

## FlashIDA usage guide

### General description

FlashIDA runs from a command line. When active it connects to the instrument and controls the acquisition of mass spectrometry data. The currently running MS method (if any) will be suppressed by the FlashIDA's one. The acquisition parameters are controlled by the method file in XML format (more information about the method is in the corresponding section below), during the run the software produces some text output to the console and two log files (the corresponding session describes how logging can be modified).

Below is the description of available command line parameters

-h, --help	Display a short usage information and exit
-v, --version	Display version number and exit
-o, --nooc	Normally, the software will wait for the contact closure signal from the instrument before starting method execution, using this option one can override the contact closure to start the method immediately.
-t, --test	Run the software in the test mode. Do not connect to the instrument and read the spectral information from the text file. This option is only used for internal testing and, thus, won't be discussed in detail.
-m, --method=VALUE	Location of the method file, the value can be absolute or relative (in respect to the executable) path. If no value is specified "method.xml" file in the executable folder will be used.
-r, --rawname=VALUE	By default, the log files are created as "FlashLog_{TIMESTAMP}" and "IDALog_{TIMESTAMP}". This parameter allows customizing created log files for easier analysis afterward. The value can be a text (NAMEID) or a file path, in the latter case only the last part of it without the extension will be used as NAMEID. The log files will be named as "FlashLog_{NAMEID}" and "IDALog_{NAMEID}"

### Structure of the method file

Method file contains parameters of MS1 and MS2 scans acquired by the instrument and FlashIDA parameters. It is a plain XML file and can be modified by the user, if necessary, please, note, however, that the XML element names and allowed values are case sensitive and must match exactly to the backend representation. The software comes with the exemplary method file – "method.xml". Below, is the description of all allowed XML elements.

### Method parameters

These parameters are relevant for the method as a whole

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Name	Description
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<b>TopN</b>	The maximal number of fragment scans that will be scheduled for each MS1 scan, should be a positive counting number
<b>Duration</b>	The length of the method in minutes should be a positive real number. The method starts either by contact closure signal or immediately if the corresponding argument is provided (see above), after a set number of minutes, FlashIDA will exit and return the current method (if any) will continue.
<b>MS1</b>	Parameters of MS1 scans, see available member elements below
<b>MS2</b>	Parameters of MS2 scans, see available member elements below
<b>IDA</b>	Parameters of FlashIDA, see available member elements below

### Scan parameters

These parameters can be relevant both for MS1 and MS2 scans, see the **Level** column.

<b>Name</b>	<b>Level</b>	<b>Description</b>
<b>Analyzer</b>	MS1/MS2	Mass analyzer used for this scan, allowed values are <b>IonTrap</b> and <b>Orbitrap</b>
<b>FirstMass</b>	MS1/MS2	The lowest m/z for the scan range should be a positive real number, compatible with your instrument properties
<b>LastMass</b>	MS1	The highest m/z for the scan range should be a positive real number, compatible with your instrument properties. Highest m/z for MS2 scans adjusted according to the m/z and charge of the precursor.
<b>OrbitrapResolution</b>	MS1/MS2	The resolution of an Orbitrap scan should be a positive counting number, compatible with the instrument. The official set of allowed values is <b>7500, 15000, 30000, 50000, 60000, 120000, 240000, 500000</b> , however, there are some reports, that other values will in the same range, for example, 33000, will work as well.
<b>AGCTarget</b>	MS1/MS2	AGC Target used for a scan should be any positive counting number compatible with the selected mass analyzer.
<b>MaxIT</b>	MS1/MS2	Maximum injection time (in ms) used for a scan, should be a positive real number
<b>Microscans</b>	MS1/MS2	The number of microscans to average should be positive counting number
<b>DataType</b>	MS1/MS2	Type of the spectral data to acquire, allowed values are <b>Profile</b> and <b>Centroid</b>

