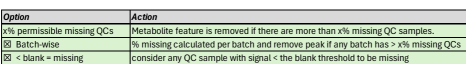
## **Cheat Sheet**





## Configuration file

| Name             | Options  | Explanation  |
|------------------|--|--|
| LogTransform     | "true"/"false"                                   | Perform Log10 transformation on data before correction (to ensure normal distribution)   |
| RemoveZeros      | "true"/"false"                                   | There is no such thing as zero. Samples is either detected (value) or not detected (missing). Zeros mess up statistics.  |
| OutlierMethod    | "None","Percentile","Linear","Quadratic","Cubic" | The 'linear'/quadratic/cubic' options fit a simple polynomial function to the QC data using robust least squares regression. Points outside the population confidence interval are deemed outliers. The 'percentile' option implements the standard non-parametric < Q1 – 1.5 IQR or > Q3 + 1.5 IQR outlier detection method on the QC samples.                                    |
| OutlierCI        | value between between 0.9 and 0.99               | Set the Confidence interval for the ploynomial methods above. 90%-99%  |
| OutlierPostHoc   | "Ignore","MPV","NaN"                             | For the corrected data. Either ignore outliers or replace with the QC median peak value (MPV) or remove (replace with "not a number" NaN).   |
| IntraBatchMode   | "Mean","Linear","Spline"                         | Three correction modes. "Spline" is the default QCRSC algorithm that requires optimisation of the smoothing parameter. "Linear" is a simple Robust (bisquare) linear regression based on the QC values & requires no smoothing optimisation. "Mean" equalises the QC mean across batches & ignores within batch systematic change.   |
| InterBatchMode   | "QC","Reference"                                 | QC' = both the intra- and inter-batch based on the QC samples;<br>'Reference' = intra-batch correction based on the QCs but the inter-<br>batch correction uses the samples labelled as 'Reference'.   |
| QCRSCgammaRange  | "x:y:z"  | The range of values (x to z) to search to determine to optimal correction curve (in increments of y). A value >=4 results in a curve equivalent to a linear regression. A value < 0 results in a highly nonlinear curve.   |
| QCRSCcvMethod    | "3-Fold","5-Fold","7-Fold","Leaveout"            | Type of cross-validation used for optimising the smoothing parameter value.  |
| QCRSCmcReps      | integer  | The number of Monte Carlo (random) resamples of the k-fold cross validation. The resulting cvMSE the mean of the generated set of cvMSEs.  |
| CorrectionType   | "Subtract","Divide"                              | Subtract or divide the correction curve from the raw data. Optional depending on whether you believe the bias to be additive or multiplicative. Recommendation: If multiplicative & Log10 transformation is used then Subtract.  |
| BlankRatioMethod | "QC","Median","Percentile"                       | BlankRatio = 100*max(BlankValue)/SampleReferenceValue. 'QC' sets the SampleReferenceValue as the median QC value; 'Median' sets the SampleReferenceValue as the median Sample value; 'Percentile' sets BlankRatio = % of Samples < max(BlankValue)*RelativeLOD (RelativeLOD is a user defined constant - default 1.5). In this instance missing values are considered Blank values |
| RelativeLOD      | value  | The RelativeLOD is a multiplier (relative to the Blank) used to set a value below which a peak is considered background noise. i.e., "estimated LOD = max(Blank) x relativeLOD"  |
| StatsParametric  | "true"/"false"                                   | Choose whether summary statistics for each corrected feature are calculated using parametric (mean, standard deviation etc) or non-parametric methods (median, median absolute deviation etc).   |
| ParallelProcess  | "true"/"false"                                   | Switch on the parallel processor to use mulitple processor cores to speed up the QCRSC engine.   |



## Clean & Explore

| Name                          | Explanation  |
|-------------------------------|--|
| Filter:                       |  |
| batch-wise/complete           | Should the filter bank calculate its peak-wise statistics across all batches or calulate each batch individually and then summarise based on the dropdown option (e.g. "Mean" = the mean of all the batch statistics & "Max" = the poorest statistic comparing all batches). |
| % missing thesh =             | % allowed missing samples. e.g. if 20% then peak is removed if number of missing > 20%.  |
| Before/After                  | Compare the performance statistics/visualisations before or after correction. Use this to convince yourself of the utility of the correction algorithm.  |
| PCA preprocessing:            |  |
| log2(Fold wrt QC)             | First perform log2 fold change - with fold = sample value / mean QC value.   |
| log10 Transform               | OR log10 transformation  |
| Autoscale / Pareto scale      | Preform autoscaling (mean centre then divide by standard deviation) or Pareto Scaling (mean centre then divide by sqrt(standard deviation). Scaling is always performed after the transformation.  |
| PCA Missing value imputation: |  |
| KNN column                    | KNN missing value imputation replacing with the nearest metabolite feature   |
| KNN row                       | KNN missing value imputation replacing with the nearest sample   |
| k =                           | replaces missing values with a weighted mean of the k nearest-neighbor columns/rows.  KNN imputation always performed after transformation & scaling.  |
| blank/20%min                  | replaces missing values with the maximum blank value for that peak or if no blanks detected 20% of the lowest value. blank/20%min is always performed before transformation or scaling.  |
| PCA projection:               |  |
| Project in QC samples         | PCA model is generated using only the Sample data. This removes any possible bias from   |
| Project in Blank samples      | the QC, Blank, or Reference samples. The QC, Blank, or Reference sample data can   |
| Project in Reference samples  | applied to the PCA model (projected through) and plotted with the Sample data.   |