MeltDB 2.0–advances of the metabolomics software system

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Received on May 6, 2012; revised on July 11, 2013; accepted on July 14, 2013

1 INTRODUCTION

Metabolomics research covers all aspects of the investigation of small molecule metabolite compositions resulting from cellular processes and constitutes an integrated part of systems biology (Bino et al., 2004). Like transcriptomics and proteomics, metabolomics is capable of measuring extrinsically initiated changes in organisms. The metabolome, the entity of all small molecules in a cell, organism or tissue, is considered to be the closest to the phenotype of all ‘-omes’ (Fiehn, 2002).

Compared with other molecular levels or -omics methods, metabolomics is challenging in its high degree of interdisciplinarity, interlinking experts from research fields as diverse as engineering, physics, chemistry and biology and from cheminformatics over bioinformatics to statistics, data mining and finally visualization.

Both sample acquisition and subsequent analysis are automated in high-throughput instruments, which has continuously posed challenges on the systematic storage and computational processing of the gathered experimental datasets, starting in the early 2000s. The increasing number and quality of measurements not only raised the generated data volume but also allowed to address more complex biological questions within conducted experiments. To comprehensively address these demands, bioinformatics internet applications were developed. MeltDB, ‘a software platform for the analysis and integration of data from metabolomics experiments’, has been published by Neuweger et al. (2008). Xia et al. (2009) released MetaboAnalyst, ‘a comprehensive tool suite for metabolomic data analysis’. Carroll et al. (2010) published the MetabolomeExpress web server as ‘a public place to process, interpret and share GC/MS metabolomics datasets’.

Since around 2008, we have observed that the requirements to comprehensive metabolomics software platforms have changed: The general growth of the field of metabolomics and the increasing number of collaborations diversified the user community of researchers and their individual scientific goals. It is obvious that the success of a metabolomics study depends on an efficient and effective collaboration of this interdisciplinary research community. Thus, not only the availability and sharing of the data is important but also special functions have to be significantly extended with specific features to consider all researcher’s demands and perspectives. In addition, the ever-increasing throughput and the constant lack of time makes it immensely important that automated pre-processing methods are reliable and that analyses and manual intervention are fast and easy. Since Metabolomics approaches are applied to more and more scientific objectives, a powerful set of statistical methods is mandatory, ranging from hypothesis-driven statistical tests to less specified and untargeted data-mining methods, such as clustering and dimension reduction. Finally, the wealth of generated data poses a necessity for exploratory data analysis tools and information visualization.

To tackle these new challenges systematically, a next generation of bioinformatics tools needed to be developed, covering all of the aforementioned aspects of metabolome data analysis, ranging from processing raw data (RD) to finishing and finally the derivation of biological knowledge. During the stages of that process, one can identify four successive data categories that represent different levels of data classification and annotation as well as different levels of abstraction. First, RD, stored and organized in meaningful groups, build the basis. Then, pre-processed data (PD) is computed, where peaks and their
quantities have been detected. It follows integrated data (ID), where peaks that putatively originate from the same compound are consistently annotated over chromatograms of an experiment and thus become comparable. Last, derivative data (DD) is achieved by statistical analyses of metabolite quantities in an experiment and then visualized to allow effective exploration and to draw conclusions.

In this manuscript, we present MeltDB 2.0, which offers novel tools to challenge the rising wealth of data quality and quantity and support the analysis of all four categories RD, PD, ID and DD and includes a multitude of updates. New and improved preprocessing methods underpin the reliability of automatically created annotations. At the same time, straightforward tools for manual peak annotation simplify the curation even of large experiments. To help answering questions of different scientific objectives, the set of statistical analyses and data-mining tools has been strongly enriched. To finally nail down the quintessence of an experiment's outcome, data exploration is supported by new interactive and telling information visualizations.

