

Q Exactive™ QuickStart Guide

About This Guide

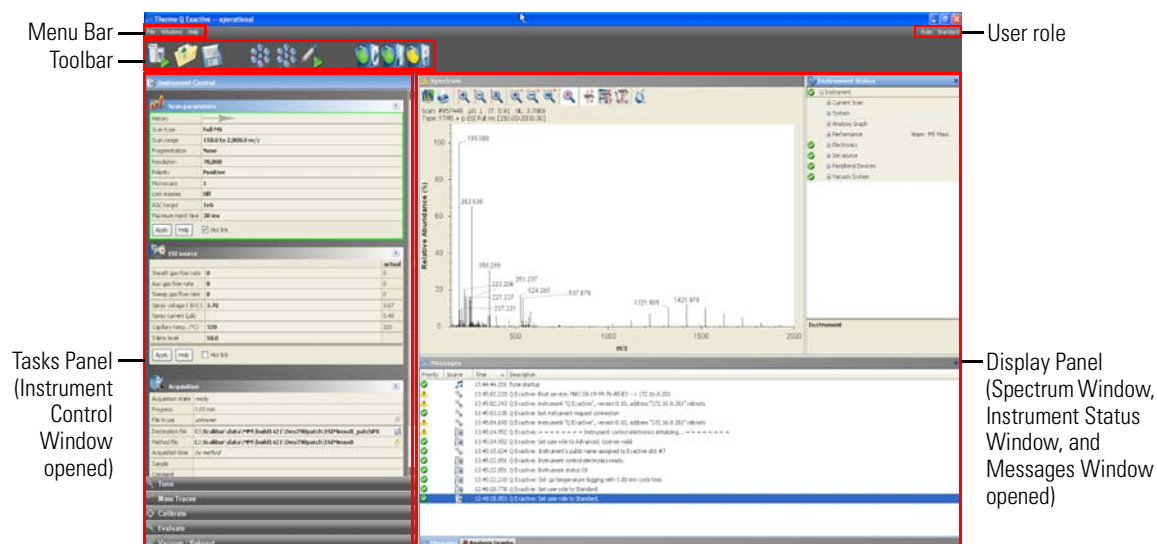
This *Q Exactive QuickStart Guide* gives an introduction on setting up and using the Thermo Scientific Q Exactive mass spectrometer. This guide is intended for all personnel who operate the Q Exactive mass spectrometer.

Please note that this guide does not contain any safety notices and precautionary statements. Thus, it is mandatory to read also the *Q Exactive Operating Manual* and the *Ion Max and Ion Max-S API Source Hardware Manual*.

Q Exactive Tune

Use Q Exactive Tune to operate the Q Exactive mass spectrometer via the data system computer. It has the following components:

- The *menu bar* provides drop-down menus with commands for operating the program. On its right side it shows the active user role.
- The *toolbar* provides symbol shortcuts for frequently used commands.
- The *tasks panel* provides six windows: Instrument Control, Tune, Mass Traces, Calibrate, Evaluate, and Vacuum/Bakeout. The *instrument control window* is divided into three additional windows for defining scan parameters, setting API source parameters, and acquiring and storing measurement data.
- The *display panel* provides various windows for showing real-time information about the current measurement and the instrument status.



❖ To display the Q Exactive Tune program

Choose **Start > Programs > Thermo Exactive Series > Tune**.

Before going on, check that the following preconditions are fulfilled:

- The main power circuit breaker switch is in the On (I) position and the electronics service switch is in the Operating Mode position, the gas pressure is within the operational limit, and the vacuum levels are sufficient for operating the instrument.
- The Ion Max API source with H-ESI probe is installed on the mass spectrometer. You have checked the proper position of the H-ESI probe in the Ion Max API source housing.
- Power supply and communication between syringe pump and mass spectrometer are established. A clean, 500 µL Unimetrics syringe with 450 µL of calibration solution for positive ion mode is placed into the syringe pump. An infusion line is installed between the syringe pump and the grounding union that is held by the grounding bar of the Ion Max API source.

Preparation





Getting Ions from Infusion Experiments

Setting up Q Exactive Tune



Refer to the *Q Exactive Operating Manual* for instructions about preparing the positive ion mode calibration solution or buying the ready-to-use solution from Thermo Fisher Scientific (product number 88323).

- The data system computer with the Q Exactive Tune software is switched on and the mass spectrometer is placed in Standby mode.

For information about performing these tasks, refer to the *Q Exactive Operating Manual*, the *Q Exactive Software Manual* or the Q Exactive Tune Help, and the *Ion Max and Ion Max-S API Source Hardware Manual*.

This topic describes how to test the instrument before you can tune and calibrate your Q Exactive mass spectrometer automatically.

❖ To set up the mass spectrometer in the Q Exactive Tune software for tuning and calibration in the H-ESI/MS mode

1. If you have not already done so, choose **Start > Programs > Thermo Exactive Series> Tune** to start the Q Exactive Tune program.
2. In the Q Exactive Tune window, on the toolbar, click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On. When you turn the mass spectrometer to On, you initiate the following events:
 - The mass spectrometer begins scanning.
 - Nitrogen flows into the H-ESI probe.
 - The Q Exactive mass spectrometer applies high voltage to the H-ESI probe.
 - The spectrum window of Q Exactive Tune shows a real-time display of the mass spectrum.
3. If necessary, open an existing tune file:
 - a. Choose **File > Load Tune File** to display the Open dialog box.
 - b. Browse for the folder C:\Xcalibur\methods\. Select a tune file, for example *ESI_installation.mstune*.
 - c. Click **Open** to open the file. Q Exactive Tune downloads the Tune Method parameters to the mass spectrometer.
4. Examine the pre-tune H-ESI source settings:
 - a. From the Instrument Control panel, open the H-ESI source window. See Figure right.
 - b. Set the *sheath gas flow rate* to 12.
 - c. Leave the auxiliary gas flow rate at 0.
 - d. Leave the sweep gas flow rate at 0.
 - e. Set the *spray voltage* to 4 kV.
 - f. Set the *capillary temperature* to 320 °C.
 - g. Leave the *S-lens level* at 50.
 - h. Set the *heater temperature* to 0 °C.
 - i. If you have made any changes in the window, click **Apply**.

Note Select the Hot link check box to apply all changes in real time. A green frame around the parameter box indicates an active hot link. ▲

		actual
Sheath gas flow rate	12	
Aux gas flow rate	0	
Sweep gas flow rate	0	
Spray voltage (kV)	4.00	
Spray current (µA)		
Capillary temp. (°C)	320	
S-lens level	50.0	
Heater temp (°C)	0	

Apply Help Hot link

Testing the Mass Spectrometer



5. Set the scan parameters for tuning and calibration:
 - a. From the Instrument Control panel, open the Scan parameters window. See Figure right.
 - b. Set *Scan type* to *Full MS*.
 - c. Enter a *scan range* between *m/z 150 and 2000*.
 - d. Set *Fragmentation* to *None* to specify that the ion source fragmentation option is turned off.
 - e. Select *17500* for *Resolution*.
 - f. Set the *ion polarity mode* to *Positive*.
 - g. Set the total number of *microscans* to *1*.
 - h. Set *Lock masses* to *Off*.
 - i. Select *1e6* for *AGC target*.
 - j. Specify a *maximum injection time* of *30 ms*.
 - k. If you have made any changes in the window, click **Apply** or select the Hot link check box.

