Gas Chromatograph Mass Spectrometer

GCMS-QP Series
GCMS-TQ Series

Compatible with GCMSsolution Ver. 4.4 or later

Operation Guide

Basic Operation Guide

Read the instruction manual thoroughly before you use the product. Keep this instruction manual for future reference.
This page is intentionally left blank.
Read this Instruction Manual thoroughly before using the product.

Thank you for purchasing the GCMS-QP series and GCMS-TQ series gas chromatograph mass spectrometer.

This manual is intended to explain basic operations to first-time users. Read this manual thoroughly before using the product and operate the product in accordance with the instructions in this manual.

Also, keep this manual for future reference.

This manual assumes that the reader is knowledgeable of basic operations of Windows. For the operation of Windows, refer to the instruction manual that comes with that product.

Important

• If the user or installation location changes, ensure that this Instruction Manual is transferred with the product.
• If this manual is lost or damaged, immediately contact your Shimadzu representative to request a replacement.
• To ensure safe operation, read all Safety Instructions before using the product.
• To ensure safe operation, contact your Shimadzu representative if product installation, adjustment, or reinstallation (after the product is moved) is required.
Introduction

• All rights are reserved, including those to reproduce this manual or parts thereof in any form without written permission from Shimadzu Corporation.
• Information in this manual is subject to change without notice and does not represent a commitment on the part of the vendor.
• Any errors or omissions which may have occurred in this manual despite the utmost care taken in its production will be corrected as soon as possible, although not necessarily immediately after detection.
• Shimadzu Corporation is not responsible for errors or injuries resulting from following the instructions in this document.
• Shimadzu Corporation is not responsible for errors or injuries resulting from the use of equipment by the customer.
• The contents of PC hard drives may be lost due to unforeseen circumstances. In order to protect important data, be sure to make backup copies on a regular basis.
• Microsoft, Windows and Excel are registered trademarks of Microsoft Corporation in the United States and/or other countries.
Other company names and product names mentioned in this manual are trademarks or registered trademarks of their respective companies.
The TM and R symbols are omitted in this manual.
• Replacement parts for this product will be available for a period of seven (7) years after the product is discontinued.
Product Warranty

Shimadzu Corporation provides the following warranty for this product.

Details

1. Period: Please contact your Shimadzu representative for information about the period of this warranty.

2. Description: If a product/part failure occurs for reasons attributable to Shimadzu during the warranty period, Shimadzu will repair or replace the product/part free of charge. However, in the case of products which are usually available on the market only for a short time, such as personal computers and their peripherals/parts, Shimadzu may not be able to provide identical replacement products.

3. Limitation of Liability
   1) In no event will Shimadzu be liable for any lost revenue, profit or data, or for special, indirect, consequential, incidental or punitive damages, however caused regardless of the theory of liability, arising out of or related to the use of or inability to use the product, even if Shimadzu has been advised of the possibility of such damage.
   2) In no event will Shimadzu's liability to you, whether in contract, tort (including negligence), or otherwise, exceed the amount you paid for the product.

4. Exceptions: Failures caused by the following are excluded from the warranty, even if they occur during the warranty period.
   1) Improper product handling
   2) Repairs or modifications performed by parties other than Shimadzu or Shimadzu designated companies
   3) Product use in combination with hardware or software other than that designated by Shimadzu
   4) Computer viruses leading to device failures and damage to data and software, including the product’s basic software
   5) Power failures, including power outages and sudden voltage drops, leading to device failures and damage to data and software, including the product’s basic software
   6) Turning OFF the product without following the proper shutdown procedure leading to device failures and damage to data and software, including the product’s basic software
   7) Reasons unrelated to the product itself
   8) Product use in harsh environments, such as those subject to high temperature or humidity levels, corrosive gasses, or strong vibrations
   9) Fires, earthquakes or any other act of nature, contamination by radioactive or hazardous substances, or any other force majeure event, including wars, riots, and crimes
   10) Product movement or transportation after installation
   11) Consumable items
       Note: Recording media, such as floppy disks and CD-ROMs are considered consumable items.

* If there is a document such as a warranty provided with the product, or there is a separate contact agreed upon that includes warranty conditions, the provisions of those documents shall apply.

The warranty period for products with special specifications or for system products is specified separately.
About This Operation Guide

## Notation

This operation guide uses the notation described below.

<table>
<thead>
<tr>
<th>Notation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="CAUTION" /></td>
<td>Indicates a potentially hazardous situation which, if not avoided, may result in minor to moderate injury or equipment damage.</td>
</tr>
<tr>
<td><img src="image" alt="NOTE" /></td>
<td>Indicates additional information that is provided to ensure the proper use of this product.</td>
</tr>
<tr>
<td><img src="image" alt="Reference" /></td>
<td>Indicates the location of related information.</td>
</tr>
<tr>
<td><img src="image" alt="Hint" /></td>
<td>Indicates information provided to improve product performance.</td>
</tr>
<tr>
<td>![ ]</td>
<td>Indicates items displayed on the screen, such as buttons, menu selections, settings, windows, and icons. Example: Click [OK].</td>
</tr>
</tbody>
</table>
To ensure safe product operation, read these important safety instructions carefully before use and follow all DANGER, WARNING and CAUTION instructions given in this section.

**Safety Precautions**

**WARNING**

- For repairs, contact Shimadzu or your Shimadzu representative. Not doing so could cause a fire, electrical shock, or injury.

- Do not modify or disassemble the instrument without the express approval of an authorized Shimadzu representative. Doing so could cause an accident from electric shock or a short circuit. It could also cause injury or instrument failure.

- Read the instruction manual thoroughly before handling or operating the equipment, and be sure to following the procedures described. Not handling the equipment as described is potentially dangerous.

**Installation Site Precautions**

**WARNING**

- The solvents used with the gas chromatograph mass spectrometer may be flammable or toxic. Install the product in a well-ventilated room. Otherwise, solvent vapors may cause poisoning, or ignite and cause a fire.

- Do not install this instrument in a location with flammable or explosive gases or liquids. Because this product is not designed to be explosion proof, doing so could cause a fire or explosion.

- Do not place flammable materials near the column oven exhaust at the back of the instrument, as they could ignite and cause a fire.

- The lab table or other surface on which this instrument is installed should be level, stable, and sufficiently strong to support the instrument's weight. Otherwise, the unit could tip over or fall off the surface.

**NOTE**

Do not install the instrument in a location with corrosive gases, gases containing organic solvents, halogen compounds, or siloxanes, oil mist, or high levels of debris/dust. The instrument performance could be affected and its service life hortened.

Do not operate the instrument in an environment where condensation may form. Doing so could cause it to malfunction.
High-Pressure Gas Precautions

**WARNING**

- A high-pressure gas cylinder will be used to supply the carrier gas. When handling the gas cylinders, observe the following suggestions.
  - Keep gas cylinders in a well-ventilated area outside of the instrument installation site. Avoid exposure to direct sunlight. Use lines to transport the gas from the cylinders to the instrument. For flammable gases, this precaution is required by law.
  - Do not place the high-pressure gas cylinder in a location where the temperature can exceed 40 °C.
  - Choose an instrument installation site with sufficient ventilation, and include checking for gas leaks using an electronic gas leak detector in your daily inspection procedure. Do not smoke or use open flames within 5 m of the instrument when using highly combustible gases, such as acetylene and hydrogen, or potentially combustible gases, such as oxygen and nitrous oxide. Install and maintain effective fire extinguishers.
  - Secure the high-pressure gas cylinder with a cylinder stand or chain, etc., so that it does not fall over.
  - Be sure to use a pressure release valve specified as "not to be used with oil." Also, do not use a pressure release valve having piping, that is in contact with gas, whose inner surface is oily.
  - When finished using the gas, immediately close the main cylinder valve.
  - Verify that the pressure gauges are functional at least once every three months.
  - Warning signs (adhesive aluminum plates) are available to indicate hydrogen gas use. Ask your Shimadzu representative for more details. Signs are supplied free of charge to sites in which they are mandatory.
  - Legal authorization is required to use cylinders with a capacity of 300 m3 or greater.

Operation Precautions

**WARNING**

- Always wear safety glasses or goggles when handling solvents. If solvent gets into the eyes, blindness could result. Should solvent get into the eyes, immediately flush with large amounts of water and seek medical attention.
- Do not place solvents near PCs, printers or other instruments, as fire or instrument damage could result.
- Do not use flammable sprays (hair sprays, insecticide sprays, etc.) near this instrument, as they could ignite and cause a fire.
Handling Emergencies

The following measures should be taken in the event of an emergency such as a malfunction of the gas chromatograph mass spectrometer.
Take adequate precautions and contact your Shimadzu representative as necessary before resuming use of the instrument.

**Emergency Shutdown Procedure**

1. Turn OFF the gas chromatograph mass spectrometer.
2. Turn OFF all accessories.
3. Close the valves for the pipes supplying carrier gas, CID gas, hydrogen, and air.
4. Disconnect the power supply.
   - If the power cable is attached to a switchboard, turn OFF the switchboard.
   - If the power cable is plugged into an outlet, unplug the cable.
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Two operation guides, Basic Operation Guide and Method Development Guide, are available for the GCMS-QP series and TQ series models. This document covers the basic operations related to the QP series and TQ series.

<table>
<thead>
<tr>
<th>Name</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic Operation Guide</td>
<td>Covers the procedures to perform analysis using the existing method files for the GCMS-QP series or TQ series models and to create new method files (for Scan mode or SIM mode).</td>
</tr>
<tr>
<td>Method Development Guide</td>
<td>Mainly covers the procedures to create method files (for SIM mode or MRM mode, etc.) using the Smart database.</td>
</tr>
</tbody>
</table>

**NOTE**

Icons and windows for functions that can only be used on the TQ series, QP2100 Ultra, QP2100 SE or QP2020 will not be displayed on the software if the GCMS model used is QP2010, QP2010 Plus or PARVUM2.

![TQ]

Indicates the procedure that can be used on the GCMS-TQ series models.

![QP]

Indicates the procedure that can be used on the GCMS-QP2010 Ultra, GCMS-QP2010 SE or GCMS-QP2020 models.