2 IMPLEMENTATION AND METHODS
The first version of the MeltDB software platform, a three-tiered web application and database server published in 2008 (Neuweger et al., 2008), provides means for the standardization, systematic storage and analysis of gas chromatography–mass spectrometry (GC-MS) metabolomics experiments. Within a powerful project and user management, raw chromatograms of various file formats can be uploaded and organized into chromatogram groups (e.g. replicates, factor levels) and experiments. A flexible processing pipeline allows to find, quantify and identify peaks in the raw chromatograms. Subsequently, a set of statistical tools and visualizations can be applied to analyze the gathered data tables. This fast growing, free online platform today hosts >25 distinct projects conducted by >150 registered users from around the world. More than 17,000 chromatograms have been uploaded and analyzed yet.

In the following, all major improvements to the entire process from RD to DD will be described in more details. Figure 1 summarizes the four stages of data processing and associates visualizations and data mining methods that can be performed in MeltDB 2.0 to each stage.

2.1 From RD to PD: improved pre-processing
In metabolomics data analysis, pre-processing is a critical step, as ID and DD build on PD. To ensure a reliable data basis for statistical data exploration, MeltDB 2.0 is equipped with several new and updated algorithms for the early steps of experiment data analysis.

The growing list of pre-processing methods now includes support for the centWave algorithm by Tautenhahn et al. (2008) for chromatographic peak detection, which features a high sensitivity, and updates of the XCMS package (Smith et al., 2006) for chromatogram alignment and profiling analyses. In addition, the ChromA (Hoffmann and Stoye, 2009) software is added to the list of supported chromatogram alignment tools. ChromA computes pairwise alignments of chromatograms without a priori knowledge, but it is capable of optionally using previously matched or identified peaks as anchor points, which speeds up the process.

The calculation of retention time indices in GC-MS measurements is improved and can now also be performed manually using the web interface. Peaks of added substances can be assigned with retention indices and will be used as anchors for interpolating other peaks retention indices (Ettre, 1994), which...
support subsequent peak identification (Kopka et al., 2005). The
detection of alkanes as retention markers can be automated.

Furthermore, peak identification itself is facilitated with a
powerful feature: MeltDB 2.0 offers a new Reference list tool
to save peaks of measured reference substances as Reference in
the MeltDB database. The stored data comprises retention
indices, quantification masses and mass spectra of reference com-
ounds. This helps to generate project specific databases that
complement the Golm Metabolite Database (http://gmd.
mpimp-golm.mpg.de/) (Kopka et al., 2005) or the National
Institute of Standards and Technology standard reference database
1A (http://www.nist.gov/srd/nist1a.cfm). The tool allows to
aggregate References and to use their underlying mass spectra for
efficient peak identification and comparison.

2.2 From PD to ID: profiling methods
To complete the first step towards ID, peaks in different chro-
matograms that derive from the same small molecule have to be
named consistently and need to be associated to each other.
Thus, a new support for GC-MS-based metabolite profiling ex-
periments has been implemented. The focus for the profiling
approach in MeltDB 2.0 is to combine the results from chem-
ometrics approaches with further identification.

The generic MeltDB approach can be applied on netCDF,
mzXML (Pedrioli et al., 2004) and mzDATA (Orchard et al.,
2007) measurements from any supported analytical system. The
novel tool registers peaks for non-targeted metabolite profiling
based on multiple criteria. These are similarity of mass spectra,
retention time difference and the existence of common extracted-
ion chromatogram (EIC) peaks above a given signal-to-noise
threshold. It allows to annotate completely unknown peaks that
are consistently detectable in several measurements of a metab-
olomics experiment. Thus, these potentially interesting peaks can
be subjected to further statistical analyses in MeltDB and become
accessible for profiling experiments, where the aim is in general to
find differences in the metabolic composition of two or more
sample groups.

Parameterizations of all pre-processing tools can be custom-
ized freely from within the web interface. Parameterizations are
persisted project wide so that other users from the same
project—who typically use similar instrumental setups—can reuse
them. As pre-processing methods are usually too long run-
ning tasks as to complete during a web request or to even allow
interactivity, these jobs are forwarded to our compute cluster and
run decoupled from the web server.