Scan parameters	
History	→
Scan type	Full MS
Scan range	150.0 to 2,000.0 m/z
Fragmentation	None
Resolution	17,500
Polarity	Positive
Microscans	10
Lock masses	Off
AGC target	1e6
Maximum inject time	30 ms
<input type="button" value="Apply"/> <input type="button" value="Help"/> <input type="checkbox"/> Hot link	

The mass spectrometer is now properly set up for tuning and calibration in the H-ESI/MS mode.

In this section, you can test whether the mass spectrometer is operating properly. Infuse a low concentration calibration solution for positive ion mode—containing n-butylamine, caffeine, MRFA, and Ultramark 1621—into the H-ESI source using the syringe pump. Monitor the real-time display of the mass spectrum to ensure that a stable spray of solution enters the mass spectrometer.

Refer to the *Q Exactive Operating Manual* for instructions about preparing the positive ion mode calibration solution or buying the ready-to-use solution from Thermo Fisher Scientific (product number 88323).

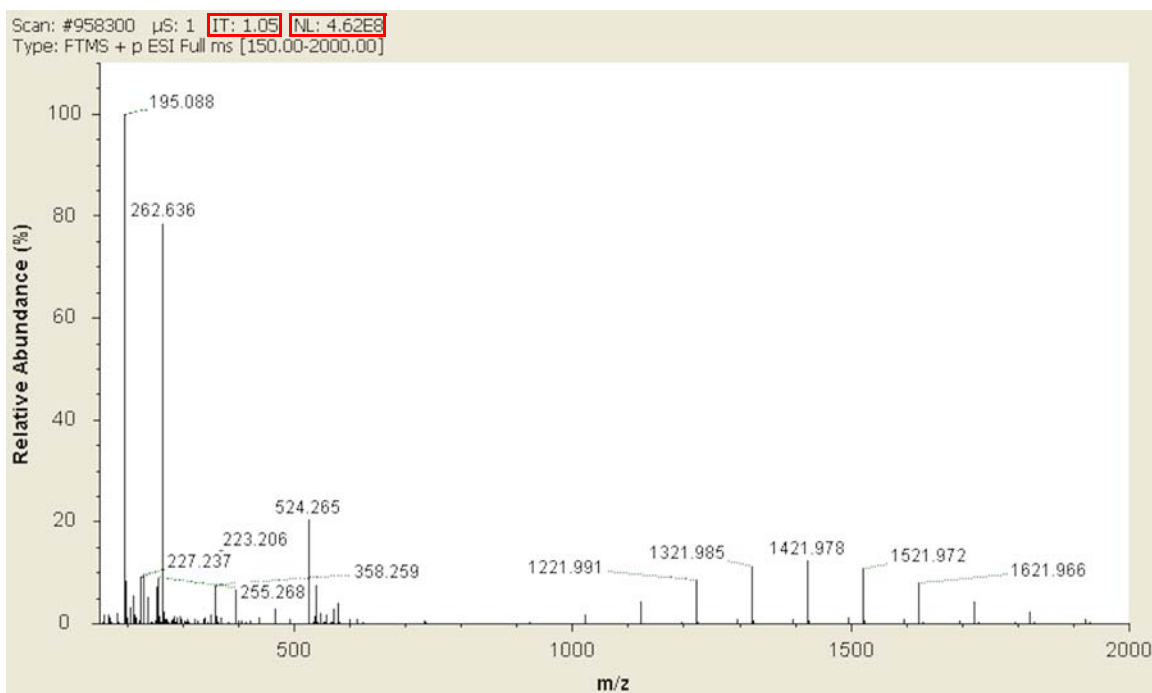
❖ To test the operation of the mass spectrometer in H-ESI/MS mode

1. Click the arrow of the **Syringe Pump** button to display the Syringe Pump dialog box. See Figure right.
2. Turn on the syringe pump and set an infusion flow rate of 5 $\mu\text{L}/\text{min}$, as follows:
 - a. Select *Unimetrics* to specify the proper *syringe type*.
 - b. Select *500* in the *Volume* list box to set the proper syringe volume. Note that the Q Exactive mass spectrometer automatically sets the syringe ID to its proper value of 3.257 mm.
 - c. Specify a *flow rate* of *5 $\mu\text{L}/\text{min}$* .
 - d. Click **Apply** to apply the syringe parameters.
 - e. Click **Start** to start the syringe pump.

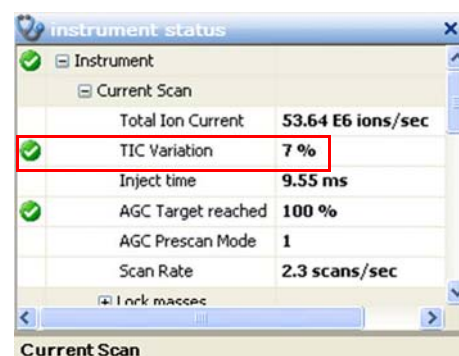
Syringe type	Unimetrics
Volume (μL)	500
Syringe inner diameter (mm)	3.257
Flow rate ($\mu\text{L} / \text{min}$)	5.000
<input type="button" value="Apply"/> <input type="button" value="Help"/>	
<input type="button" value="Manual control"/> <input type="button" value="Start"/> <input type="button" value="Stop"/>	

Note Once you have set the syringe parameters, just click the **Syringe Pump** button to switch on or off the syringe pump. ▲

3. Test the efficiency and stability of the spray of calibration solution into the mass spectrometer. Monitor the data for the calibration solution, as follows:
 - a. In the spectrum window of Q Exactive Tune, observe the mass spectra of the singly-charged ions of the positive ion mode calibration solution. See Figure below. The ions are as follows:
 - n-Butylamine: *m/z* 74
 - Caffeine: *m/z* 138 (fragment), 195
 - MRFA: *m/z* 524
 - Ultramark 1621: *m/z* 1022, 1122, 1222, 1322, 1422, 1522, 1622, 1722, 1822
 - b. At the top of the spectrum window, notice the values for the inject time (IT) and normalized target level (NL). See Figure below.



- c. Observe the values for NL and IT in the spectrum window. As calibration solution infuses, and the readback values fluctuate, check the status of the ion current signal:
- Is the signal present?
 - Is the signal stable? Check in the instrument status window that the TIC variation has stabilized at about 15% or less from scan to scan (green check mark). See Figure right.



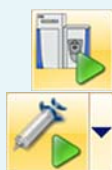
If you answered “yes” to the questions in [step 3c](#), then your mass spectrometer is operating properly.

If you answered “no” to either of these questions, try the following troubleshooting measures:

- Ensure that the H-ESI probe is properly assembled and that the needle position is correct.
- Ensure that the entrance to the ion transfer capillary is clean.
- Ensure that the solution entering the probe is free of air bubbles and that the tubing and connectors are free of leaks and that they are not clogged.
- Ensure that spray voltage is appropriate.
- Ensure that discharge does not take place.