### 1.1 Programs

GCMSsolution is made up of the programs described below. Select the program that is appropriate for the purpose (e.g., analysis or data processing).

<table>
<thead>
<tr>
<th>Icon</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GCMS Real Time Analysis</td>
<td>GCMS Real Time Analysis</td>
<td>Used to start up and shut down the instrument, make configuration settings, and perform analysis.</td>
</tr>
<tr>
<td>GCMS Analysis Editor</td>
<td>GCMS Analysis Editor</td>
<td>Used to create and edit method files and batch files during analysis.</td>
</tr>
<tr>
<td>GCMS Postrun Analysis</td>
<td>GCMS Postrun Analysis</td>
<td>Used to perform qualitative and quantitative processing, print reports, and perform other tasks involving data processing.</td>
</tr>
</tbody>
</table>
1.2 Routine Analysis Operation Flowchart

1. Prepare for analysis
   • Maintenance
   • Auto tuning
   • Create folder

2. Acquire data

3. Analyze data
   • Qualitative analysis
   • Quantitative analysis
1.3 Flowchart of Qualitative and Quantitative Analyses

Start up GCMS system
- Switch the power ON
- Specify maintenance and configuration settings, etc.
- Start up vacuum system
- Check for vacuum leaks

Execute autotuning

Qualitative Analysis

1. Create method file
   - Specify instrument parameters
     - Specify (Q3) Scan mode
     - Specify similarity search parameters

2. Acquire data

3. Analyze data
   - Display mass spectrum
   - Perform similarity search
   - Register displayed spectrum

4. Print report
   - Edit report
   - Output report

Quantitative Analysis

1. Perform qualitative analysis (standard sample)
   - Use standard samples to confirm retention times for target compounds

2. Create method file
   - <Quantitative analysis using (Q3) Scan mode>
     - Create compound table
   - <Quantitative analysis using (Q3) SIM mode>
     - Create compound table
     - Create SIM table

3. Acquire data

4. Analyze data
   - Check and correct calibration curve
   - Calculate quantitative values for samples with unknown concentrations

5. Print report
   - Edit report
   - Output report

Shut down the instrument
- Shut down the vacuum system
- Switch the power OFF
2.1 Turning ON the Power

Switch ON any peripheral or accessory equipment connected to the system, before switching ON the main GCMS system.

Switch ON any sample pretreatment unit connected to the system before switching ON the GCMS system.

When using a TQ series model, open the CID gas (argon gas) supply valve and supply the CID gas required for measurements in MRM mode.

1. Turn ON the power to the GC.

2. Turn ON the power to the MS.

3. Turn ON the power to the PC, printer, and display.

4. Double-click the (GCMS Real Time Analysis) icon.
   The [GCMS Real Time Analysis] program starts.
Click [OK].

2.2 Layout of Operating Areas

<table>
<thead>
<tr>
<th>No.</th>
<th>Name</th>
<th>GCMS Program</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Title Bar</td>
<td>Real Time Analysis, Analysis Editor, Postrun Analysis, and Browser</td>
<td>Displays the name of the program, process, and file currently running or being processed.</td>
</tr>
<tr>
<td>2</td>
<td>Menu Bar</td>
<td>Real Time Analysis, Analysis Editor, Postrun Analysis, and Browser</td>
<td>Displays command menus corresponding to the window currently open.</td>
</tr>
<tr>
<td>3</td>
<td>Toolbar</td>
<td>Real Time Analysis, Analysis Editor, Postrun Analysis, and Browser</td>
<td>Displays command tool buttons corresponding to the window currently open.</td>
</tr>
<tr>
<td>4</td>
<td>Assistant Bar</td>
<td>Real Time Analysis, Analysis Editor, Postrun Analysis, and Browser</td>
<td>Command icons are arranged in order of typical operation sequence. The assistant bar is named according to the window that is currently open. For example, when the [Batch] window is open, the assistant bar is named the [Batch] assistant bar.</td>
</tr>
<tr>
<td>5</td>
<td>Instrument Monitor</td>
<td>Real Time Analysis</td>
<td>Displays analytical instrument parameter values in real time.</td>
</tr>
</tbody>
</table>

GCMS Real Time Analysis

GCMS Postrun Analysis
2.3 Inspecting Consumable Items and Maintenance Parts

Check the state of the GCMS consumable items using the procedure described below.

1. Move the mouse pointer over the icon for a consumable item in the instrument monitor to display the current state and the recommended replacement point for the corresponding item.

When a consumable item approaches its recommended maximum usage frequency, the background of the corresponding icon turns black to alert the user.

This note is shown when mouse pointer is moved over the septum icon. This means that the septum has been used 13 times out of a maximum 100 times.

NOTE
When replacing the analysis column, or when a consumable item has passed its recommended replacement point, perform maintenance with reference to "Appendix K Maintenance" P.104.

Depending on the analysis content, the appropriate replacement frequency may be greater than the recommended frequency.
2.4 System Configuration

Check and set the modules used for analysis using the procedures described below. If the column has been changed, enter the column information using the procedure in the [MS Navigator] window.

2.4.1 Setting the Modules Used for Analysis

1. Click the [System Configuration] icon on the [Real Time] assistant bar.

2. Check that the components shown in the [Modules Used for Analysis] area correspond to the actual modules in the GC/MS system that are to be used for the analysis.

If the modules to be used for current analysis do not correspond to the modules shown in this window, set as shown in the following example:

1. Select [AOC-20i+s] in the [Available Modules] area if for example, AOC-20i with AOC-20s are to be used for analysis.

2. Click to register the module in [Modules Used for Analysis].

3. Click [Set].
Setting the CID Gas

For the GCMS-TQ series model, the default is set to use CID gas (argon gas). Turn OFF the CID gas to perform analysis using only the Q3 Scan or Q3 SIM mode.

1. Double-click the (MS) icon under [Modules Used for Analysis].
2. Deselect the [Use CID Gas] checkbox.
3. Click [OK]. Returns to the [System Configuration] sub-window.
4. Click [Set]. The configuration has now been set to use no CID gas. From now on, parameters regarding CID gas will not be displayed.
2.5 Vacuum System Startup

Open the carrier gas cylinder valve to supply carrier gas.
If carrier gas is being controlled by accessory/peripheral equipment, use that equipment to supply carrier gas before starting the vacuum system.

1. Click the [Vacuum Control] icon on the [Real Time] assistant bar.
   The [Vacuum Control] window opens.

2. Click [Auto Startup].
   The vacuum system starts.

3. When [Completed] is displayed, click [Close].
2.6 Checking for Vacuum Leakage

1. Wait for 10 minutes after starting up the vacuum system.

2. Click the [Tuning] icon on the [Real Time] assistant bar.
   
   The [Tuning] window opens.

3. Click the [Peak Monitor View] icon on the [Tuning] assistant bar.
   
   The [Peak Monitor] window opens.
2.6 Checking for Vacuum Leakage

Check for vacuum leaks.

1. Click the arrow button in [Monitor Group] setting, and select [Water, Air] from the list.

2. Click (Filament ON/OFF) to turn ON the filament. Peaks will be displayed in the three windows.

3. Change the detector voltage gradually by clicking the up or down arrow buttons so that the peak height for $m/z$ 18 (water) corresponds to half the height of the display window.

4. Compare the peak heights for $m/z$ 18 (water) and $m/z$ 28 (nitrogen). Check that the peak height for $m/z$ 28 (nitrogen) is not more than twice that for $m/z$ 18 (water).

**NOTE**

If the peak height for $m/z$ 28 (nitrogen) is more than twice that for $m/z$ 18 (water), it is possible that there is an air leak. Search for the location of the leak. Refer to the System User's Guide for details on how to check for vacuum leaks.

5. Click (Filament ON/OFF) to turn OFF the filament.

5. Click button in the top-right corner to close the [Tuning] window.

The message [Save current tuning file?] is displayed. Click [No].
2 Starting GCMS

6 Click the [Top] icon on the assistant bar.

2.7 Autotuning

Wait for approximately 2 hours (before starting qualitative analysis) or 4 hours (before starting quantitative analysis) after starting up the vacuum system and then perform autotuning using the procedures described below.

Perform autotuning periodically, even with the vacuum system operating.

\[\textbf{NOTE}\]
Create calibration curves again after performing autotuning.

2.7.1 Setting Analysis Conditions

If no analysis conditions have been created, start from "2.7.2 Executing Autotuning" P.13.

If a method file is already created, parameters can be specified in the instrument according to the following procedure.

\[\textbf{NOTE}\]
However, parameters for an accessory or peripheral equipment, except for AOC-20 auto-injector/auto-sampler, cannot be specified by using the following procedure. When using an accessory/peripheral equipment, set the parameters on the equipment itself, or by using the software specific to that equipment/device.

1 Click the [Data Acquisition] icon on the [Real Time] assistant bar.
   The [Acquisition] window opens.

2 Click \(\text{file}\) (Open) on the toolbar.
   The [Open Method File] sub-window opens.
2.7 Autotuning

3 Select the method file to load, then click [Open].

The method file is loaded.

![Open Method File Window]

NOTE

If the message “The hardware configuration for this method is different from the current instrument configuration. The measurement condition in the method file is modified according to the current instrument configuration.” appears, click [OK] and click  (Save) on the toolbar.

4 Select [Download Initial Parameters] on the [Acquisition] menu.

The set parameters are transferred to the instrument. When the parameter values become equal to the settings, [GC: Ready] and [MS: Ready] are displayed.

![Download Initial Parameters Window]

2.7.2 Executing Autotuning

1 Click the [Tuning] icon on the [Real Time] assistant bar.

The [Tuning] window opens.
2 Click the [Peak Monitor View] icon on the [Tuning] assistant bar.
The [Peak Monitor] window opens.

The [Select Tuning Mode] sub-window opens.

**Select Tuning Mode appropriate for the application**

When using a TQ series, QP2010 Ultra, QP2010 SE or QP2020 model and creating a new tuning file, choose the tuning mode appropriate for the concentration level of target compounds being measured. Since the tuning file is created with an emission current corresponding to the selected mode, it enables measuring samples with an appropriate dynamic range.