As soon as pre-processing is completed at least to the point of
profiling, data tables can be exported as XLS or TSV files. This
allows to subsequently use the MeltDB 2.0 processing results in
other external programs of choice.

2.3 Analyzing PD: efficient mass decomposition
To elucidate single mass signals in a spectrum, e.g. to identify an
unknown compound or to explain a certain fragmentation, po-
tential sum formulas explaining this signal may be insightful.
Thus, a combination of the efficient mass decomposition algo-

rithm described by Böcker et al. (2006) with the filter criteria
described by Kind and Fiehn (2007) has been implemented.

![Fig. 2. The improved algorithm was compared with the original mass decomposition approach. For compounds of molecular mass close to 1000 Da, 7-fold runtime improvements can be observed. As presented in the graph, the relation of the runtimes of both methods is not linear. Especially for large compounds, the improved variant becomes most beneficial](image-url)
measurements. Several filters can be activated to reduce the number of computed sum formulas. As the implementation does automatically extract and sort, the dominant ions found in the mass spectrum, the access to the mass decomposition is greatly simplified for the researcher.

After the efficient generation and filtering step, the sum formulae are compared with the KEGG compound database. Matching sum formulae are highlighted, and both compound name and synonyms are presented to the user.

2.4 From RD to DD: new user interfaces

Visual inspection and the possibility for manual curation is important at all data abstraction levels, i.e. from RD to DD. Intuitive and responsive data visualizations are required as well as intelligent tools that allow to solve common tasks, such as the inspection of processing results or the navigation from an experiments overview to the spectrum of a single peak, in a few steps.

Introducing Asynchronous Javascripting and XML to MeltDB 2.0 using the jQuery library (http://jquery.com) allows a new quality of interactive and dynamic data display (The user can interact with visualizations to cause small changes to the data representation. These changes are performed in place without requiring a reload of the visualization) in terms of speed, responsiveness and user guidance. This was utilized to improve the Experiment Total Ion Current view [cmp. (Fig. 3b)] with peak specific tooltips containing information about the associated compounds, latest annotations, and quantities (Fig. 3b.1). On demand, the complete peak object can be loaded and displayed inside the experiment TIC view, giving i.a. access to its spectrum, the complete list of annotations and observations.

One additional major improvement that benefits from the employment of Asynchronous Javascripting and XML is the dynamic manual annotation dialog [cmp. (Fig. 3c)]. The interactive annotation functionality for whole experiments has been improved so that aligned peaks with high mass spectral similarity can be annotated in parallel. The streamlined web interface helps to annotate peaks across chromatograms in a consistent manner, when researchers annotate manually, correct errors of automated annotation tools or correct data that have been imported beforehand. This consistency is ensured by an autocompletion of compound names according to the KEGG database and is important for statistical analyses and comparison of results among different experiments.

2.5 From ID to DD: extended statistics and data mining

In metabolomics, data analysis and mining can be driven by various intentions and will strive for DD with different purposes. One way to group these aims is to relate them to one of the following questions: (i) What are the significant features of a sample group separating it from other sample groups? (ii) Do groups of samples form clusters according to their features and quantities? (iii) Can a sample be assigned to a class based on its spectral features? For each of these questions, a variety of statistical analysis methods exists.

