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# **toffee Documentation**

***Release stable***

**ProCan Software Engineering at Children's Medical Research Inst**

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# CHAPTER 1

## OpenMSToffee: C++

### Contents

- *OpenMSToffee: C++*
  - *OpenSwathWorkflow*
  - *Internal Class Structure*
  - \* *Extracting data from mzML and mzXML files*

TODO...

## 1.1 OpenSwathWorkflow

```
1 $ OpenSwathWorkflow --helphelp
2 OpenSwathWorkflow -- Complete workflow to run OpenSWATH
3 Version: 2.3.0 Jun 21 2018, 07:51:05, Revision: 763e76a
4 To cite OpenMS:
5   Rost HL, Sachsenberg T, Aiche S, Bielow C et al.. OpenMS: a flexible open-source
6   software platform for mass spectrometry data analysis. Nat Meth. 2016; 13, 9: 741-
7   748. doi:10.1038/nmeth.3959.
8 Usage:
9   OpenSwathWorkflow <options>
10 Options (mandatory options marked with '*'):
11   -in <files>*
12   ↳ Input files separated by blank (valid formats: 'mzML', 'mzXML', 'sqMass')
13   -tr <file>*
14   ↳ Transition file ('TraML', 'tsv', 'pqp') (valid formats: 'traML', 'tsv', 'pqp')
15   -tr_type <type>
16   ↳ Input file type -- default: determined from file extension or content (continues on next page)
```

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14    ↵(valid: 'traML', 'tsv', 'pqp')                                     ↵
15      -tr_irt <file>                                              ↵
16      ↵Transition file ('TraML') (valid formats: 'traML')           ↵
17      -rt_norm <file>                                              ↵
18      ↵normalization file (how to map the RTs of this run to the ones stored in the library). If set, tr_irt may be omitted. (valid formats: 'trafoXML')       ↵
19      -swath_windows_file <file>                                         ↵
20      ↵Optional, tab separated file containing the SWATH windows for extraction: lower_offset upper_offset \newline 400 425 \newline ... Note that the first line is a header and will be skipped.          ↵
21      -sort_swath_maps                                              ↵
22      ↵Sort input SWATH files when matching to SWATH windows from swath_windows_file           ↵
23      -use_ms1_traces                                              ↵
24      ↵Extract the precursor ion trace(s) and use for scoring           ↵
25      -enable_uis_scoring                                           ↵
26      ↵Enable additional scoring of identification assays             ↵
27      -out_features <file>                                            ↵
28      ↵Output file (valid formats: 'featureXML')                      ↵
29      -out_tsv <file>                                               ↵
30      ↵TSV output file (mProphet compatible TSV file) (valid formats: 'tsv')           ↵
31      -out_osw <file>                                              ↵
32      ↵OSW output file (PyProphet compatible SQLite file) (valid formats: 'osw')           ↵
33      -out_chrom <file>                                             ↵
34      ↵Also output all computed chromatograms output in mzML (chrom.mzML) or sqMass (SQLite format) (valid formats: 'mzML', 'sqMass')           ↵
35      -min_upper_edge_dist <double>                                         ↵
36      ↵Minimal distance to the edge to still consider a precursor, in Thomson (default: '0' ↵
37      ↵)
38      -rt_extraction_window <double>                                         ↵
39      ↵Only extract RT around this value (-1 means extract over the whole range, a value of 600 means to extract around +/- 300 s of the expected elution). (default: '600')           ↵
40      -extra_rt_extraction_window <double>                                         ↵
41      ↵Output an XIC with a RT-window that by this much larger (e.g. to visually inspect a larger area of the chromatogram) (default: '0' min: '0')           ↵
42      -mz_extraction_window <double>                                         ↵
43      ↵Extraction window used (in Thomson, to use ppm see -ppm flag) (default: '0.05' min: '0')           ↵
44      -ppm                                                               M/
45      ↵z extraction_window is in ppm                                         ↵
46      -sonar                                                       ↵
47      ↵Data is scanning SWATH data                                         ↵
48      -min_rsq <double>                                              ↵
49      ↵Minimum r-squared of RT peptides regression (default: '0.95')           ↵
50      -min_coverage <double>                                         ↵
51      ↵Minimum relative amount of RT peptides to keep (default: '0.6')           ↵
52      -split_file_input                                              ↵
53      ↵The input files each contain one single SWATH (alternatively: all SWATH are in separate files)           ↵
54      -use_elution_model_score                                         ↵
55      ↵Turn on elution model score (EMG fit to peak)           ↵
56      -readOptions <name>                                              ↵
57      ↵Whether to run OpenSWATH directly on the input data, cache data to disk first or to perform a datareduction step first. If you choose cache, make sure to also set tempDirectory (default: 'normal' valid: 'nor ↵
58      ↵', 'cache', 'cacheWorkingInMemory', 'workingInMemory')           ↵