- TQ series, QP2010 Ultra or QP2020: High concentration (20 µA), standard (60 µA, default), or high sensitivity (150 µA)
- QP2010 SE: High concentration (20 µA) or standard (60 µA, default)

**Perform Autotuning Even with CID Gas OFF**

When using a TQ series model, a tuning method can be chosen depending on the measurement mode.
2.7 Autotuning

Table: Estimated Time Required for Autotuning

<table>
<thead>
<tr>
<th>System Configuration</th>
<th>Tuning Condition</th>
<th>Acquisition Mode</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use CID Gas</td>
<td>Perform Autotuning Even with CID Gas OFF</td>
<td>Q3 Scan</td>
<td>Q3 SIM</td>
</tr>
<tr>
<td></td>
<td>Possible</td>
<td>Possible</td>
<td>Impossible</td>
</tr>
<tr>
<td></td>
<td>Possible *1</td>
<td>Possible *1</td>
<td>Possible</td>
</tr>
<tr>
<td></td>
<td>Possible</td>
<td>Possible</td>
<td>Possible</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(Recommended time)</td>
</tr>
</tbody>
</table>

*1 : High sensitivity analysis

For GCMS-QP series models, autotuning takes about 3 minutes.

4 Select the filament to be used.

5 Click the [Start Auto Tuning] icon on the [Tuning] assistant bar.

6 Enter a file name and click [Save].

Autotuning starts.

When autotuning is completed, a report is printed.
7 Click (Save) on the toolbar.

**NOTE**

If [Perform Autotuning Even with CID Gas OFF] is not selected, select [ON] in the [CID Gas] drop-down list to view the tuning results.

<table>
<thead>
<tr>
<th>Tuning Date</th>
<th>CID Gas</th>
</tr>
</thead>
<tbody>
<tr>
<td>2/5/2014 15:57:14</td>
<td>OFF</td>
</tr>
<tr>
<td>Acquisition Mode</td>
<td>Q3 Scan</td>
</tr>
<tr>
<td>Q1 Resolution</td>
<td>OFF</td>
</tr>
</tbody>
</table>
2.7.3 Checking Autotuning Results

Check the results of autotuning.

1. Check that the FWHM (full width at half maximum) values are in the range 0.5 to 0.7.
2. Check that the detector voltage does not exceed 2 kV.
3. Check that the relative intensity ratio for \( m/z \) 502 is at least 2% (for QP2010S and SE : 1%).
4. Check that the peak intensity for \( m/z \) 69 is at least twice that for \( m/z \) 28.

**NOTE**

If any irregularities are discovered above, possible causes could include a vacuum leak, poor column connections, or contaminated ion source.

See "Appendix K Maintenance" P.104 to implement corrective measures.
This chapter describes a series of routine analysis operations using the existing method files and batch files.

3.1 Preparing for Analysis

TQ QP

When the ecology mode is set in the [GCMS Real Time Analysis] program, click [Cancel].

1 Inspect consumable items and maintenance parts.
When a consumable item has passed its recommended replacement point, or when the column requires maintenance, perform maintenance with reference to "Appendix K Maintenance" P.104.

2 Perform autotuning.
Perform autotuning periodically (once every two weeks, for example) even with the vacuum system operating, referring to "2.7.2 Executing Autotuning" P.13. When quantitative analysis is to be performed, create calibration curves again after autotuning.

3 Create a new folder.
[Create New Project (Folder)] can be used to create a new folder at the same directory level as currently open in Data Explorer and to copy the target files.
3.2 Editing a Batch File

For routine analyses, it may be more convenient to load an exiting batch file and partially edit it. The following procedure describes how to edit information in specified rows(s) collectively.

1. **Click the [Batch Processing] icon on the [Real Time] assistant bar.**
   The [Batch Table] window opens.

2. **Click the [Batch Processing] icon on the [Real Time] assistant bar.**
   The [Batch Table] window opens.

   (Example of a batch table for qualitative analysis)

<table>
<thead>
<tr>
<th>Folder</th>
<th>VisList</th>
<th>Sample Name</th>
<th>Sample ID</th>
<th>Sample Type</th>
<th>Analysis Type</th>
<th>Method File</th>
<th>Data File</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>Methylene Chloride</td>
<td>0 Unknown</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>STD 50ppb</td>
<td>STD-3001</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>STD 10ppb</td>
<td>STD-3002</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>STD 100ppb</td>
<td>STD-3003</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>STD 1000ppb</td>
<td>STD-3004</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
</tbody>
</table>

   (Example of a batch table for quantitative analysis)

<table>
<thead>
<tr>
<th>Folder</th>
<th>VisList</th>
<th>Sample Name</th>
<th>Sample ID</th>
<th>Sample Type</th>
<th>Analysis Type</th>
<th>Method File</th>
<th>Data File</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>Methylene Chloride</td>
<td>0 Unknown</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>STD 50ppb</td>
<td>STD-3001</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>STD 10ppb</td>
<td>STD-3002</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>STD 100ppb</td>
<td>STD-3003</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>STD 1000ppb</td>
<td>STD-3004</td>
<td>T OC</td>
<td>Methylene Chloride</td>
<td>20131127</td>
<td></td>
</tr>
</tbody>
</table>
Add or delete rows depending on the number of samples being analyzed.

1. Click on the row number to be edited to highlight the whole row.

<table>
<thead>
<tr>
<th>Menus</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copy Row</td>
<td>Copies the selected row.</td>
</tr>
<tr>
<td>Add Row</td>
<td>Adds a row to the end.</td>
</tr>
<tr>
<td>Insert Row</td>
<td>Inserts a new row above the selected row.</td>
</tr>
<tr>
<td>Paste Row</td>
<td>Pastes the copied row.</td>
</tr>
<tr>
<td>Delete Row</td>
<td>Deletes the selected row.</td>
</tr>
</tbody>
</table>

2. Right-click on the selected row, and select the appropriate editing command from the menu that is displayed.

Drag the mouse from the row of the unknown sample to the row specified with serial numbers.

Select [Fill Down] on the [Edit] menu. The entire content of the first row is copied.
6. Select [Fill Series] on the [Edit] menu. Edited parameters will be appended with serial numbers. Modify the sample name, etc. if necessary.

![Fill Series and Fill Down options]

To collectively edit specified rows without changing other rows, right-click the cell in the first row and click [Fill Down] or [Fill Series].

7. Select [Save Batch File As] on the [File] menu.

8. Open the folder where the method file is saved, enter a name, and save the file.

9. Set the syringe rinse solvent and samples in the autosampler.
Click the [Start] icon on the [Batch] assistant bar.
Analysis starts. When the "Do you want to go into the ecology mode after batch processing ends?" message appears, click [Yes].

- To abort batch processing, click the (Stop) icon on the [Batch] assistant bar.
- To execute only specified rows, select the rows by clicking or dragging the mouse, then start the analysis.

For details on how to create data file names automatically or how to change settings for continuous data acquisition during the measurement, refer to "Appendix D Batch File" P.74.

3.3 Dana Analysis

Reference
Refer to "4.5 Analyzing Data" P.34 when performing qualitative analysis.
Refer to "5.3 Analyzing Data" P.54 when performing quantitative analysis.
Refer to "Appendix L Quantitative Browser" P.110 when quantitating multiple samples.
4 Qualitative Analysis

4.1 Selecting a Folder

1. Start up the [GCMS Real Time Analysis] program.

2. Click the (Data Explorer) icon on the toolbar to display Data Explorer.

3. Click (Project (Folder) Selection).
   The [Project (Folder) Selection] window opens.

4. Click the folder to be used.

5. Click [Close].
Creating a Folder

1. Click the desired hard drive or folder and then click [New Folder].
   The [Create New Folder] window opens.

2. Type a folder name and click [OK].
   A folder is created in the drive or folder selected in step 1, and the [Project (Folder Selection)] window returns.
4.2 Creating a Method File

Set the instrument (i.e., autosampler, GC, MS) parameters and similarity search parameters using the procedure described below.

1. Click the [Data Acquisition] icon on the [Real Time] assistant bar.
   The [Acquisition] window opens.


4.2.1 Setting Autosampler Parameters

1. Click the [Sampler] tab and specify the number of rinses appropriate for the sample.

4.2.2 Setting GC Parameters

1. Click the [GC] tab and set the analysis conditions.

2. Input an initial temperature for the column oven (40 to 100 °C).

3. Input an injection temperature based on consideration of the boiling point of the target compound (200 to 300 °C).

4. Select [Split] or [Splitless].

- **NOTE**

  Selecting Injection Mode
  
  - Split: Select this mode if the concentration of the target compound is high. As a rough guideline, select this mode when the target compound concentration is greater than 10 ng/uL.
  
  - Splitless: Select this mode if the concentration of the target compound is low. As a rough guideline, select this mode when the target compound concentration is less than 10 ng/uL.

5. Select [Pressure] when the method calls for a constant pressure mode, and select [Linear Velocity] when the method calls for a constant linear velocity mode for the carrier gas. When no reference method is available, select [Linear Velocity].

6. When no reference method is available, refer to the table "Typical Pressure Settings for Carrier Gas" to set an initial value for the pressure. The linear velocity will be set automatically.

   **Typical Pressure Settings for Carrier Gas**

<table>
<thead>
<tr>
<th>Middle bore capillary column (I.D. 0.25 mm)</th>
<th>Semi-wide bore capillary column (I.D. 0.32 mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 m</td>
<td>60 m</td>
</tr>
<tr>
<td>75 to 150 kPa</td>
<td>100 to 250 kPa</td>
</tr>
</tbody>
</table>

7. If "Split" is selected as the injection mode, enter a split ratio. If "Splitless" is selected, enter "-1.0".

8. Set appropriate conditions for separating the target compound from other peaks.
4.2.3 Setting MS Parameters

1 Click the [MS] tab and set the analysis conditions.

![Image of MS parameters setting](image)

1 Input [Interface Temp.] (200 to 300 °C).

2 Input [Start Time] and [End Time] according to the note below.

**NOTE**

In the absence of information about the elution time of the solvent peak, set [Start Time] to zero minutes, and set [End Time] to the [GC Program Time] value. After one analysis of a standard sample or the solvent, and obtaining the solvent peak profile, change the [Start Time] to a time after the end of the solvent peak (see the figure shown on page 24).

