

Agilent MassHunter Workstation Software – Data Acquisition for 6400 Series Triple Quadrupole LC/MS

Familiarization Guide

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Use the exercise in this guide to learn to use the Agilent 6400 Series Triple Quad LC/MS. You can do this exercise with the demo data files, SulfaDrugs, shipped with the system (in the **Data** folder of your Qualitative Analysis installation disk), or with data you acquire.



With this exercise, you learn how to determine the best acquisition settings for analyzing your compounds of interest. These instructions help you understand not only how to set up a worklist to optimize instrument parameters for best sensitivity in acquisition, but also how to use the Qualitative Analysis program to identify parameter values producing optimum signal response. You can also learn about the Qualitative Analysis program by using the *Qualitative Analysis Familiarization Guide or the online Help for the Qualitative Analysis program*.

In this exercise, you create an acquisition method for a mixture of four sulfa drugs, optimizing both the fragmentor and collision energy voltages to maximize sensitivity. One of the ways to optimize parameters is to create a worklist, or sequence, of data file acquisitions, each using a different method. This exercise uses this protocol for method development.

Another way to develop a method is to use the manual tune capability to optimize various parameters, including collision energy, to obtain the optimal signal response for each multiple reaction monitoring (MRM) transition. A third technique has you set up an acquisition method that directs the instrument to make multiple injections of the sample from an autosampler vial and acquire the data to a single data file. The method contains multiple time segments, one for each injection, with an incremental change made to a particular parameter (e.g., collision energy) in each segment.

This exercise uses the first protocol for method development.

NOTE See the *Concepts Guide* to learn more about how the triple quadrupole mass spectrometer works and why the fragmentor and collision energy voltages are important. For background information, see Chapter 3, "Agilent Triple Quad MS and Sensitivity", in the *Concepts Guide*. See the online Help for detailed information on how the program works.

Each task is presented in a table with three columns:

- Steps Use these general instructions to proceed on your own to explore the program.
- Detailed Instructions Use these if you need help or prefer to use a step-by-step learning process.
- Comments Read these to learn tips and additional information about each step in the exercise.

Before you begin

Before you begin, you need to check that your system is ready. If you plan to acquire data, you also need to set up the instrument.

Prepare your system

- 1 Check that:
 - The Data Acquisition program has been installed.
 - The LC modules and the 6400 Series Triple Quad LC/MS have been configured.
 - The performance has been verified.
 - The system has been turned on.

If these actions have not yet been done, see the *Installation Guide* for your instrument.

2 Copy the data files to your PC.

Copy the folder named **SulfaDrugs** in the **Data** folder on your Qualitative Analysis installation disk to any location on your hard disk. This folder contains all the data files needed for this exercise.

NOTE Do not re-use the sulfa drug data files already on your system unless you know that you copied them from the originals on the disk and you are the only one using them.

Do not use these sample data files to look at sample information or print a report.

Prepare to acquire data

If you do not intend to acquire data but want to learn how to use the Qualitative Analysis program for method development, you can skip this step, which tells you how to prepare the demo sample. You then do those tasks that show you how to use the Qualitative Analysis program with the sulfa drug data files shipped with the system.

Parts List The exercise in this guide uses this equipment and materials:

- Agilent 1100 or 1200 LC modules: well-plate sampler, binary pump, thermostatted column compartment, DAD
- Zorbax column (see Table 1 on page 4)
- A 10-µL sulfa mix sample (prepared in this step)

Table 1 2	orbax column	s
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Triple Quadrupole	Column Description	Film Thickness	Pore Size	Part Number
6410	SB-C18 2.1mm x 30mm	3.5 µm	100Å	873700-902
6430 or 6460	SB-C18 2.1mm x 50mm	1.8 µm	80Å	822700-902

1 Prepare the LC solvent.

In 1-liter reservoirs of HPLC-grade water and acetonitrile (ACN), add 1 mL of 5M $\rm NH_4HCO_2$ (Ammonium Formate) each to make 5mM $\rm NH_4HCO_2$ in water and acetonitrile and use for the A and B channels, respectively.

- **2** Prepare the sample.
 - **a** Add 10 μ L sulfa mix from one of the ampoules (500 μ L) to 990 μ L of solvent A in an Eppendorf vial so that the final concentration is 1 ng/ μ L.
 - **b** Place a sample vial containing an injectable amount of the prepared sample in the autosampler.
- **3** Set up the LC column.

Use the appropriate Agilent column from Table 1.

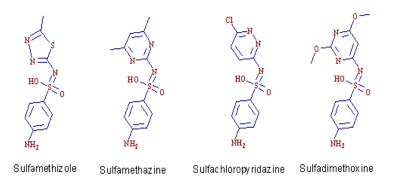
4 Set the column temperature.

For the Agilent 6460 with Agilent Jet Stream Technology, set the column temperature to 60° C.

For the Agilent 6410 series, set the column temperature to 40° C. This exercise can also run at room temperature.

The Electrospray LC Demo Sample (P/N 59987-20033) contains five ampoules with 100 ng/ μ L each of sulfamethizole (M+H)⁺ = 271, sulfamethazine (M+H)⁺ = 279, sulfachloropyridazine (M+H)⁺ = 285, and sulfadimethoxine (M+H)⁺ = 311.

Before you begin Prepare to acquire data



NOTE Determining optimal parameter values for acquiring sample compound data requires that the Agilent Triple Quad instrument already be tuned on the Tuning Mix calibrant ions. Before proceeding with this exercise, make sure you have used Checktune or Autotune to verify that calibrant ions each have the proper mass assignment, peak width, and signal intensity.

See the *Quick Start Guide* or online Help for instructions on tuning the instrument.

Task 1. Enter acquisition parameters and acquire data

Exercise 1 – Develop an acquisition method for the 6400 Series

For this exercise you analyze a mixture of four sulfonamide compounds.

Task 1. Enter acquisition parameters and acquire data

In this exercise, you enter the conditions for the analysis of the sulfa drug mix.

Steps	Detailed Instructions	Comments	
1 Enter LC parameters appropriate for sulfa drug mix.	 a Double-click the Data Acquisition icon. b Make sure that Acquisition appears as 	• The Data Acquisition window appears. See Figure 1.	
See Table 2.	 the selection in the Context text box. If Tune is the selection, click Acquisition from the Context dropdown menu in the Combo bar. c Enter the LC parameters listed in the Table 2. 		