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37   -mz_correction_function <name>
38   ↵Use the retention time normalization peptide MS2 masses to perform a mass_
39   ↵correction (linear, weighted by intensity linear or quadratic) of all spectra._
40   ↵(default: 'none' valid: 'none', 'unweighted_regression'
41
42   ↵'weighted_regression', 'quadratic_regression', 'weighted_quadratic_regression',
43   ↵'weighted_quadratic_regression_delta_ppm', 'quadratic_regression_delta_ppm')
44   -irt_mz_extraction_window <double>
45   ↵Extraction window used for iRT and m/z correction (in Thomson, use ppm use -ppm_
46   ↵flag) (default: '0.05')
47   -ppm_irtwindow
48   ↵IRT m/z extraction_window is in ppm
49   -tempDirectory <tmp>
50   ↵Temporary directory to store cached files for example (default: '/tmp/')
51   -extraction_function <name>
52   ↵Function used to extract the signal (default: 'tophat' valid: 'tophat', 'bartlett')
53   -batchSize <number>
54   ↵The batch size of chromatograms to process (0 means to only have one batch,_
55   ↵sensible values are around 500-1000) (default: '0' min: '0')
56
57   Common UTIL options:
58   -ini <file>
59   ↵Use the given TOPPINI file
60   -log <file>
61   ↵Name of log file (created only when specified)
62   -instance <n>
63   ↵Instance number for the TOPPINI file (default: '1')
64   -debug <n>
65   ↵Sets the debug level (default: '0')
66   -threads <n>
67   ↵Sets the number of threads allowed to be used by the TOPP tool (default: '1')
68   -write_ini <file>
69   ↵Writes the default configuration file
70   -write_ctd <out_dir>
71   ↵Writes the common tool description file(s) (Toolname(s).ctd) to <out_dir>
72   -no_progress
73   ↵Disables progress logging to command line
74   -force
75   ↵Overwrite tool specific checks.
76   -test
77   ↵Enables the test mode (needed for internal use only)
78   --help
79   ↵Shows options
80   --helphelp
81   ↵Shows all options (including advanced)
82   --log_arguments
83   ↵Print out all the command line arguments
84
85   Debugging:
86   -Debugging:irt_trafo <text>
87   ↵Transformation file for RT transform
88
89   Parameters for the RTNormalization for iRT peptides. This specifies how the RT_
90   ↵alignment is performed and how outlier detection is applied. Outlier detection can_
91   ↵be done iteratively (by default) which removes one outlier per iteration or using_
92   ↵the RANSAC algorithm.:
93   -RTNormalization:alignmentMethod <choice>
94   ↵How to perform the alignment to the normalized RT space using anchor points continues on next page
95   ↵': perform linear regression (for few anchor points). 'interpolated': Interpolate_
96   ↵between anchor points (for few, noise-free

```

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```

65      e_
66      ↵anchor points). 'lowess' Use local regression (for many, noisy anchor points). 'b_
67      ↵spline' use b splines for smoothing. (default: 'linear' valid: 'linear',
68      ↵'interpolated', 'lowess', 'b_spline')
69      -RTNormalization:outlierMethod <choice>
70      ↵Which outlier detection method to use (valid: 'iter_residual', 'iter_jackknife',
71      ↵'ransac', 'none'). Iterative methods remove one outlier at a time. Jackknife_
72      ↵approach optimizes for maximum r-squared improvem
73
74      ↵ent while 'iter_residual' removes the datapoint with the largest residual error_
75      ↵(removal by residual is computationally cheaper, use this with lots of peptides)._
76      ↵(default: 'iter_residual' valid: 'iter_residu
77
78      ↵', 'iter_jackknife', 'ransac', 'none')
79      -RTNormalization:useIterativeChauvenet
80      ↵Whether to use Chauvenet's criterion when using iterative methods. This should be_
81      ↵used if the algorithm removes too many datapoints but it may lead to true outliers_
82      ↵being retained.
83      -RTNormalization:RANSACMaxIterations <number>
84      ↵Maximum iterations for the RANSAC outlier detection algorithm. (default: '1000')
85      -RTNormalization:RANSACMaxPercentRTThreshold <number>
86      ↵Maximum threshold in RT dimension for the RANSAC outlier detection algorithm (in_
87      ↵percent of the total gradient). Default is set to 3% which is around +/- 4 minutes_