Your mass spectrometer is operating properly in the H-ESI mode. You are now ready to tune and calibrate the mass spectrometer.

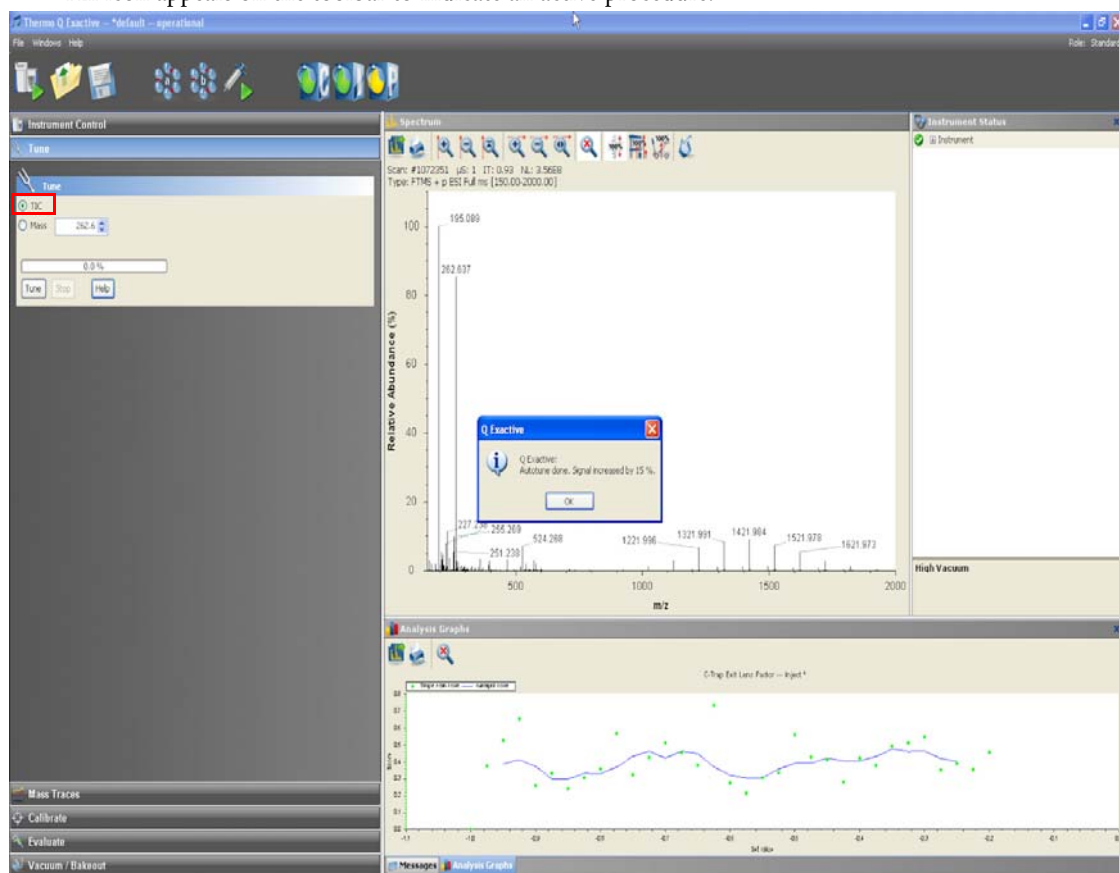
Tuning



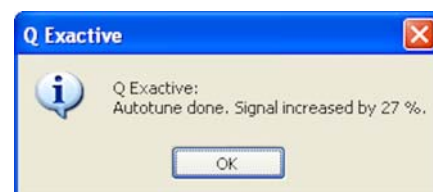
❖ To tune the Q Exactive mass spectrometer

1. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
2. Click the **Syringe Pump** button to start the syringe pump.
3. In the Tune window, select the TIC option button to specify that the Q Exactive mass spectrometer optimizes on the total ion current (TIC) intensity. See Figure on next page.

4. Ensure that *NCE fragmentation is not activated* in the Fragmentation dialog box. (See [page 10.](#)) Otherwise, the tuning procedure comprises only HCD relevant parameters.
5. Click **Tune** to start the automatic tuning procedure:
 - The Tune window indicates the completed percentage of the tuning process by a progress bar.
 - A message box informs you about the active tuning procedure and the currently tuned parameter.
 - The analysis graphs window displays a graphic representation of the tuning process for the currently tuned parameter.
 - The messages window shows a record of the tuning steps.
 - An icon appears on the toolbar to indicate an active procedure.



6. A message box informs you that the tuning procedure is completed. The message box also contains information about the factor by which the signal was increased. See Figure right. Click **OK** to close the message box.
7. Choose **File > Save Tune File as** to save the adjusted parameters in a Tune Method:
 - a. In the displayed dialog box, enter a name for the new tune file.
 - b. Click on **Save** to save the new tune parameters and to close the dialog box.



The new tune file is specific to your particular analyte and solvent flow rate. (In this example, you save settings obtained using calibration solution.) You can call the Tune Method again and use it as a starting point for optimization of the mass spectrometer for a different analyte of interest or with a different flow rate.

After having tuned the Q Exactive mass spectrometer, it should be calibrated to ensure the mass accuracy of the results. Usually, it is sufficient to perform a mass calibration only. Before starting the calibration, check again that you have a stable spray: Focus on the injection time (IT) and on the normalized level (NL) in the header information on the spectrum window.

Note If the system was in Off mode before, it is necessary to put the instrument into On mode for at least 90 minutes before a mass calibration is performed. ▲