Table 2 LC parameters for sulfa drug mix

Parameter	6410	6430 or 6460				
PUMP						
• Flowrate	800 µL/min	800 µL/min				
Solvent A	$5 \text{ mM NH}_4\text{HCO}_2 \text{ in H}_2\text{O}$	$5 \text{ mM NH}_4\text{HCO}_2 \text{ in H}_2\text{O}$				
• Solvent B	5 mM NH ₄ HCO ₂ in ACN 90:10 acetonitrile:water	5 mM NH ₄ HCO ₂ in 90:10 acetonitrile:water				
Gradient (min - %B)	0 min - 13%	0 min - 13%				
	1.80 min - 60%	1.80 min - 60%				
	2.50 min - 60%	2 min - 60%				
Stop Time	2.50 min	2.0 min				
Post Time	2.50 min	2.0 min				
NJECTOR						

Task 1. Enter acquisition parameters and acquire data

Parameter	6410	6430 or 6460
• Inj. Vol.	1 µL	2.0 µL
 Injection 	Standard	With needle wash
Draw Position	3.0 mm	0.0 mm
UV DETECTOR		
• Ch A	254 nm (4 nm BW on DAD)	254 nm (4 nm BW on DAD)
REF A (DAD only)	400 nm (80 nm BW)	400 nm (80 nm BW)
COL THERM		
• Temp	40°C	60 °C

Table 2 LC parameters for sulfa drug mix (continued)

Task 1. Enter acquisition parameters and acquire data

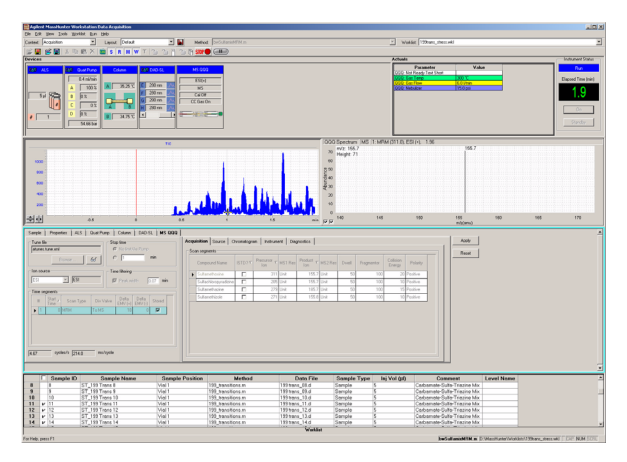


Figure 1 Agilent MassHunter Workstation Software – Data Acquisition window

Task 1. Enter acquisition parameters and acquire data

Steps		D	etailed Instructions	Comments	
2	Enter MS parameters appropriate for sulfa drug mix and save the method as <i>iii</i> MS2Scantest.m, where <i>iii</i> are your initials.	-	Click the MS QQQ tab in the Method Editor pane. Click the Scan Type cell, and click MS2Scan from the list.		
		C	Enter the other MS parameters as		
	See Table 3.		listed in Table 3. These parameters are in either the Acquisition or the Source tabs.		
		d	Save the method as <i>iii</i> MS2Scantest.m, where <i>iii</i> are your initials.		

Table 3MS parameters for sulfa drug mix

Parameter	Value	
• Inlet	ESI (positive polarity)	ESI (positive polarity) with Agilent Jet Stream Technology
• Gas Temp	350 °C	350 °C
Scan Type	MS2Scan	MS2Scan
• Nebulizer	50 psi	35 psi (nitrogen)
• Dry Gas	12 L/min	10 L/min
Range	100 to 400	100 to 400
Sheath Gas Temperature	not applicable	400 °C
Sheath Gas Flow	not applicable	12 L/min
• Nozzle Voltage	not applicable	0 V
Capillary voltage positive	4000 V	4000 V
• Delta EMV pos	400 V	200 V

Task 1. Enter acquisition parameters and acquire data

	Stop time No limit/As Pump	Acquisition Source	Chromatogram	Instrument	Diagnostic	5		1	Apply
Browse 60'	C ☐ min Time filtering IV Peak width 0.07 min ■ Delta Delta EMV (+) EMV (+) Stored	Sean segment Name	Start Mass 1	End Mass 1000	Scan Time 500	Fragmentor 135	Polarity Positive		Reset
1 0 US2 Searce To MS MS2 SIM MS2 SIM MSP MRM Product Ion Procusor Ion Nextra I cos Versita Gram 225 cycles/s 443.8 ms/cycl	0 0 17		0.1 Profile	• amu					

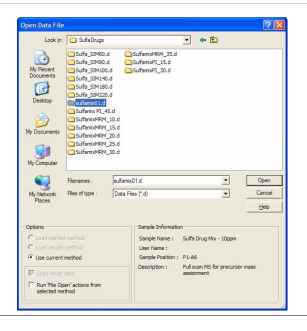
Figure 2 Select Scan Type of MS2 Scan in the MS QQQ tab

Steps	Detailed Instructions	Comments
 3 Acquire data (optional). Set up a one-line worklist with the method you just created. Name the data file <i>iiisulfamix01.d</i>, where <i>iii</i> are your initials. Designate a directory path to hold your data files and method. 	 a If necessary, click the Worklist icon to display the Worklist pane. b Click Worklist > Worklist Run Parameters. Verify that the parameters are set properly. Click OK. c Click Worklist > Add Multiple Samples. d Type <i>iii</i>sulfamix01.d and <i>iii</i>MS2Scantest.m as the data file name and the method name, respectively. e Click the Sample Position tab. f Select None as the autosampler. g Type 1 as the Number of samples. h Click OK. i In the Worklist pane, mark the check box to the left of the sample as shown below. 	 You have just acquired a full scan MS data file to see what ions are being formed from the sample. This step is optional because you can perform the next step with an example data file that comes with the program. If you prefer, you can create your own data file as described in this step.
	Sample Name Sample Position	Method Data File S
	Sample1 Vial 1 pfhMS25	Scantest.m pfhsulfamix01.d San
	j Click the Start Worklist Run icon in the main toolbar or click the Run > Worklist command.	

Task 2. Determine precursor ion masses

In this exercise, you determine the precursor ions for each of the sulfa drugs in the acquired data file.

Steps	Detailed Instructions	Comments	
 Open the acquired data file. In the Qualitative Analysis program, open either the example file, sulfamix01.d, or the data file you created in "Task 1. Enter acquisition parameters and acquire data" on page 6. 	a Double-click the Qualitative Analysis icon.	 When you open the sulfa drug directory after installation, the Load result data (lower left corner) check box is grayed out. If you see the check box marked, this means that the data file(s) already contains results. Clear this check box before opening the file. 	



Task 2. Determine precursor ion masses

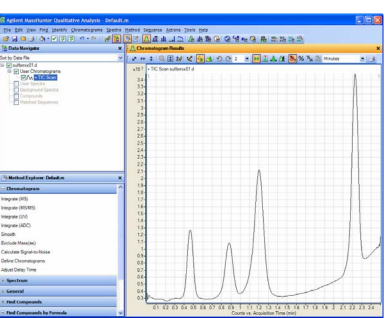
Steps	Detailed Instructions	Comments		
	 b Do one of the following: Select the example data file sulfamix01.d, and click Open. Select the data file you created in "Task 1. Enter acquisition parameters and acquire data" on page 6, and click Open. By default, the system displays the Total Ion Chromatogram (TIC). 	 The figure below shows the default layout. This is what you want to see. The Qualitative Analysis program displays a newly opened data file with the same layout and display settings used for the previous data file. Therefore, you MUST make sure to return to the default settings for this exercise. 		

Before you begin, make sure that all previous settings are returned to their default values:

- Restore default layouts
 - Click View > Window Layouts > Restore Default Layout.
- Make sure the method is default.m. (see title bar)
 - Click Method > Open.
 - Select default.m, and click Open.
- Return display options to default settings.
 - Click Tools > Plot Display Options
 - Click **Default**, and then **OK**.