<b>RFLens</b>	MS1	RF Lens value in %, should be a positive real number
<b>SourceCID</b>	MS1	In-source CID fragmentation energy in eV, should be a positive real number
<b>IsolationMode</b>	MS2	Select the device used for precursor isolation, allowed values are <b>Quadrupole</b> and <b>IonTrap</b>
<b>Activation</b>	MS2	Activation/Fragmentation method, the following methods are supported in the current version <b>CID, HCD, ETD</b> . EThcD scans can be made selecting ETD and non-zero Collision Energy (see below), the latter will be used as the value for the supplemental activation
<b>CollisionEnergy</b>	MS2	Normalized collision energy both for HCD and CID activation, should be positive counting number
<b>ReactionTime</b>	MS2	Reaction time for ETD fragmentation, for other fragmentation methods any value will be ignored, should be a positive real number
<b>ReagentMaxIT</b>	MS2	Maximum injection time for reagent ion in ETD, for other fragmentation methods any value will be ignored, should be a positive real number
<b>ReagentAGCTarget</b>	MS2	AGC target for reagent ion in ETD, for other fragmentation methods any value will be ignored, should be positive counting number compatible with the instrument

#### FlashIDA parameters

These parameters are relevant for the FlashIDA algorithm

<b>Name</b>	<b>Description</b>
<b>MaxMs2CountPerMs1</b>	, should be a positive counting number
<b>QScoreThreshold</b>	The quality score threshold for precursors to be considered for fragmentation should be a real number
<b>MinCharge</b>	The minimal charge to consider for spectral deconvolution should be a positive countable number
<b>MaxCharge</b>	The maximal charge to consider for spectral deconvolution should be a positive countable number
<b>MinMass</b>	The minimal precursor mass to consider for spectral deconvolution should be a positive real number

<b>MaxMass</b>	Maximal precursor mass to consider for spectral deconvolution should be a positive real number
<b>Tolerances</b>	Tolerance for mass deconvolution in ppm should be an array of two real numbers – upper and lower tolerances, represented as two <b>&lt;double&gt;</b> elements in XML
<b>RTwindow</b>	The expected retention time window (in seconds) for elution of the same proteoform, should be a positive real number

## Changing logging

FlashIDA uses the **log4net** library to create logs. The configuration used by **log4net** is saved in the corresponding section of the **.config** file. This section will not cover the complete configuration of **log4net** logging, please, refer to the [log4net manual](#) for it.

Logging messages generated by the software are forwarded into three different places (appenders in log4net terms) – the console window, and two text files. It is relatively easy to change how many details are logged in each of the appenders, by changing **<threshold value>** element of an appender. For example, the following change

Before

```
<appender name="ConsoleAppender" type="log4net.Appender.ColoredConsoleAppender" >
...
  <threshold value="INFO" />
...
</appender>
```

After

```
<appender name="ConsoleAppender" type="log4net.Appender.ColoredConsoleAppender" >
...
  <threshold value="WARN" />
...
</appender>
```

will limit console output only to Warning and Error messages, Debug and Info messages will be still written to other appenders.

The general level of logging can be modified by changing **<level value>** existing **<logger>**s. Two loggers are used in FlashIDA – “**General**” for general info, and “**IDA**” for messages related to FlashIDA core. By default, both report all messages – i.e. Debug and higher level. The following example shows, how to exclude debug messages from logging

Before

```
<logger name="General">
  <level value="DEBUG" />
...
</logger>
<logger name="IDA">
```

```

    <level value="DEBUG" />
...
</logger>

```

After

```

<logger name="General">
  <level value="INFO" />
...
</logger>
<logger name="IDA">
  <level value="INFO" />
...
</logger>

