# LCMS Troubleshooting Tips



Causes	Solutions
MS	<ul> <li>If all peaks have changed in sensitivity check maintenance requirements; and clean the ion source and if required ion optics.</li> <li>Ensure appropriate and accurate acquisition parameters are being used.</li> <li>Check for adducts such as Na+, NH4+, K+. Consider different glassware, make up new mobile phases.</li> <li>Increase additive or buffer concentration if required</li> <li>Ensure probe is at correct distance from orifice.</li> <li>Check for matrix effects such as ion suppression. Consider different LC methods and sample preparation procedures.</li> <li>Replace capillary and desolvation line.</li> <li>Perform and check MS tune.</li> </ul>
Sample	<ul> <li>Check for sample degradation. Prepare a fresh sample.</li> <li>Check sample preparation and standard/QC concentrations.</li> <li>Include an internal standard with known concentration during sample preparation.</li> <li>Incorrect sample diluent used.</li> <li>Check injection volume.</li> </ul>
Loss of column performance	<ul> <li>Check the peak widths and resolution. Test the performance of the column using your standard test for loss in performance.</li> <li>Replace analytical column.</li> </ul>
LC leaks	- Check for loose fittings post injector on the system.
Mobile phase	<ul> <li>Check concentration of additives.</li> <li>If suppressing mobile phase components were used in previous method, clean MS source and flush out LC system.</li> </ul>

#### **Low Pressure**

Causes	Solutions
Partial leak in system	- Check all connections and retighten any which have leaks.
Flow rate	<ul> <li>Check the method has the correct flow rate.</li> <li>Test the flow rate accuracy using a calibrated flow rate meter or collect a specific volume and monitor the time required.</li> <li>Replace worn out or damaged pump seals.</li> </ul>
Method	<ul><li>Check if method is using correct temperature and correct solvents.</li><li>If a column section valve is used, check correct column selected.</li></ul>
Incorrect column	- Use correct column with correct dimensions and particle geometry.
Column temperature too high	<ul> <li>Set adequate column temperature and check no column damage if exceeded column temperature limit.</li> </ul>
Airlock in LC tubing	<ul> <li>Remove tubing from degasser and ensure flow under gravity. Reconnect and purge pumps in isopropanol.</li> </ul>
Stuck check valve	<ul><li>Purge LC system using isopropanol and ensure check valves are working correctly.</li><li>Sonicate the check valves in isopropanol.</li></ul>



**Sensor malfunction** - Repair or replace pressure sensor.

Causes	Solutions
MS out of tune	- Perform and check system tune.
TOF Calibration	- Perform TOF calibration.
Calibration performed incorrectly	- Ensure sample analytes are within calibration range and adjust if required.
Detector saturation	- Dilute sample or adjust injection volume.

#### Changes in MS Resolution

Causes	Solutions
MS out of tune	- Perform and check tune.

#### **Undesired Fragmentation**

Causes	Solutions	
lon source setting too harsh	<ul><li>Check source temperatures are appropriate for analyte.</li><li>Check ionisation voltage is appropriate for analyte.</li></ul>	
Collision energy too high / low	<ul> <li>Check and optimise collision cell gas pressure and collision energy.</li> </ul>	
lon optics	- Ensure correct voltage is applied to desolvation line and QArray.	

# **Basic Steps**

Follow these steps to isolate where the problems is.

Check the obvious explanations first and change only one thing at a time!



• LC pump pressures

 Argon gas cylinder (level and pressure)

(pressure readings)

Roughing pump

MS vacuum

Gas generator

• Ion source maintenance

(oil level & gas ballast)

#### Check the Basics:

- Power & electrical connections
- Communication cables
- Sample preparationAnalytical conditions
- Mobile phase preparation
- Needle rinse & seal washes
   Salvant Parising /
- Solvent flowing / no air bubbles

#### Identify the Cause:

- Clearly define the problem
- Review sample and maintenance logs to identify trends in the data or possible problem indicators
- Use a logical sequence of events to isolate possible causes

#### **Document Everything:**

- Document typical values such as analyte retention times and normal initial LC operating pressures. Use as a benchmark to indicate deterioration in system performance
- Document all troubleshooting steps and results. This may help you identify and solve the next problem faster
- Always inject a known sample and compare to previous data as a reference to ensure restored performance