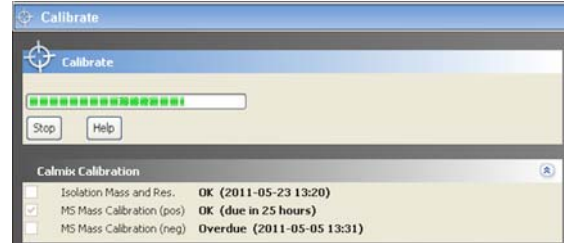
Mass Calibration

Positive Ion Mode Calibration



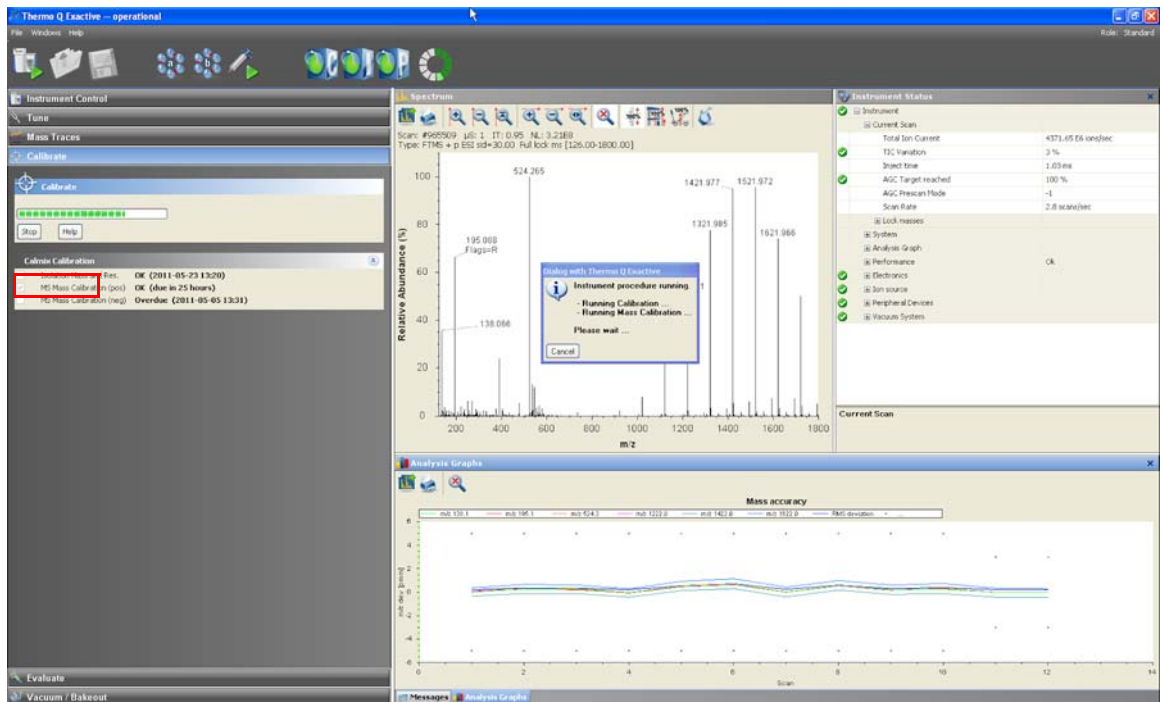
❖ To calibrate the Q Exactive mass spectrometer for positive ion mode

1. Place a clean, 500 μ L Unimetrics syringe with 450 μ L of calibration solution for the positive ion mode into the syringe pump.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
3. Click the **Syringe Pump** button to start the syringe pump.
4. Click the title bar of the Calibrate window to display it.
5. In the Calmix Calibration sub-window, select the MS Mass Calibration (pos) check box to specify a mass calibration for the positive ion mode. See Figure right.
6. In the Calibrate window, click **Calibrate** to start the automatic calibration procedure:

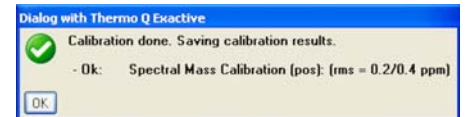


- The Calibrate window indicates the completed percentage of the calibration process by a progress bar.
- Q Exactive Tune displays a message box that informs about the active calibration procedure and the currently calibrated parameter.
- The analysis graphs window displays a graphic representation of the calibration process for the currently tuned parameter.
- The messages window shows a record of the calibration.

7. An icon appears on the toolbar to indicate an active procedure.



8. A message box appears when the calibration procedure is completed. See Figure right. The message box also informs you that Q Exactive Tune has automatically saved the calibration results. Click **OK** to close the message box.



Before you can start measuring samples, you must ensure that the quadrupole isolation performance for the narrow isolation window is sufficient. Therefore, evaluate the isolation calibration for the positive ion mode as described on [page 8](#).

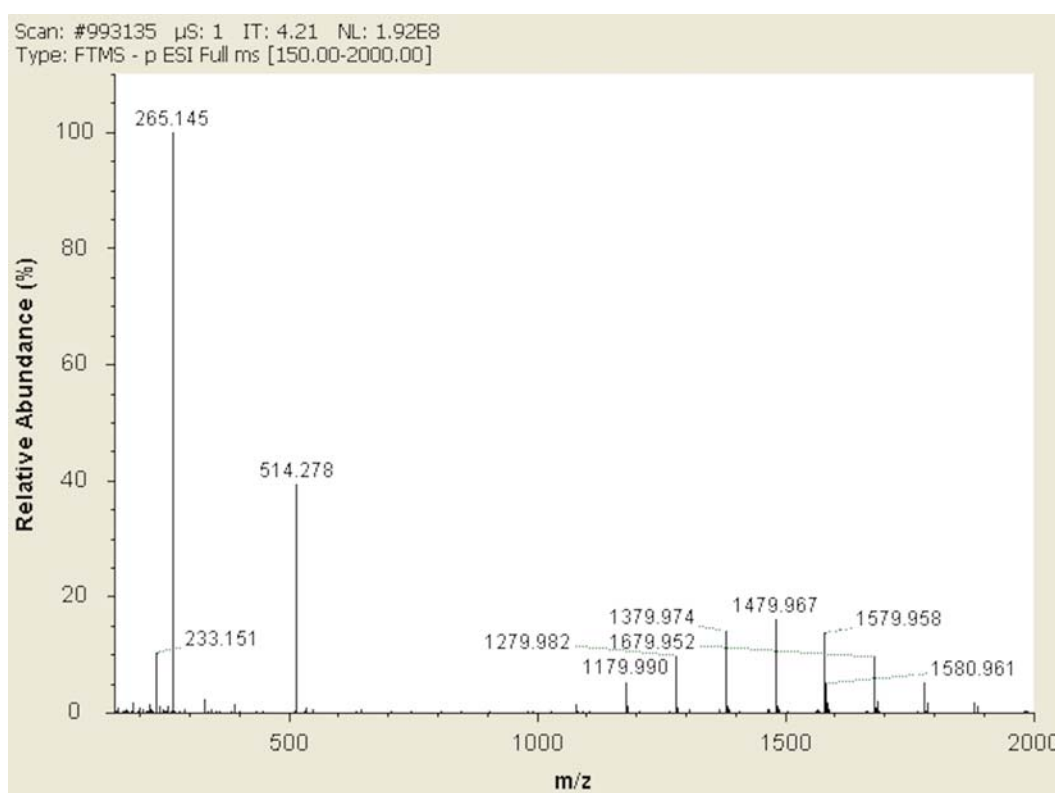
Negative Ion Mode Calibration



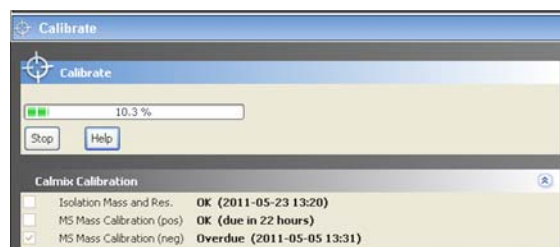
Before performing measurements in the negative ion mode, the mass spectrometer must be calibrated with a calibration solution containing sodium dodecyl sulfate, sodium taurocholate, and Ultramark 1621. Refer to the *Q Exactive Operating Manual* for instructions about preparing the negative ion mode calibration solution or buying the ready-to-use solution from Thermo Fisher Scientific (product number 88324).