```

Logging can be suppressed completely or in part, by disabling corresponding appenders in logger configuration. For example, the following change (marked lines) will disable text file logging and will leave only console one.

```

<logger name="General">
  <level value="DEBUG" />
  <appender-ref ref="ConsoleAppender" />
  <!-- <appender-ref ref="GeneralFile" /> -->
</logger>
<logger name="IDA">
  <level value="DEBUG" />
  <appender-ref ref="IDAFile" />
  <!-- <appender-ref ref="IDAInfoForward" /> -->
</logger>

```

## Running FlashIDA acquisition

FlashIDA controls only the MS part of the acquisition. It can be used on its own, for example, during direct infusion experiment, or “on top” of the existing LC-MS method, in the latter case, it will override any existing MS method. A single instance of FlashIDA has to be executed for each LC-MS run, although it is possible (no hardware damage should occur) to run multiple instances, the result will be unpredictable, since different instances, will schedule their task to the instrument concurrently. FlashIDA will exit and return the control to Tune or Xcalibur after the FlashIDA method has finished. Please, note, that in the very beginning of the FlashIDA method, one might expect, so-to-say, transition period, when the instrument will execute scans both from the Xcalibur method and FlashIDA, this happens due to the way the instrument work, and should not last longer than few seconds.

## Setting up FlashIDA-only experiment

This type of experiment is relevant for the cases when you do not need/want to use any LC in front of the instrument, for example when performing the direct injection, or you want to run FlashIDA manually for some time.

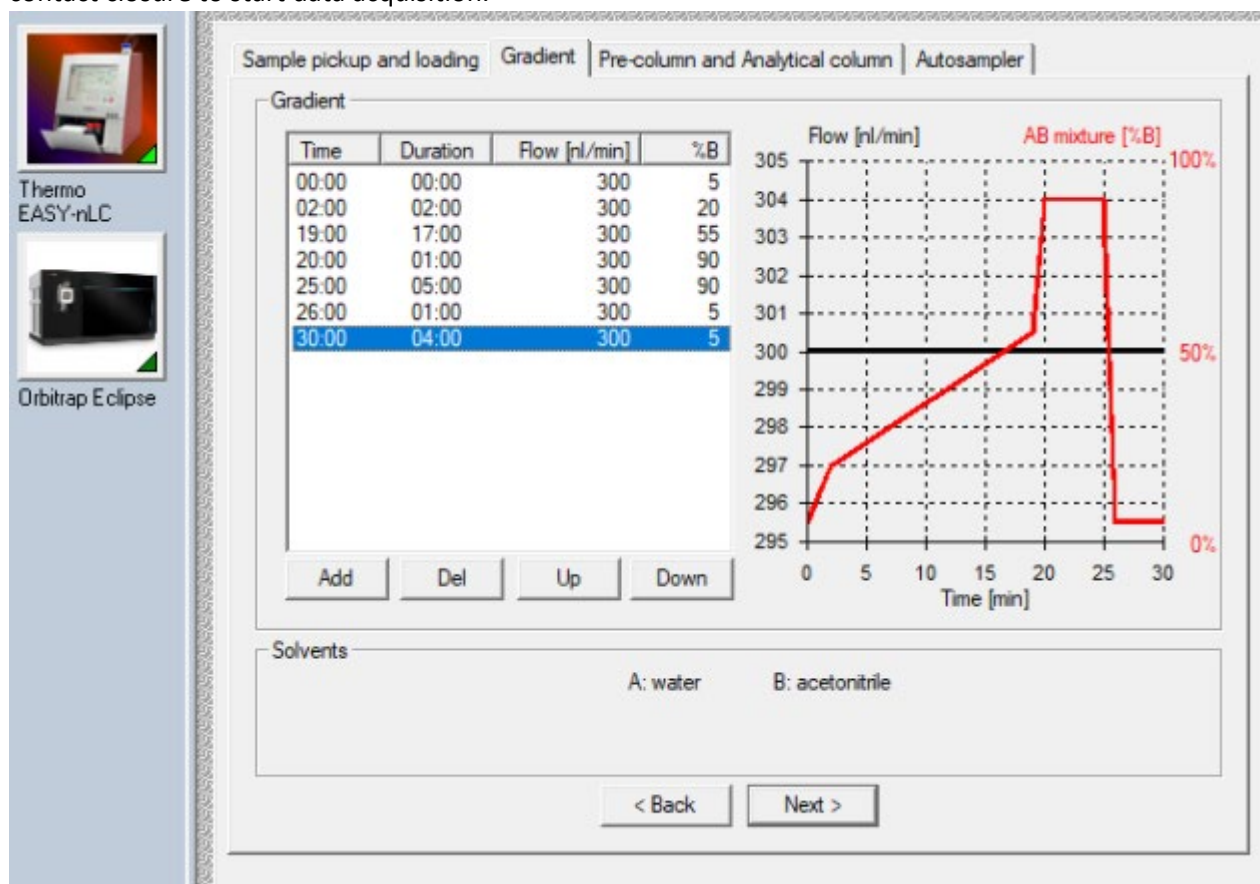
Make sure that all other parts of the experiment are running fine, for example, you have a stable spray, the molecule of interest is entering the mass spectrometer, and start FlashIDA executable with -o, --noacc command line parameter to start the acquisition ASAP. You will need to make sure that either **method.xml** file is appropriate to the experiment you want to execute, or provide -m, --method parameter pointing to

