An R package to analyse LC/MS metabolomic data: MAIT (Metabolite Automatic Identification Toolkit)

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1 INTRODUCTION

Liquid chromatography and mass spectrometry (LC/MS) is an analytical technique used widely in metabolomics to detect molecules in biological samples (Theodoridis et al., 2012). A wide array of software tools is available for LC/MS profiling data analysis, including commercial, programmatic and online tools. A commercial example is Analyst®, whereas some open-source packages permit programmatic processing, such as the R package XCMS (Smith et al., 2006) to detect peaks or CAMERA (Kuhl et al., 2012) and AStream (Alonso et al., 2011) for peak annotations. There have been efforts on just peak annotation using JAVA (Brown et al., 2011). MZmine and mzMatch are modularized tools coded in JAVA that are focused on LC/MS data preprocessing and visualization (Katajamaa et al., 2006; Pluskal et al., 2010; Scheltema et al., 2011). Online tools permit sample processing through a web Graphical User Interface, such as XCMSOnline (http://xcmsonline.scripps.edu) or MetaboAnalyst (Xia et al., 2009). Refer to Supplementary Table S1 for a comparative between the capabilities for some of the main available tools. In this context, we introduce a new R package called Metabolite Automatic Identification Toolkit (MAIT) for automatic LC/MS analysis. The goal of the MAIT package is to provide an array of tools that makes programmable metabolomic end-to-end statistical analysis possible (see Section 3 of the Supplementary Material for details about the MAIT modularity). MAIT includes functions to improve peak annotation through the process called biotransformations and to assess the predictive power of statistically significant metabolites that quantify class separability.

2 METHODS

MAIT includes the stages peak detection, peak annotation, statistical analysis and table and plots creation (Fig. 1). The peak detection stage detects the peaks in the LC/MS sample files. The peak annotation stage improves the identification of the metabolites in the metabolomic samples by increasing the chemical and biological information in the dataset. A statistical analysis reveals the significant sample features and measures their predictive power. MAIT uses the R package XCMS to detect and align peaks. For the peak annotation step, MAIT uses three steps:

- First, MAIT uses the CAMERA package to perform the first annotation step (Kuhl et al., 2012). In this stage, MAIT uses a peak correlation distance and a retention time window to find which peaks came from the same source metabolite based. The peaks within each peak group are annotated following a reference adduct/fragment table and a mass allowance window.
- Biotransformations could be related to specific in-source mass losses. Therefore, in the second annotation step, they are detected using a mass allowance window inside the peak groups (Breitling et al., 2006). For this search, MAIT already includes a biotransformations table (here Human biotransformations). User-defined biotransformation tables can be set as input, following the procedure defined in Supplementary Text (Section 6.6).
- Finally, a predefined metabolite database is mined for significant masses. This identifies metabolites with the help of the Human Metabolome Database (Wishart et al., 2009), 2009/07 version.

The objective of analysing the metabolomic profiling data is to obtain the statistically significant features (SSF) that contain the highest amount of class-related information. To gather these features, MAIT can apply statistical tests such as ANOVA or Student’s t-test to every feature,
selecting the significant set of features given a threshold P-value. A validation test is included to quantify SSF class separability by a repeated random subsampling cross-validation using three methods: partial least squares and discriminant analysis, support vector machines and K-nearest neighbours (Hastie et al., 2009). MAIT computes overall and class-related classification ratios to evaluate the SSF class-related information.

3 RESULTS

The example data files are a subset of the data used in the reference (Saghatelian et al., 2004), which are distributed freely through the faahKO package (Smith, 2012). MAIT was used to read and analyse these samples using the functions depicted in Figure 1 (see the tutorial in the Supplementary Information). The significant features for each class are found using statistical tests and analysed through the different plots that MAIT produces. Using the following function call, 2640 peaks were detected:

R> MAIT <- sampleProcessing(dataDir = 'DataXcms', project = 'MAIT_Demo', snThres = 2, rtStep = 0.03)

At this point, the first annotation stage is launched:

R> MAIT <- peakAnnotation(MAIT.object = MAIT)

Next, we gather the significant features from the peaks detected. After the Welch’s tests, 106 of these features were found to be significant through the spectralSigFeatures function. Statistical plots such as heat maps, boxplots and principal component analysis score plots can be generated (Supplementary Figs S3 and S4). Significant features are annotated after checking for certain neutral losses (biotransformations).

R> MAIT <- spectralSigFeatures(MAIT, P = 0.05)

By using only the SSF, a validation stage is launched, obtaining a classification ratio of 100% with three training samples for all classifiers. These results suggest that the significant variables separate both classes completely.

R> MAIT <- Validation(MAIT, Iterations = 20, trainSamples = 3)

Finally, the database is mined to identify the significant features.

R> MAIT <- identifyMetabolites(MAIT, peakTolerance = 0.005)

4 CONCLUSIONS

MAIT provides a set of tools and functions to perform an automatic end-to-end analysis of LC/MS metabolomic data, putting special emphasis on peak annotation and metabolite identification. In addition, MAIT validation functions make it possible to estimate predictive power for significant variables.

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