3 Click [Relative to the Tuning Result].

If peak intensity is too low, change the value within the range +0.1 to +0.3., as necessary.

Increasing the detector voltage by 0.1 kV increases the peak intensity by 2 to 4 times.

4 Input a value that is 0.5 minutes less than the [Start Time] setting. (If the resulting value is less than zero, enter "0".)

Relationship between Start Time and Solvent Elution Time

5 Select [Scan]. (For TQ series models, select Q3 scan.)

6 Enter the mass range to be measured, where [Start m/z] is the lower mass limit, and [End m/z] is the upper mass limit. The typical value for [Start m/z] is 35, and the typical value of [End m/z] is the highest molecular weight of the target compounds in the sample plus some margin of error (+15).
4 Qualitative Analysis

4.2.4 Setting Similarity Search Parameters


2. Click the [Similarity Search] tab and set the search conditions.

   3. Click .
   4. Click .

   2. Open the library to be used.
2. To remove a library from the selection, highlight the library file name by dragging the mouse over it, then press the [Delete] key.
3. Select [Do not include duplicate hits].
4. After completing the settings, click [OK] to return to the original window.

### 4.2.5 Saving the Method File


![Image of Save Method File As dialog box]

2. Enter a file name and click [Save].

![Image of file selection dialog box]

### 4.3 Repeating Autotuning

If autotuning has not been performed under the analysis conditions, perform the procedures described under "2.7 Autotuning" P.12.
4.4 Sequential Analysis

Create a batch file necessary for qualitative analysis and perform sequential analysis using the procedures described below.

4.4.1 Creating a Batch File

   The [Batch Table] window opens.


   The [Batch Table Wizard] window opens.
4 With the Batch Table Wizard, make the appropriate settings and create a batch table.

1 Click [Unknown Only].
2 Click and specify the method file to be used.
3 Deselect both [Data Processing] items.
4 Click [Next].

5 Input [Vial #] and [Sample Count].
6 Input [Injection Volume].
7 Click [Next].
8 Enter [Data File Name].
   If the file name ends with a number, the files are named sequentially.
9 Click [Finish]. The batch table is displayed. Edit the batch table as required.

- Insert a row above the first row.
- Copy a row for the unknown sample and paste it on the first row.
- Edit the vial #, sample name and data filename.

It is recommended to measure the blank (solvent, etc.) before starting analysis.
4.4.2 Saving Batch Files

1. Select [Save Batch File As] on the [File] menu.

![Image of Save Batch File As dialog box]

2. Open the folder where the method file is saved, enter a name, and save the file.

![Image of folder and Save Batch File As dialog box]

4.4.3 Executing Sequential Analysis

1. Set the syringe rinse solvent and samples in the autosampler.

2. Click the [Start] icon on the [Batch] assistant bar.

Analysis starts.
4.5 Analyzing Data

Use the procedure described below to perform basic qualitative data processing for data measured in Scan mode, for examples, to display mass spectra, perform background subtraction, and perform similarity search.

1. Double-click the (GCMS Postrun Analysis) icon. The [GCMS Postrun Analysis] program starts.

2. Click the [Qualitative] icon on the [Postrun] assistant bar.

4.5.1 Loading Data Files

1. Double-click the data file to analyze. The data file opens. If the required folder is not found, refer to "4.1 Selecting a Folder" P.23.
4.5.2 Displaying Mass Spectra

1. Specify a range in the TIC window by dragging the mouse so that both the peak top and baseline are highlighted.

   Drag the mouse so that both the peak top and baseline are displayed. To undo the zoom, right-click in the MC window and select [Undo Zoom] on the pop-up menu.

2. Move the mouse pointer to the peak top and double-click.

3. Click (Spectrum Subtraction) on the toolbar.

4. Double-click at the background processing position.

With the following types of peaks, process the parts indicated by arrows as background.

Example 1) Example 2) Example 3)

If a red peak appears on a mass spectrum, it indicates that the peak is saturated. This spectrum has a different pattern (intensity ratio) from the mass spectrum for the target compounds. In such a case, click the left and right arrow buttons to select a mass spectrum that shows no red peaks.

Background spectrum can be subtracted from one of positions.
4.5.3 Searching for Similarity

1. Click the [Similarity Search] icon on the [Qualitative] assistant bar.
   The [Similarity Search Results] window opens.

2. Check the similarity search results.
   
   1. Use to switch between the mass spectra for the compounds found.
   2. Select the checkbox for the applicable compound to enter a compound name in the spectrum table.
   3. After checking the mass spectra, click (Register Target Spectrum to Spectrum Process Table).
      The mass spectrum is registered.
      By registering the target mass spectrum in the spectrum process table, you can re-check the similarity search results or output them in reports.
4.5.4 Editing the Spectrum Process Table

1. Click the [Qualitative Table] icon on the [Qualitative] assistant bar. The [Qualitative Table] window opens.

2. Click (Maximize).

To sort the spectrum process table in chronological order, click [Sort Table] on the [Edit] menu.

4

After check is complete, close the [Similarity Search Results] window.

- To edit a compound name, select the desired row and click [Edit Compound Name] on the [Edit] menu. Enter the compound name in the [Edit Compound Name] window displayed and click [OK].

- To delete a row in the spectrum processing table, select the desired row and click [Delete Rows] on the [Edit] menu.

5

Close the [Qualitative Table] window.

4.5.5 Saving Data Files

1

Click (Save) on the toolbar.
The qualitative table is saved in the data file.
4.6 Printing Qualitative Analysis Reports

It is convenient to use a template to create a report of analyzed data. Depending on the data, edit the area of the chromatogram to display in the report, or edit the number of compounds to be displayed in the report of similarity search results.

4.6.1 Loading Report Formats

1. Click the [Report] icon on the [Qualitative] assistant bar.
   The [Data Report] window opens.

   The [File New] window opens.

3. Select [Use Template] and select the format [Qualitative Analysis Report].

4. Click [OK].
   The [Qualitative Analysis Report] format opens.
4.6.2 Editing Report Formats


2. Click the [Chromato] tab.

3. In the [Area] area, deselect [Auto] for the X-axis and enter the time range.
4.6 Printing Qualitative Analysis Reports

4. Click [OK].

5. Click the next page icon on the toolbar to display the second page.

   The [GCMS Library Properties] window opens.

7. Click the [Result] tab.
Enter the [Maximum Compound Number] (maximum number of search results to display).

- Selecting [Print the Hit Compounds] prints the report for the maximum number of search results to display in order starting with the highest similarity.
- Selecting [Print Only Specified Compound] prints the report for the compounds selected for registration in the similarity search.

Click [OK].
4.6.3 Outputting Reports

1. Click the [Preview] icon on the [Data Report] assistant bar. The print preview window opens.

2. After checking the report content, click [Print] to print the report.

3. Click [Save] on the toolbar. The report is saved as part of the data file.
5.1 Creating a Method File

With reference to "4 Qualitative Analysis" P.23 analyze standard samples (including internal standard substances when using the internal standard method for analysis) and register the retention times and mass spectra of the target compounds in the spectrum process table.

5.1.1 Creating a Compound Table

1. Start the [GCMS Postrun Analysis] program and click the [Compound Table] icon on the [Postrun] assistant bar. The [Create Compound Table] window opens.

2. From Data Explorer, double-click the data file in which the spectrum process table for the target compounds was saved.

3. Click the [Wizard (New)] icon on the [Compound Table] assistant bar. The [Compound Table Wizard] window opens.
5.1 Creating a Method File

4. Select [Use current Spectrum Process Table] and then click [Next] in the [Compound Table Wizard 1/7] window.

5. Click [Next] in the [Compound Table Wizard 2/7] window.

6. Select a row in the table, check the mass spectrum for each compound, and click [Next] in the [Compound Table Wizard 3/7] window.

7. Specify the calibration curve type, the quantitative method, and other parameters as required, and click [Next].

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Explanation</th>
</tr>
</thead>
</table>
| 1   | Quantitative Method   | • External Standard: Quantitation is performed using a calibration curve obtained from the absolute quantity (concentration) and the area or height value of the target compound in a standard sample.  
    |                       | • Internal Standard: An internal standard is added to the sample, the sample is analyzed, and quantitation is performed using the relationship between the relative sensitivity and the quantitative ratio with respect to the internal standard compound. |
| 2   | Calculated by         | Select [Area] or [Height]. Normally, select [Area].                         |
| 3   | # of Calib. Levels    | Input the number of concentration levels of the calibration curve.          |
| 4   | Unit                  | Set the concentration unit used for reports.                               |
| 5   | Format of Concentration| Set the number of digits used to indicate concentrations.                  |
8 Make the appropriate settings for concentrations and measurement ions, and click [Next].

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Standard</td>
<td>Set the concentrations of the standard samples. If the concentration varies with the compound, make the necessary corrections after completing the wizard procedure.</td>
</tr>
<tr>
<td>2</td>
<td>Internal Standard</td>
<td>Set the concentration of the internal standard.</td>
</tr>
<tr>
<td>3</td>
<td># of Reference Ions</td>
<td>Input the number of reference ions used to perform peak identification.</td>
</tr>
<tr>
<td>4</td>
<td>Decimal for mass</td>
<td>Determine the number of decimal places for target ion and reference ion m/z values. Selecting [1 Decimal] increases the sensitivity level.</td>
</tr>
</tbody>
</table>

9 Set the type, compound name, target ion, and reference ion for each substance. After entering the required information for all the compounds, click [Next].

1 Change the compound displayed by changing the ID number.
2 Select [Target] in the [Type] list. Select [I.S.] when setting for an internal standard.
3 Change the type and m/z value.
   - To change the type, click cell for the type to be changed and select "Target Ion", "Ref. Ion", or "Not used".
   - To change the mass value, click in the cell containing that value, click the resulting arrow button, point your cursor at the desired peak on the mass spectrum, and then double-click it.