Fig. 3. This figure presents representative screenshots from the MeltDB 2.0 interface, which are part of an efficient workflow for manual data curation and quality improvement. (a) The Show Absent/Present Compounds view lists which compounds were identified in which chromatogram groups in how many chromatograms (pie charts). Checked fields tell that the respective compound was identified in all chromatograms of that group. The darker column in the right represents the entire experiment. (b) The Experiment Total Ion Chromatogram view depicts the total intensities over time for all chromatograms of an experiment. Detected peaks are marked with colored dots. Blue: Detected peak; Green: peak that was consistently detected throughout chromatograms; Red: Peak that was identified using any database. Hovering a peak highlights other peaks (link-and-brush) with the same annotation and description and shows a tooltip with detailed information (b.1). From this view, semi-automated tools for annotation (c) can be accessed. The Annotate tool (screenshot shown) allows to manually define a description and select a KEGG compound. Similar peaks from other chromatograms are suggested and can be annotated in a batch. Alternatively, the MeltDB-own Warped Peak Detection can be applied to find and quantify peaks, which are similar to ones that are already annotated in other chromatograms using the tool ReQuantify Peaks for Existing Annotations. Sample names in the screenshots were modified.
A binding to the R software (R Development Core Team, 2011) makes numerous statistical tools available from within the MeltDB software (Neuweger et al., 2008). Its database objects are converted to R objects in a standardized manner. Information about chromatogram associations to chromatogram groups is pertained in the data representation. This ID conversion avoids the cumbersome process of converting data tables from proprietary software to a format a statistics software package can interpret and analyze. By default, data are gathered from the database on the fly, but snapshots can be stored to speed up statistical analyses vastly, especially for large experiments.

Statistical analyses and data mining tools in MeltDB are accessed through a standardized parameterization and data selection form. Where appropriate, this basic form is extended with specific options and parameters. The basic form consists of a list of all chromatograms of the experiment from which the user may select. Additionally, features can be selected to be considered in the analysis. Features can either be identified compounds or unidentified features that have been detected consistently among most chromatograms. A feature or compound can be chosen as reference to normalize to. When available, ribitol is preselected. One can select whether peak intensities or peak areas will be used for quantitation. Quantitations can be scaled linearly or logarithmically. Missing values can be handled in different ways.

To determine the significance and variances of features (see question a), the t-test of the Perl CPAN package Statistics::TTest (Juan, 2003) and Statistics::KruskalWallis (Lee, 2003) is offered as well as analysis of variance using the aov method of the R statistical software (R Development Core Team, 2011), which fits a linear model. For all of these, Bonferroni, Holm and Benjamini & Hochberg corrections are calculated (Benjamini and Hochberg, 1995; Holm, 1979). For each feature (metabolite) presented in the analysis of variance and Kruskal–Wallis test results, a boxplot view and the extracted ion chromatograms of all samples can directly be accessed. In another view, the m-values (log-2 signal ratios) of features of all chromatogram groups of an experiment in reference to the same features in a user-selected chromatogram group are displayed tabularly. Volcano plots can be created plotting either the a-values (average log-2 signal values) or t-test values (as negative decadic logarithm of the P-value) against m-values. Variable importance estimation via the random forest algorithm from the caret R package (Kuhn et al., 2011) can be applied to find differing features in groups. The metabolite set enrichment analysis published by Persicke et al. (2011) is another powerful tool in MeltDB for the identification of differentially regulated metabolic pathways.

Samples may aggregate to clusters according to their features quantities (see question b), regardless of the groups they nominally belong to. To visualize these clusters, MeltDB provides the dimensionality reduction methods principal component analysis (PCA, prcomp method in R, cmp. Fig. 4b), independent component analysis (ICA, fastICA package for R) and partial least squares discriminant analysis (caret package for R) (Kuhn et al., 2011; Marchini et al., 2010; R Development Core Team, 2011). Hierarchical clustering allows to display dendrograms of chromatograms and is made available using the hclust method in R, which can be applied with different linkage methods (cmp. Fig. 4c). The heatmap method of R is used to show false color maps of feature signals in chromatograms, sorting columns and
MeltDB 2.0 presents random forest can be trained with chromatogram groups representing which the variable importance estimation has been aforementioned. MeltDB with the powerful R package caret (Kuhn et al., 2011) of which the variable importance estimation has been aforementioned. Now, classification algorithms (support vector machine, random forest) can be trained with chromatogram groups representing different classes \( \omega_0, \ldots, \omega_{c-1} \) and then be applied to other chromatograms of samples that have not yet been assigned to any class \( \omega_k \). For evaluation purposes, the user may opt to partition chromatograms into training and testing groups randomly. Additionally, MeltDB uses caret to compute and evaluate the classification performances of the algorithms random forest, \( k \) nearest neighbors, support vector machine, neural networks and partial least squares, to estimate which classification algorithm performs best on a problem.