❖ To calibrate the Q Exactive mass spectrometer for negative ion mode

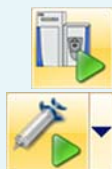
1. Place a clean, 500 μL Unimetrics syringe with 450 μL of calibration solution for the negative ion mode into the syringe pump.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
3. Click the **Syringe Pump** button to start the syringe pump.
4. In the spectrum window of Q Exactive Tune, observe the mass spectra of the singly-charged ions of the negative ion mode calibration solution. See Figure below. The ions are as follows:
 - Sodium dodecyl sulfate: m/z 265
 - Sodium taurocholate: m/z 514
 - Ultramark 1621: m/z 1280, 1380, 1480, 1580, 1680, 1780



5. Click the title bar of the Calibrate window to display it.
6. In the Calmix Calibration sub-window, select the MS Mass Calibration (neg) check box to specify a mass calibration for the negative ion mode.
7. In the Calibrate window, click **Calibrate** to start the automatic calibration procedure.
8. The Calibrate window indicates the completed percentage of the calibration process by a progress bar. See Figure right.
9. A message box informs you when the calibration procedure is completed. Click **OK** to close the message box.



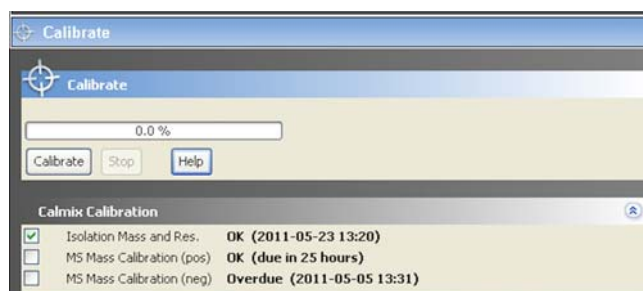
Isolation Mass and Resolution Calibration



If the evaluation of the quadrupole isolation as described on [page 8](#) fails you have to calibrate isolation mass and resolution of the quadrupole. This ensures that the quadrupole isolation performance for the narrow isolation window is sufficient for measurement. The following example describes this calibration for the positive ion mode.

❖ To calibrate isolation mass and resolution of the quadrupole

1. Place a clean, 500 μ L Unimetrics syringe with 450 μ L of calibration solution for the positive ion mode into the syringe pump.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
3. Click the **Syringe Pump** button to start the syringe pump.
4. Click the title bar of the Calibrate window to display it.
5. In the Calmix Calibration sub-window, select the Isolation Mass and Res. check box. See Figure right.
6. In the Calibrate window, click **Calibrate** to start the automatic calibration procedure.
7. The Calibrate window indicates the completed percentage of the calibration process by a progress bar.
8. A message box informs you when the calibration procedure is completed. Click **OK** to close the message box.



The Q Exactive mass spectrometer is now ready to perform your measurements.

Evaluation

Evaluation of Positive Mass Calibration

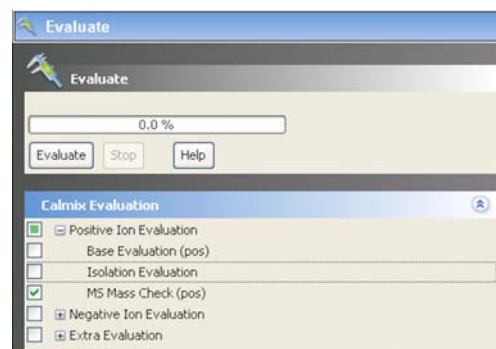


Use the Evaluate window of the Tasks panel to perform an automatic check of the instrument calibration.

The following example describes an evaluation of the positive mass calibration.

❖ To evaluate the mass calibration for positive ion mode

1. Place a clean, 500 μ L Unimetrics syringe with 450 μ L of calibration solution for the positive ion mode into the syringe pump.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
3. Click the **Syringe Pump** button to start the syringe pump.
4. Click the title bar of the Evaluate window to display it.
5. In the Calmix Evaluation sub-window, select the MS Mass Check (pos) check box to evaluate the mass calibration for the positive ion mode. See Figure right.
6. In the Evaluate window, click **Evaluate** to start the automatic evaluation procedure.
7. The Evaluate window indicates the completed percentage of the evaluation process by a progress bar.
8. A message box informs you when the evaluation procedure is completed. Click **OK** to close the message box.



If this evaluation fails, calibrate the Q Exactive mass spectrometer as described on [page 6](#).

This example describes the evaluation of the isolation calibration for the positive ion mode.

❖ To evaluate the isolation calibration for the positive ion mode

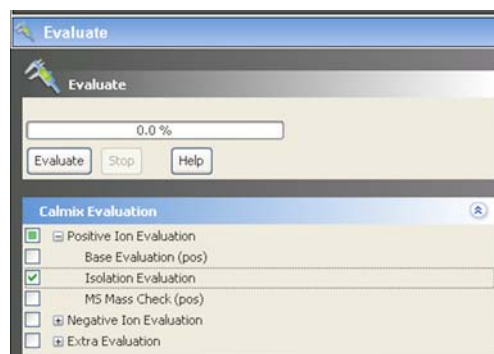
1. Place a clean, 500 μ L Unimetrics syringe with 450 μ L of calibration solution for the positive ion mode into the syringe pump.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.

Evaluation of Isolation Calibration





3. Click the **Syringe Pump** button to start the syringe pump.
4. Click the title bar of the Evaluate window to display it.
5. In the Calmix Evaluation sub-window, select the Isolation Evaluation check box. See Figure right.
6. In the Evaluate window, click **Evaluate** to start the automatic evaluation procedure.
7. The Evaluate window indicates the completed percentage of the evaluation process by a progress bar.
8. A message box informs you when the evaluation procedure is completed. Click **OK** to close the message box.
Q Exactive



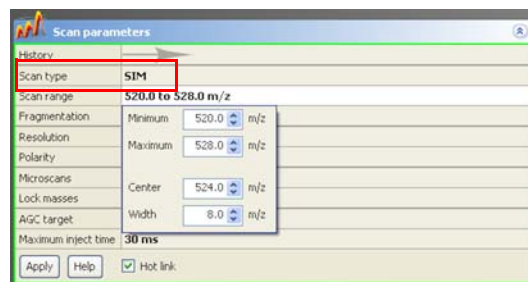
If this evaluation fails, please perform an “Isolation Mass and Resolution” calibration as described on [page 8](#).

In the previous examples, the Q Exactive mass spectrometer operates in the full scan mode. The following examples show the mass spectrometer operating in other scan modes.

In the following example, the quadrupole of the Q Exactive mass spectrometer works as mass filter and is used to perform a Selected Ion Monitoring (SIM) scan of a specified m/z range.