10 Click [Finish].
   A compound table is created. Correct the contents of the compound table as required.

11 Click \[\text{3D View}\] to set the compound table to the display mode.

\[\text{NOTE}\]
   To correct the compound table again, enter edit mode by clicking \[\text{Edit}\] at the top-right corner of the table.

12 Click the [Save Compound Table] icon on the [Compound Table] assistant bar.
   The method file that was used to acquire the data will be selected automatically.
Click [Save].
This completes the procedure for creating a quantitative method for Scan mode.

If greater sensitivity is required, use the following procedure to create a quantitative analysis method for the SIM mode.

5.1.2 Creating a SIM Table

1. Click the [Create MS Table [COAST]] icon on the [Compound Table] assistant bar.
The [Select Method File] window opens.

2. Enter a file name and click [Save].
The [Creation of Automatic MS Table [COAST]] window opens.
Click (Maximize) in the [Creation of Automatic MS Table [COAST]] window.

A SIM table is created automatically. Check the chromatogram and SIM table and, if necessary, modify the table with reference to the following procedure.

1. To ensure sufficient sensitivity:
   To ensure sufficient sensitivity, it is best to specify no more than 20 m/z values per row (i.e. per group).
   If necessary, modify the SIM table.

2. To edit table rows (i.e. groups):
   To edit table rows (i.e. groups), right-click on the desired row and select the following on the menu that appears.
   • Add Row : Adds a row to the bottom of the table.
   • Insert Row : Inserts a new row above the selected row.
   • Delete Row : Deletes the selected row.

3. To split groups:
   To split groups, use the following procedure. (Example: Splitting Group 3 into two groups)
   1. Click the third row of the SIM table.
   2. Right-click on the table and select [Insert Row].
   3. Click the inserted row and drag the mouse on the chromatogram to specify and enlarge the desired area.
   4. Click near the center of peaks labeled with compound names.
      Group 3 is divided into two groups.

When finished, click [OK].
A method is created for SIM mode quantitative analysis.
5.2 Sequential Analysis

Create a batch file necessary for quantitative analysis and perform sequential analysis using the procedure described below.

5.2.1 Creating a Batch File

   The [Batch Table] window opens.


   The [Batch Table Wizard] window opens.
4 Make the appropriate settings with the Batch Table Wizard and thereby create a batch table.

1. Select [Standard & Unknown].
2. Click and specify the method file to be used.
3. Select [Quantitative].
4. Click [Next].

5. Input [Vial #].
   The number of calibration points is loaded automatically from the method.
6. Input [Injection Volume].
7. Input [Average Count] (i.e., the number of repetitions).
8. Click [Next].
9 Enter [Data File Name].
   If the file name ends with a number, the files are named sequentially.
10 Click [Next].

11 Input [Vial #].
12 Input [Sample Count].
13 Input [Injection Volume].
14 Click [Next].
5.2 Sequential Analysis

15 Enter [Data File Name].
   If the file name ends with a number, the files are named sequentially.

16 Click [Finish].
   The batch table is displayed.

   It is recommended to measure the blank (solvent, etc.) before starting analysis.
   1. Insert a row above the first row.
   2. Copy a row for the unknown sample and paste it on the first row.
   3. Edit the vial #, sample name and data filename.

5.2.2 Saving Batch Files

1 Select [Save Batch File As] on the [File] menu.
5. Quantitative Analysis

2. Open the folder where the method file is saved, enter a name, and save the file.

5.2.3 Executing Sequential Analysis

1. Set the syringe rinse solvent and samples in the autosampler.

2. Click the [Start] icon on the [Batch] assistant bar.
   Analysis starts.

5.3 Analyzing Data

5.3.1 Checking and Correcting Calibration Curves

   The [Calibration Curve] window opens.
Double-click the method file used in analysis from Data Explorer.

Select a compound in the compound table and click the calibration curve level.
Check the calibration curve created and the chromatogram.

Reference
If no peaks are identified or detected, perform identification or peak integration with reference to "Manual Identification and Manual Peak Integration" P.57.
To change the method used to plot calibration curves, see "Appendix I Editing Parameters for Quantitative Analysis" P.96.
Only after correcting the calibration curves, click (Save) on the toolbar to save the method file.

Peaks that are detected in the chromatograms after automatic peak integration, will have peak detection marks (↑↓).

The detected peaks are subjected to identification based on the retention times and ion ratios (▼ peak identification mark).

<table>
<thead>
<tr>
<th>Chromatogram</th>
<th>Countermeasure</th>
</tr>
</thead>
<tbody>
<tr>
<td>No peaks are detected.</td>
<td>Perform manual peak integration. (P.58)</td>
</tr>
</tbody>
</table>
5.3 Analyzing Data

Manual Identification and Manual Peak Integration
If no peaks are identified or detected, perform identification or peak integration using the procedure described below.

Manual Identification

1. Right-click in a chromatogram and select [Manual Identification] from the displayed menu.

A bar is displayed.

<table>
<thead>
<tr>
<th>Chromatogram</th>
<th>Countermeasure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peaks are detected but different peaks are identified.</td>
<td>Perform manual identification. (P.57)</td>
</tr>
<tr>
<td><img src="image1" alt="Chromatogram" /></td>
<td></td>
</tr>
<tr>
<td>Peaks are detected and identified but peak integration is not performed</td>
<td>Perform manual peak integration. (P.58)</td>
</tr>
<tr>
<td>properly.</td>
<td></td>
</tr>
<tr>
<td><img src="image2" alt="Chromatogram" /></td>
<td></td>
</tr>
</tbody>
</table>
2 Click the peak to be identified.
The peak is identified.

Manual Peak Integration

1 Right-click in a chromatogram and select [Manual Peak Integrate...] from the displayed menu.
A bar is displayed.

2 Drag the mouse from the start point to the end point of the peak.
The [Select Base Line] window opens.
3 Select the baseline and click [OK].
The peak is integrated and identified.

![Baseline selection interface]

NOTE
The same process can be accomplished by performing the following operations on the chromatogram.

<table>
<thead>
<tr>
<th>Process</th>
<th>Operation</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Identification</td>
<td>[Shift] + [Ctrl] + right-click</td>
<td>Identifies integrated peaks.</td>
</tr>
<tr>
<td>Manual Peak Integration</td>
<td>[Shift] + right-click-drag</td>
<td>Connects start point and end point as baseline.</td>
</tr>
</tbody>
</table>

5.3.2 Re-quantifying after Correcting a Calibration Curve
After correcting a calibration curve, re-quantify the data for samples with unknown concentrations.

1 Click the [Batch Processing] icon on the [Postrun] assistant bar.
The [Batch Table] window opens.

3 Click the [Select Data File] icon on the [Batch] assistant bar. The [Select Data File] window opens.

4 Click the data file for sample with unknown concentrations, for which re-quantification is to be performed and click (Add). The data file is selected.

5 Click [OK].

6 A batch table is displayed. Assign a name to the batch file and save it.
5.3 Analyzing Data

5.3.3 Checking and Correcting Quantitation Results

Check the quantitation results for the samples with unknown concentrations.


2. Double-click the data file to be checked from Data Explorer.

   The data file being checked opens.

3. Click the [Results] tab in the [Compound Table View].

   The quantitation results are displayed.
4 Display the standard spectra sub-window and reference data sub-window in the [Quantitative View] area.

If necessary, see "Displaying Standard Spectra" P.63, "Displaying Reference Data" P.64 to display information about identified compounds.

5 Click on a compound name in the compound table and check the chromatogram in the [Quantitative View].

Check the results while viewing the peak identification/detection marks and baseline in the chromatogram.

Reference

If necessary, perform identification or peak integration with reference to "Manual Identification and Manual Peak Integration" P.57.

The same process can be accomplished more easily by performing the following operations on the chromatogram.

<table>
<thead>
<tr>
<th>Process</th>
<th>Operation</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Identification</td>
<td>[Shift] + [Ctrl] + right-click</td>
<td>Identifies integrated peaks.</td>
</tr>
<tr>
<td>Manual Peak Integration</td>
<td>[Shift] + right-click-drag</td>
<td>Connects start point and end point as baseline.</td>
</tr>
</tbody>
</table>
When peaks are integrated for quantitation, concentrations calculated from the calibration curve are displayed. However, if no concentration is displayed, character strings shown below are displayed according to the cause.

<table>
<thead>
<tr>
<th>Displayed Character String</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>No peak is detected.</td>
<td>Quantitative peak integration resulted in no peaks detected.</td>
</tr>
<tr>
<td>No peak is found in Window/Band range.</td>
<td>No peaks were detected within the retention time range specified for identification.</td>
</tr>
<tr>
<td>Ratio of reference ion does not match.</td>
<td>Peak is not identified due to the difference between specified and measured reference ion ratio values exceeding the allowable range.</td>
</tr>
<tr>
<td>Under the minimum similarity index.</td>
<td>Peak is not identified due to the measured similarity being less than the specified similarity setting, when mass pattern matching is specified in identification parameters.</td>
</tr>
<tr>
<td>No peak is identified.</td>
<td>Automatic identification results were manually deleted.</td>
</tr>
<tr>
<td>---</td>
<td>When the calibration curve is quadratic and the area is larger than the local maximum value (or smaller than the local minimum value), &quot;---&quot; is displayed since the concentration cannot be calculated. The target component may be out of the measurement range. Confirm the peak.</td>
</tr>
</tbody>
</table>

**NOTE**
When peaks are integrated for quantitation, concentrations calculated from the calibration curve are displayed. However, if no concentration is displayed, character strings shown below are displayed according to the cause.

6 After checking the results, click (Save) on the toolbar.
The data file is saved.

- **Displaying Standard Spectra**
Data can be analyzed more easily by comparing the displayed spectrum with a standard spectrum. When scan mode is used for measurements, data can be analyzed more easily by comparing the displayed spectrum with a standard spectrum.

1 Click [Spectrum View] - [Display Setting] on the [View] menu. If the [Spectrum Graph Display Setting] window is displayed, select [Display Standard Spectrum]. The standard spectrum is displayed.
The standard spectrum can be hidden by repeating step 1 above.
When the measured spectrum is enlarged by dragging, the standard spectrum is enlarged correspondingly.
Compounds can be identified from the shape of chromatograms, retention times, and other information obtained by referencing measurement data of standard samples or spiked samples.