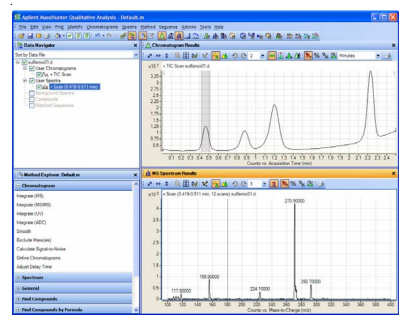
0r...

- Restore the General layout.
 - Click Tools > Configure for Workflow > General.
 - Click OK.
 - (optional) You may be asked to save method changes.
- Return display options to default settings.
 - Click Tools > Plot Display Options
 - Click **Default**, and then **OK**.



Task 2. Determine precursor ion masses

Steps				D	etailed Instructions	Comments		
2	Determine precurso for all four peaks. • You have determ correctly if you fi are similar to the this table:	ined the	em values	a b	In the Chromatogram Results window, make sure that the Range Select icon in the toolbar \bowtie is On. Click the left mouse button and drag the cursor across the first peak to produce a shaded region, as in the figure below.	•	The system displays an averaged spectrum across the peak in the MS Spectrum Results window. The precursor mass of the first compound, sulfamethizole, is determined to be m/z 270.9. To obtain a single scan, double-	
Compound RT m/z		C	Right-click the shaded area, and click		click the apex of the peak.			
	Sulfamethizole Sulfachloropyridazine	0.47 0.88	270.9 284.9		Extract MS Spectrum from the shortcut menu.			



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d Repeat step a through step c for the other compounds.
 The precursor ion masses should

match those in the table in step 2.

- e Click File > Close Data File.
- f When asked if you want to save the results, click **No**.
- Some compounds form sodium (Na) and/or potassium (K) adducts as well, corresponding to M + 23and M + 39 masses respectively. Seeing these masses along with the M + H can make for an easy confirmation of which ion is the pseudo-molecular ion (M + H)+.

Sulfamethazine

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Sulfadimethoxine

1.20

2.23

If you acquired the data file

may be different.

using the Agilent Jet Stream

Technology, the retention times

Close the data file after finding the precursor ion masses.

279.0

311.0

Task 3. Find optimum fragmentor voltage for maximum response

Task 3 shows you how to carry out the optimization for fragmentor voltage by creating selected ion-monitoring experiments for each compound within a method and setting up multiple methods with varying fragmentor voltages.

Steps	Detailed Instructions	Comments
 Set up six methods for six different fragmentor voltages. Change to a SIM experiment. 	a In the Scan Type dropdown MS2 SIM.	list, click
 Use 60, 80, 100, 140, 180 and 220 volts as the fragmentor voltages for the six methods. Save the methods as <i>iii</i>MS2SIM<i>xxx.m</i>, where <i>iii</i> are your initials and <i>xxx</i> is the voltage. 	Tune file atunes.tune.xml Browse Browse 6d Ion source ESI Time segments # Start / Scan Type Ø 1 0 MS2 SIM To MS MRM Product Ion Precursor Ion Neutral Loss Neutral Loss 1.23	EMV (+) EMV (-)

eps	De	etailed Instruction	S	Com	nents			
	C d f g	In the Acquisition Compound Name (precursor ion ma sulfadimethoxine Right-click anywh segments section Type the Compou Mass for sulfach Repeat steps c ar sulfamethazine at Save the method where <i>iii</i> are your Change the fragm and save the met <i>iii</i> MS2SIM060, w initials. Repeat step g for and 220, saving th <i>iii</i> MS2SIM080, <i>iii</i> <i>iii</i> MS2SIM080, <i>iii</i> <i>iii</i> MS2SIM080, <i>iii</i>	e and M iss) for here in t a, and c nd Nan foropyri ad d for nd sulfa as <i>iii</i> hod as where <i>iii</i> voltage he meth <i>i</i> MS2S ad <i>iii</i> MS	the Scan lick Add F ne and the dazine. amethizolo S2SIM14 s. fare your are your s 80, 100, sods as M100, S2SIM220	di th • Th Au Bow. e> • m fra e> • 0.m, 60, 180	fferent set e Acquisiti ne Instrume cquisition p periment f ass, startin	of column on windov ent Contro program cr or each co ng with a d oltage of 1	l and Data eates a SIM mpound
	1		hromatogra	am Instrum	ent Diagnostics			
		Scan segments Compound Name	ISTD?	Mass 🗸	MS2 Res	Dwell	Fragmentor	Polarity
		sulfadimethoxine			Unit	200	-	Positive
			Г	285	Unit	200	140	Positive
		sulfachloropyridazine			Unit Unit	200		Positive Positive

Task 3. Find optimum fragmentor voltage for maximum response

	s Detailed Instructions						Comments				
 Set up and run the worklist optional). Set up six samples with Sample Name SulfaDrugMix to inject 1ul from vials 1-6 or the ones you choose. Specify the data files as <i>iii</i>SulfaSIMxxx.d, where <i>iii</i> are your initials and xxx is the voltage. 	b c	maka Click new the I To see left c Wor Type files. Type run. Click samµ five s volta Marl	k F water k F water k C k C k C k C k C k C k C k C k C k C	sure the work ile > New rorklist. You to worklist. You to worklist. up the run, rner of the ist Run Par he paths for he informat Vorklist > 1 worklist > 1 wor	t icon if necessa orklist is visible. Worklist to s do not need to right-click the u worklist, and cli rameters. The method and tion for the 60 vo Add Sample. And to the Worklist. And to the Worklist for ox to the left of the r each of the six	tart a save upper ck I data oltage other Add	can use data fi	tional because you les shipped with th orm many of the cercise.			
	4	-			0 I D **			0.1.7			
				ample Name ulfaDrugMix	Sample Position Vial 1	Acq Metho MS2SIM060.m		Sample Type Sample			
		2 ¥	S	ulfaDrugMix	Vial 1	MS2SIM080.m	d:\Sulfa_SIM080.d	Sample			
				ulfaDrugMix	Vial 1	MS2SIM100.m		Sample			
				ulfaDrugMix	Vial 1	MS2SIM140.m		Sample			
				ulfaDrugMix	Vial 1	MS2SIM180.m		Sample			
		6 ¥	' Si	ulfaDrugMix	Vial 1	MS2SIM220.m	n d:\Sulfa_SIM220.d	Sample			

a checkmark.

ps	Detailed Instructions	Comments
 Set up a qualitative method to view the EIC data automatically. Open the data file Sulfa_SIM60.d or your own <i>iii</i>Sulfa_SIM60.d, where <i>iii</i> are your initials. In the Method Editor, add in the EICs corresponding to the precursor ion masses of 271, 279, 285, and 311. Save the method as <i>iii</i>Exercise1, where "<i>iii</i>" are your initials. 	Image: Sector Activity Image: Sector Activity Stort by Data Rise Image: Sector Activity Image: Sector Activity Image: Sector Activity Image: Sector Activity	Conta 1940 Conta
	Calculate Signal to Noise Define Chromatograms MS level: MS	Add Change Deter Deter Consolidation Deter Change Deter Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Deter Change Cha