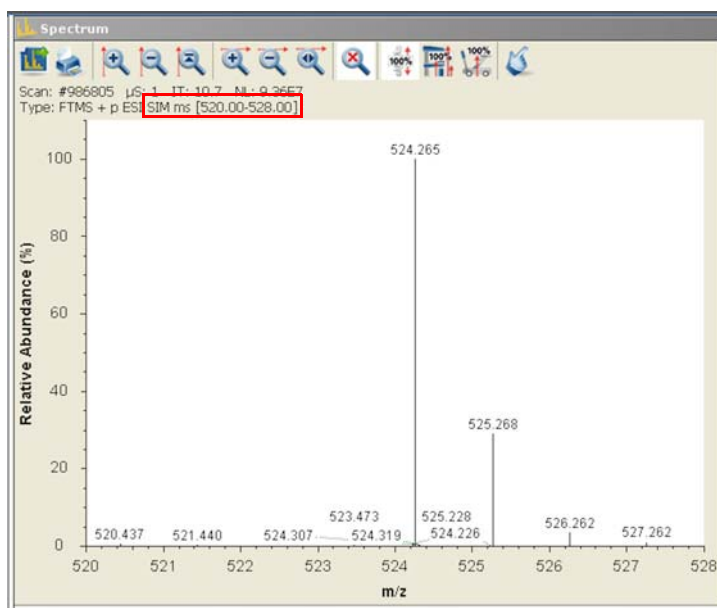
❖ To perform a SIM scan

1. If necessary, tune and calibrate your Q Exactive mass spectrometer as described above.
2. Set the scan parameters in the Scan parameters window:
 - a. Click into the Scan Range field to display the Scan Range dialog box.
 - b. Enter a value for the Center mass (m/z 524, for example). See Figure right.
 - c. Enter a width not larger than 10 amu (m/z 8, for example).Note that the scan type changes to SIM.
- d. If necessary, adjust the other scan parameters according to your requirements.
- e. Click **Apply** or select the Hot link check box.



3. Perform the data acquisition as described on [page 12](#).

The Figure below shows the typical spectrum of a SIM view (m/z 524 example).



Other Experiment Types

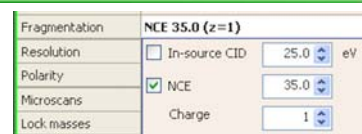
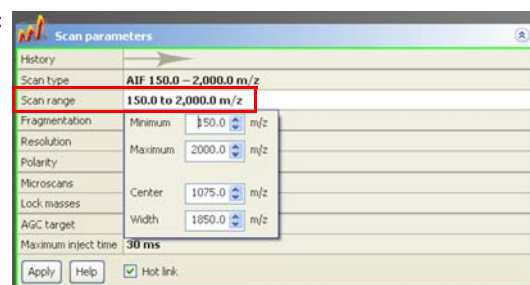
Performing SIM Scans

Performing AIF Experiments

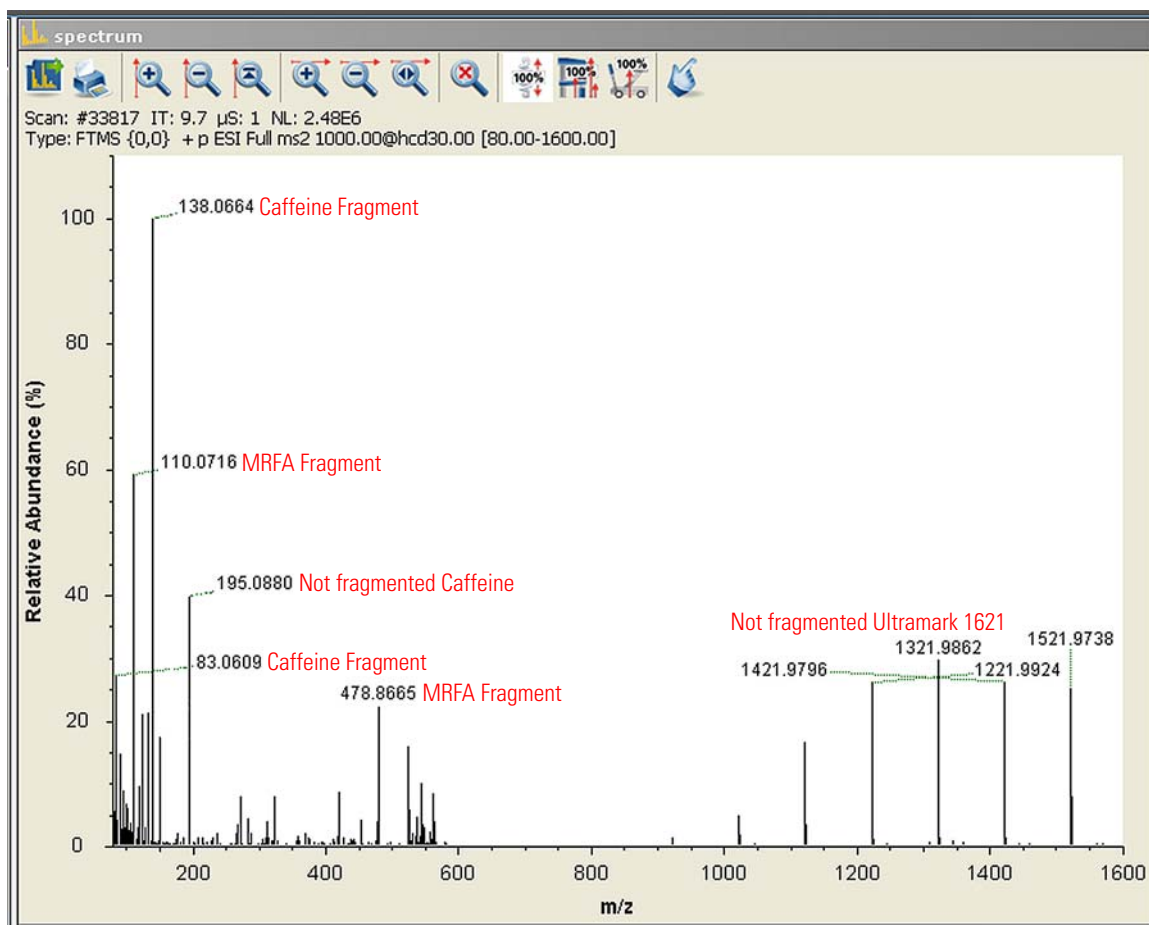
The Q Exactive mass spectrometer allows performing All Ion Fragmentation (AIF) measurements with in-source CID and with normalized collision energy (NCE). The following example describes an AIF fragmentation experiment.

❖ To set the scan parameters for an AIF fragmentation experiment with NCE

1. If necessary, tune and calibrate your Q Exactive mass spectrometer as described above.
2. Set the scan parameters in the Scan parameters window:
 - a. In the Scan type field, select **AIF-MS/MS**.
 - b. Click into the Scan Range field to display the Scan Range dialog box.
 - c. Adjust the scan range according to your requirements (from m/z 150 to m/z 2000, for example). See Figure right.
 - d. Click into the Fragmentation field to display the Fragmentation dialog box.
 - e. Select the NCE check box and enter the value for the normalized collision energy (35, for example) into the spin box. See Figure right.
 - f. Leave the value for Charge at 1.
 - g. If necessary, adjust the other scan parameters according to your requirements.
 - h. Click **Apply** or select the Hot link check box.
3. Perform the data acquisition as described on [page 12](#).



The Figure below shows a typical spectrum of an AIF fragmentation experiment with HCD.