1. Select [Open Reference Data File] on the [File] menu to open the data file being referenced. The reference data is displayed.

### Closing Reference Data


**NOTE**

Up to three reference data files can be displayed. Reference data peaks cannot be integrated.
5.4 Printing Quantitative Analysis Reports

It is convenient to use a template to create a report of analyzed data.

5.4.1 Creating and Outputting Quantitative Analysis Reports

   The [Data Report] window opens.

   The [File New] window opens.
3. Select [Use Template] and select the format [Quantitative Analysis Report].

4. Click [OK].
   The [Quantitative Analysis Report] format opens.

5. Click the [Preview] icon on the [Data Report] assistant bar.
   The print preview window opens.
5.4 Printing Quantitative Analysis Reports

6 After checking the report content, click [Print] to print the report.

7 Click (Save) on the toolbar.

The report is saved as a data file.
6 Shutting Down GCMS

6.1 Vacuum System Shutdown

1. Click the [Vacuum Control] icon on the [Real Time] assistant bar.
   The [Vacuum Control] window opens.

2. Click [Auto Shutdown].
   The vacuum system shuts down.

3. When [Completed] is displayed, click [Close].
6.2 Turning OFF the Power

Turn OFF the power by performing the procedure for turning ON the power in reverse. If accessory/peripheral equipment is connected, switch OFF the accessory/peripheral equipment power last.

Reference
See "2.1 Turning ON the Power" P.4 for details on how to turn ON the power.

1. Quit the [GCMS Real Time Analysis] program and all other programs that are running.
2. Turn OFF the power to the PC, printer, and display.
3. Turn OFF the power to the MS unit.
4. Turn OFF the power to the GC unit.
GCMSsolution uses the file formats described below.

<table>
<thead>
<tr>
<th>File type</th>
<th>Icon</th>
<th>Extension</th>
<th>File contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data file</td>
<td><img src="image" alt="Icon" /></td>
<td>.qgd</td>
<td>In addition to the raw data acquired (e.g., chromatograms and spectra), the following information is saved.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Calculation results such as area values and concentrations</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Status information such as the oven temperature and error status at the time data is acquired</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Contents of method files used in analysis (including configuration settings used for analysis)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Contents of report format file (when reports are output)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Contents of batch files (when batch processing is performed)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Contents of tuning file used in analysis</td>
</tr>
<tr>
<td>Method file</td>
<td><img src="image" alt="Icon" /></td>
<td>.qgm</td>
<td>Analysis conditions, peak integration parameters, compound tables, etc. are saved.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Because the configuration settings are saved when the method is edited, the configuration settings are checked when the method file is loaded to ensure that they agree with the current settings. Created calibration curves are also saved in the method file.</td>
</tr>
<tr>
<td>Report format file</td>
<td><img src="image" alt="Icon" /></td>
<td>.qgr</td>
<td>The report format information used to output a report, such as layout information and detailed settings, is saved. Once a report format file has been created, it can be used repeatedly to output reports of the same format.</td>
</tr>
<tr>
<td>Batch file</td>
<td><img src="image" alt="Icon" /></td>
<td>.qgb</td>
<td>Batch tables used to perform automatic sequential processing are saved. The same files can be used in both the [GCMS Real Time Analysis] program and the [GCMS Postrun Analysis] program.</td>
</tr>
<tr>
<td>Tuning file</td>
<td><img src="image" alt="Icon" /></td>
<td>.qgt</td>
<td>The conditions used to perform instrument adjustment (tuning) and the tuning results are saved.</td>
</tr>
<tr>
<td>Library file</td>
<td><img src="image" alt="Icon" /></td>
<td>.lib</td>
<td>These files are used to register the compound information and spectral data used to perform similarity searches. The libraries consist of public libraries (e.g., NIST and Wiley) and private libraries.</td>
</tr>
</tbody>
</table>
Appendix

Viewing Help

If you do not know how to perform a procedure, refer to Help using one of the procedures described below.

B.1 Displaying Help from the Menu Bar

1. Click [Contents] on the [Help] menu displayed in the menu bar of a window to display the [GCMS Help window].

Searching from the [Contents] Tab

1. Double-click the applicable topic.

Searching from the [Index] Tab

1. Type the applicable word.
2. Select the applicable topic and click [Display].

Searching from the [Search] Tab

1. Type the applicable word and click [Search].
2. Select the applicable topic and click [Display].

B.2 Displaying Help with the F1 Key

1. Press the [F1] key on the keyboard.
   Help for the open window is displayed.
Use the procedure described below when analyzing samples one-by-one using the autosampler or when performing analysis using manual injection.

1. **Start the [GCMS Real Time Analysis] program, then click the [Data Acquisition] icon on the [Real Time] assistant bar.**
   The [Acquisition] window opens.

2. **Double-click the method file to be used in Data Explorer.**

3. **Click the [Sample Login] icon on the [Acquisition] assistant bar.**
The [Sample Login] window opens.

1 Enter [Sample Name] and [Data File].
2 When using an autosampler, input [Vial #] in which the sample is set and [Injection Volume].
3 Click [OK].

**Hint**
[Tuning File] is not set usually. If it is left blank, the tuning file saved in the previous tuning will be used.

4 **When using an autosampler, set syringe rinse solvent and samples in the specified positions.**

5 **Click the [Download] icon on the [Acquisition] assistant bar.**
The method file settings are transferred to the instrument.
When preparation for GC and MS has been completed, the [Start] icon turns green, indicating that it can be selected.
If using autosampler model AOC-20i, the analysis starts automatically.

**Hint**
- In manual injection mode, inject the sample and then press [START] on the GC unit keyboard.
- If using accessory/peripheral equipment, start such equipment first, then click the (Start) icon.
- To abort analysis before completion, click the (Stop) icon on the [Acquisition] assistant bar.
D.1 Generating Filenames Automatically

In the [Settings] window displayed when clicking [Settings] on the [Batch] assistant bar, data filenames can be generated automatically. The settings are saved in the batch file.


2. Specify the format of the data file names to be automatically created.

   1. Click the [Data Filename] tab.
   2. Select the [Create filenames automatically with] checkbox.
   3. Add or delete items in the [Selected Items] box.
   4. Click [OK].

When the settings are complete, the [Data File] column in the batch table is highlighted in yellow.

Example: Automatically generating data filenames by putting [Batch Start Date] and [Sample Name] in the [Selected Items] box.
D.2 Editing a Batch File

1. Modify a batch file.
   
   Symbols that cannot be used in a data filename, such as "/" should not be included in the sample names.

<table>
<thead>
<tr>
<th>Batch Start Date</th>
<th>Sample Name</th>
<th>Datafile (.qgd)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20131220</td>
<td>Standard 1ppb</td>
<td>20131220_Standard 1ppb_1</td>
</tr>
<tr>
<td></td>
<td>Standard 10ppb</td>
<td>20131220_Standard 10ppb_2</td>
</tr>
<tr>
<td></td>
<td>Standard 100ppb</td>
<td>20131220_Standard 100ppb_3</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>20131220 Unknown_4</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>20131220 Unknown_5</td>
</tr>
</tbody>
</table>

NOTE
Symbols that cannot be used in a data filename, such as "/" should not be included in the sample names.

When analyses are in progress, both the [Batch Table] window and [Acquisition] window are displayed. To switch to the [Batch Table] window,

1. Click on the table.
2. Click the [Pause/Restart] icon on the [Batch] assistant bar.
   
The [Batch Table] window opens, allowing unexecuted rows to be edited.

NOTE
Analysis of rows currently being analyzed will continue to be executed.
2 Edit the batch table.
Right-click on the row to be edited, then select [Add Row], [Delete Row], or other action on the menu that appears.
The vial number, data file name, or other information can be changed as well.

3 Click (Save) on the toolbar.

4 Click the [Pause/Restart] icon on the [Batch] assistant bar.
The analysis restarts.

📝 NOTE
Some accessory/peripheral equipment may prevent using this function.
D.3 Adding Batch Files (Batch Queue)

D.3.1 Creating Batch Files to Add

1. Double-click the (GCMS Analysis Editor) icon.
   The [GCMS Analysis Editor] program starts.

2. Click the [Batch Processing] icon on the [Real Time] assistant bar.
   The [Batch Table] window opens.

3. Create the batch file to be added.

4. Name and save the batch file.

   NOTE
   • The analysis will not start if the same data file name is used more than once or the specified method file
does not exist.
   • The batch queue is not activated until the [GCMS Analysis Editor] program is closed.

5. Quit the [GCMS Analysis Editor] program.
D.3.2 Adding Batch Files

1. Start the [GCMS Real Time Analysis] program.

2. Click the [Batch Table] window.
   The content of the toolbar, menu bar, and assistant bar changes.

   The [Batch Queue] window opens.

4. Click [Add] to open the batch file to be added.

   If multiple batch files were added, change their order by clicking the desired batch file, then clicking [Move Up] or [Move Down]. Then analysis starts in that order from the top.

5. When finished editing, click [OK].
Reducing the carrier gas flow rate after analysis is finished is recommended to reduce carrier gas consumption.

## Appendix E

### Reducing the Carrier Gas Flow Rate During Standby

Reducing the carrier gas flow rate during standby for analysis is recommended to reduce carrier gas consumption.

#### E.1 Ecology Mode

Using the ecology mode reduces power consumption and carrier gas consumption during standby for analysis.

To cancel the ecology mode, click [Cancel] in the [Ecology Mode] window.

When the ecology mode is canceled, settings before switching to the ecology mode are restored.

#### E.1.1 Setting the Mode Manually

1. **Click the [Ecology Mode] icon in the instrument monitor.**
   
   A message window opens.

2. **Click [Yes].**
   
   The [Ecology Mode] window opens and the mode switches to the ecology mode. After switching to the ecology mode, the column oven temperature and the total carrier gas flow rate decrease. (For the TQ series model, CID gas supply stops.)
E.1.2 Setting the Mode Using Batch Processing

This allows switching the instrument to the ecology mode after the entire sequential analysis is finished.