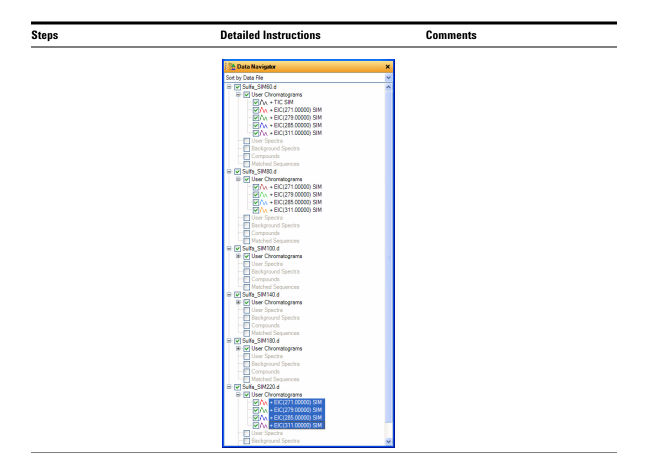
teps	Detailed Instructions	Comments
	 d If necessary, click Define Chromatograms in the Chromatogram section of the Method Explorer. e To delete the BPC chromatogram, click Delete. f Select EIC for the Chromatogram Definition Type, g In the MS Chromatogram tab, make sure MS Level is set to All and Scans is set to All Scan Types. h Clear the Do cycle sum check box. i Type 271 as the m/z value. j Click Add. k Repeat steps i and j for the other precursor ions, 279, 285 and 311. l Click Method > Save As. The system opens the Save As dialog box m Save the method as <i>iii</i>Exercise 1.m. n Click Method > Save As. 	 The default Method Editor list selection after installation is Integrate. You can also select Define Chromatograms from the Method Items list.
	Method Editor: Define Chromatograms	×
	Method Editor: Define Chromatograms Method Editor: Define Chromatograms Method Rems• B	×

Steps	Detailed Instructions	Comments
 Extract the chromatogram for the data file and view the results. Make sure you can see all five chromatograms, the TIC and four EICs. 	a Click the Run button on the Method Editor toolbar.	 You can also click the Chromatograms > Extract Defined Chromatograms command to extract the defined chromatograms.
	 b To see the TIC and four EICs, click the arrow next to the Maximum Number of List Panes icon in the Chromatogram Results toolbar, as shown in the example below. c Select 5 to view five chromatograms simultaneously. The system displays chromatogram results as shown below. 	
	A Chromatogram Results	×
	x10 ⁶ + TIC SIM Sulfa_SIM60.d 1 + TIC SIM Sulfa_SIM60.d 3 + EIC(271 00000) SIM Sulfa_SIM60.d 7 + EIC(279 00000) SIM Sulfa_SIM60.d 1 + EIC(285 00000) SIM Sulfa_SIM60.d 2 + EIC(285 00000) SIM Sulfa_SIM60.d 1 + EIC(311.00000) SIM Sulfa_SIM60.	X 36 % % Minutes

St	teps	Detailed Instructions		Comments				
	 teps Extract the remaining ion chromatograms automatically. Extract Defined Chromatograms should be the default action for Assign File Open Actions. Open the remaining data files, Sulfa_SIM80.d through Sulfa_SIM220.d. Close the Method Explorer. 	 a Select File Open Actions from the General section in the Method Explorer. b Make sure that Actions to be run list only contains Extract Defined Chromatograms. i Method Editor: Assign Actions to Run Opening a Data Chromatograms. i Method Editor: Assign Actions to Run Opening a Data Chromatograms. i Method Editor: Assign Actions to Run Opening a Data Chromatograms. i Method Editor: Assign Actions to Run Opening a Data Compound Automation Analysis Automation Analysis Automation Find Compounds by Auto MS/MS Find Compounds by Auto MS/MS Find Compounds by Formula Export mzData Generate Compound Report 		•	The Qualitative Analysis Method Editor lets you define actions to be performed automatically upon opening a data file(s).			
			Generate Analysis Report Generate Formulas from Compound					
		d	Click File > Open Data File. The system displays the Open Data File dialog box. Select the data files to be opened, Sulfa_SIM80.d through Sulfa_SIM220.d. Mark the Run 'File Open' actions from selected method checkbox. (lower left corner)					

Task 3. Find optimum fragmentor voltage for maximum response

g To close the Method Explorer and Method Editor, click the **X** in the upper right corner of each window.



Task 3. Find optimum fragmentor voltage for maximum response

Steps	Detailed Instructions	Comments
 Select the fragmentor voltage that produces the maximum response for each of the precursor ions. Close the data files after you determine the optimum voltage. 	 a In the Data Navigator window, highlight the EICs for 271.0 m/z. b Click the Show only the highlighted items icon, 2. Only the 271 m/z check boxes are now marked. c Look at the relative intensities of each peak to determine which fragmentor voltage setting will be best to use for the 271 precursor. 	A fragmentor voltage of 100 should
	Image: Sort by Data Navigator × Image: A chromatogra Sort by Data Fle ✓ Image: A chromatogra	
	✓/ + EIC(271.00000) SIM ✓	You can overlay the chromatograms by clicking the Overlaid mode icon in the Chromatogram Results toolbar.
	d Repeat step a through step c for the other three base peaks or precursor	 Click the different EICs in the Data Navigator window to change which chromatogram is labeled in the
	ions. e Click File > Close Data File.	chromatogram is labeled in the Chromatogram Results window.
	 f Click Close when the Close Data File dialog box appears. 	When the color of the label of the chromatogram matches the color of the chromatogram that has the

highest intensity, you use the fragmentor voltage that was used

for that file.

Task 4. Determine product ion masses

Task 4. Determine product ion masses

In this part of the method development, we will use three collision energies to determine the best fragment ions to use for the eventual Multiple Reaction Monitoring (MRMs).

Steps	Detailed Instructions	Comments		
 Set up three product ion acquisition methods and acquire data. Use the MS parameters in the example below, but change the Fragmentor voltage to the optimum voltage you determined in the previous task. 	 a Click the MS QQQ tab in the Method Editor pane. b Select Product Ion in the Scan Type combo box to scan each precursor ion for all its product ions. c Enter all MS parameters as listed in the example below, making sure the Collision Energy is set to 15 and the 			
 Save methods as <i>iii</i>Sulfamix PI_xx.m, where <i>iii</i> are your initials and xx is the collision energy. 	Fragmentor voltage is set to 15 and the optimum voltage determined in Task 3. d Save the method as <i>iii</i> Sulfamix PI 15.m.			
unurgy.	e Repeat step c and step d for collision energies of 30 and 45.			

I une rile	stop ame		equivition Source ci	moniatogram	matrument	Diagnos	lica				
atunes.tune.xml	No limit/As Pump	Шr	Scan segments								
Browse 65	C 1 min		Segment Name	Precursor Ion ∇	MS2 From	MS2 To	Scan Time	Fragmentor	Collision Energy	Polarity	
			Sulfadimethoxine	311	50	320	250	140	15	Positive	
Ion source	Time filtering		Sulfachloropyridazine	285	50	320	250	140	15	Positive	
ESI ESI	Peak width 0.07 min		Sulfamethazine	279	50	320	250	140	15	Positive	
Time comonte			 Sulfamethizole 	350	50	320	250	140	15	Positive	
# Start Scan Type Div V. 1 0 Product Ion To MS 1 0 Product Ion To MS	Start / Scan Type Div Valve Delta Delta Stored										
1.1 cycles/s 913.1 ms/	cycle										

- 2 Set up and run the worklist (optional).
 - Specify the data files as *iii*Sulfamix Pl_xx.d, where *iii* are your initials and xx is the collision energy.
- **a** Scroll down if necessary to make sure the worklist is visible.
- b Add three samples to the worklist for collision energies 15, 30 and 45.
- c Mark the checkbox to the left of the Sample Name for each sample you are adding.
- d Click Run > Worklist.