88      ↵on a 120 gradient. (default: '3')
89      -RTNormalization:RANSACSamplingSize <number>
90      ↵Sampling size of data points per iteration for the RANSAC outlier detection_
91      ↵algorithm. (default: '10')
92      -RTNormalization:estimateBestPeptides
93      ↵Whether the algorithms should try to choose the best peptides based on their peak_
94      ↵shape for normalization. Use this option you do not expect all your peptides to be_
95      ↵detected in a sample and too many 'bad'
96
97      ↵peptides enter the outlier removal step (e.g. due to them being endogenous peptides_
98      ↵or using a less curated list of peptides).
99      -RTNormalization:InitialQualityCutoff <value>
100     ↵The initial overall quality cutoff for a peak to be scored (range ca. -2 to 2)
101     ↵(default: '0.5')
102     -RTNormalization:OverallQualityCutoff <value>
103     ↵The overall quality cutoff for a peak to go into the retention time estimation_
104     ↵(range ca. 0 to 10) (default: '5.5')
105     -RTNormalization:NrRTBins <number>
106     ↵Number of RT bins to use to compute coverage. This option should be used to ensure_
107     ↵that there is a complete coverage of the RT space (this should detect cases where_
108     ↵only a part of the RT gradient is actually
109
110    ↵covered by normalization peptides) (default: '10')
111    -RTNormalization:MinPeptidesPerBin <number>
112    ↵Minimal number of peptides that are required for a bin to counted as 'covered'
113    ↵(default: '1')
114    -RTNormalization:MinBinsFilled <number>
115    ↵Minimal number of bins required to be covered (default: '8')
116
117    RTNormalization:lowess:
118      -RTNormalization:lowess:span <value>
119      ↵Span parameter for lowess (default: '0.6666666666666667' min: '0' max: '1')
120
121    RTNormalization:b_spline:

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86      -RTNormalization:b_spline:num_nodes <number>
87      ↳Number of nodes for b spline (default: '5' min: '0')                                     ▾

88  Scoring parameters section:
89      -Scoring:stop_report_after_feature <number>
90      ↳Stop reporting after feature (ordered by quality; -1 means do not stop). (default:
91      ↳'-1')
92      -Scoring:rt_normalization_factor <value>
93      ↳The normalized RT is expected to be between 0 and 1. If your normalized RT has a
94      ↳different range, pass this here (e.g. it goes from 0 to 100, set this value to 100)           ▾
95      ↳(default: '100')
96      -Scoring:quantification_cutoff <value>
97      ↳Cutoff in m/z below which peaks should not be used for quantification any more             ▾
98      ↳(default: '0' min: '0')
99      -Scoring:write_convex_hull
100     ↳Whether to write out all points of all features into the featureXML
101     -Scoring:uis_threshold_sn <number>
102     ↳N threshold to consider identification transition (set to -1 to consider all)           ▾
103     ↳(default: '0')
104     -Scoring:uis_threshold_peak_area <number>
105     ↳Peak area threshold to consider identification transition (set to -1 to consider         ▾
106     ↳all) (default: '0')
107     -Scoring:scoring_model <choice>
108     ↳Scoring model to use (default: 'default' valid: 'default', 'single_transition')          ▾

109    Scoring:TransitionGroupPicker:
110        -Scoring:TransitionGroupPicker:stop_after_feature <number>
111        ↳Stop finding after feature (ordered by intensity; -1 means do not stop). (default:
112        ↳'-1')
113        -Scoring:TransitionGroupPicker:min_peak_width <value>
114        ↳Minimal peak width (s), discard all peaks below this value (-1 means no action).       ▾
115        ↳(default: '14')
116        -Scoring:TransitionGroupPicker:peak_integration <choice>
117        ↳Calculate the peak area and height either the smoothed or the raw chromatogram data.      ▾
118        ↳ (default: 'original' valid: 'original', 'smoothed')
119        -Scoring:TransitionGroupPicker:background_subtraction <choice>
120        ↳Remove background from peak signal using estimated noise levels. The 'original'         ▾
121        ↳method is only provided for historical purposes, please use the 'exact' method and