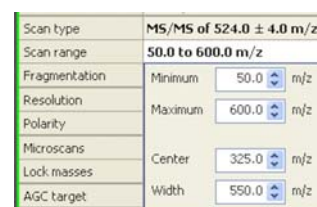
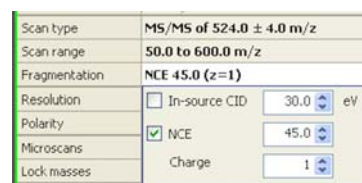
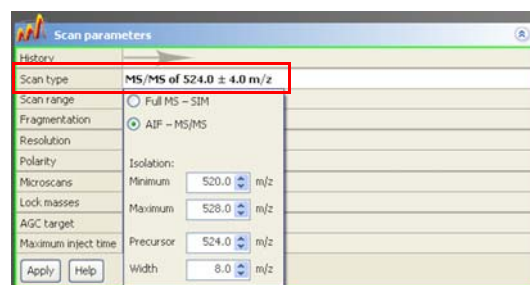
Performing MS/MS Scans

The following example shows an MS/MS experiment. Precursor ions that were selected by using the quadrupole are sent to the HCD collision cell of the Q Exactive mass spectrometer. Here, they are fragmented with normalized collision energy (NCE).

❖ To perform an MS/MS experiment

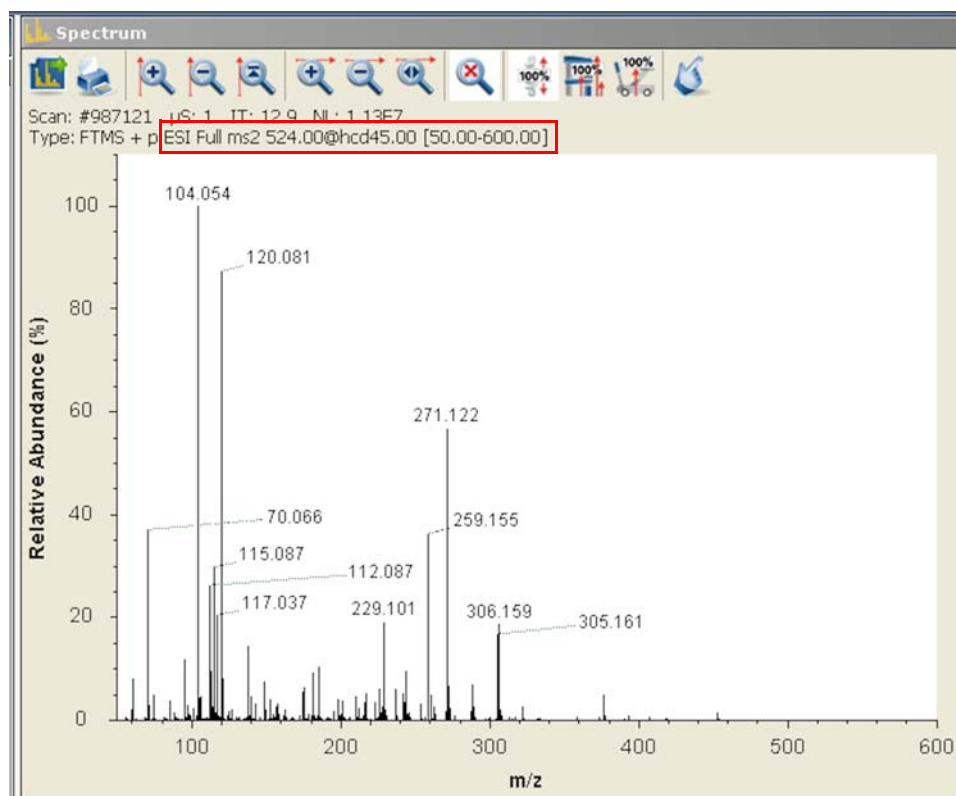
1. If necessary, tune and calibrate your Q Exactive mass spectrometer as described above.
2. Set the scan parameters in the Scan parameters window:

- a. In the Scan type field, select **AIF-MS/MS**. See Figure right.
 - b. In the Isolation area, enter a value for the precursor mass (m/z 524, for example).
 - c. Enter a width not larger than 10 amu (m/z 8, for example).
- Note that the scan type changes to MS/MS.
- d. Click into the Fragmentation field to display the Fragmentation dialog box.
 - e. Select the NCE check box and enter the desired value for the collision energy (45, for example) into the spin box. See Figure right.
 - f. Leave the value for Charge at 1.
 - g. Click into the Scan Range field to display the Scan Range dialog box.
 - h. Adjust the scan range according to your requirements (from m/z 50 to m/z 600, for example). See Figure right.
 - i. If necessary, adjust the other scan parameters according to your requirements.
 - j. Click **Apply** or select the Hot link check box.



3. Perform the data acquisition as described on [page 12](#).

The Figure below shows an example of a typical MS/MS spectrum.



Data Acquisition




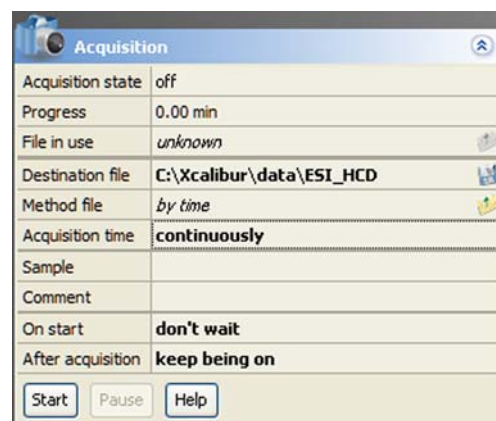
Setting up and Running Methods

Creating the Instrument Method

After you have entered the scan parameters of your experiment, you can finally start to acquire data.




❖ To perform data acquisition

1. Specify the acquisition parameters in the Acquisition window:
 - a. In the Destination file field, type the full path (Drive:\path\file name) of the raw file that is used to store the acquired data. Alternatively, click the  button to the right of the text box to open a dialog box to browse your directories. See Figure right.
 - b. Click into the Acquisition time field to open the dialog box. Select the continuously option button to specify that the acquisition will continue until you click **Stop**. The Acquisition time field shows *continuously*.
2. Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
3. Click the **Syringe Pump** button to start the syringe pump.
4. In the Acquisition window, click **Start** to initiate the data acquisition. The spectrum window shows a real-time display of the mass spectrum. The data are saved in the selected raw file.
5. Click **Stop** to stop the acquisition.

