probe, and try another probe of the same type. On the Tune window Diagnostics page, ensure that the Source ID (V) value is 1.5 to 2.5 V for ESI and 2.5 to 3.5 V for APCI.

8.4.14 Ripple

The peaks and baseline appear to vary cyclically in intensity.



Warning: To avoid high-pressure liquid jet spray, wear safety goggles when inspecting the sample capillary, injection loop, or tubing from the LC system.

Possible Cause	Corrective Action
Erratic LC solvent flow.	Troubleshoot the LC system.
Poor nebulization due to incorrect temperature and gas flow settings.	Adjust the temperature and gas flow settings. Liquid in the source enclosure indicates that the temperature is too low.
Vibration from the rotary pump or even other equipment in the same building.	Look for and eliminate excessive bench top and instrument vibration.

8.4.15 Loss of Communication with the Instrument

Possible Cause	Corrective Action
The instrument to MassLynx host communication has failed.	Reset the workstation and, when rebooted, reboot the embedded PC from the front panel using a short length of PEEK tubing to operate the reset switch (see Figure 1-3 on page 5). Wait 3 minutes for the audible signal indicating that the embedded PC has booted from the Quattro Premier before starting MassLynx.

8.4.16 IEEE Communication Errors

Possible Cause	Corrective Action
The instrument's components have powered up in the wrong sequence.	Power down the system components and start up the system components in the correct order: <ol style="list-style-type: none">1. Workstation2. Quattro Premier3. Inlet modules Wait 3 minutes for the audible signal indicating that the embedded PC has booted from the Quattro Premier before starting MassLynx.
There is a wrong or conflicting IEEE address.	Check the system's IEEE settings and enter the correct addresses.
There is a faulty IEEE cable in the IEEE chain.	Systematically replace the IEEE cables until the problem cable is located.
Network cables are confused with the site network.	Ensure that the network cable for the instrument is connected to the correct network card in the PC. Ensure that the network card with the BNC connector is configured to the site network.

8.5 High Noise Levels in MRM Analyses

The background noise in MRM analysis can be either electronic or chemical. To distinguish between the two:

1. Start an acquisition.
2. During the acquisition, set Ion Energy 1, and Ion Energy 2 fully negative in the Tune window Analyser page.

A significant decrease in signal when the ion energies are set negative implies that the major contribution to the overall noise is chemical.

Any residual noise is electronic.

8.5.1 Chemical Noise

Possible Cause	Corrective Action
High background noise due to carry-over after tuning with strong concentrations.	Repeat injections of 10% formic acid and/or isopropanol.
Contaminated injector. (The signal changes upon injection of mobile phase).	Repeat injections of 10% formic acid and/or isopropanol.
Contaminated tubing.	Replace the tubing.
Contaminated probe.	Flush with methanol at 0.5 mL/min until the background noise level falls. Replace the ESI or APCI sample capillary (see Sections 7.12 and 7.13).
Contaminated HPLC system.	Infuse mobile phase from the solvent reservoir using a syringe pump. Compare the MRM background levels. Confirm the purity of solvents and replace them if necessary. Ensure that all solvents are HPLC grade.
Contaminated glassware.	Ensure that glassware is not cleaned with commercial surfactants.

8.5.2 Electronic Noise

Corrective Action
Ensure that the valleys of peak-peak noise, when ion energies are fully negative, touch the baseline. Increase the Ion Counting Threshold to suit; ensure that this does not reduce the sensitivity on low level peaks too much (see Section 5.3.4).

8.6 Contacting Waters

You can easily correct many problems with the Quattro Premier. However, if this is not the case, you must contact Waters.

Customers in the USA and Canada should report maintenance problems they cannot resolve to Waters Technical Service (800 252-4752). All others should visit <http://www.waters.com> and click Offices, or phone their local Waters subsidiary or Waters corporate headquarters at 34 Maple Street, Milford, MA 01757, USA.

When contacting Waters, have the following information available:

- The nature of the symptom
- The Quattro Premier serial number

Depending on the nature of the fault, it may also be useful to have the following information available:

- Details of the flow rate, mobile phases, and sample concentrations
- Details of the gas cell operating pressure
- The Tune window settings
- The software version update reference

Appendix A

Starting Up and Shutting Down the Instrument

A.1 Starting Up the Instrument

1. Switch on the power switch located on the lower-right side corner of the instrument's front panel (see [Section 1.6.14](#)).
2. Allow 3 minutes for the embedded PC to initialize. An audible alert is given when the PC is ready.
3. Start the MassLynx software. The MassLynx window appears and the word "Ready" appears in the status bar at the bottom (see [Figure A-1](#)).
4. Click the Instrument shortcut bar MS Tune icon to open the Tune window (see [Figure A-2](#)).
5. Select Options > Pump.



Caution: The instrument may be damaged if the parameters in the Tune window Diagnostics page are modified by an unqualified user; the parameters on this page should only be modified by a Waters field service engineer.

6. Click the Diagnostics tab.
Note: If the Diagnostics tab is not present, select Options > View Diagnostics Page.
7. Monitor the Turbo Speeds. These should reach 98 to 100% within approximately 5 minutes of Options > Pump being selected.
8. Ensure that the instrument has pumped sufficiently such that the Vacuum LED on the front panel is steady green (see [Section 1.5.5](#)). The mass spectrometer is sufficiently evacuated to enable operation in 20 minutes.

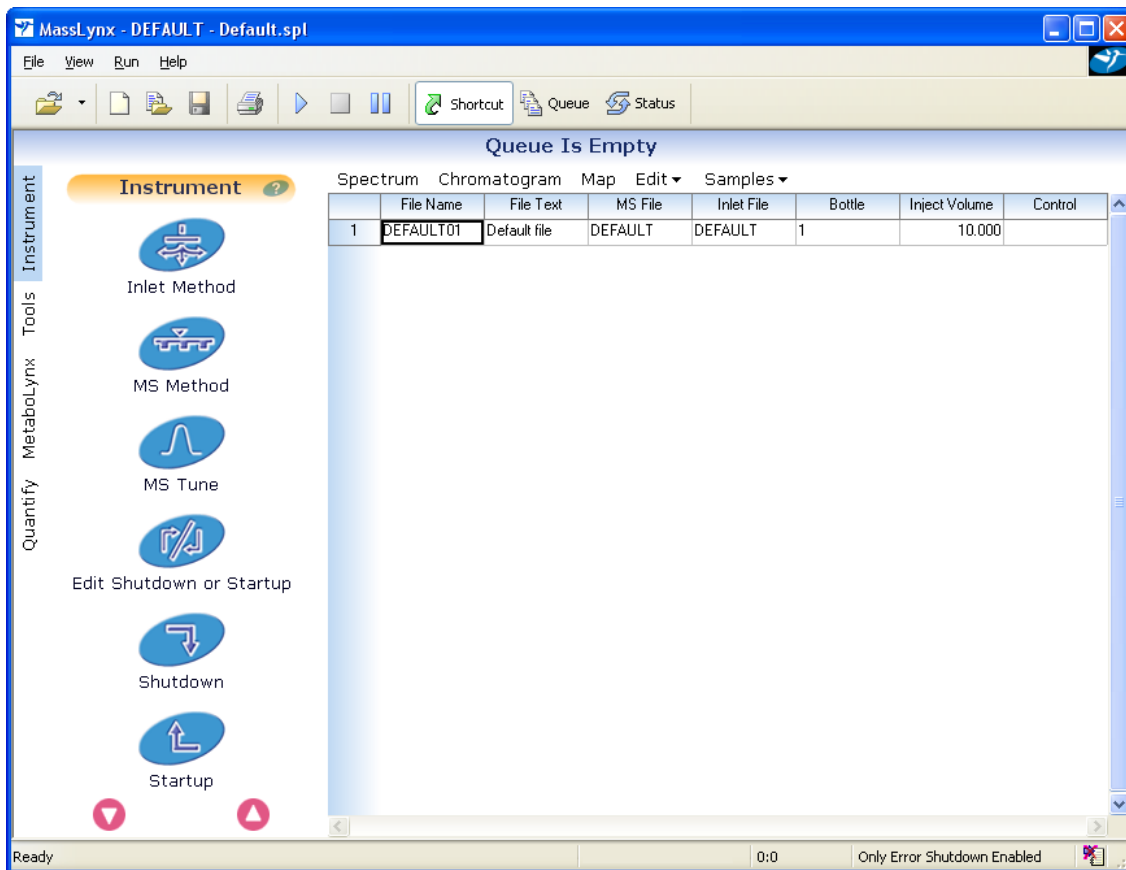


Figure A-1 MassLynx Window

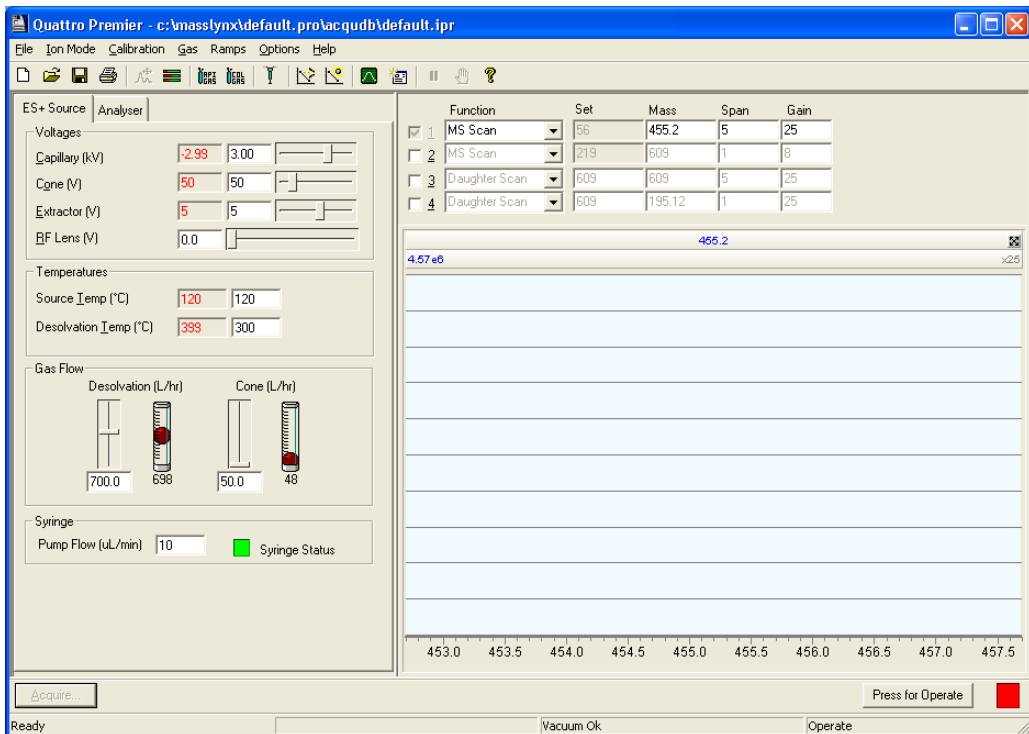


Figure A-2 Tune Window

A.2 Shutting Down the Instrument

A.2.1 Emergency Shutdown



Warning: *The power switch does not isolate the instrument from the mains power supply. Unplug the mains power from the rear of the instrument to isolate the instrument from the mains power supply.*

If you must shut down the instrument in an emergency, switch off the power at the wall mounted isolation switch(es), if fitted. If not, switch off the power switch located on the lower-right side corner of the instrument front panel (see [Section 1.5.1](#)), and turn off all peripherals.