- This step is optional because you can determine the product ion masses from the data files shipped with the system.
- Use the instructions in Step 2 of Task 3 to set up the worklist.

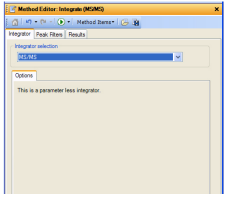
$\label{eq:Exercise1-Develop} Exercise 1-Develop an acquisition method for the 6400 Series$

Task 4. Determine product ion masses

Steps	Detailed Instructions	Comments
 Set up a qualitative method to integrate and extract product ion spectra. Use the data files SulfamixPl_xx.d, where xx is the collision energy, or your own data files, <i>iii</i>SulfamixPl_xx.d. Open Method Explorer and Method Editor. Use TICs set up for MS/MS, product ion and each of the precursor ions 271, 279, 285, 311. Make sure the MS/MS integrator has been selected and the maximum number of peaks has been limited to the largest 100 peaks. Add the ability to integrate and extract peak spectra to the file actions run upon data opening. Save the changes to the current method. 	 a Click the Open Data File icon in the toolbar. b Select SulfamixPl_15.d. c Make sure that the Run File Open Actions from Specified Method check box is clear, and click Open. d Make sure the Method Explorer and the Method Editor windows are displayed; otherwise, click the Method Explorer and then Method Editor icons. Refer to the Method Editor icons. e In the Chromatograms Section in the Method Explorer window, select Define Chromatograms. f Delete any existing chromatograms list. g Select TIC from the Chromatogram Definition list. h For MS Level, select MS/MS. i Mark the Do cycle sum check box. j For Scans, select Product ion. k For Precursor ion m/z, type 271. l Click Add. m Repeat steps j and k for each ion. 	 The Qualitative Analysis program should already be open and contair <i>iii</i>exercise 1.m as the method.
	🚰 Method Editor: Define Chromatograms	×
	Image: Construction of the second	

Task 4. Determine product ion masses

Steps	Detailed Instructions	Comments	
	 n From the Method Explorer in the Chromatogram section, click Integrate (MS/MS). o Select the MS/MS Integrator, if necessary. 	 These data files contain MS/MS data, so you need to modify the parameters in the Integrate (MS/MS) section. If the data file contained only MS data, you would need to modify the parameters in the Integrate (MS) section. 	



- Figure 3 Integrate (MS/MS) > Integrator Tab
- p Click the Peak Filters tab. Make sure that the Limit to the largest check box is marked and set to the value 100 (peaks) as shown below.

Integrator A Peak Filters Results								
Filter on	O Peak height		 Peak are 	a				
Height filte		> =	10000	counts				
Relativ	e height	>=	5.000	% of largest peak				
Area filters				_				
Absolu	te area	>=	10000	counts				
Relativ	e area	>=	1.000	% of largest peak				
	e area number of peaks	>=	1.000	% of largest p				

Figure 4 Integrate (MS/MS) > Peak Filters tab

Task 4. Determine product ion masses

Steps	Detailed Instructions	Comments
	 q Click General in Method Explorer, an then click File Open Actions. r Select Integrate and extract peak spectra from the Available actions li and click r to add this to Action to be run. 	st
	🗄 🚰 Method Editor: Assign Actions to Run Opening a Data File	×
	🛃 🔄 🕶 🖓 🔸 🔁 👘	
	Available actions Compound Automation without Report Cromatogram Pack Survey without Analysis Report Find Compounds by Marcel MS/MS Find Compounds by Formula Export Generate Compound Report Generate Analysis Report Generate Analysis Report Compound Strong Compound Fingerate And Extract Peak Spectra Actions to be run Extract Peak Spectra Integrate And Extract Peak Spectra	× • •

Figure 5 General > File Open Actions tab

	S	To apply the changes to the current method, <i>iii</i> exercise1.m, click the Save Method icon.	
4 Run the qualitative r current data file.	nethod on the •	In the Method Editor toolbar, click the Run button, . When the Assign Actions to Run Opening A Data File section is displayed, the Actions to be run list is done.	 The program first extracts the product ion chromatograms for each precursor ion in the data file. Next, it finds the largest peak in the total ion chromatograms, and integrates and extracts peak spectra from each integrated peak. See Figure 6 on page 28.

Task 4. Determine product ion masses

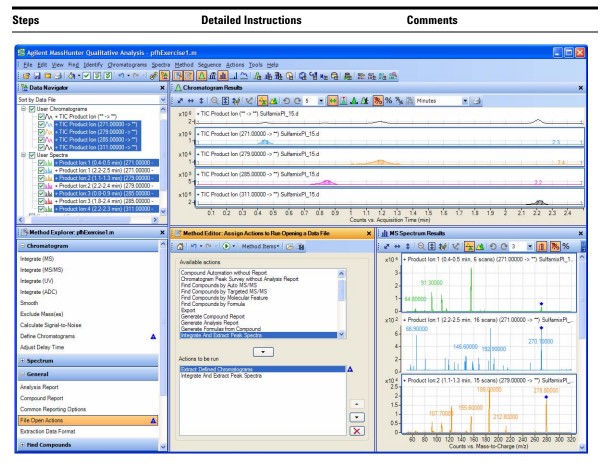


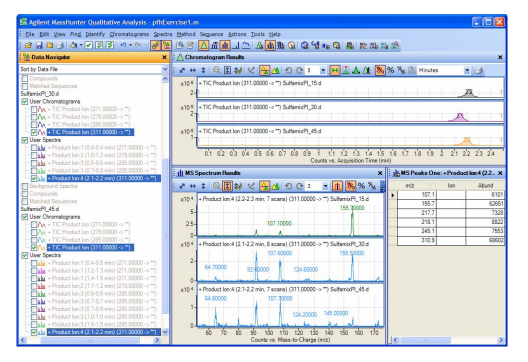
Figure 6 Results for integration and extraction of peak spectra.

5 Run the 'File Open' actions on the remaining product ion data files.

- Use either the example files, Sulfamix Pl_xx.d, or the data files you acquired in step 2.
- a Click File > Open Data File. The system displays the Open Data File dialog box.
- b Hold the Ctrl key and do one of these:
 Select the two data files Sulfamix
 - PI 30.d, and Sulfamix PI 45.d.
 - Select the data files you acquired in step 2.
- c Mark the **Run 'File Open' actions from** selected method check box in the Open Data File dialog box, and click **Open**.
- After the data files open, the Qual method first extracts the product ion chromatograms for each precursor ion.
- Next, it finds the largest peak in the total ion chromatograms, and integrates and extracts peak spectra from each integrated peak.