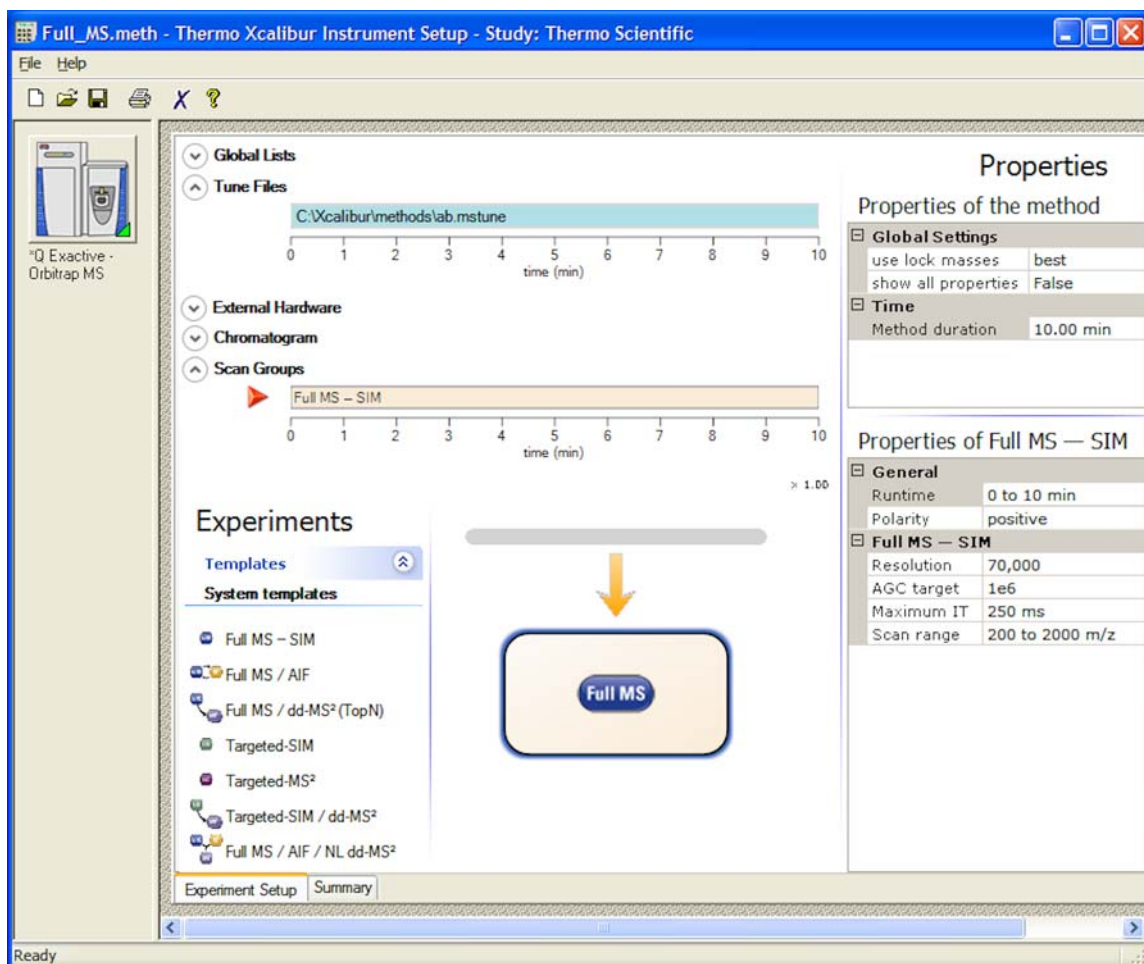


The following example describes how to create an instrument method in the Instrument Setup view of Xcalibur. You can then use the instrument method to acquire data with the Q Exactive mass spectrometer. The tune file used in the instrument method is used for data acquisition.

❖ To set up an instrument method

1. Display the Instrument Setup page in Xcalibur:
 - a. Choose **Start > Programs > Thermo Xcalibur > Xcalibur** to display the Xcalibur Home Page window.
 - b. Choose **GoTo > Instrument Setup** to display the Instrument Setup window.
 - c. If necessary, click **Q Exactive - Orbitrap MS** in the View bar to display the Method Editor.
2. In the Method Editor, drag an experiment symbol (for example, Full MS - SIM) from the Workflows pane (bottom left) to the gray bar in the Graph pane. A corresponding time bar is displayed in the Scan Groups pane.
3. Edit the fields of the Properties pane (right) to set up the experiment parameters. Refer to the *Q Exactive Software Manual* or the Q Exactive Tune Help for a description of the available parameters.
4. Assign a tune file:
 - a. On the External Hardware pane, click the  **Tune Files** icon to expand the Tune Files pane.
 - b. In the Properties of Tunefiles pane, click the Base Tunefile field.
 - c. Click the  button to open a dialog box.
 - d. Browse for the tune file (for example, ab.mstune) and assign it to the experiment.
5. Click  on the toolbar to save the new instrument method:
 - a. In the Save As dialog box, enter the name (Full_MS.meth, for example) and the location for your instrument method. Then click **Save**.
 - b. In the File Summary Information Dialog Box, enter a comment as header information for your instrument method. Then click **OK** to close the dialog box.

The instrument method can now be used to acquire data with your Q Exactive mass spectrometer, either from the Xcalibur Sequence Setup or from the Q Exactive Tune window. The Figure on the next page shows the example of an instrument method that was set up as described above.




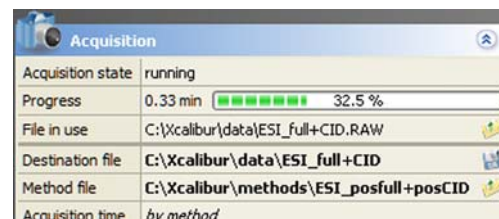
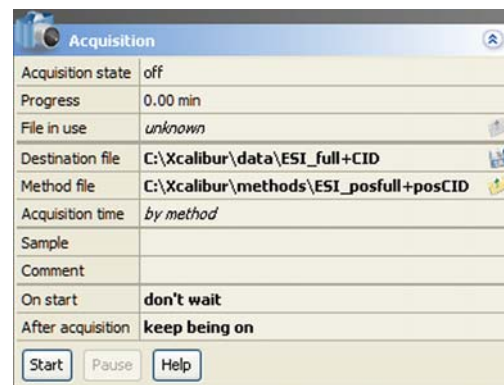
Running the Q Exactive Mass Spectrometer



This section shows an example of how to use an instrument method to run the Q Exactive mass spectrometer.

❖ To run the Q Exactive mass spectrometer with an instrument method


- Specify the acquisition parameters in the Acquisition window of Q Exactive Tune, as follows:
 - In the Destination file field, type the full path (Drive:\path\file name) of the raw file that is used to store the acquired data. See Figure right.
 - In the Method file field, click the  button to the right of the field to browse the computer for the instrument method file that was created in the previous topic. The Acquisition time field shows *by method*.
- Click the **On/Off/Standby** button to take the mass spectrometer out of the Standby mode or Off mode and turn it On.
- Click the **Syringe Pump** button to start the syringe pump or start a separate pump for infusion experiments.
- In the Acquisition window, click **Start** to initiate the data acquisition:
 - The Start button is replaced by a Stop button.
 - The Acquisition state changes to *running*.
 - A progress bar indicates the completed percentage of the acquisition. See Figure right.
 - Icons appear on the toolbar to indicate an active data acquisition.



After Operation

- If necessary, Q Exactive Tune appends a time stamp to the name of the destination raw file to prevent overwriting existing files.

When the Scan parameters window and the Spectrum window are monitored, it is possible to view how Q Exactive Tune changes the values according to the settings defined in the instrument method.

The acquisition ends when the acquisition time defined in the instrument method has expired. To display the raw file with the acquired data in Qual Browser, click the  button in the File in use field.

When the measurement is finished, Thermo Fisher Scientific recommends leaving the Q Exactive mass spectrometer in Standby mode and not in Off mode to provide the best mass accuracy for the next analysis.