Note: *Data may be lost.*

A.2.2 Overnight Instrument Shutdown

When the instrument is to be left unattended for a long time, for example, overnight or at weekends, proceed as follows:

1. On the MassLynx Tune window Source page, set the Voltages, Temperatures, Gas Flow, and Syringe parameters to zero.
2. If required, click Press for Standby to switch the instrument out of Operate.

Note: *It is not necessary to switch the instrument out of Operate. However, this is acceptable provided that the instrument warm-up time is considered when restarting analysis.*

A.2.3 Complete Instrument Shutdown

1. In the Tune window, click Press for Standby.
2. Select Options > Vacuum.
3. Select Vent Instrument.

A message confirms the vent command.

4. Click OK. The turbomolecular pumps are switched off. When the turbomolecular pumps have run down to half their normal operating speed, the vent valves are opened and the instrument is automatically vented. The Vacuum LED will switch off after changing from green to amber.
5. Exit MassLynx.

6. Shut down the PC.
7. Switch off all peripherals.
8. Switch off the power switch located on the lower-right side corner of the instrument front panel (see [Section 1.5.1](#)).

Note: *The power switch does not isolate the instrument; fans may be heard running, even when the instrument is “off”.*

Appendix B

Setting-Up the Syringe Pump

This appendix describes how to set-up the syringe pump, which is used for infusing the standard solution during the tuning process.



Warning: To avoid electric shock, clip the ground cable, attached beneath the lower right of the front panel, onto the syringe needle.

1. Clip the ground cable (with a plug-in clip), attached beneath the lower right of the front panel, onto the syringe needle.
2. Mount the syringe onto the pump (Figure B-1).

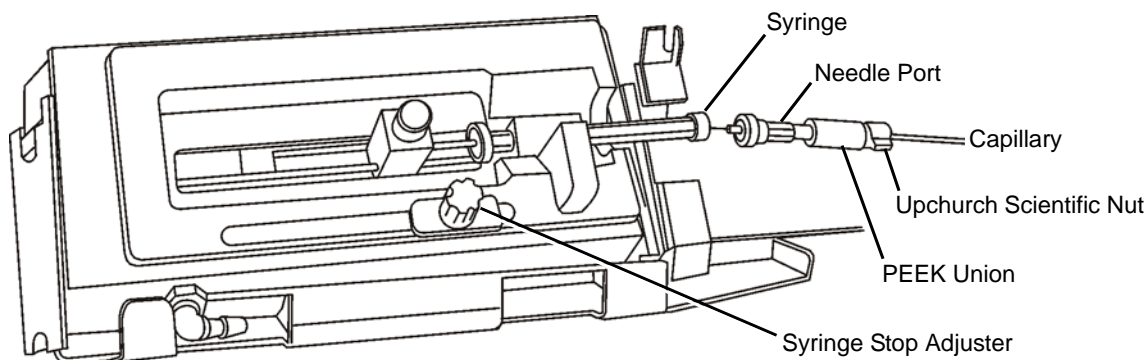


Figure B-1 Syringe Pump



Caution: The syringe pump has a positive syringe stop to prevent certain syringe types from breaking. Nevertheless, as added protection against syringe breakage, setting the syringe stop adjuster is recommended. This prevents the syringe plunger from travelling its full stroke inside the syringe barrel, thereby reducing the likelihood of breakage.

3. Set the syringe stop and syringe stop adjuster appropriately.
4. Screw the Rheodyne 9013 needle port fitting into the PEEK union and tighten it so that it will not leak.



Caution: *The sample capillary is fragile. Take great care to avoid damaging it while performing this procedure.*

5. Use a ceramic silica cutter to make a square, even cut on both ends of the sample capillary (supplied in the ESI probe installation kit) before installing. Examine new cuts for squareness using an eye glass. When cutting the capillary, allow enough length to form loops at angles and corners.
6. Feed the capillary through the hole at the top of the molding, the capillary will emerge from a hole at the top right side of the syringe pump area (see [Figure 1-3 on page 5](#)).
7. Connect the capillary to the PEEK union, using an Upchurch[®] Scientific nut, ferrule, and PTFE tubing.
8. Before using the syringe pump, ensure that the type of syringe used is selected in the MassLynx software (see [Appendix C, The MassLynx Tune Window](#)).

Appendix C

The MassLynx Tune Window

Note: For the highest mass accuracy, tune and calibrate the instrument using a suitable reference compound before sample data are acquired (see [Chapter 2](#), [Chapter 3](#) and [Chapter 6](#).)

C.1 Opening the MassLynx Tune Window

Click the MassLynx window Instrument shortcut bar MS Tune icon to open the Tune window. [Figure C-2](#) summarizes the Tune window layout.

C.2 Selecting the Ionization Mode

Select the required ionization mode from the Ion Mode menu ([Figure C-1](#)). The current mode has a check mark next to it.

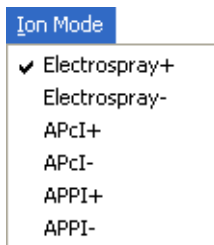


Figure C-1 Ion Mode Menu

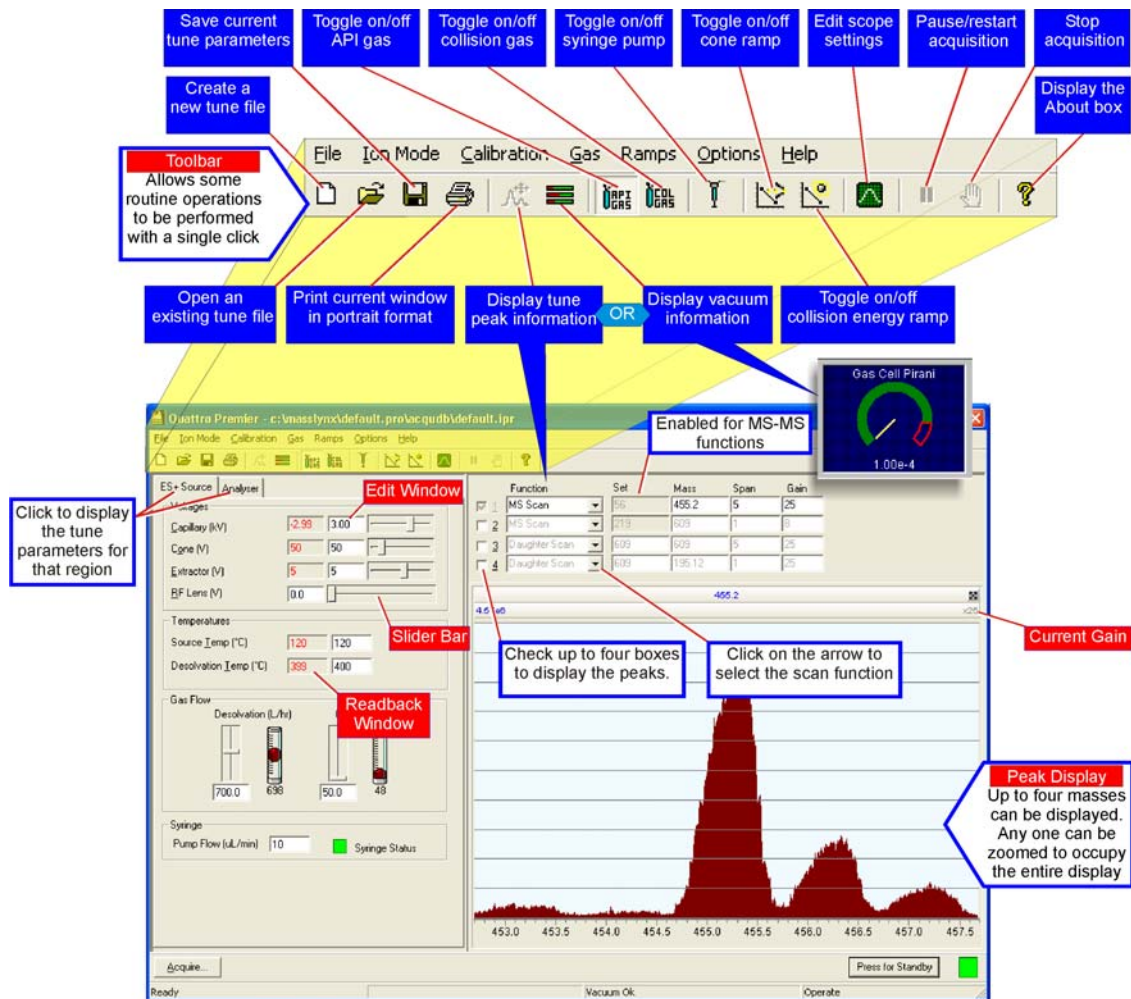




Figure C-2 Tune Window

C.3 Controlling Gas Flows


C.3.1 Controlling the Nebulizer, Desolvation, and Cone Gas Flows

To toggle the nebulizer, desolvation, and cone gasses on and off, click  or select Gas > Gas.

C.3.2 Controlling the Collision Gas Flow

To toggle the collision gas on and off, click  or select Gas > Collision Gas.

C.4 Controlling the Syringe Pump

To toggle the syringe pump on and off, click .

C.5 Selecting the Syringe Type

1. Select Options > Syringe Type to open the Syringe Selection dialog box ([Figure C-3](#)).

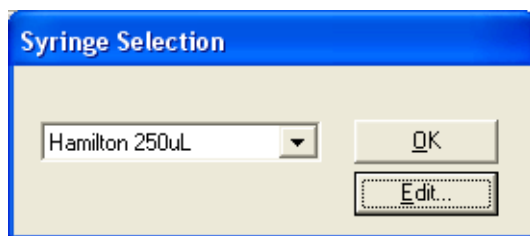



Figure C-3 Syringe Selection Dialog Box

2. Choose the syringe type from the drop-down list.
If the required syringe type is not available in the drop-down list, click Edit to open the syringe list, and then add the required details.
3. Click OK.

C.6 Selecting the Scan Time and Inter Scan Delay

Enter the Scan Time and Inter Scan Delay in the Scope Setup dialog box (Figure C-4).

This is opened by clicking  or selecting Options > Scope Parameters.