122        ↳set parameters using the PeakIntegrator:
123
124        ↳settings. The same original or smoothed chromatogram specified by peak_integration
125        ↳will be used for background estimation. (default: 'none' valid: 'none', 'original',
126        ↳'exact')
127        -Scoring:TransitionGroupPicker:recalculate_peaks <choice>
128        ↳Tries to get better peak picking by looking at peak consistency of all picked peaks.      ▾
129        ↳ Tries to use the consensus (median) peak border if theof variation within the
130        ↳picked peaks is too large. (default: 'true'
131
132        ↳valid: 'true', 'false')
133        -Scoring:TransitionGroupPicker:use_precursors
134        ↳Use precursor chromatogram for peak picking
135        -Scoring:TransitionGroupPicker:recalculate_peaks_max_z <value>
136        ↳Determines the maximal Z-Score (difference measured in standard deviations) that is
137        ↳considered too large for peak boundaries. If the Z-Score is above this value, the
138        ↳median is used for peak boundaries (defau
139
140        ↳value 1.0). (default: '0.75')                                              lt

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108     -Scoring:TransitionGroupPicker:minimal_quality <value>
109     ↪Only if compute_peak_quality is set, this parameter will not consider peaks below
110     ↪this quality threshold (default: '-1.5')
111     -Scoring:TransitionGroupPicker:resample_boundary <value>
112     ↪For computing peak quality, how many extra seconds should be sample left and right
113     ↪of the actual peak (default: '15')
114     -Scoring:TransitionGroupPicker:compute_peak_quality <choice>
115     ↪Tries to compute a quality value for each peakgroup and detect outlier transitions.
116     ↪The resulting score is centered around zero and values above 0 are generally good
117     ↪and below -1 or -2 are usually bad. (defa
118     ↪ult: 'true' valid: 'true', 'false')
119     -Scoring:TransitionGroupPicker:compute_peak_shape_metrics
120     ↪Calulates various peak shape metrics (e.g., tailing) that can be used for
121     ↪downstream QC/QA.
122     -Scoring:TransitionGroupPicker:boundary_selection_method <choice>
123     ↪Method to use when selecting the best boundaries for peaks. (default: 'largest'
124     ↪valid: 'largest', 'widest')

125 Scoring:TransitionGroupPicker:PeakPickerMRM:
126     -Scoring:TransitionGroupPicker:PeakPickerMRM:sgolay_frame_length <number>
127     ↪The number of subsequent data points used for smoothing.

128     ↪This number has to be uneven. If it is not, 1 will be added. (default: '11')
129     -Scoring:TransitionGroupPicker:PeakPickerMRM:sgolay_polynomial_order <number>
130     ↪Order of the polynomial that is fitted. (default: '3')
131     -Scoring:TransitionGroupPicker:PeakPickerMRM:gauss_width <value>
132     ↪Gaussian width in seconds, estimated peak size. (default: '30')
133     -Scoring:TransitionGroupPicker:PeakPickerMRM:use_gauss <choice>
134     ↪Use Gaussian filter for smoothing (alternative is Savitzky-Golay filter) (default:
135     ↪'false' valid: 'false', 'true')
136     -Scoring:TransitionGroupPicker:PeakPickerMRM:peak_width <value>
137     ↪Force a certain minimal peak_width on the data (e.g. extend the peak at least by
138     ↪this amount on both sides) in seconds. -1 turns this feature off. (default: '-1')
139     -Scoring:TransitionGroupPicker:PeakPickerMRM:signal_to_noise <value>
140     ↪Signal-to-noise threshold at which a peak will not be extended any more. Note that
141     ↪setting this too high (e.g. 1.0) can lead to peaks whose flanks are not fully
142     ↪captured. (default: '0.1' min: '0')
143     -Scoring:TransitionGroupPicker:PeakPickerMRM:write_sn_log_messages
144     ↪Write out log messages of the signal-to-noise estimator in case of sparse windows
145     ↪or median in rightmost histogram bin
146     -Scoring:TransitionGroupPicker:PeakPickerMRM:remove_overlapping_peaks <choice>
147     ↪Try to remove overlapping peaks during peak picking (default: 'true' valid: 'false',
148     ↪'true')
149     -Scoring:TransitionGroupPicker:PeakPickerMRM:method <choice>