   The [Batch Table] window opens.

2. Create and save a batch file.

E.2 Reducing the Carrier Gas Flow Rate During Standby

For models QP2010, QP2010 Plus and QP2010S, perform the following operations.

E.2.1 Creating a Method File That Reduces the Carrier Gas Flow Rate

As an example, the following describes how to create a method file that reduces the total flow rate to 20 mL/min.

1 Start the [GCMS Real Time Analysis] program, then in Data Explorer, double-click the method file to be used for sequential analysis.

4 Click the [Start] icon on the [Batch] assistant bar.

5 When the ecology mode confirmation message appears, click [Yes].

The mode switches to the ecology mode after the sequential analysis is completely finished, including the batch queue.

NOTE
The setting can be canceled by repeating step 3, but leave the setting as it is.
2. Change [Total Flow] to 20 mL/min, then name and save the method file.

E.2.2 Creating Batch Files

2 In Data Explorer, double-click the batch file to be used for sequential analysis.

![Data Explorer Batch Window]

3 Right-click on the batch table and select [Table Style] on the menu that appears. The [Table Style] window opens.

![Table Style Window]

4 Click [Run Mode] in the [Hide Items] list, then click [Add>>] and [OK]. A [Run Mode] column is added to the end of the batch schedule.
Edit the batch file.
Add a row at the end and select a method file created in "Appendix E.2.1 Creating a Method File That Reduces the Carrier Gas Flow Rate" P.81.
Vial number, level number, and injection volume settings do not need to be changed from their default values. Enter a data file name that is not the same as any other row.

Click the [Run Mode] cell for the row that specifies the method file that reduces the flow rate, then click the arrow button that appears.
The [Run Mode] window opens.

Configure [Run Mode] settings as shown below, then click [OK].

Name and save the batch file, then click the [Start] icon on the [Batch] assistant bar.
During this process, the method file for reducing the carrier gas flowrate is loaded when the final row is reached, and continuous data acquisition ends when the flowrate reaches 20 mL/min.
Appendix F

Peak Integration for Total Ion Current Chromatogram (TIC)

When qualitatively analyzing multiple components, perform peak integration as described below to make the analysis operation easier.

1. Double-click the (GCMS Postrun Analysis) icon. The [GCMS Postrun Analysis] program starts.

2. Click the [Qualitative] icon on the [Postrun] assistant bar.

3. Open the data file.
**F Peak Integration for Total Ion Current Chromatogram (TIC)**

**4 Set the peak integration parameters and perform peak integration for the entire TIC.**

1. Click the [Peak Integration for All TICs] icon on the [Qualitative] assistant bar.

2. Click the [Peak Integration] tab in the [Quantitative Parameters] window.

3. Click [Auto(Area)].

4. Set a value in [# of Peaks].

5. Click [OK].

**5 Open the TIC peak table and check the peaks detected.**

1. Click the [Qualitative Table] icon on the [Qualitative] assistant bar. The [Qualitative Table] window opens.

2. Click the [TIC] tab. The results for peak integration now can be checked.
6 Click [Select All] on the [Edit] menu.

7 Click [Edit] menu again and then click [Register to Spectrum Process Table].

8 Perform similarity search for every row.

1 Click the [Spectrum Process] tab in the [Qualitative Table] window.
2 Click [Search All Table] on the [Similarity Search] menu. [Done] is displayed in the [Search] cell.
Double-click the first row and check the similarity search results in order.

<table>
<thead>
<tr>
<th>No.</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Similarity: The closer this value is to 100, the greater the similarity in mass spectra.</td>
</tr>
<tr>
<td>2</td>
<td>To enter a compound name in the spectrum table, select the box for the applicable compound.</td>
</tr>
<tr>
<td>3</td>
<td>Click to copy the selected compound names to the spectrum table.</td>
</tr>
<tr>
<td>4</td>
<td>Use to switch between the mass spectra for the compounds found.</td>
</tr>
<tr>
<td>5</td>
<td>Hit numbers for the compounds found.</td>
</tr>
<tr>
<td>6</td>
<td>Allows switching between search results for each row in the spectrum process table.</td>
</tr>
</tbody>
</table>
Modifying the TIC Table

If peaks cannot be detected properly by automatic peak integration, take the following actions.

<table>
<thead>
<tr>
<th>Phenomena</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>No peak is detected.</td>
<td>Click [Manual Peak Integrate] - [TIC] on the [Qualitative] menu to perform peak integration of the target compounds.</td>
</tr>
<tr>
<td>Overlapped multiple peaks are identified as a single peak.</td>
<td>Click [Sprit Peak] on the [Qualitative] menu and move the mouse pointer to the desired position for peak splitting.</td>
</tr>
<tr>
<td>Peaks are detected and identified, but peak integration has not been done correctly.</td>
<td>Delete the row(s) from the TIC peak table.</td>
</tr>
</tbody>
</table>

Copying the Hit #1 Compound Name in the Spectrum Process Table

If [Copy Compound Name of Hit #1 to Spectrum Process Table] is selected, the name of the compound that was hit first in the similarity search can be automatically entered. However, it is necessary to confirm that the first hit compound is actually the target compound because it might not be.
Index Searches

It is possible to search for information related to the target compounds (e.g., spectra and information on the structure) in the library.

1. Click the [Library Editor] icon on the [Postrun] assistant bar.
   The [Library Editor] window opens.

2. Click [Open Library] on the [File] menu to open the library to be used.
   The library opens.

3. Click the cell in the [Index] column to select an item.

4. Enter information for the index item in the [Parameter] column for the row where the index item was selected.
5 Click [Start] on the [Index Search] menu.
The results are displayed.

6 Confirm the applicable information (e.g., spectrum or structure).
Displaying the appropriate mass chromatogram while analyzing data for qualitative analysis makes analysis easier.

### Confirming the Purity of Peaks

Displaying mass chromatograms can be used to check the presence of two or more overlapping peaks, or in other words, to check the purity of a peak in the chromatogram.

![Mass Chromatogram Example](image)

### Looking for Target Compound Peaks Among Multiple Peaks

In some cases, peaks for target compounds cannot be confirmed in a total ion chromatogram (TIC). If characteristic mass spectral peaks (i.e., m/z) of the target compounds are known, displaying the mass chromatograms makes it easier to check the position of the target compound's peaks in the chromatogram.

![Mass Chromatogram Example](image)

Perform the operation as described below.

1. Perform "Index Searches" P.90 to check the mass spectra of the target compounds.
2. Check one to three ions in the high m/z region.
3. Enter m/z values in the fragment table and display the mass chromatograms.
4. Perform similarity searches for the mass spectra of the target peaks.
H.1 Displaying Chromatograms from Mass Spectra

1. In the mass spectrum, specify and enlarge the range containing the desired peaks by dragging the mouse.

![Mass Spectrum Image]

2. Move the mouse pointer to the spectral peak to be displayed and double-click.

A mass chromatogram is displayed in the MC window, enlarged by an automatically set enlargement rate.

![Mass Chromatogram Image]

- To hide the mass chromatogram, deselect the applicable cell in the [Disp.] column in the [MC Fragment Table] window.
- To undo enlarging, right-click on the mass spectrum and select [Undo Zoom] on the menu that appears.
H.2 Displaying Chromatograms from Fragment Tables

1. Click the [Fragment Table] icon on the [Qualitative] assistant bar. The [MC Fragment Table] window opens.

2. Enter the applicable values in the [m/z] and [Factor] columns, select the corresponding cells in the [Disp.] column, and click [OK]. A mass chromatogram is displayed in the MC window.
H.2 Displaying Chromatograms from Fragment Tables

The display can be changed as shown below by enabling/disabling [Base Shift] in the table.

- **With Base Shift**

![With Base Shift](image1)

- **Without Base Shift**

![Without Base Shift](image2)
Appendix

Editing Parameters for Quantitative Analysis

Change quantitative processing parameters as necessary.

1. Start the [GCMS Postrun Analysis] Program and open the method file.

2. Click the [Quantitative Parameters] icon on the [Calibration] assistant bar.
   The [Quantitative Parameters] window opens.

3. Click the [Quantitative] tab.
Change the [Curve Fit Type], [Zero], and [Weighted Regression] settings, as necessary.

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Explanation</th>
</tr>
</thead>
</table>
| 1   | Curve Fit Type     | Specifies how to plot the calibration curve.  
• Linear: Determines the calibration curve as a straight line from the obtained values.  
• Point to point: Points are connected by a broken line. No formula is displayed for point to point calibration curves.  
• Quadratic: Curve is fit to each point using a quadratic equation. This requires at least three points on the calibration curve. For two points or less, the curve is calculated as linear.  
• Mean RF: First, it determines straight lines passing through the origin and each point. Then it finds the simple average of the slopes for each line. |
| 2   | Zero               | Select either [Not Forced] or [Force Through]. Normally, select [Not Forced].                                                               |
| 3   | Weighted Regression| A typical least squares method of plotting calibration curves could result in a quantitation error that is larger the lower the concentration at the calibration point. In general, when the calibration curve has a large dynamic range (maximum concentration is at least 50 times higher than the minimum quantitation limit), formulas are weighted to reduce the weight of higher concentration points of the calibration curve. Typically, formulas are optimized by checking the correlation coefficient and contribution ratio.  
• [1/C²]: Formulas are weighted by the inverse of the concentration value squared.  
• [1/C]: Formulas are weighted by the inverse of the concentration value.  
• [1/A²]: Formulas are weighted by the inverse of the area value squared (or height value when a height is specified for the data used).  
• [1/A]: Formulas are weighted by the inverse of the area value squared (or height value when a height is specified for the data used). |

After finishing making changes, click [OK]. Calibration curves are corrected according to the changed parameters.
Reports can be output from GCMSsolution using the two methods described below.

- **Image printing**: The image in the displayed window is automatically converted to a report.
- **Report creation**: A report format is set and output manually.

### J.1 Printing Images (Printing Spectra and Chromatograms Displayed in Windows)

1. Call up the applicable data in the [Data Analysis] window in the qualitative or quantitative processing modes of the [GCMS Postrun Analysis] program.