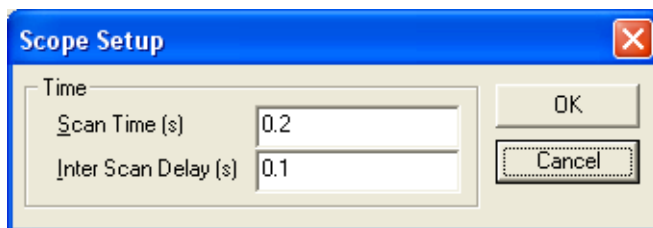


Figure C-4 Scope Setup Dialog Box

The Scan Time(s) and Inter Scan Delay (s) parameter values control the speed with which the tune peak display is updated.

Tuning is more responsive when these parameters are set to low values.

C.7 Setting the Ramp Controls

C.7.1 Creating a Cone Voltage Ramp

1. Select Ramps > Cone Ramp Gradient to open the Cone Ramp dialog box (Figure C-5).

Two values of cone voltage (Cone Start Volts and Cone End Volts) are defined at two particular masses (Start Mass and End Mass). These values define a gradient for the cone voltage, which is then extrapolated to cover the full mass range.

2. Make any changes required, and click OK to exit.

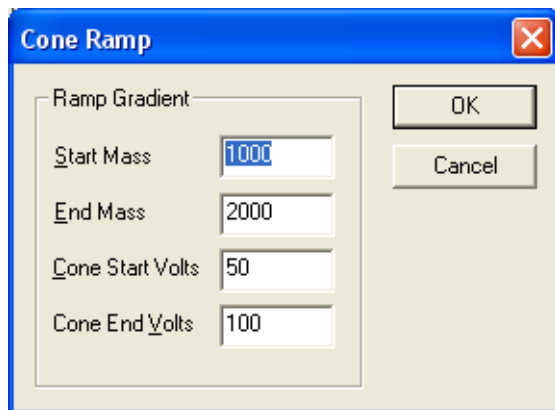



Figure C-5 Cone Ramp Dialog Box

C.7.2 Controlling the Cone Voltage Ramp

To toggle the cone voltage ramp on and off, click  or select Ramps > Use Cone Ramp.

C.7.3 Creating a Collision Energy Ramp

1. Select Ramps > Collision Energy Ramp Gradient to open the Collision Ramp dialog box (Figure C-6).

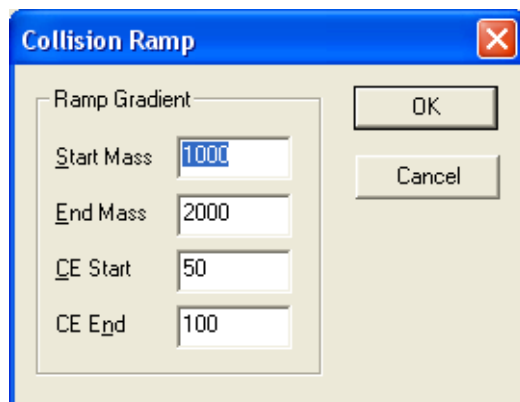



Figure C-6 Collision Ramp Dialog Box


Two values of collision energy (CE Start and CE End) are defined at two particular masses (Start Mass and End Mass). These values define a gradient for the collision energy voltage, which is then extrapolated to cover the full mass range.

2. Make any changes required, and click OK to exit.

C.7.4 Controlling the Collision Energy Ramp

To toggle the collision energy ramp on and off, click  or select Ramps > Use Collision Energy Ramp.

C.8 Resetting the Zero Level

The zero level (or baseline) can be repositioned by clicking  or by selecting Options > Reinitialize.

This command causes the instrument control system to measure the position of the noise signal so that any baseline offset caused by the electronics or instrumentation can be compensated for.

You should reset the zero level whenever the multiplier voltage is changed.

C.9 Controlling the Display of Readback Windows

The Readbacks dialog box (Figure C-7) is opened by selecting Options > Readbacks.

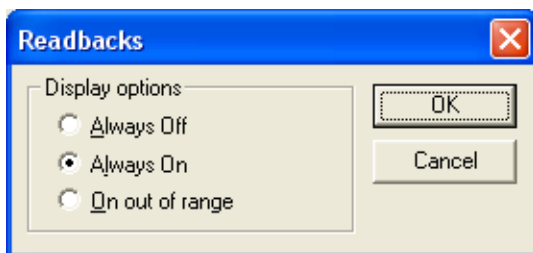


Figure C-7 Readbacks Dialog Box

There are three options for displaying system readbacks in the Tune window:

- Readbacks displayed continuously (Always On).
- Readbacks hidden (Always Off).
- Readbacks displayed only when differing from their defined values by more than 10% (On out of range).

A number of the readbacks are for indication purposes only and are not true (calibrated) records of the actual voltages on the instrument. The acceptable variation between the set value and the readback value varies depending on the particular tune parameter. If you are concerned about any reading, contact Waters for advice (see [Section 8.6](#)).

C.10 Changing Tune Parameter Settings

You can modify most parameters in the following ways:

- Use the mouse to drag the slider bar.
- Click the slider bar and use the left and right arrow keys to change the value by one increment. The edit window is updated as the slider bar is activated.
- Type a new value into the edit window, then press the Enter key.

Other parameters have only an edit window and are changed by direct typing.


The speed with which the system responds to changes depends on the speed with which the peak display is refreshed. For the fastest response, set the Scope Setup dialog box's Scan Time (s) and Inter Scan Delay (s) values as short as possible (see [Figure C-4](#)).

C.11 Saving Instrument Tune Parameters

Instrument tuning parameters can be saved in an instrument parameter file (.ipr), which can be recalled later.

An instrument parameter file contains all the parameters for all supported ionization modes, not just the ionization mode currently selected. Instrument parameter files also contain settings for the analyzer, inlet set points, and peak display.

C.11.1 Creating a New Instrument Parameter File

To create a new instrument parameter file, click  or select File > New.

C.11.2 Saving Tune Parameters in an Instrument Parameter File

To save the current tune parameters with the existing instrument parameter file name, click


 or select File > Save.

To save the current tune parameters with a new file name:


1. Select File > Save As to open the Save As dialog box.
2. Enter a new file name or select an existing file from the displayed list.
3. Click Save.
4. If the selected file already exists on disk, a warning is displayed. Click Yes to overwrite the existing information, or No to enter a different file name.

C.11.3 Opening an Existing Instrument Parameter File

To open an existing instrument parameter file, i.e., restore a saved set of tune parameters:

1. Click  or select File > Open to open the Open dialog box.
2. Select the required instrument parameter file, either by typing its name or selecting one from the list.
3. Click Open.

C.12 Printing Tune Information

To print a report containing a copy of the on-screen tune peak information and a record of each parameter setting, click  or select File > Print.

This report is not configurable by the user.

C.13 Using the EasyTune Source Page

The Tune window EasyTune Source page replaces the Source and Analyser pages. It displays the tune parameters commonly used during instrument tuning, while the more rarely used parameters are hidden.



By default, the MassLynx Tune window opens showing the Source and Analyser pages. To open the Tune Window EasyTune Source page (Figure C-8), select Options > EasyTune.

Using the EasyTune Source page, the instrument is tuned in a similar manner to that described in Chapters 2 and 3.

To close the EasyTune Source page and return to the Source and Analyser pages, select Options > Advanced.

C.14 Changing Between the Peak and Vacuum Displays

Peak information, or vacuum gauge readbacks, can be displayed in the top-right corner of the Tune window.

- To display the vacuum information, select  or select Options > Vacuum Monitor.
- To display the peak information, select  or Options > Peak Editor.

C.15 Setting-Up Peaks for Tuning

C.15.1 Selecting Peaks for Tuning

1. With the Tune window set to display peak information (see Section C.14), select the peaks to be displayed by selecting the appropriate numbered box(es) in the top-right of the Tune window.
2. For each active peak, enter the required values for the Mass, Span, and Gain.

Note: For MS/MS functions, Set is also enabled, allowing the mass of the parent, daughter, neutral loss, or neutral gain ion (as appropriate) to be entered.

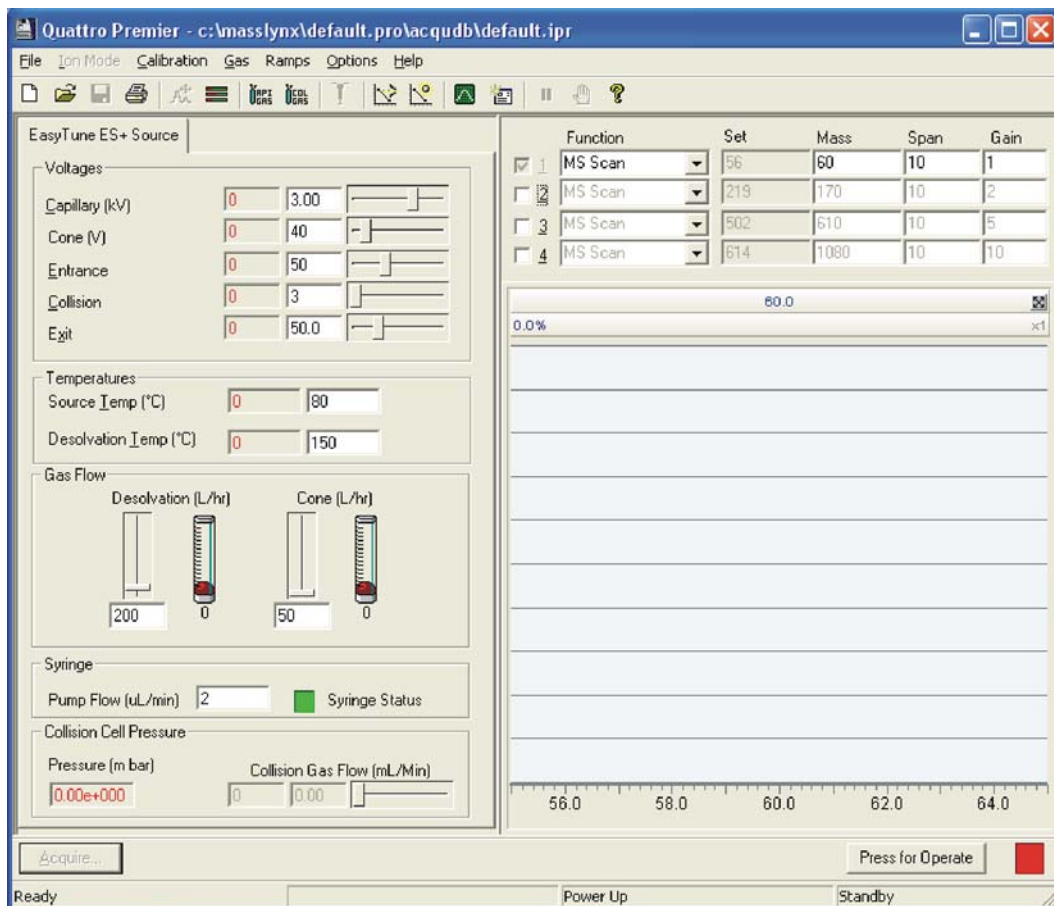


Figure C-8 Tune Window EasyTune Source Page


C.15.2 Selecting the Operating Mode for a Peak

To select the operating mode (scan function), select the required mode for the peak from the Function drop-down list.