150     ↪Which method to choose for chromatographic peak-picking (OpenSWATH legacy on raw
151     ↪data, corrected picking on smoothed chromatogram or Crawdad on smoothed
152     ↪chromatogram). (default: 'corrected' valid: 'legacy',
153     ↪'corrected', 'crawdad')

154 Scoring:TransitionGroupPicker:PeakIntegrator:
155     -Scoring:TransitionGroupPicker:PeakIntegrator:integration_type <choice>
156     ↪The integration technique to use in integratePeak() and estimateBackground() which
157     ↪uses either the summed intensity, integration by Simpson's rule or trapezoidal
158     ↪integration. (default: 'intensity_sum' valid:
159     ↪'intensity_sum', 'simpson', 'trapezoid')

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131   -Scoring:TransitionGroupPicker:PeakIntegrator:baseline_type <choice>
132     ↳The baseline type to use in estimateBackground() based on the peak boundaries. A
133     ↳rectangular baseline shape is computed based either on the minimal intensity of the
134     ↳peak boundaries, the maximum intensity or
135
136   ↳the average intensity (base_to_base). (default: 'base_to_base' valid: 'base_to_base'
137   ↳', 'vertical_division', 'vertical_division_min', 'vertical_division_max')
138
139 Scoring:DIAScoring:
140   -Scoring:DIAScoring:dia_extraction_window <value>
141     ↳DIA extraction window in Th or ppm. (default: '0.05' min: '0')
142   -Scoring:DIAScoring:dia_extraction_unit <choice>
143     ↳DIA extraction window unit (default: 'Th' valid: 'Th', 'ppm')
144   -Scoring:DIAScoring:dia_centroided
145     ↳Use centroided DIA data.
146   -Scoring:DIAScoring:dia_byseries_intensity_min <value>
147     ↳DIA b/y series minimum intensity to consider. (default: '300' min: '0')
148   -Scoring:DIAScoring:dia_byseries_ppm_diff <value>
149     ↳DIA b/y series minimal difference in ppm to consider. (default: '10' min: '0')
150   -Scoring:DIAScoring:dia_nr_isotopes <number>
151     ↳DIA number of isotopes to consider. (default: '4' min: '0')
152   -Scoring:DIAScoring:dia_nr_charges <number>
153     ↳DIA number of charges to consider. (default: '4' min: '0')
154   -Scoring:DIAScoring:peak_before_mono_max_ppm_diff <value>
155     ↳DIA maximal difference in ppm to count a peak at lower m/z when searching for
156     ↳evidence that a peak might not be monoisotopic. (default: '20' min: '0')
157
158 Scoring:EMGScoring:
159   -Scoring:EMGScoring:max_iteration <number>
160     ↳Maximum number of iterations using by Levenberg–Marquardt algorithm. (default: '10')
161
162 Scoring:Scores:
163   -Scoring:Scores:use_shape_score <choice>
164     ↳Use the shape score (this score measures the similarity in shape of the transitions
165     ↳using a cross-correlation) (default: 'true' valid: 'true', 'false')
166   -Scoring:Scores:use_coelution_score <choice>
167     ↳Use the coelution score (this score measures the similarity in coelution of the
168     ↳transitions using a cross-correlation) (default: 'true' valid: 'true', 'false')
169   -Scoring:Scores:use_rt_score <choice>
170     ↳Use the retention time score (this score measure the difference in retention time)
171     ↳(default: 'true' valid: 'true', 'false')
172   -Scoring:Scores:use_library_score <choice>
173     ↳Use the library score (default: 'true' valid: 'true', 'false')
174   -Scoring:Scores:use_intensity_score <choice>
175     ↳Use the intensity score (default: 'true' valid: 'true', 'false')
176   -Scoring:Scores:use_nr_peaks_score <choice>
177     ↳Use the number of peaks score (default: 'true' valid: 'true', 'false')
178   -Scoring:Scores:use_total_xic_score <choice>
179     ↳Use the total XIC score (default: 'true' valid: 'true', 'false')
180   -Scoring:Scores:use_sn_score <choice>
181     ↳Use the SN (signal to noise) score (default: 'true' valid: 'true', 'false')
182   -Scoring:Scores:use_dia_scores <choice>
183     ↳Use the DIA (SWATH) scores. If turned off, will not use fragment ion spectra for
184     ↳scoring. (default: 'true' valid: 'true', 'false')
185   -Scoring:Scores:use_ms1_correlation
186     ↳Use the correlation scores with the MS1 elution profiles
187   -Scoring:Scores:use_sonar_scores
188     ↳Use the scores for SONAR scans (scanning swath)