the location of the method file. If you do not have the acquisition running already, you will need to start it manually.

#### Setting up FlashIDA experiment through Xcalibur

This way of operation is handy when you want to use LC in front of the instrument, or want to acquire several samples in a sequence. Acquisition of the raw files is handled by Xcalibur, while FlashIDA is used to substitute the MS part of the experiment. FlashIDA can monitor the contact closure signal from the instrument and start the method using this signal.

1. Start setting up your LC-MS experiment as usual, i.e. create the LC part relevant to your needs. If your LC is not controlled by the Xcalibur software, you still can set up the MS-only method and use contact closure to start data acquisition.



2. Make sure that the **Global Parameters** of the MS method such as spray voltage, gas flows in the source, are set to the necessary values. There are no special requirements for the **Scan Parameters** part of the method since all settings here will be overridden by FlashIDA. It is advised, however, to set the total

duration of the method to match the LC method duration.

3. Adapt the existing **method.xml** file to your needs or create the new one with the settings of your experiment. Make sure that the duration of the experiment in the method file does not significantly exceed the duration of the LC run. If the duration of the FlashIDA experiment is shorter, the Xcalibur method will be used for the remaining time.

4. A new instance of FlashIDA must be started before acquiring each raw file since the FlashIDA tool will exit after the method is over. The **Pre Acquisition** command of Xcalibur is handy to perform this task automatically. Simply add the command to execute in the corresponding window, for example

```
D:\FlashIDA\Flash.exe -m D:\methods\FlashIDA\30_min_method.xml
```

Assuming you have FlashIDA executable in D:\FlashIDA and the method file in

D:\methods\FlashIDA\30\_min\_method.xml

5. Xcalibur allows sending some metadata of the running sample to command-line tools. In particular, *%R: Provides the current raw data file*, this can be used together with the `-r`, `--rawname` parameter of FlashIDA to name the log files according to the raw file name. For example,

```
D:\FlashIDA\Flash.exe -m D:\methods\FlashIDA\30_min_method.xml -r %R
```

If any of the log files exist, a timestamp will be added to the filename. By default, log files will use just a timestamp. For the complete list of available metadata refer to the **Run Sequence** section of Xcalibur help.

### Understanding log-files

FlashIDA uses two log files per run, **FlashLog\_** one is used for most of the information, while, **IDALog\_** one only for the information from FlashIDA core, i.e. the information related to the targets found in each scan. By default, both logs are set for **Debug**-level output, the section above explains how to change the debugging level or disable it if desired.