C.15.3 Selecting the Tune Mass for a Peak

Either:


1. In the appropriate peak window, on the right side of the Tune window, click and drag the mouse within the bounds of the axis to draw a “rubber band” around the region of interest.

2. Release the mouse button. This range is redisplayed to fill the window. The value displayed in the Mass box is the mass at the center of the window. This operation can be repeated as often as required.
3. Clicking  once displays the previous magnification range and mass; clicking it again returns to the default settings.


Or:

1. Enter a value in the Mass box for the required peak.
2. Click Return.


This becomes the default, so if the range is then altered using the mouse and you click

 twice, Mass returns to this value.

Or:

1. Position the cursor at the top of the peak window, just below the line displaying the gain value (e.g., 2×).
2. When  appears, click and drag the mouse until the required mass is displayed in the Mass box and at the top of the window.


This becomes the default, so if the range is altered using the mouse and you click

 twice, Mass returns to this value.

C.15.4 Selecting the Span of a Displayed Peak

Either:


1. Click and drag the mouse, horizontally, from one end to the other of the region of interest. As the mouse is dragged, a “rubber band” indicates the selected range. Do not go beyond the bounds of the axis.
2. Release the mouse button to redisplay the selected range so that it fills the current window. This operation can be repeated as often as required.

Clicking  once displays the previous magnification range, clicking it again returns to the default settings.

Or:

1. Enter a value in the Span box for the required peak.

2. Click Return.

This becomes the default, so if the range is altered with the mouse and you click  twice, Span returns to this value.

C.15.5 Changing the Gain of a Displayed Peak

Either:

1. Double-click the line displaying the gain value (e.g., 2×) above the peak, to double the gain applied to that peak.
2. Double-click below the peak display to halve the gain.

Or:

1. Click and drag the left mouse button, vertically, from one end to the other of the region of interest.

As the mouse is dragged, a marquee indicates the selected range.

Do not go beyond the bounds of the axis.

2. Release the mouse button to redisplay the selected range so that it fills the current window.

Or:

1. Enter a value in the Gain box for the required peak.
2. Click Return.

C.16 Customizing the Peak Display

C.16.1 Opening the Peak Display Menu

Use the Peak Display menu ([Figure C-9](#)) to customize the Peak Display. Open it by right-clicking in the appropriate peak display window in the Tune window.

Note: *The display window for each peak can be individually customized, e.g., the peak color for peak 1 can be red, for peak 2 green, etc.*



Figure C-9 Peak Display Menu

C.16.2 Customizing the Colors and Numbers of Displayed Traces

To change the color of the background and traces, and to change the number of traces displayed, select Customise > Plot Appearance from the Peak Display pop-up menu to open the Customise Plot Appearance dialog box (Figure C-10).

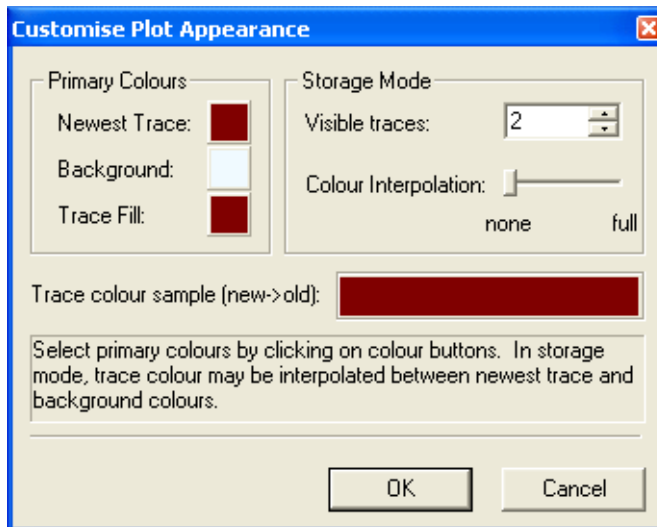


Figure C-10 Customise Plot Appearance Dialog Box

To change the colors on the display, click the color box adjacent to Newest Trace, Background, or Trace Fill as required, and select a new color from the Color dialog box.

To change the number of displayed traces, enter the required value in the Visible traces box, within the range 2 to 20.

If more than one trace is displayed, the older traces can be displayed in different color shades to the newer ones. Drag the Colour Interpolation slider toward the full position. The colors of the older traces appear in the Trace colour sample (new->old) field.

C.16.3 Customizing the Peak Trace Line Appearance

Each trace may be displayed as:

- Outline only – select Trace > Outline from the Peak Display pop-up menu.
- With the area below the line filled – select Trace > Fill from the Peak Display pop-up menu.
- Maximum and minimum points only – select Trace > Min/Max from the Peak Display pop-up menu.

The selected option displays a check mark in the Peak Display pop-up menu.

C.16.4 Customizing the Peak Intensity Display

To display the peak intensities as absolute values (counts/second), select Intensity > Absolute Intensity from the Peak Display pop-up menu.

To display the peak intensities as percentage values relative to the intensity of the highest peak, select Intensity > Relative Intensity from the Peak Display pop-up menu.

The Peak Display pop-up menu Intensity > Normalise Data option can be selected in conjunction with either of the above options. It controls the way in which the peak display is scaled. When enabled, the display scales to the value of the intensity of the highest peak; when disabled, the display scales to the default value set in the MassLynx software. It is recommended that the Intensity > Normalise Data option is normally enabled.

The selected options display a check mark in the Peak Display pop-up menu.

C.16.5 Customizing the Peak Display Grid

The Peak Display vertical and horizontal grid lines may be independently displayed or hidden.

To display the horizontal grid lines, select Grid > Horizontal from the Peak Display pop-up menu.

To display the vertical grid lines, select Grid > Vertical from the Peak Display pop-up menu.

The selected options display a check mark in the Peak Display pop-up menu.

C.17 Selecting the Instrument Name

1. Select Options > Instrument Name to open the Instrument Name dialog box.
2. Enter the required name in the ID text box.
3. Click OK.

C.18 The Diagnostics Page



Caution: *The instrument may be damaged if the parameters in the Tune window Diagnostics page are modified by an unqualified user. The parameters on this page should only be modified by a Waters field service engineer.*

The Tune window Diagnostics page, opened by selecting Options > View Diagnostics Page, is intended for use by Waters field service engineers only.

C.19 Manually Controlling the T-Wave Optics

The T-Wave devices have been optimized for use under normal operating conditions and their control should normally be left to MassLynx. Nevertheless, manual control of the optics may be desirable under certain conditions, for example, at high gas cell pressures ($>7 \times 10^{-3}$ mbar).

Note: *Inappropriate settings will result in poor performance in terms of sensitivity and/or high levels of cross-talk.*

The source T-Wave is only applied when using the optional MUX-technology interface (see the *Waters Micromass Quattro Premier MUX-technology Interface Operator's Guide* for details). The collision cell T-Wave is applied for MS methods that require collision gas in the cell.

To select manual control of the T-Wave devices, select Options > View T-Wave Manual Controls in the Tune window. This opens the T-WAVE page (Figure C-11). Separate control of the source and/or collision cell T-Wave pulse voltage and velocity is enabled by selecting the appropriate Enable Manual Controls check box(es).

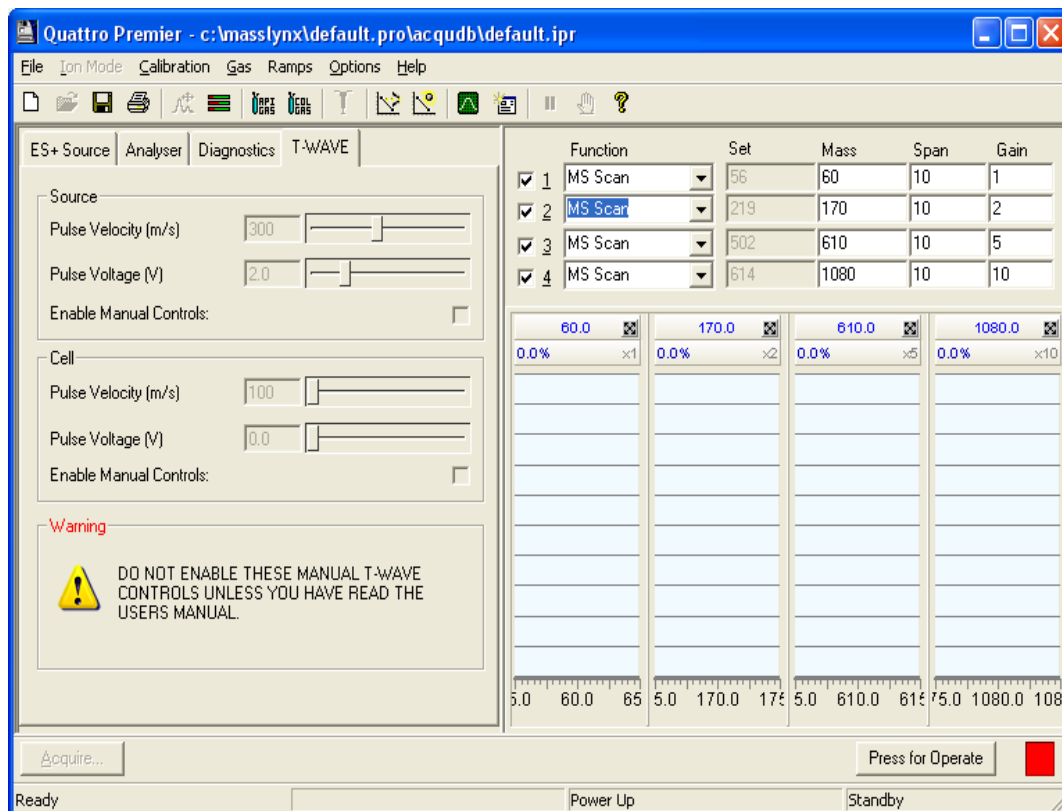


Figure C-11 Tune Window T-WAVE Page

Each T-Wave device has two adjustable parameters: the Pulse Velocity (m/s) and Pulse Voltage. The standard setting for the Pulse Velocity is 300 m/s. Setting a velocity that is too low or too high may increase cross-talk between MRM channels. The Pulse Voltage is set to different values depending of the MS method being employed; the default values are shown in Table C-1. If the pulse voltage is set too low, cross-talk will increase and ions will slip over the pulse instead of “surfing” in front of it. Setting the pulse voltage to too high a value will result in a decrease in sensitivity.