2. Display the chromatogram and mass spectrum in the window in the way desired for the report.

4. Adjust the size as necessary.

5. After editing, click the [Print] icon on the [Report] assistant bar. The report is output.

6. After outputting the report, close the [Report] window.
J.2 Creating Reports

With report creation, reports are output after setting report formats or using previously created templates.
Process and save the results to be output (such as spectral information) in advance.

1. Open the applicable data in the [GCMS Postrun Analysis] - [Data Analysis] window.
The same report is output for both the qualitative and quantitative windows.

2. Click the (Report) icon on the [Qualitative] or [Quantitative] assistant bar.
The [Data Report] window opens.

J.2.1 Using Templates


2. Select [Use Template], select the applicable template, and click [OK].

If this selection window is not displayed, select [Option] on the [Tool] menu to display the [Setting Options] window and, on the [File New] tab, select [Prompt on File New] for the report format file.
J.2 Creating Reports

J.2.2 Using Previously Created Report Files

1 In Data Explorer, double-click the report file to be used.

J.2.3 Manually Setting Report Content

1 Click the buttons on the toolbar for the information to be printed or select the desired items on the [Item] menu.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Name</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Sample information" /></td>
<td>Sample information</td>
<td>Select to print sample information.</td>
</tr>
<tr>
<td><img src="image" alt="Method" /></td>
<td>Method</td>
<td>Select to print methods.</td>
</tr>
<tr>
<td><img src="image" alt="Peak table" /></td>
<td>Peak table</td>
<td>Select to print the peak tables in qualitative tables.</td>
</tr>
<tr>
<td><img src="image" alt="Chromatogram" /></td>
<td>Chromatogram</td>
<td>Select to print the chromatograms (TIC, MIC, and MC).</td>
</tr>
<tr>
<td><img src="image" alt="Spectrum graph" /></td>
<td>Spectrum graph</td>
<td>Select to print the mass spectra registered in spectrum processing tables.</td>
</tr>
<tr>
<td><img src="image" alt="Mass table" /></td>
<td>Mass table</td>
<td>Select to print the mass tables for the spectra registered in spectrum processing tables.</td>
</tr>
<tr>
<td><img src="image" alt="Quantitative graph" /></td>
<td>Quantitative graph</td>
<td>Select to print the chromatograms and quantitative values obtained in quantitative results.</td>
</tr>
<tr>
<td><img src="image" alt="Quantitative table" /></td>
<td>Quantitative table</td>
<td>Select to print the tables obtained in quantitative results.</td>
</tr>
<tr>
<td><img src="image" alt="Calibration curve" /></td>
<td>Calibration curve</td>
<td>Select to print calibration curves.</td>
</tr>
<tr>
<td><img src="image" alt="Tuning" /></td>
<td>Tuning</td>
<td>Select to print the tuning results obtained when data acquisition is executed. Select [GC/MS Tuning] or [GC/MS/MS Tuning] icon.</td>
</tr>
<tr>
<td><img src="image" alt="Library search" /></td>
<td>Library search</td>
<td>Select to print the library search results obtained for the mass spectra registered in spectrum tables. • Searches must be performed in the spectrum tables.</td>
</tr>
</tbody>
</table>
2 Drag the mouse in the layout view to specify the print range. The properties window for the item being laid out opens.

3 Set [Properties] and click [OK].

Reference
Refer to Help for details on property settings.

To display a properties window again, double-click on the corresponding item.
4 Click the [Preview] icon on the [Data Report] assistant bar and check the contents of the report being output.

5 After the checking the report content, click [Print] to output the report.

6 Select [Save Format File As] on the [File] menu to name and save the report file. This allows loading the report format in the future to create reports easily.
K.1 Maintenance

Replace or clean the consumable items and maintenance parts as necessary, referring to the [MS Navigator] window using the procedure described below.

1. Double-click the (GCMS Real Time Analysis) icon.
   The [GCMS Real Time Analysis] program starts.

   The [MS Navigator] window opens.

3. Click on the instrument for which maintenance will be performed.
4 Read the precautionary information carefully and then click the applicable item under the maintenance menu.

Perform maintenance by following the instructions displayed on the screen. Click an image to enlarge it. Click [Back] in the enlarged window to return to the original window.

To perform another maintenance item, click [Back] and repeat the procedure from step 3.
After completing maintenance, close the [MS Navigator] window.

After performing maintenance, reset the usage frequencies and usage times using the procedure described in "Appendix K.3 Changing Replacement Guidelines for septa and Glass Inserts" P.108.
K.2 Easy sStop

Using Easy sTop allows replacing septa and glass inserts without stopping the vacuum system. Therefore, it significantly reduces the time required for stabilizing the system after replacement and eliminates the need for autotuning.

To protect columns, Easy sTop keeps the temperature of the sample injection unit, column oven, and interface at 70 °C or below. Consequently, it can take about 30 minutes depending on the settings until glass inserts and septa can be displayed.

1. **Double-click one of the icons for consumables in the instrument monitor.**

   ![Consumeable Tab Page](image)

2. **Click [Easy sTop].**
   The [Easy sTop] window opens and the injection unit, column oven, and interface temperatures decrease. When each temperature reaches 70 °C or lower, the “Push the replace button” status is displayed in the [Easy sTop] window.

   ![Easy sTop Window](image)
3. Click [Replace], then replace septa or glass inserts in the sample injection unit. For replacement procedures, refer to the septum replacement procedure or insert replacement procedure in the [MS Navigator] window.

4. After replacement, click [Complete] in the [Easy sTop] window. If there is no air leaking in, the sample injection unit, column oven, and interface temperatures return to their previous temperatures before Easy sTop started.

5. Reset the usage counter for the septum and glass insert. For instructions on how to reset usage counters, see the procedure on page 108, starting with step 3.
K.3 Changing Replacement Guidelines for septa and Glass Inserts

For septa, replacement frequency varies depending on the syringe needle diameter. The septum can be used about 100 times with the recommended syringe, and about 30 times with a gastight syringe before replacement.

Glass insert replacement frequency varies depending on the sample. Set replacement guidelines based on the sample.

1. Click the [System Configuration] icon on the [Real Time] assistant bar.
   The [System Configuration] window opens.

2. Double-click [SPL1] under [Modules Used for Analysis].
   The [Modules of Analytical Line #1] window opens.
3 Click [Injection Port Maintenance].
   The [Injection Port Maintenance (SPL1)] window opens.

4 Input [Septum Used Counts] and [Insert Used Counts] settings.
   To restore default settings, click [Default].

5 Click [OK].
   The [Modules of Analytical Line #1] window returns.

6 Click [OK].
   The [System Configuration] window returns.

7 Click [Set].
   The replacement guidelines for septa and glass inserts are changed.
L.1 Data Analysis using Quantitative Browser

Using this browser allows multiple samples to be quantitatively processed at one time.

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[Quantitative Result View]</td>
<td>Use to check the quantitative calculation results (area, concentrations, etc.) of multiple data files. Click ⬤ to switch between compounds to display.</td>
</tr>
<tr>
<td>2</td>
<td>[Compound Table View]</td>
<td>Click the [Results] tab to check the quantitative values of each compound in the data file selected in [Quantitative Result View].</td>
</tr>
<tr>
<td>3</td>
<td>[Calibration Curve View]</td>
<td>Displays a calibration curve of the ID selected in [Compound Table View].</td>
</tr>
<tr>
<td>4</td>
<td>[Chromatogram View]</td>
<td>Displays chromatograms of the compounds that are in the data files selected in [Quantitative Result View] and also selected in [Compound Table View].</td>
</tr>
<tr>
<td>5</td>
<td>Sample Type Toolbar</td>
<td>Data files for the specified sample type can be displayed in [Quantitative Result View].</td>
</tr>
<tr>
<td></td>
<td></td>
<td>All: All sample types, Std: Standard samples, Unk: Unknown samples</td>
</tr>
</tbody>
</table>
L.1.1 Loading Data Using Quantitative Browser

1. Double-click the (GCMS Browser) icon on the desktop.

2. Click the [Quant Browser] icon on the [Browser] assistant bar.

3. Click the [Batch] tab in Data Explorer.

4. Drag and drop the batch file used for analysis into [Quantitative Result View].

   Individual data files can be dragged and dropped to open. In addition, right-click a desired row and then click [Delete] to delete a data file.
L.1.2 Checking and Correcting Calibration Curves

1. Click the [Modify Calibration Curve] icon on the [Quant Browser] assistant bar.

2. Check the calibration curve and make necessary corrections.

   ![Reference](Image)
   
   Refer to "5.3.1 Checking and Correcting Calibration Curves" P.54 for the operation procedure.

3. Click the (Top) icon on the [Calibration] assistant bar.

   When the [Quant Browser] window appears, quantitative recalculation is done based on the corrected calibration curve.

L.1.3 Checking and Correcting Quantitative Results of Unknown Samples

![Reference](Image)

Reference

If necessary, perform identification or peak integration with reference to "Manual Identification and Manual Peak Integration" P.57.
The same process can be accomplished more easily by performing the following operations on the chromatogram.

<table>
<thead>
<tr>
<th>Process</th>
<th>Operation</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Identification</td>
<td>[Shift] + [Ctrl] + right-click</td>
<td>Identifies integrated peaks.</td>
</tr>
<tr>
<td>Manual Peak Integration</td>
<td>[Shift] + right-click-drag</td>
<td>Connects start point and end point as baseline.</td>
</tr>
<tr>
<td>Delete Identification</td>
<td>[Shift] + [Ctrl] + right-double-click</td>
<td>Voids identification and removes quantitative calculation results.</td>
</tr>
</tbody>
</table>

The intensity axis in [Chromatogram View] can be fixed by moving the mouse pointer to the desired data file in [Quantitative Result View], right-clicking it, and then clicking [Fix the Intensity Axis to this Data].

**L.2 Saving Data Files**

1. **Click** (Save) on the toolbar.
   The data file is saved.

2. **Click [Save Browsing File As] on the [Layout] menu.**
   Enter a name and save the file. The browsing file (that stores information on the loaded data files) is saved.
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