Task 4. Determine product ion masses

Steps	Detailed Instructions	Comments	
 6 Identify product ions. View each set of TICs and spectra individually (e.g., 271 m/z first). Close the data files. 	 a In the Data Navigator, select the TICs and spectra for the 271 m/z precursor ion. b Click the Show only the highlighted items icon, . c Click View > MS Spectrum Peak List 1. d Examine the spectra to see which fragment ions are produced at which collision energies. e Repeat steps a-c until all the product ions are identified. 	 The m/z 155.7 product ion is the most abundant of any product ion and the highest signal is recorded at 15 V. This means that a good choice for the MRM for sulfamethizole would be 271.0 > 155.7 when the collision energy is around 15 V. The peak may not be labeled if the peak is too wide. 	



f Click the Close Data File icon in the main toolbar, and click Close when the dialog box containing the list of data files pops up.

The product ions appear to be: Sulfamethizole-271.0 > 155.7 Sulfamethazine-279.0 > 185.8 Sulfachloropyridazine-285.0 > 155.7 Sulfadimethoxine-311.0 > 155.7

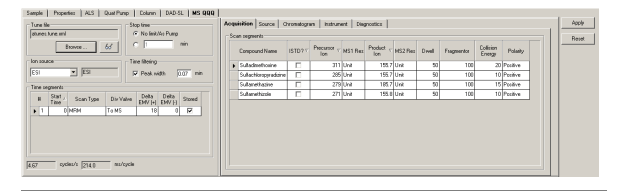
•

Task 5. Find optimum collision energy for MRM acquisition

Task 5. Find optimum collision energy for MRM acquisition

In this task, you set up MRM acquisition methods for the sulfa drugs for different collision energies. By examining the spectra and comparing peak intensities, you determine the optimal collision energy settings for the compounds.

Steps	Detailed Instructions	Comments	
 Set up three MRM acquisition methods. Use all the MS parameters in the example below except for the collision energy value. Use collision energies of 10, 15 and 20. Save methods as <i>iii</i>Sulfamix MRM_xx.m, where <i>iii</i> are your initials and xx is the collision energy. 	 a Click the MS QQQ tab. b Set Scan Type to MRM. c Enter all MS parameters shown in the example below except for the collision energy value. d In the collision energy column, type 10 for each compound. e Save the method as <i>iii</i>Sulfamix MRM_10.m. f Repeat step d and step e for collision energies of 15, 20, saving the methods as <i>iii</i>Sulfamix MRM_xx.m, where <i>iii</i> are your initials and xx is the collision energy. 	 Because the largest peaks were produced with a collision energy of 15 in the previous exercise, you will look at only those collision energies to either side of 15. 	



Task 5. Find optimum collision energy for MRM acquisition

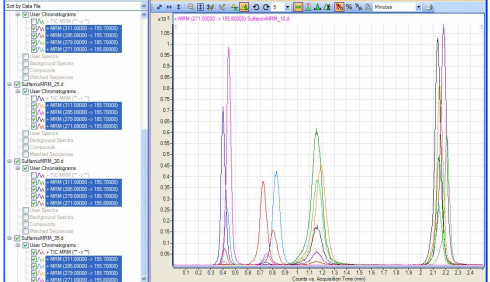
OK

Cancel

Steps	Detailed Instructions	Comments		
 2 Set up and run the worklist (optional). Specify the data files as <i>iii</i>Sulfamix MRM_<i>xx.</i>d, where <i>iii</i> are your initials and <i>xx</i> is the collision energy. 	 a Click the Worklist icon if necessary to make sure the worklist is visible. b Add three samples to the worklist for collision energies 10, 15, 20. c Mark the checkbox to the left of the Sample Name for each of the three samples. d Click Run > Worklist. 	• This step is optional because you can use the six example data files in the next step.		
 3 Compare the compound transition intensities at different collision energies. • Open the MRM data files: SulfamixMRM_10.d SulfamixMRM_15.d SulfamixMRM_20.d • Set the MRM chromatogram extraction parameters as shown at right for all transitions. • Disable the TICs for clarity and examine the peak intensities. • Compare the intensities of each compound transition obtained at one collision energy with the same compound transition obtained at one collision energy with the same compound transition obtained at another collision energy. (Do this in Overlaid Mode with all the MRM chromatograms.) • Close the data files but don't save results. • Refer to Table 4 on page 33 for optimal method settings for each compound. 		 Why a spectrum for MRM? It's a feature of the program to show spectra even for MRM experiments and can be quite handy for comparing relative intensities of product ions generated from the same precursor. 		

Task 5. Find optimum collision energy for MRM acquisition

Steps	Detailed Instructions	Comments		
	 h Click the Overlaid Mode icon, i Compare peak intensities for each compound transition in each data file in the Chromatogram Results window. 	 Compare the colors shown in Chromatogram Results with the color next to the MRM transition name in the Data Navigator. You can also right-click the Chromatogram Results window header and compare the colors of the chromatograms to the colors of the titles in the shortcut menu. 		
🏠 Data Navigator	× Chromatogram Results	×		
Sort by Data File		s 💌 🛃		



Unless you decide to acquire MRMs at lower collision energies, you should find that the optimal method settings are as shown in Table 4.

- j Click the **Close Data File** icon in the main toolbar, and click **Close** when the Close Data File dialog box appears.
- You now have all the information you need to do an MRM acquisition experiment of the sulfa drug mixture. Consider doing at least one more run with those settings.

•

Compounds	MRM Transition	Fragmentor	Collision
Sulfamethizole	271.0 > 155.8	100 V	10
Sulfamethazine	279.0 > 185.7	100	15
Sulfachloropyradizine	285.0 > 155.7	100	10
Sulfadimethoxine	311.0 > 155.7	100	20

Task 1. Create a batch file from an existing MRM data file

Exercise 2 – Develop a Dynamic MRM acquisition method from an MRM acquisition data file

The purpose of this exercise is to create a Dynamic MRM method from an acquired MRM data file for PM-34 Pesticides with the correct retention times for Dynamic MRM using the Quantitative Analysis program. All transitions in the MRM method must have the same polarity.

For this exercise, you have three main tasks:

- "Task 1. Create a batch file from an existing MRM data file" on page 34
- "Task 2. Print a report in the Quantitative Analysis program" on page 38
- "Task 3. Create a Dynamic MRM method using the results from the report" on page 39

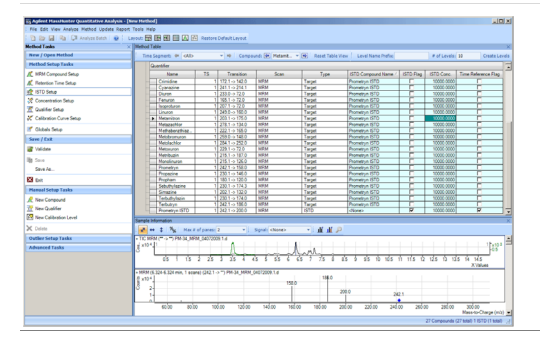
Task 1. Create a batch file from an existing MRM data file

In this exercise, you create a batch and a method from an existing MRM data file.