Table C-1 Default Pulse Voltage Values

MS Mode	Pulse Voltage (V)	
	Collision Cell	Source
MS1	0 (T-Wave is "OFF")	2*
MS2	0 (T-Wave is "OFF")	2
Daughter	2	2
MRM	2	2
Parent	5	2
Neutral Loss/Gain	5	2
Survey	2	2

*** Note:** The source pulse voltage is only applied when using the optional MUX-technology Interface.

Appendix D

Calibration Reference Information

D.1 Overview


Calibration reference files consist of two columns of numbers separated by any number of spaces or tab characters. The first column contains the reference peak masses, and the second column contains the reference peak intensities.

The reference files listed in this appendix have all ion intensities set to 100%. Actual ion intensities are not, in fact, all 100%, but the calibration software does not take account of the ion intensities and this is therefore a convenient way of storing the reference files in the required format. However, if required, realistic intensity values can be entered to improve the appearance of the reference spectra.

Most samples can be purchased from the Sigma chemical company. To order, contact Sigma at <http://www.sigma.sial.com>. This site contains a list of worldwide Sigma offices, many with local toll-free numbers.

D.2 Editing a Calibration Reference File

Calibration reference files can be created or edited using any Windows text editor. To read a reference file into the Notepad text editor:

1. In the Tune window, select Calibration > Calibrate Instrument to open the Calibration dialog box.
2. Select the required reference file from the drop-down list box.
3. Click  or select Edit > Reference File to open the reference file in Notepad.

To save the reference file after editing, select File > Save to save the file under the current name, or select File > Save As to save as a new reference file with a new name.

Textual information or comments can be stored in the reference file. Lines that are textual information or comments must start with the semicolon (;) character.

D.3 Positive Ion Calibration Reference Files

The following table lists typical reference files that can be used for positive ion calibration.

Ref. File Name	Chemical Name [Sigma Code #]	Molecular Mass	m/z	Uses
PEGNH4	Ammoniated polyethylene glycol		50 to 2000	General calibration
NAICS	Sodium iodide/cesium iodide mixture		20 to 4000	General, ES+ calibration
UBQ	Bovine ubiquitin [U6253]	8564.85	650 to 1500	General
HBA	Human α globin [H753]	15126.36	700 to 1500	Hb analysis
SOD	Superoxide dismutase [S2515]	15591.35	900 to 1500	Hb (internal calibration)
HBB	Human β globin [H7379]	15867.22	800 to 1500	Hb analysis
MYO	Horse heart myoglobin [M1882]	16951.48	700 to 1600	General
PEGH1000	Polyethylene glycol + ammonium acetate mixture PEG 200+400+600+1000		80 to 1000	ES+ and APCI+ calibration
PEGH2000	Polyethylene glycol + ammonium acetate mixture PEG 200+400+600+1000+1450		80 to 2000	ES+ calibration
NAIRB	Sodium iodide/rubidium iodide mixture		20 to 4000	ES+ calibration

D.3.1 Polyethylene Glycol

PEG + NH₄⁺

Reference File: PEGHNH4.REF

Calculated m/z Value								
89.06		459.28		872.54		1268.78		1665.01
133.09		503.31		916.57		1312.80		1709.04
177.11		564.36		960.60		1356.83		1753.07
221.14		608.39		1004.62		1400.86		1797.09
239.15		652.41		1048.65		1444.88		1841.12
283.18		696.44		1092.67		1488.91		1885.15
327.20		740.46		1136.70		1532.94		1929.17
371.23		784.49		1180.73		1576.96		1973.20
415.25		828.52		1224.75		1620.99		2017.22

D.3.2 Sodium Iodide and Cesium Iodide Mixture

Reference File: NAICS.REF

Calculated m/z Value								
22.9898		772.4610		1671.8264		2571.1918		3470.5572
132.9054		922.3552		1821.7206		2721.0861		3620.4515
172.8840		1072.2494		1971.6149		2870.9803		3770.3457
322.7782		1222.1437		2121.5091		3020.8745		3920.2400
472.6725		1372.0379		2271.4033		3170.7688		
622.5667		1521.9321		2421.2976		3320.6630		

D.3.3 Sodium Iodide and Rubidium Iodide Mixture

Reference File: NAIRB.REF

Calculated m/z Value								
22.9898		772.4610		1671.8264		2571.1918		3470.5572
84.9118		922.3552		1821.7206		2721.0861		3620.4515
172.8840		1072.2494		1971.6149		2870.9803		3770.3457
322.7782		1222.1437		2121.5091		3020.8745		3920.2400
472.6725		1372.0379		2271.4033		3170.7688		
622.5667		1521.9321		2421.2976		3320.6630		

D.4 Negative Ion Calibration Reference Files

Note: A positive ion calibration is sufficient for most applications. A specific negative ion calibration is not required when using the Quattro Premier.

Ref. File Name	Chemical Name [Sigma Code #]	Molecular Mass	m/z	Uses
NAINEG	Sodium iodide/cesium iodide (or rubidium iodide) mixture		200 to 3900	ES- calibration
SUGNEG	Sugar mixture of maltose [M5885], raffinose [R0250], maltotetraose [M8253], and corn syrup [M3639]		100 to 1500	Low mass range
MYONEG	Horse heart myoglobin [M1882]	16951.48	700 to 2400	General

You can use rubidium iodide to obtain a peak at m/z 85 ($^{85}\text{Rb}^+$) with an intensity of about 10% of the base peak at m/z 173. Rubidium iodide has the advantage that no rubidium clusters that can complicate the spectrum are formed. Note that rubidium has two isotopes (^{85}Rb and ^{87}Rb) in the ratio 2.59:1, giving peaks at m/z 85 and 87.

Use reference file NAIRB.REF.

D.5 Preparing Reference Sample Solutions

D.5.1 Preparing the PEGNH₄ Reference Sample Solution

Add 1 ng of each of the following to 50:50 acetonitrile/10 mM aqueous ammonium acetate:

- PEG 200
- PEG 400
- PEG 600
- PEG 1000

D.5.2 Preparing the Sodium Iodide and Cesium Iodide Mixture Sample Solution

Mix a 2 ng/ μ L solution of sodium iodide with a 50 ng/ μ L solution of cesium iodide.

Appendix E

Performance Specifications

Note: For the purpose of the following specifications, signal-to-noise is calculated as the ratio of the chromatographic peak height to twice the standard deviation of the noise.

E.1 Electrospray Positive Ion

The measured signal-to-noise ratio obtained from the chromatogram monitoring the transition m/z 609 to m/z 195 on injection of 5 pg of reserpine is $\geq 500:1$. This is based on a 5- μL injection of a 1 pg/ μL reserpine solution in 70:30 acetonitrile/water (no additives) at a flow rate of 200 $\mu\text{L}/\text{min}$ in MRM mode, 0.5 second dwell, span 0 Da.

The resolution of the precursor and product ions is <1 Da peak width at half-height.

E.2 Electrospray Negative Ion

The measured signal-to-noise ratio obtained from the chromatogram monitoring the transition m/z 503 to m/z 179 on injection of 25 pg of raffinose is $\geq 50:1$. This is based on a 5- μL injection of a 5 pg/ μL raffinose solution in 70:30 acetonitrile/water (no additives) at a flow rate of 200 $\mu\text{L}/\text{min}$ in MRM mode, 0.5 second dwell, span 0 Da.

The resolution of the precursor and product ions is <1 Da peak width at half-height.

E.3 MS Resolution

The resolution is demonstrated using a 1 $\mu\text{g}/\mu\text{L}$ solution of PPG 2000 in 50:50 acetonitrile/water containing 1-mM ammonium acetate. The peaks at m/z 2009.5 and 2010.5 should be resolved with a valley between them of no more than 15% of the height of the 2010.5 peak. It is recommended that fifteen 1-second scans are summed and the resulting spectrum smoothed (two passes, 0.5 Da SG).

E.4 Mass Measurement Accuracy

The mass measurement accuracy is measured from the mean of five repeat analyses of the $[M+NH_4]^+$ peak at m/z 1004.622 from 1 $\mu\text{g}/\mu\text{L}$ PEG 1000 in 50:50 acetonitrile/water containing 2 mM ammonium acetate. The mean measured mass will be 1004.622 ± 0.05 Da. The standard deviation of the mean will be ≤ 0.05 Da.

A mass calibration will be performed using the $[M+H]^+$ peaks from a separate analysis over the mass range m/z 700 to 1300 and the resolution on the $[M+H]^+$ peak at m/z 1031.62 must be between 0.3 and 0.4 Da wide at half height after smoothing.

E.5 APCI Positive Ion

Measured signal-to-noise ratio obtained from the chromatogram monitoring the transition m/z 331.2 to m/z 109.1 on injection of 50 pg of 17- α -hydroxyprogesterone will be $\geq 70:1$, using a 5 μl injection of a 10 $\text{pg}/\mu\text{l}$ 17- α -hydroxyprogesterone solution in 70:30 acetonitrile/water (no additives) at a flow rate of 1 mL/min in MRM mode, 0.2 second dwell, span 0 Da.

The resolution of the precursor and product ions will be < 1 Da peak width at half height.

Appendix F

Theory and Principles of Operation

F.1 Ionization Techniques

Two atmospheric pressure ionization techniques are available:

- Electrospray ionization (see [Section F.1.1](#)).
- Atmospheric pressure chemical ionization (see [Section F.1.2](#)).

F.1.1 Electrospray Ionization (ESI)

Electrospray ionization takes place as a result of imparting a strong electrical field to the eluent flow as it emerges from the nebulizer, producing an aerosol of charged droplets. These undergo a reduction in size by solvent evaporation until they have reached a charge density sufficient to allow sample ions to be ejected from the droplet's surface ("ion evaporation").

A characteristic of ESI spectra is that ions may be singly- or multiply-charged. Since the mass spectrometer filters ions according to their mass-to-charge ratio, compounds of high molecular weight can be determined if multiply-charged ions are formed.

Eluent flows up to 1 mL/min can be accommodated, although it is often preferable to split the flow such that 100 to 200 $\mu\text{L}/\text{min}$ of eluent enters the mass spectrometer.

F.1.2 Atmospheric Pressure Chemical Ionization (APCI)

Atmospheric pressure chemical ionization generally produces protonated or deprotonated molecular ions from the sample via a proton transfer (positive ions) or proton abstraction (negative ions) mechanism. The sample is vaporized in a heated nebulizer before emerging into a cloud of solvent ions formed within the atmospheric source by a corona discharge. Proton transfer, or abstraction, then takes place between the solvent ions and the sample. Eluent flows up to 2 mL/min can be accommodated without splitting the flow.

F.2 Ion Optics

Figure F-1 shows the Quattro Premier ion optics.