```

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```
159 -Scoring:Scores:use_ms1_fullscan
  ↵Use the full MS1 scan at the peak apex for scoring (ppm accuracy of precursor and↑
  ↵isotopic pattern)
160 -Scoring:Scores:use_uis_scores
  ↵Use UIS scores for peptidoform identification↑
```

## 1.2 Internal Class Structure

```
class OpenMSToffeeWorkflow : public TOPPBase
```

### Public Types

```
enum FileFormat
```

Values:

TSV

SQLITE

### Public Functions

```
OpenMSToffeeWorkflow(bool testing = false)
```

```
struct FileArguments
```

### Public Members

```
std::string toffeeFilePath
```

input tof file path

```
std::string srlFilePath
```

input srl file path (tsv or pqp)

```
std::string alignmentTSVFilePath
```

input alightment tsv file path

```
std::string outputPath
```

output file path

```
std::string inputRTTrafoXML
```

if not empty, use this to specify RT norm (trafoXML)

```
std::string outputRTTrafoXML
```

if not empty, save RT norm to here (trafoXML)

```
FileFormat format
```

defines if using TSV or SQLite

### 1.2.1 Extracting data from *mzML* and *mzXML* files

```
class HDF5ChromatogramConsumer : public IMSDataConsumer
```

## Public Types

```
using MapType = OpenMS::PeakMap
using SpectrumType = MapType::SpectrumType
using ChromatogramType = MapType::ChromatogramType
```

## Public Functions

```
HDF5ChromatogramConsumer (const std::string &h5FilePath)
~HDF5ChromatogramConsumer ()
void consumeSpectrum (SpectrumType &s)
void consumeChromatogram (ChromatogramType &c)
void setExpectedSize (size_t expectedSpectra, size_t expectedChromatograms)
void setExperimentalSettings (const OpenMS::ExperimentalSettings &exp)
```

## Public Static Functions

```
void chromatogramToHDF5 (const std::string &mzMLFilePath, const std::string &h5FilePath)
Save an mzML chromatogram file to a much more (>100x) compressed HDF5 file This data is saved as a series of 1D vectors:
```

- **names** gives the transition ids of the chromatograms
- **offset** gives the index into the retention time and intensity vectors for the corresponding transition id
- **size** gives the size of the data in the retention time and intensity vectors for the corresponding transition id. I.e. its data resides at [offset, offset + size)
- **retentionTime** all retention time data concatenated into a single vector
- **intensity** all intensity data concatenated into a single vector

### Parameters

- mzMLFilePath: path to the mzML file generated by OpenSwath
- h5FilePath: output file path

## class RTNormalisation

Calculate the world to iRT normalisation transformation using raw SWATH-MS data contained in a toffee file, and a list of precursor and product ions that can be used for alignment

## Public Functions

```
RTNormalisation (const std::string &toffeeFilePath, const std::string &alignmentTSVFilePath)
```

### Parameters

- toffeeFilePath: the toffee file for which we wish to calculate the world to iRT normalisation

- alignmentTSVFilePath: the path to a TSV file of retention time alignment precursor and product ions

```
void updateNormalisationParams (const OpenMS::Param &param)  
    Update the default parameters to input into the normalisation algorithms.
```

```
void updateMzCorrectionFunction (const std::string &mzCorrectionFunction)  
    Update the default method for correcting the mass over charge.
```

```
void updateMinRSquared (double minRSquared)  
    Update the default minimum R-squared value in regression fitting method.
```

```
void updateMinCoverage (double minCoverage)  
    Update the default minimum coverage of the fit regression method.
```

```
void updateChromExtractParams (const OpenMS::ChromExtractParams &param)  
    Update the chromatogram extraction configuration.
```

```
void updateFeatureFinderParams (const OpenMS::Param &param)  
    Update the feature finding configuration.
```

```
OpenMS::TransformationDescription calculateNormalisation () const  
    Calculate the world to iRT normalisation.
```

```
OpenMS::TransformationDescription calculateNormalisationFromFile (const std::string  
    &inputTrafoPath)  
const  
    Calculate the world to iRT normalisation.
```

```
void saveToFile (const OpenMS::TransformationDescription &transform, const std::string  
    &trafoXMLFilePath) const  
    Save a previously calculated iRT normalisation to file.
```

**Warning** this function does not check if the transformation is in world to iRT coordinates, or its inverse.  
It relies on the user to take care!