Steps		Detailed Instructions		C	Comments	
1	Open the Quantitative Analysis program and create a batch file with one sample file, PM-34_MRM_04072009.1.d. • Copy the data file PM-34_MRM_04072009.1.d from the installation disk to the \MassHunter\Data\MRM_to _DMRM folder.	b c d	Double-click the QQQ Quantitative Analysis icon. Click File > New Batch. Navigate to the \MassHunter\Data\ MRM_to_DMRM folder. Type MRM_to_DMRM in the Batch Name text box. Click OK. Click File > Add Samples. Select the file PM-34_MRM_04072009.1.d. Click OK.	•	The file PM-34_MRM_04072009.1.d is on the installation disk in the \Support\Data folder. Copy this entire folder to the \MassHunter\Data\ MRM_to_DMRM folder.	
2	Create a method for that batch using MRM data.		Click Method > New > New Method from acquired MRM data. Select the PM-34_MRM_04072009.1.d data file. Click OK.			

Task 1. Create a batch file from an existing MRM data file

Steps	Detailed Instructions	Comments	
 Add the compound Prometryn ISTD with the following information:. Time Segment: 1 Transition: leave blank Scan: MRM Type: ISTD Precursor lon: 242.1 Product lon: 200.0 Retention time: 6.324 minutes 	 a Select New Compound in the Manual Setup Tasks section in the Method Tasks pane. b Enter the Name, Time Segment, Scan, and Type for this new compound. c Select MRM Compound Setup under Method Setup Tasks in the Method Tasks pane. d Enter the Precursor Ion, Product Ion and RT (Retention Time). e Select ISTD Setup in the Method Setup Tasks section in the Method Tasks pane. f Mark ISTD Flag and Time Reference Flag for Prometryn ISTD. g Select Prometryn ISTD as the ISTD Compound Name for all compounds except for Prometryn ISTD. h Type 10000 as the ISTD Conc. 		



Exercise 2 – Develop a Dynamic MRM acquisition method from an MRM acquisition data file

Task 1. Create a batch file from an existing MRM data file

Exercise 2 – Develop a Dynamic MRM acquisition method from an MRM acquisition data file

Task 1. Create a batch file from an existing MRM data file

teps		Detailed	Instructi		Comments				
						•			
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P ISTD Setup	- Diuron	1 233.0->72.0	MRM	Target	Linear	Include	None		
	Fenuron	1 165.1 -> 72.0	MRM	Target	Linear	Include	None	_	
oncentration Setup	- Isoproturon	1 207.1 -> 72.0 1 249.0 -> 160.0	MRM	Target	Linear	Include	None	_	
Cuslifier Setup	Linuron Metamitron	1 203.1 -> 175.0	MRM	Target Target	Linear	Include	None	_	
Calibration Curve Setup	Metazachlor	1 278.1 -> 134.0	MRM	Target	Linear	Include	None	_	
Camadon Carle Seup	Methabenzthiaz	1 222.1 -> 165.0	MRM	Target	Linear	Include	None	-	
ኛ Globals Setup	Metobromuron	1 259.0 -> 148.0	MRM	Target	Linear	Include	None	_	
Save / Exit	Metalachior	1 284.1 -> 252.0	MRM	Target	Linear	Include	None		
	Metoxuron	1 229.1 -> 72.0	MRM	Target	Linear	Include	None		
Validate	Metribuzin	1 215.1 -> 187.0	MRM	Target	Linear	Include	None		
bi Save	Monolinuron	1 215.1 -> 126.0	MRM	Target	Linear	Include	None	_	
	Prometryn	1 242.1 -> 158.0	MRM	Target	Linear	Include	None	_	
Save As	Propazine Propham	1 230.1 -> 146.0 1 180.1 -> 120.0	MRM	Target Target	Linear	Include	None	_	
X Exit	Sebuthylazine	1 230.1 -> 174.3	MRM	Target	Linear	Include	None	_	
	Simazine	1 202.1 -> 132.0	MRM	Target	Linear	Include	None	-	
Manual Setup Tasks	Terbuthylazin	1 230.1 -> 174.0	MRM	Target	Linear	Include	None		
R New Compound	Terbutryn	1 242.1 -> 186.0	MRM	Target	Linear	Include	 None 		
New Qualifier								-	
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New Calibration Level New Calibration Level Outlier Setup Tasks	* TIC MRM ("->") PM-34_MRM_0 2 ×10 *]1 3 2.5-		45 5 55 6		8.5 9 9.5 10	10.5 11 11.5 12 12		-2	
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New Calibration Level Delete Outlier Setup Tasks	* TIC MRM (* -> *) PM-34_MRM (* * 10 4 1 3 2.5 0 * MRM (\$ 324-6 324 min, 1 scans) (* * 10 4 * 10 5 * 1 5	2'5 3 3'5 4		6.5 7 7.5 8 18 6 .0	85 9 95 10	10.5 11 11.5 12 12		-2	
New Calibration Level Colote Outlier Setup Tasks	TIC MRM (* -> *) PM-34_MRM (* * * *) PM-34_MRM (* * * *) PM-34_MRM (* * * * * * * * * * * * * * * * * * *	2'5 3 3'5 4	_04072009.1.d	6.5 7 7.5 8 186.0 0	8.5 9 9.5 10			-2	
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New Calibration Level Delete utlier Setup Tasks	+ TIC MRM ("→ ") PM-34_MRM (2'5 3 3'5 4 242.1 -> "") PM-34_MRM	_04072009.1.d	65 7 75 8 1860	200.0	242.1	X	-2 (5 falues	
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New Calibration Level Delete utlier Setup Tasks	+ TIC MRM ("→ ") PM-34_MRM (2'5 3 3'5 4 242.1 -> "") PM-34_MRM	_04072009.1.d	65 7 75 8 1860	200.0	242.1 240.00 260.00	280.00 300.0	-2 is plues -Charge (m/2)	

- errors, if necessary.
 - c Click Method > Save As.
 - d Enter MRM to DMRM.
 - e Click OK.
 - f Click Method > Exit.
 - g Click Yes to apply the method to the batch.
- 6 Analyze and save the batch.
- a Click Analyze > Analyze Batch. **b** Click **File > Save Batch**.

Task 2. Print a report in the Quantitative Analysis program

Task 2. Print a report in the Quantitative Analysis program

In this task, you print a report using the MRM_TO_DMRM.xltx template that is included on the Acquisition for Triple Quad installation disk, in the $\Support\Data$ folder.

Steps		D	etailed Instructions	Comments				
1	Print a report using the template MRM_to_DMRM.xltx.	a b c d e f g h	Click Report > Generate . The system displays the Report dialog box. Click Generate report results file . Specify the default destination directory for saving Excel reports in the Report folder text box; for example, \ <i>MassHunter\Data\</i> <i>MRM_to_DMRM\QuantReports\</i> <i>MRM_to_DMRM</i> . Click Add. Select MRM_to_DMRM.xltx. Click Open. Click OK.	•	Copy the MRM_to_DMRM.xltx template from the \Support\Data folder on the installation disk. For this report, you do not need to print the report, so Printer is set to <none> and Publish Format is <none>.</none></none>			
2	Check the status of the report using the Queue Viewer program.	a b c						

determine the optimal collision energy settings for the compounds.

Task 3. Create a Dynamic MRM method using the results from the report

In this exercise, you print a report using the MRM_TO_DMRM.xltx template that is included on the Acquisition for Triple Quad installation disk, in the $\Support\Data$ folder.

determine the optimal collision energy settings for the compounds.