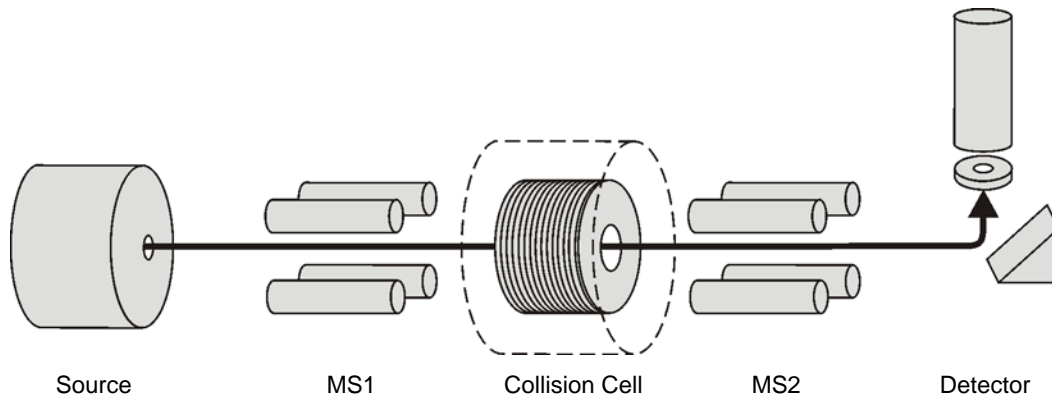


Figure F-1 Ion Optics

F.3 MS Operating Modes

Table F-1 shows the MS operating modes.

Table F-1 MS Operating Modes

Operating Mode	MS1	Collision Cell	MS2
MS1	Resolving (scanning)	Pass all masses	
MS2	Pass all masses		Resolving (scanning)
SIR	Resolving (static)	Pass all masses	

The MS1 mode, in which MS1 is used as the mass filter, is the most common and most sensitive method of performing MS analysis. This is directly analogous to using a single quadrupole mass spectrometer.

The MS2 mode of operation is used, with collision gas present, when switching rapidly between MS and MS/MS operation (for example, survey scan mode). It also provides a useful tool for instrument tuning and calibration before MS/MS analysis, and for fault diagnosis.

The SIR (Selected Ion Recording) mode of operation is used as a quantitation mode when no suitable fragment ion can be found to perform a more specific MRM analysis (see [Section F.4.3](#)).

F.4 MS/MS Operating Modes

The four common MS/MS operating modes are summarized in [Table F-2](#).

Table F-2 MS/MS Operating Modes

Operating Mode	MS1	Collision Cell	MS2
Daughter (Product) Ion Spectrum	Static (at parent mass)	Pass all masses	Scanning
Parent (Precursor) Ion Spectrum	Scanning		Static (at daughter mass)
Multiple Reaction Monitoring (MRM)	Static (at parent mass)		Static (at daughter mass)
Constant Neutral Loss Spectrum	Scanning (synchronized with MS2)		Scanning (synchronized with MS1)

F.4.1 Daughter (Product) Ion Mode

The daughter (product) ion mode is shown in [Figure F-2](#). It is the most commonly used MS/MS operating mode.

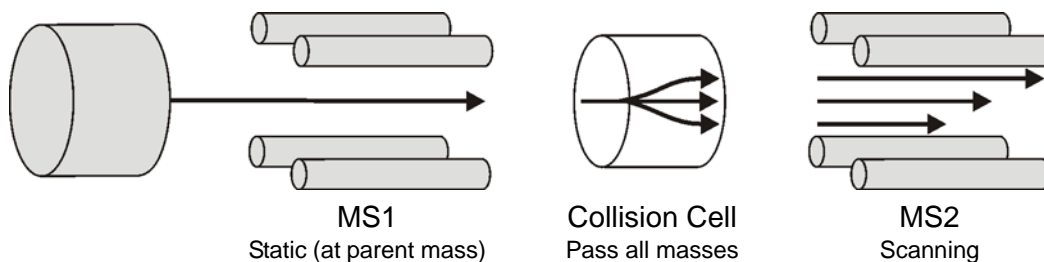


Figure F-2 Daughter (Product) Ion Mode

Typical Applications

- Structural elucidation (for example, peptide sequencing)
- Method development for MRM screening studies:
 - Identification of daughter ions for use in MRM transitions.

- Optimization of CID tuning conditions to maximize the yield of a specific daughter ion to be used in MRM analysis.

Figure F-3 shows an example of daughters of the specific parent at m/z 609 from reserpine in Electrospray positive ion mode.

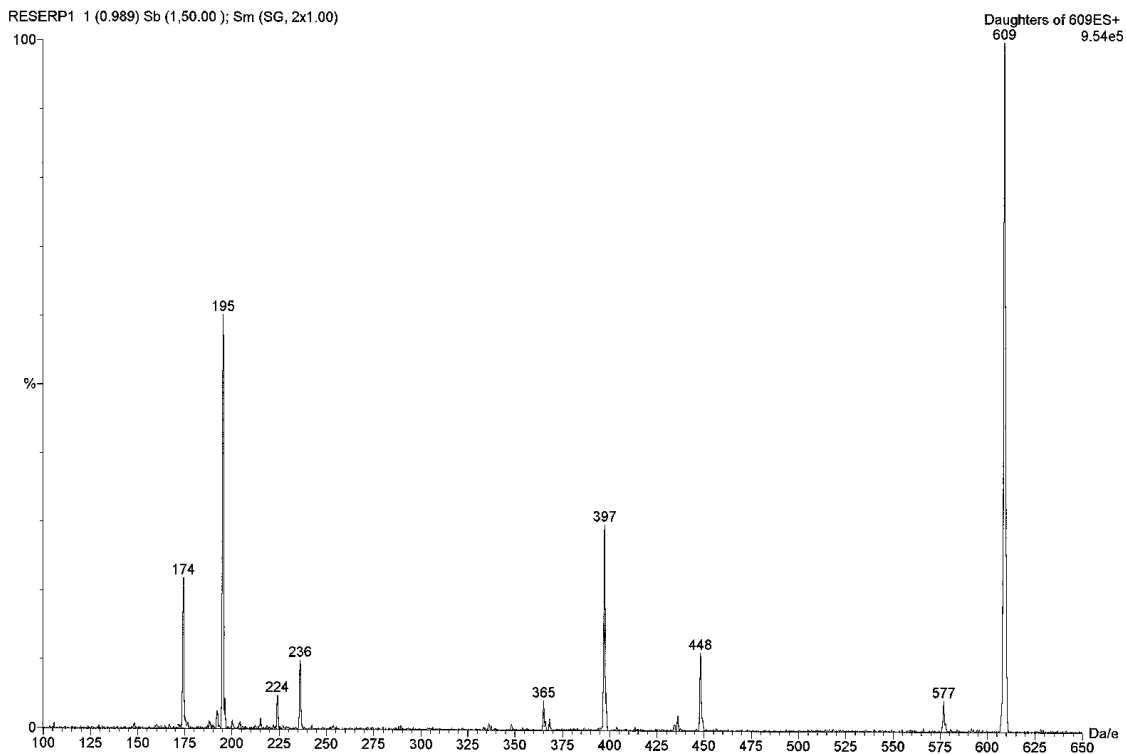


Figure F-3 Daughters of the Specific Parent at m/z 609 from Reserpine in Electrospray Positive Ion Mode

F.4.2 Parent (Precursor) Ion Mode

The parent (precursor) ion mode is shown in [Figure F-4](#).

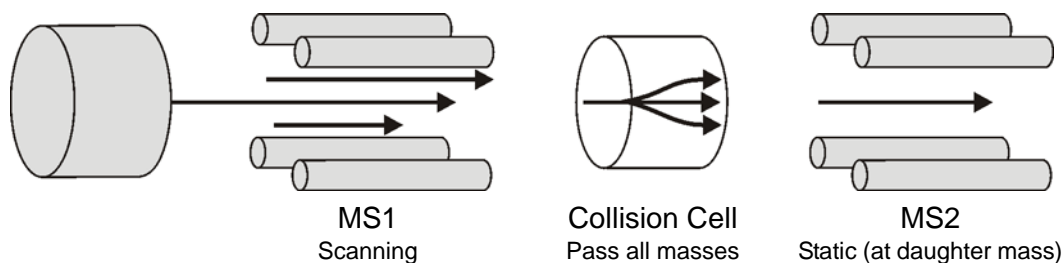


Figure F-4 Parent (Precursor) Ion Mode

A typical application is for structural elucidation, that is, complementary or confirmatory information (for daughter scan data).

[Figure F-5](#) shows an example of parents of the specific daughter ion at m/z 195 from reserpine in electrospray positive ion mode.

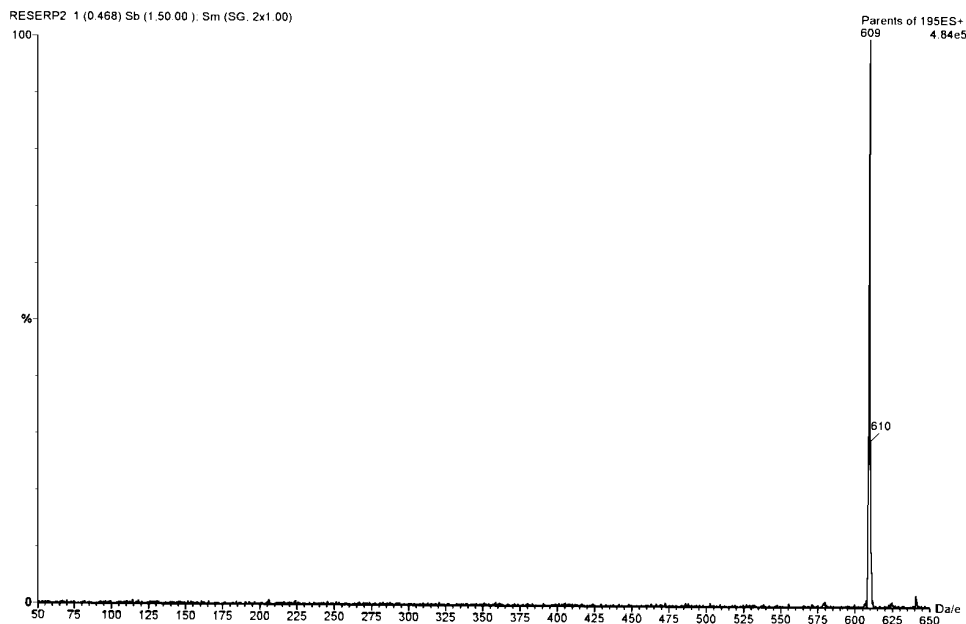


Figure F-5 Parents of the Specific Daughter Ion at m/z 195 from Reserpine in Electrospray Positive Ion Mode

F.4.3 Multiple Reaction Monitoring (MRM) Mode

The MRM mode (Figure F-6) is a highly selective MS/MS equivalent of SIR. As both MS1 and MS2 are static, greater dwell time on the ions of interest is allowed, and therefore better sensitivity compared to scanning MS/MS.