```
void calculateAndSaveToFile (const std::string &trafoXMLFilePath) const  
    Calculate the world to iRT normalisation and save it to file.
```

## Public Static Functions

```
OpenMS::Param defaultNormalisationParams ()  
    Default parameters to input into the normalisation algorithms. These can be updated, see below.
```

```
static std::string defaultMzCorrectionFunction ()  
    Default method for correcting the mass over charge. This can be updated, see below.
```

```
static double defaultMinRSquared ()  
    Default minimum R-squared value in regression fitting method. This can be updated, see below.
```

```
static double defaultMinCoverage ()  
    Default minimum coverage of the fit regression method. This can be updated, see below.
```

# CHAPTER 2

## OpenMSToffee: Python

### Contents

- *OpenMSToffee: Python*
  - *Working with toffee files*
    - \* *Calculating RT normalisation*
  - *Converting SRLs*
  - *Internal Class Structure*

## 2.1 Working with *toffee* files

### 2.1.1 Calculating RT normalisation

```
1 $ toffee_openms_rt_normalisation --help
2 usage: toffee_openms_rt_normalisation [-h]
3                                     toffee_filename alignment_filename
4                                     transformation_xml_filename
5
6 Calculate the retention time normalisation for a toffee file using OpenMS as
7 the wrapper
8
9 positional arguments:
10    toffee_filename      The output filename (*.tof)
11    alignment_filename   The input alignment library (iRT) filename (*.tsv)
12    transformation_xml_filename
13                                The output transformation XML file (*.trafoXML)
```

(continues on next page)

(continued from previous page)

```

15 optional arguments:
16   -h, --help           show this help message and exit

```

## 2.2 Converting SRLs

```

1 $ srl_peakview_to_openms --help
2 usage: srl_peakview_to_openms [-h] [-output_dir OUTPUT_DIR]
3                               [-minimum_number_of_transitions MINIMUM_NUMBER_OF_
4 →TRANSITIONS]
5                               [-maximum_number_of_transitions MAXIMUM_NUMBER_OF_
6 →TRANSITIONS]
7                               [-mz_cutoff MZ_CUTOFF]
8                               [-precursor_product_limit PRECURSOR_PRODUCT_LIMIT]
9                               [--drop_modifications] [--make_pqp]
10                              [--keep_intermediate_files] [--debug]
11                             sciex_filename
12
13 Convert a Sciex ProteinPilot/PeakView/OneOmics SRL into a format that can be
14 used by OpenMS
15
16 positional arguments:
17   sciex_filename      The input filename (*.txt)
18
19 optional arguments:
20   -h, --help           show this help message and exit
21   -output_dir OUTPUT_DIR
22                           The directory to save the output files. By default it
23                           will be the same as the input file.
24   -minimum_number_of_transitions MINIMUM_NUMBER_OF_TRANSITIONS
25                           Define the minimum number of transitions for any PSM
26   -maximum_number_of_transitions MAXIMUM_NUMBER_OF_TRANSITIONS
27                           Define the maximum number of transitions for any PSM
28   -mz_cutoff MZ_CUTOFF  Filter out any precursor ions with a mass over charge
29                           below this threshold
30   -precursor_product_limit PRECURSOR_PRODUCT_LIMIT
31                           Filter out any product ions with a mass over charge
32                           with this many Da of the precursor ion
33   --drop_modifications Remove all modifications from the final SRL
34   --make_pqp            Create the new PQP sqlite format in addition the TSV
35   --keep_intermediate_files
                           Remove intermediate files that were created
                           --debug                Switch on more detailed logging

```

## 2.3 Internal Class Structure

```
OpenMSToffee.log.set_stream_logger(name='OpenMSToffee', level=20, format_string=None,
                                    fname=None)
```

Add a stream handler for the given name and level to the logging module. By default, this logs all boto3 messages to stdout.

```
>>> import OpenMSToffee as omt
>>> omt.log.set_stream_logger(name='OpenMSToffee', level=logging.INFO)
```

### Parameters

- **name** (*string*) – Log name
- **level** (*int*) – Logging level, e.g. `logging.INFO`
- **format\_string** (*str*) – Log message format

```
class OpenMSToffee.srl_peakview_to_openms.OpenSwathLibraryFromPeakview(peakview_fname,
                                                               out-
                                                               put_basename=None,
                                                               out-
                                                               put_dir="",
                                                               min-
                                                               i-
                                                               num_number_of_transitions
                                                               max-
                                                               i-
                                                               num_number_of_transitions
                                                               mz_cutoff=400.0,
                                                               pre-
                                                               cur-
                                                               sor_product_limit=10.0,
                                                               mod-
                                                               ifi-
                                                               ca-
                                                               tions_to_keep='CAM',
                                                               make_pqp=False,
                                                               clean_up_files=True,
                                                               de-
                                                               bug=True)
```

Convert SRL file formats between PeakView and OpenMS. Furthermore, run the decoy generation on the OpenMS data once it has been generated

```
DROP_TEXT = 'DROP'