S	Steps		etailed Instructions	Comments				
1	Open the file MRM_to_DMRM.xlsx in Excel.	a b	Navigate to the QuantReports folder in the batch. Right-click the new file and click Open .	•	In this example, the batch is in the \ MassHunter\Data \ MRM_to_DMRM folder.			
2	 Modify the results file. Delete the column Dwell. Delete the column QualFlag. Delete the empty row. Delete the second header row 	a b c d e f g h i	the program. Click the Dwell column header. Right-click the header and click Delete . Click the QualFlag column header. Right-click the header and click Delete . Click any empty row. Right-click the row and click Delete .					
3	 Copy the results to the clipboard. Copy only the results. Do not copy the header information. 	a b	Select the results from A3 to M55 . Right-click the selected area and click Copy to Clipboard .					

Exercise 2 – Develop a Dynamic MRM acquisition method from an MRM acquisition data file

Task 3. Create a Dynamic MRM method using the results from the report

;			D	etailed	Instruc	tions	;	Comments							
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7 Chloroxuron	FALSE	291.1	unit	72.0	unit	_	120	2	25	7.372	10.00	Positive	15792	4021	
Chlorpropham	FALSE	214.1	unit	172.0	unit	X	Cut			7.604	10.00	Positive	701	253	
Crimidine	FALSE	172.1	unit	142.0	unit					0.975	10.00	Positive	429	285	
0 Cyanazine	FALSE	241.1	unit	214.1	unit		Copy			5.550	10.00	Positive	14858	3644	
1 Diuron	FALSE	233.0	unit	72.0	unit	1	Paste			6.336	10.00	Positive	4236	955	
2 Fenuron	FALSE	165.1	unit	72.0	unit		Paste Special			3.654	10.00	Positive	29280	4087	
3 Isoproturon	FALSE	207.1	unit	72.0	unit		Insert			6.391	10.00	Positive	40483	10488	
4 Linuron	FALSE	249.0	unit	160.0	unit					7.132	10.00	Positive	1695	461	
5 Metamitron	FALSE	203.1	unit	175.0	unit		Delete			3.495	10.00	Positive	8127	1068	
6 Metazachlor	FALSE	278.1	unit	134.0	unit		Clear Contents			6.708	10.00	Positive	31176	7606	
7 Methabenzthiazuron	FALSE	222.1	unit	165.0	unit		Filter			5.981	10.00	Positive	20503	5612	
8 Metobromuron	FALSE	259.0	unit	148.0	unit		-			6.436	10.00	Positive	1744	498	
9 Metolachior	FALSE	284.1	unit	252.0	unit		Sort			7.975	10.00	Positive	12218	3131	
0 Metoxuron	FALSE	229.1	unit	72.0	unit		Insert Comment			5.108	10.00	Positive	14192	3558	
1 Metribuzin	FALSE	215.1	unit	187.0	unit	_				5.458	10.00	Positive	10617	2647	
2 Monolinuron	FALSE	215.1	unit	126.0	unit	2	Eormat Cells			6.199	10.00	Positive	2416	734	
3 Prometryn	FALSE	242.1	unit	158.0	unit		Pick From Drop-down Li	t		6.304	10.00	Positive	126951	29244	
4 Propazine	FALSE	230.1	unit	146.0	unit		Name a Bange			6.996	10.00	Positive	48092	11249	
5 Propham	FALSE	180.1	unit	120.0	unit		Hyperlink			6.570	10.00	Positive	2096	537	
6 Sebuthylazine	FALSE	230.1	unit	174.3	unit	20			_	7.157	10.00	Positive	134285	33543	
7 Simazine	FALSE	202.1	unit	132.0	unit		120		20	5.376	10.00	Positive	13297	3317	
8 Terbuthylazin	FALSE	230.1	unit	174.0	unit		120		15	7.157	10.00	Positive	134285	33543	
9 Terbutryn	FALSE	242.1	unit	186.0	unit		120		15	6.336	10.00	Positive	191300	43892	
0 Atrazine	FALSE	216.1	unit	132.0	unit		120		20	6.233	10.00	Positive	6405	1696	
1 Atrazine-desethyl	FALSE	188.1	unit	104.0	unit		120		25	4.047	10.00	Positive	4013	753	
2 Atrazine-desethyl-desisoprop		146.0	unit	68.0	unit		120		20	0.477	10.00	Positive	300	142	
3 Chlorotoluron	FALSE	213.1	unit	140.0	unit		120		20	6.056	10.00	Positive	1093	263	
4 Chloroxuron	FALSE	291.1	unit	218.0	unit		120		25	7.372	10.00	Positive	853	229	
5 Chlorpropham 6 Cyanazine	FALSE	214.1	unit	154.0	unit		80		15		10.00	Positive	692	230	
6 Cyanazine 7 Diuron	FALSE	241.1	unit	104.0	unit		120		25	5.550	10.00	Positive	2809	713	
8 Fenuron	FALSE	233.0 165.1	unit	160.0 120.0	unit		120		20	6.336 3.654	10.00	Positive Positive	389 1519	91 255	
9 Isoproturon	FALSE	207.1	unit	120.0	unit		120		10	6.391	10.00		5374	1454	
	FALSE	207.1	unit	165.0			120		15	6.391	10.00	Positive	1473	402	
0 Linuron 1 Metamitron					unit		120		20	7.132		Positive			
	FALSE	203.1	unit	104.0	unit						10.00	Positive	4051	539	
2 Metazachlor	FALSE	278.1	unit	210.0	unit		80		5	6.708	10.00	Positive	18841	4618	
3 Methabenzthiazuron	FALSE	222.1	unit	150.0	unit		120		20	5.981	10.00	Positive	2733	802	
4 Metobromuron 5 Metolachlor	FALSE	259.0	unit	170.0	unit		120		15	6.436 7.975	10.00	Positive	1538	412	
+ H Report 1 Report 2	91	764 1	1000	1.00.0	1000		1.00			7 975	10.00	Programming a	7784		b l

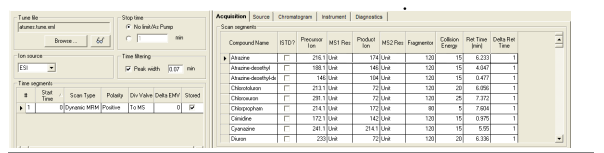
- 4 Switch to the Acquisition Program and paste the results into the Scan Segments table.
 - Verify that you only have one time segment.
- **a** Select Dynamic MRM as the Scan Type.
- **b** Select Positive as the Polarity.
- c Select ToMS as the Divert Valve.
- d Enter 200 for the Delta EMV value.
- e Mark the Stored check box.
- f Verify that only one time segment exists.
- **g** Select the first row in the Scan Segments Table on the
- h Right-click in the table and click **Paste** from clipboard.
- i Verify that the table was added properly.

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Exercise 2 – Develop a Dynamic MRM acquisition method from an MRM acquisition data file

Task 3. Create a Dynamic MRM method using the results from the report

Steps Detailed Instructions Comments



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In This Book

This exercise helps you use the Agilent 6400 Series Triple Quadrupole LC/MS system. In this guide, you acquire data and then analyze the results using the Qualitative Analysis program to learn how to develop an acquisition method.

If you have comments about this guide, please send an e-mail to feedback_lcms@agilent.com.

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m C}$ Agilent Technologies, Inc. 2009

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