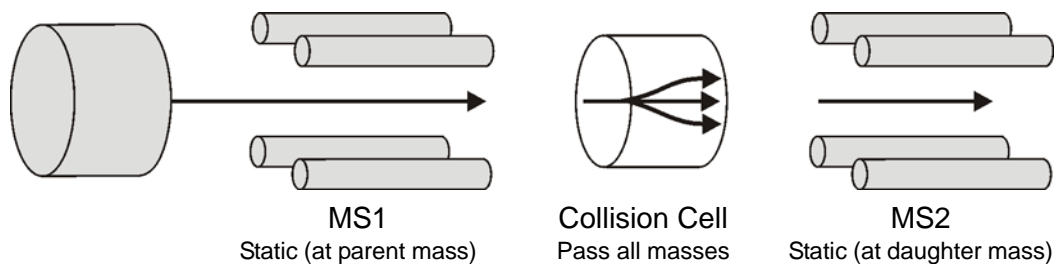


Figure F-6 MRM Mode

A typical application is for quantification of known analytes in complex samples:

- Drug metabolite and pharmacokinetic studies.
- Environmental, for example, pesticide and herbicide analysis.
- Forensic or toxicology, for example, screening for target drugs in sport.
- MRM does not produce a spectrum as only one transition is monitored. As in SIR, a chromatogram is produced.

F.4.4 Constant Neutral Loss Mode

The constant neutral loss mode is shown in [Figure F-7](#). It detects the loss of a specific neutral fragment or functional group from an unspecified parent or parents.

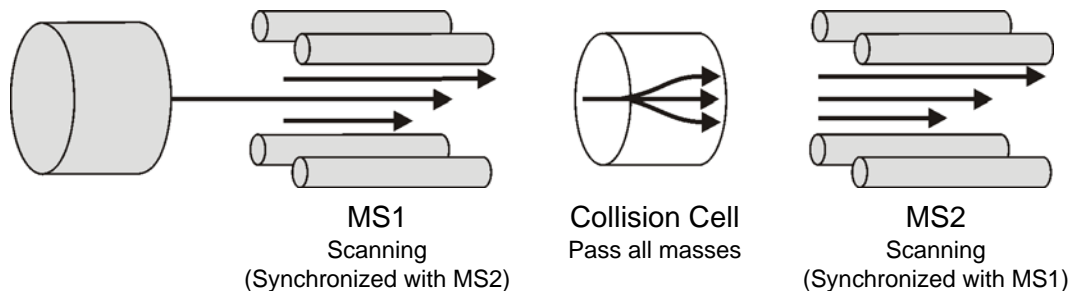


Figure F-7 Constant Neutral Loss Mode

A typical application is for screening mixtures for a specific class of compound that is characterized by a common fragmentation pathway, indicating the presence of compounds containing a common functional group.

The scans of MS1 and MS2 are synchronized. When MS1 transmits a specific parent ion, MS2 “looks” to see if that parent loses a fragment of a certain mass. If it does, it registers at the detector.

The result is that the spectrum shows the masses of all parents that actually lost a fragment of a certain mass.

F.4.5 Source and Collision Cell T-Wave Devices

The T-Wave devices are stacked ring electrode ion guides with opposite phases of radio frequency voltage applied to adjacent plates to confine ions radially (Figure F-8).

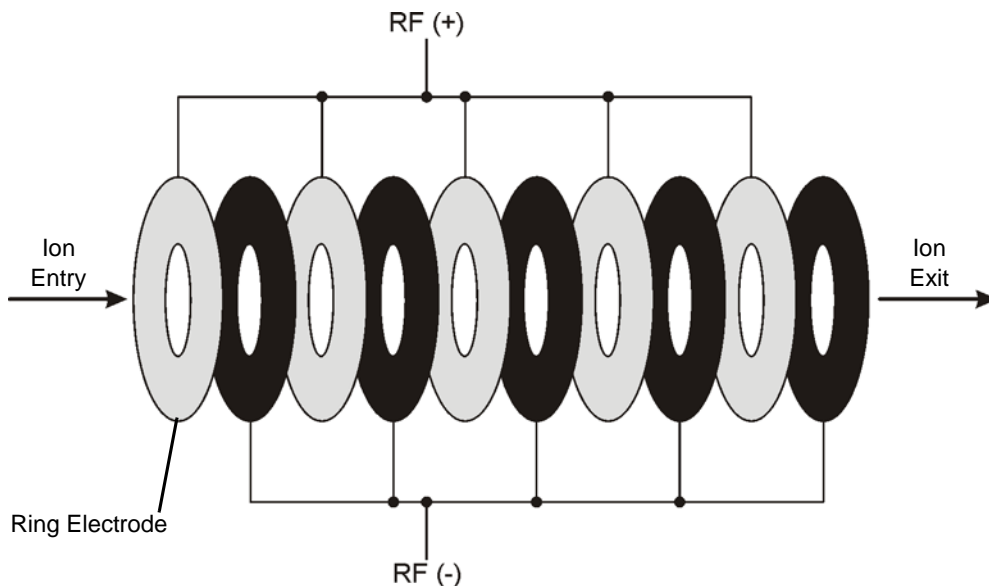


Figure F-8 T-Wave Schematic Diagram

The presence of gas in these devices causes an axial slowing of ions, which can have adverse effects for fast acquisitions such as reduced sensitivity resolution and increased cross-talk. To reduce the residence time of ions in these optics, a travelling voltage wave (T-Wave) moves along the device by application of a transient d.c. voltage to successive ring electrodes. Through appropriate choice of wave amplitude and velocity, ions “surf” on the front of this wave, reducing their transit time (Figure F-9).

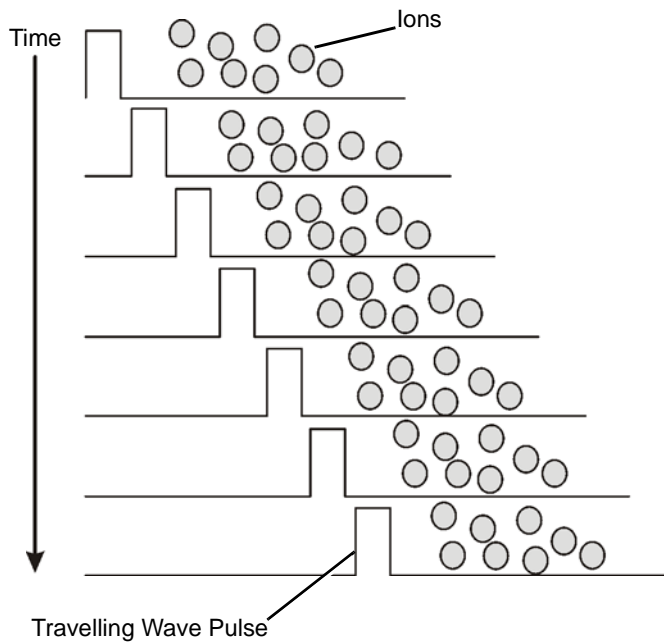


Figure F-9 Travelling Wave Pulse

Under normal operating conditions, the wave parameters are fixed by MassLynx to give optimal performance. However, under certain operating conditions, manual control may be necessary (see [Section C.19](#)).

Index

A

- a.c. power connection [14](#)
- acquisition parameters [124](#)
- Acquisition Setup dialog box [57](#)
- adduct list
 - creating [109](#)
 - opening [111](#)
 - saving [110](#)
- analog data [69](#)
 - acquiring [75](#)
- Analog Data dialog box [76](#)
- Analogue Channel [10](#)
- APCI [250](#)
- APCI probe [3](#)
 - heater
 - not working [209](#)
 - replacing [197](#)
 - installing [38](#)
 - removing [17](#)
 - stainless steel capillary, replacing [187](#)
 - tip, cleaning [173](#)
- APPI electrical connection [7](#)
- assembling
 - source ion block [168](#)
 - source T-Wave assembly [169](#)
- atmospheric pressure chemical
 - ionization [250](#)
- automated quantification [63](#)
- Automatic Calibration Check dialog
 - box [119](#)
- automatic calibration check parameters [119](#)
- Automatic Calibration dialog box [123](#)
- Aux O/P connection [12](#)

B

- backing pump [3](#)
- baseline, resetting [228](#)

C

- calibration
 - checking [129](#)
 - failure [127](#)
 - peak matching, manually editing [131](#)
 - performing [123](#)
 - removing [118](#)
 - report [129](#)
 - saving [132](#)
 - scan speed compensation [123](#)
 - scanning [123](#)
 - selecting the reference file [118](#)
 - setting general parameters [120](#)
 - setting parameters [119](#)
 - starting [125](#)
 - static [123](#)
 - verifying [132](#)
- Calibration Acquisition Setup dialog
 - box [124](#)
- Calibration Options window [126](#)
- Calibration Parameters dialog box [120](#)
- calibration reference
 - files
 - editing [241](#)
 - negative ion [244](#)
 - positive ion [242](#)
 - information [241](#)
- calibration report [126, 129](#)
- CE Int connection [12](#)
- CE Profile dialog box [105](#)

- centroid data [68, 80](#)
- Charge State Mass dialog box [107](#)
- chemical noise [213](#)
- chromatogram, viewing in real-time [65](#)
- cleaning
 - APCI probe tip [173](#)
 - cone gas cone [165](#)
 - corona discharge pin [172](#)
 - ESI probe tip [171](#)
 - extraction cone [167](#)
 - gas exhaust port [165](#)
 - ion block [167](#)
 - isolation valve stem [168](#)
 - sample cone [165](#)
 - source T-Wave assembly [166](#)
- collision energy
 - options [102](#)
 - profile table [105](#)
 - adding new entries [107](#)
 - opening [108](#)
 - saving [108](#)
 - ramp
 - applying [86](#)
 - controlling [228](#)
 - setting-up [227](#)
- Collision Gas In connection [14](#)
- collision gas, controlling [225](#)
- Collision Ramp dialog box [86, 227](#)
- Com1 connection [12](#)
- Comm For EPC connection [12](#)
- communications status [70](#)
- complete shutdown [218](#)
- cone gas
 - connection [6](#)
 - controlling [225](#)
- cone gas cone, cleaning [165](#)
- Cone Ramp dialog box [79, 226](#)
- cone voltage ramp
 - applying [79](#)

- controlling [227](#)
 - setting-up [226](#)
- connections [10](#)
 - front panel [4](#)
 - rear panel [10](#)
 - vacuum [14](#)
- constant neutral loss mode [256](#)
- contact closure connections [11](#)
- contacting Waters [214](#)
- continuum data [80](#)
- controlling
 - collision energy ramp [228](#)
 - cone gas [225](#)
 - cone voltage ramp [227](#)
 - desolvation gas [225](#)
 - gas flows [225](#)
 - nebulizer gas [225](#)
 - syringe pump [225](#)
- controls
 - embedded PC reset switch [4](#)
 - front panel [4](#)
 - Inject switch [9](#)
 - Load switch [9](#)
 - power switch [4](#)
- corona discharge pin
 - cleaning [172](#)
 - installing [37](#)
 - removing [19](#)
 - replacing [172](#)