OPENSWATH_INDEX_COLS = ['TransitionGroupId']
OPENSWATH_SORT_ASCENDING = [True, False]
OPENSWATH_SORT_COLS = ['TransitionGroupId', 'LibraryIntensity']
PEAKVIEW_HEADERS = ['Q1', 'Q3', 'iRT', 'stripped_sequence', 'relative_intensity', 'uni']
PEAKVIEW_RTCAL_PROTEIN = '[ RT-Cal protein ]'
PV_TO_OS_COL_MAPPING = {'Annotation': 'Annotation', 'CE': 'CollisionEnergy', 'Decoy': 'Decoy'}
convert()

convert_peakview_to_openswath(df)
    Internal class method, exposed only for testing

classmethod create_calibration(df_irt, basename)
    Internal class method, exposed only for testing

create_decoys(df_library, basename)
    Internal class method, exposed only for testing

filter_and_normalise_openswath(df)
    Internal class method, exposed only for testing
```

```
open_peak_view()
    Internal class method, exposed only for testing

remove_minimum_fragments(df)
    Internal class method, exposed only for testing

rename_modifications(mod_peptide_col)
    Internal class method, exposed only for testing. Convert the modification format of PeakView into UniMod
    format

classmethod split_to_srl_and_alignment(df)
    Internal class method, exposed only for testing
```

|| Master: || Dev:

OpenMSToffee is a wrapper around OpenSwath that allows us to use the `toffee` file format. It is available as a Docker image at [cmriprocan/openms-toffee](#).

We follow the [OpenVDB style guide](#) for the C++ and PEP-8 for our python code, so please aim to stay consistent with the rest of the code base. Contributions will be pass through peer review and style will be one element that is reviewed.

# CHAPTER 3

---

Changes

---



# CHAPTER 4

---

## Change Log

---

### 4.1 0.14

#### 4.1.1 0.14.1

- Changed license to MIT and fixed documentation for <https://openms-toffee.readthedocs.io>
- Bumped version of `toffee` to 0.13.1, also MIT licensed

### 4.2 0.13

#### 4.2.1 0.13.12

- Bumped version of `toffee` to 0.12.16, which now includes the mzML to toffee to mzML conversions
- Removed all mzML to toffee conversions from this package and deleted tests

#### 4.2.2 0.13.11

- Bumped version of `toffee` to 0.12.9, which now includes the toffee to mzML conversion (PD-793)

#### 4.2.3 0.13.10

- Bumped version of `toffee` to 0.12.4
- Many fixes to accommodate change in structure of the dia-test-data repository

#### **4.2.4 0.13.9**

- Added ability to convert toffee files to Spectra HDF5 files (PD-793)

#### **4.2.5 0.13.8**

- Bumped version of `toffee` to 0.11.1

#### **4.2.6 0.13.6**

- Bumped version of `toffee` to 0.10.6 (that fixes PD-749) and added unit test

#### **4.2.7 0.13.5**

- Bumped version of `toffee` to 0.10.5 (that fixes PD-735)

#### **4.2.8 0.13.4**

- Pinned version of `toffee` to 0.10.4 (that fixes PD-727)

#### **4.2.9 0.13.3**

- A few bug fixes for the mzML to toffee conversion

#### **4.2.10 0.13.2**

- Updated to version 1.2.4 of `cmriprocan/openms`.

#### **4.2.11 0.13.1**

- Significant refactor of the XML to `toffee` conversion process. This has been pulled apart into smaller modules such that individual parts can be tested. Importantly, the code that converts mzML and mzXML to HDF5 is now shared by the process that then converts these to toffee. This improves memory performance, and means that we can test both conversions.

### **4.3 0.12**

#### **4.3.1 0.12.1**

- Directly linking to upstream image in Dockerfile: `cmriprocan/openms:1.2.3` which:
  - Updated OpenMS to version CMRI-ProCan-v1.1.2
  - Updated PyProphet to 2.0.2

## **4.4 0.11**

### **4.4.1 0.11.1**

- Updated OpenMS to version CMRI-ProCan-v1.1.1 with hot-fix to solve <https://github.com/OpenMS/OpenMS/issues/3860>

## **4.5 0.10**

### **4.5.1 0.10.1**

- Updated OpenMS to version CMRI-ProCan-v1.1.0 that incorporates upstream Release 2.4.0 with our changes that enable toffee files to be used on the command line

## **4.6 License**

MIT Copyright (c) 2017-2019 Children's Medical Research Institute (CMRI)



# CHAPTER 5

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