Still having problems with your instrument? Let us know at

lcms\_support@shimadzu.co.uk

# No Peaks

Causes	Solution
MS setting issue	<ul> <li>Check the method uses appropriate MS settings for the compounds of interest.</li> <li>Ensure probe is at correct distance from orifice.</li> <li>Check protrusion of capillary from probe.</li> <li>Check spray from capillary.</li> <li>Check ion source temperatures / gas flows are as expected and stable.</li> <li>Check collision gas pressure is correct and stable.</li> <li>Ensure analyte elutes within event window set in method.</li> <li>Check for adducts such as Na+, NH<sub>4</sub>+, K+. Consider different glassware, make up new mobile phases.</li> <li>Check for matrix effects such as ion suppression. Consider different LC methods and sample preparation procedures.</li> <li>Concentration injected is below limit of detection.</li> <li>Perform and check MS tune.</li> </ul>
LC	<ul> <li>Check LC outlet tubing is connected to ion source.</li> <li>No mobile phase flow, possibly purge valve left open.</li> <li>Purge the system including injector to remove possible air bubbles in pump.</li> <li>Purge LC system using isopropanol to ensure check valves are working correctly.</li> <li>Check for crimped or damaged tubing.</li> </ul>
Compounds not retained or retained longer than run time in method conditions	<ul> <li>Check mobile phase composition is correct.</li> <li>Check correct analytical column type is being used.</li> <li>Increase run time.</li> <li>Increase solvent strength.</li> <li>Check correct flow rate is being achieved.</li> </ul>
Sample issues	<ul> <li>Prepare fresh samples.</li> <li>Review injection volume in sequence / method.</li> <li>Ensure the sample is in the correct position in the autosampler.</li> <li>Check for sample adsorption issue.</li> <li>Check for air pockets trapped in bottom of vial or well.</li> </ul>
Sample flowing	- Check divert valve settings if applicable.

## **Changing Retention Times**

to waste

Causes	Solutions
Flow rate	- Check the method uses the correct flow rate.
Insufficient equilibration	<ul> <li>The column should be equilibrated using at least 10 column volumes (check with manufacturer's manual)</li> <li>Allow more time or column volumes to equilibrate column between injections.</li> </ul>
Poor temperature control	- Ensure column oven temperature is accurate.
Change in column dimension	- Ensure the correct column including dimensions are being used.
Change in column stationary phase environment	<ul> <li>Do not use a column which has ion pairing reagent for other mobile phases due to memory effects.</li> <li>Stationary phase 'de-wetted'.</li> </ul>
Improper mobile phase	<ul> <li>Ensure the mobile phase is accurately prepared and the correct lines are being chosen on the method.</li> <li>Ensure the pump is accurately dispensing mobile phase.</li> </ul>
Instrument leaks	- Check for loose fittings throughout the system.
Air bubble in pump	- Purge pump via purge valve.
Sample diluent	- Inappropriate sample diluent for column.
Needle rinse	- Needle rinse solvent reaching the column.
Faulty or 'sticky' check valve	- Purge LC system using isopropanol to ensure check valves are working correctly.

### **Carry Over**

Causes	Solutions
Inappropriate wash settings	- Check wash solution and wash settings.
Sample concentration	- Use lower concentrated sample or inject less.
Column contamination	- Flush column; replace guard columns and analytical column if applicable.
Injector issue	<ul> <li>Changes in dispensing volume of injector, use a system suitability sample to determine volume changes.</li> <li>Check batch / method details to ensure the correct volume was programmed.</li> <li>Increase needle and loop flushing protocols to ensure no carryover from injection.</li> <li>Purging the injector metering pump.</li> </ul>
LC Gradient	<ul> <li>Insufficient time at strong solvent conditions during gradien program. Increase based on column dimensions.</li> </ul>

# **High Pressure**

Causes	Solutions
Flow rate set too high	- Reduce flow rate setting.
Blocked column	- Backflush column (if permitted) or replace column.
Incompatible mobile phase (precipitated buffer or immiscible)	<ul> <li>Use correct mobile phase; wash column and re-equilibrate.</li> </ul>
Improper column	- Use correct column with correct dimensions and particle geometry.
Injector blockage	<ul> <li>Clear blockage (review needle, loop, valve assembly and HPV outlet).</li> </ul>
Guard column / cartridge blockage	- Replace or remove guard column.
Column in-line filter blockage	- Replace or remove in-line filter.
Column temperature too low	- Set adequate column temperature.
Sensor malfunction	- Repair or replace pressure sensor.
Pump in-line filter blockage	- Replace in-line filter.
Blocked tubing	- Replace blocked tubing as necessary.