D

- data
 - analog [69](#)
 - centroid [68, 80](#)
 - continuum [80](#)
 - MCA [80](#)
 - profile [67](#)
 - SIR [68](#)

- data acquisition
 - monitoring 65
 - pausing 70
 - starting 57
 - stopping 70
- daughter (product) ion mode 252
- Daughter Scan Function Editor 85
- desolvation gas
 - connection 6
 - controlling 225
- desolvation heater not working 209
- dialog boxes
 - Acquisition Setup 57
 - Analog Data 76
 - Automatic Calibration 123
 - Automatic Calibration Check 119
 - Calibration Acquisition Setup 124
 - Calibration Parameters 120
 - CE Profile 105
 - Charge State Mass 107
 - Collision Ramp 86, 227
 - Cone Ramp 79, 226
 - Display Calibration Graphs 129
 - Exclude Mass 101
 - Include Masses 97
 - Instrument Threshold Settings 66
 - Mass Measure Parameters 121
 - Modify Charge State 106
 - Quantify Samples 63
 - Readbacks 228
 - Scope Setup 226
 - Solvent Delay 74
 - Spectrum Real-Time Update 65
 - Start Sample List Run 60
 - Syringe Selection 225
- disassembling
 - source ion block 157
 - source T-Wave assembly 155
- Display Calibration Graphs dialog box 129

divert/injection valve 8

E

- electrical connections
 - a.c. power 14
 - Analogue Channel 10
 - Aux O/P 12
 - CE Int 12
 - Com1 12
 - Comm For EPC 12
 - contact closure 11
 - Events I/P 11
 - Events O/P 11
 - Gas Fail 11
 - HV 7
 - mains power 14
 - MUX 7
 - Probe 7
 - Pump Relay 12
- electrical connections APPI 7
- electronic noise 213
- electrospray ionization 249
- embedded PC reset switch 4
- emergency shutdown 218
- ESCi multi-mode operation 3, 53
 - data acquisition 55
 - option key disk 53
 - preparing the instrument 53
 - tuning the instrument 54
- ESI 249
- ESI probe 3
 - installing 20
 - removing 35
 - stainless steel capillary, replacing 178
 - tip
 - cleaning 171
 - replacing 171
- Events I/P connections 11

- Events O/P connections 11
- Exclude List 99
 - editing 101
- Exclude Mass dialog box 101
- Exhaust connection 14
- extraction cone, cleaning 167

F

- filter elements
 - changing 139
- fitting
 - gas exhaust port 170
 - ion block 170
 - ion source enclosure 170
 - source T-Wave assembly 169
- front panel
 - connections 4
 - controls 4
- Function Editors
 - Daughter Scan 85
 - Full Scan 77
 - MRM 88
 - Parent Scan 87
 - SIR 81
 - Survey Scan 88
- function list
 - adding a new function 73
 - changing the order of functions in 74
 - copying an existing function 74
 - modifying an existing function 73
 - opening a saved list 77
 - removing a function 74
 - saving 76
 - setting a solvent delay 74
 - setting the maximum retention time 74
 - setting-up 71
- functions
 - MRM 88

- SIR 81
- survey 88

- fuses
 - main system printed circuit board 209
- mains power 14

G

- gas connections
 - Collision Gas In 14
 - cone gas 6
 - desolvation gas 6
 - Exhaust 14
 - Gas In 13
 - nebulizer gas 6
- gas exhaust port
 - cleaning 165
 - fitting 170
 - removing 150
- Gas Fail connection 11
- gas flow rates 15
- gas flows, controlling 225
- Gas In connection 13
- gas-ballasting, rotary pump 135

H

- high LC backpressure 206
- high noise levels 212
- HV electrical connection 7

I

- IEEE communication errors 212
- Include List 95
 - editing 97
- Include Masses dialog box 97
- Inject switch 9
- Inlet Configuration Wizard 33

- installing
 - APCI probe 38
 - corona discharge pin 37
 - ESI probe 20
- instrument data thresholds 66
- instrument name, selecting 237
- instrument parameter files 229
 - creating new 230
 - opening 230
- Instrument Threshold Settings dialog box 66
- insufficient vacuum 207
- intended use *xiii*
- inter scan delay, selecting 226
- ion block
 - cartridge heater, replacing 174
 - cleaning 167
 - fitting 170
 - removing 151
- ion counting threshold 68
- Ion Mode menu 223
- ion modes
 - daughter 252
 - parent 254
 - precursor 254
 - product 252
- ion optics 250
- ion source enclosure
 - fitting 170
 - removing 151
- ion source heater, not working 209
- ionization mode, selecting 223
- isolation valve 3, 23, 41, 146
 - stem, cleaning 168

K

- key disk, ESCi option 53

L

- LEDs
 - Operate 7
 - Vacuum 7
- Load switch 9
- loss of communication with instrument 211
- low intensity peaks 205
- low LC backpressure 207

M

- mains power
 - connection 14
 - fuses 14
- maintenance
 - procedures 133
 - safety 134
 - schedule 133
- mass calibration 113
 - for electrospray 114
 - scan speed 113
 - scanning 113
 - static 113
- mass flow controllers 15
- mass measure parameters 121
- Mass Measure Parameters dialog box 121
- MassLynx
 - ESCi option key disk 53
 - software 4
 - Tune window, opening 223
- maximum retention time 74
- MCA data 80
- Modify Charge State dialog box 106
- monitoring acquisitions 111
- MRM function 88
- MRM Function Editor 88
- MS Method Editor 71
 - toolbar 72

- MS operating modes 251
- MS to MS/MS switching 90
- MS/MS operating modes 252
- MS/MS scanning functions 84
- MS/MS to MS switching 93
- MS2 scan 87
- Multi Channel Analysis data 80
- multiple reaction monitoring mode 255
- multiple-sample acquisition 60
- MUX electrical connection 7

N

- nebulizer gas
 - connection 6
 - controlling 225
- neutral gain scan 87
- neutral loss scan 87
- no ion beam 203
- no peaks in the Tune window 203
- noise
 - chemical 212
 - electronic 212

O

- odor filter element, changing 139
- oil mist filter element, changing 139
- Operate LED 7
- operating modes
 - MS 251
 - MS/MS 252
- overnight shutdown 218

P

- parameters
 - acquisition 124
 - mass measure 121

- setting calibration 119
- setting general calibration 120
- parameters automatic calibration check 119
- parent (precursor) ion mode 254
- Parent Scan Function Editor 87
- Peak Display menu 234
- peak display, customizing
 - colors 235
 - display grid 236
 - intensity display 236
 - numbers 235
 - trace line appearance 236
- peaks
 - changing gain 234
 - low intensity 205
 - selecting
 - for tuning 231
 - operating mode 232
 - span 233
 - tune mass 232
 - unsteady 205
- performance specifications 247
- Pirani gauge 3
- power switch 4
- printing tune information 230
- probe
 - APCI 3
 - connections 144
 - ESI 3
 - removing 143
- Probe electrical connection 7
- profile data 67
 - removing spikes from 69
- pump
 - backing 3
 - turbomolecular 3
- Pump Relay connection 12

Q

Quantify Samples dialog box 63

R

Readbacks dialog box 228

real-time viewing, chromatogram 65

rear panel connections 10

reference file, selecting 118

reference sample solutions, preparing 245

removing

APCI probe 17

corona discharge pin 19

ESI probe 35

gas exhaust port 150

ion block 151

ion source enclosure 151

replacing

APCI probe heater 197

APCI probe stainless steel capillary 187

corona discharge pin 172

ESI probe stainless steel capillary 178

ESI probe tip 171

ion block cartridge heater 174

report, calibration 126

reset switch, embedded PC 4

rotary pump

fuse failure 209

gas-ballasting 135

oil

changing 138

checking 137

S

safety procedures, maintenance 134

sample cone

cleaning 165

fitting to instrument 170

removing 145

sample inlet 3

Sample List 60

Scan Report window 65

scan speed compensation calibration 123

scan time, selecting 226

scanning calibration 123

scans

MS2 87

neutral gain 87

neutral loss 87

Scope Setup dialog box 226

shutdown

complete 218

emergency 218

overnight 218

shutting down the instrument 218

SIR data 68

SIR function 81

SIR Function Editor 81

adding a function 83

Solvent Delay dialog box 74

source ion block

assembling 168

disassembling 157

source T-Wave assembly

assembling 169

cleaning 166

disassembling 155

fitting 169

removing 153

source, cleaning 142

Spectrum Real-Time Update dialog box 65

spectrum, viewing in real-time 65

Start Sample List Run dialog box 60

starting up the instrument 215

static calibration 123

status display LEDs 7

- survey function 88
- Survey Scan Function Editor 88
 - Adduct page 109
 - Collision Energy page 102
 - Exclude page 99
 - Include page 95
 - MS to MSMS page 90
 - MSMS Template page 89
 - Survey page 89
- switches
 - Inject 9
 - Load 9
- switching
 - MS to MS/MS 90
 - MS/MS to MS 93
- syringe pump
 - controlling 225
 - setting-up 221
- Syringe Selection dialog box 225
- syringe type, selecting 225
- System Manager window 70

T

- thresholds
 - instrument data 66
 - ion counting 68
- troubleshooting 201
 - APCI probe heater not working 209
 - chemical noise 213
 - desolvation heater not working 209
 - electronic noise 213
 - failure to recognize a probe 210
 - hardware 203
 - high LC backpressure 206
 - high noise levels 212
 - IEEE communication errors 212
 - instrument will not turn on 203
 - insufficient vacuum 207

- ion mode fault 210
- ion source heater not working 209
- isolation valve 204
- leaking nitrogen 208
- loss of communication with
 - instrument 211
- low LC backpressure 207
- no ion beam 203
- no peaks in the Tune window 203
- ripple 211
- rotary pump fuse failure 209
- rotary pump oil in exhaust tubing 208
- safety 201
- system 201
- unsteady or low intensity peaks 205
- tune information, printing 230
- tune parameters
 - changing 229
 - saving 229
- Tune window
 - Diagnostics page 237
 - EasyTune Source page 231
 - ESCi source page 54
 - T-WAVE page 238
- turbomolecular pump 3
- T-Wave
 - optics, manually controlling 237
 - theory 257

U

- unsteady peaks 205

V

- vacuum
 - connections 14
 - system 3
- Vacuum LED 7

W

Waste connection [14](#)

Waters, contacting [214](#)

Z

zero level, resetting [228](#)

ZSpray ionization source [3](#)