FlashIDA works in a loop, it receives MS scans from the instrument, analyze them, and form additional scan requests to be executed by the instrument. The loop starts with the start of the method and lasts until the duration of the method. FlashIDA can issue AGC, MS1, and MS2 scans requests. Each scan request produced by FlashIDA gets a unique numeric **Access ID** or just **ID** starting at 41. This identifier can be used to monitor the progression of each scan request. **Access ID** will be written into the trailer information structure in the raw file and can be inspected by looking at the scan header. All scan requests are stored in a FIFO queue and submitted one after another to the instrument. Scan requests are then executed by the instrument; however, it is **not** guaranteed that the instrument will execute the scans requests in the same order as they are submitted.



FlashLog will report on which MS scans are received and sent to the instrument and the state of the FIFO queue. The number in front shows a timestamp (number of ms from the starting of the application). Below are examples of log entries with an explanation.

```
786766 Sending Full Orbitrap scan [500 - 2000]; ID: 44
```

A request to perform a **Full MS1 Orbitrap** scan with mass range **500 – 2000** and Access ID **44** was sent to the instrument

```
787395 RECD FTMS MS1 Scan #8; ID: 44
```

The request was executed (Access ID **44**) and an MS1 scan with scan number **8** was received by the software

```
787399 MS1 Scan# 8 RT 0.0317269848 (Access ID 44) - 0 targets
```

The same scan has been analyzed and **0** precursor targets were generated. Scan number and retention time match the ones written to the raw file.

```
1612381 ADD m/z 739.2156/2.29 (11+) qScore: 0.9101 to Queue as #4
```

Fragmentation scan for the target(precursor) m/z **739.2156**, with isolation window **2.29** Th, was added to the FlashIDA scan request queue at position **4**. The precursor has charge **11** and quality score **0.9101**

```
1612381 QUEUE is [#0 MSn Orbitrap [729.736236572266, 22+] - #4369 Full IonTrap [500:2000] - #4370 Full Orbitrap [500:2000] - #0 MSn Orbitrap [739.215637207031, 11+] - #0 MSn Orbitrap [1445.72064208984, 9+] - #0 MSn Orbitrap [891.063781738281, 18+] - #0 MSn Orbitrap [757.001708984375, 17+] - #4369 Full IonTrap [500:2000] - #4370 Full Orbitrap [500:2000]]
```

The condition of the scan request queue, each scan is represented by the short summary with type of the scan, analyzer, precursor m/z and charge (for MSn scans), and mass range for MS1 scans. Scan added in the previous example can found in the queue at position 4.

```
1612802 POP MSn scan MZ = 739.215637207031 Z = 11 // AGC: 1, MS1: 1, MS2: 3
```

The same precursor scan request is popped from the queue, the queue contains still **1** AGC scan, **1** MS1 scan, and **3** MS2 scans

```
1612803 Sending MSn scan MZ = 739.215637207031 Z = 11; ID: 4377
```

The fragmentation scan request was sent to the instrument and received ID **4377**

IDALog will report on the result of the analysis of the scans. See examples and explanations below.

```
MS1 Scan# 3650 RT 10.2560682368 (Access ID 3686) - 4 targets
```

MS1 scan with scan number **3650**, retention time **10.2560682368** (both match to raw file information), having Access ID **3686** was processed and **4** possible precursor targets were found. Each found precursor will be represented by the following text.

```
Mass=9969.86636706996      Z=11      Score=0.819953358681964 Window=[906.9416
87011719-908.870568847656] PrecursorIntensity=1656068.5      PrecursorMassInt
ensity=3923630.359375      Features=[0.954601168632507,5.42687654495239,0.95818
1262016296,1.55814671516418,0.820968925952911,1.47546184062958] ChargeRange=[11-
14]
```

It provides, targeted mass, charge, quality score, isolation window, as well as several other parameters, much of which are only useful for the debugging of the deconvolution process.