Generally, the computed results can be downloaded as TSV, XLS, PNG or PDF files in addition to the representation in the web browser.

2.6 Diving into DD: new interactive visualizations

To ultimately grasp the gist of gathered and calculated DD information, visualizations will be powerful tools, if they are intuitive, fast, responsive, easily customizable and—most important—well represent the underlying data.

The R software is not only useful for statistical analysis computation but also for the results visualization. Parameterization of the analyzes and customization of the visualizations is realized with MeltDBs tool forms, which can be easily extended on request or requirement. Nevertheless, these visualizations are static, and even small changes require server side computing and a page reload. This makes visual data exploration a time-consuming process. Therefore, new visualizations have been developed for MeltDB 2.0, which are based on the R data output and thus are consistent with their static ‘sister’ visualizations, but prepare the results in a novel dynamic and interactive fashion.

There are several web technologies for the development of rich internet applications (RIAs) available. Oracle®’s Java™ Webstart applications, using the Java Network Launch Protocol, belong to the first RIAs created for the web (Farrell and Nezlek, 2007). The Adobe® Flash® technology and the Flash® Builder® made platform-independent RIA development even easier and provided great functionality for user interface design, backed by a large vivid community. Most recent changes in the industry, namely, the development of HTML 5, made Javascript-based RIA development applicable and seem to give insight into hidden links in small molecule patterns, which are characteristic for a group \( k' \). This propelled to extend MeltDB with the powerful R package caret (Kuhn et al., 2011) of which the variable importance estimation has been aforementioned. Now, classification algorithms (support vector machine, random forest) can be trained with chromatogram groups representing different classes \( \omega_0, \ldots, \omega_{c-1} \) and then be applied to other chromatograms of samples that have not yet been assigned to any class \( \omega_k \). For evaluation purposes, the user may opt to partition chromatograms into training and testing groups randomly. Additionally, MeltDB uses caret to compute and evaluate the classification performances of the algorithms random forest, \( k \) nearest neighbors, support vector machine, neural networks and partial least squares, to estimate which classification algorithm performs best on a problem.

Generally, the computed results can be downloaded as TSV, XLS, PNG or PDF files in addition to the representation in the web browser.

3 APPLICATION EXAMPLE AND RESULTS

This application example makes use of a dataset from another study (in preparation) in which we investigated whether it is possible to distinguish between wheat samples of different years, farming schemes and cultivars. The data shown in the following result from the GC-MS measurements of a wheat cultivar that was grown in the years 2007 and 2010 and under organic (O) and conventional (C) treatment. Consequently, samples can be divided into the four groups 2007-O, 2007-C, 2010-O and 2010-C. Each group consists of four biological replicates, which were each analyzed as two technical replicates. This results in a total of eight chromatograms per group. Chromatograms of low quality have been discarded.

The MeltDB 2.0 processing pipelines and its versatile tool box for statistical analyses was applied to reveal potential differences in the metabolic compositions of wheat from two distinct years and treatments. These insights may lead to the determination of bio markers that allow to distinguish between the respective sample groups. However, the focus of this application example is on the software and its capabilities, not on the biological interpretation of the generated results.

The MeltDB setup started by uploading the chromatograms (RD) in the netCDF file format. These were then organized into an experiment of four chromatogram groups. Peak detection was performed using the MeltDB Warped Peak Detection method. Alkanes were detected by the RISimple method, and peaks were thus provided with interpolated retention indices, completing the steps toward PD. Subsequently, Multiple Profiling was performed to consistently name peaks with similar spectra and retention indices throughout chromatograms of the experiment. Finally, to obtain ID, peaks were matched against the
The tool **Show Absent/Present Compounds** reveals a tabular presentation of pie charts, which shows in how many chromatograms of a particular chromatogram group or the entire experiment a particular compound was identified (cmp. Fig. 3a). Compounds that were identified automatically in the majority of chromatograms are likely to be found by manual inspection in the remaining measurements. This is easy owing to the mean retention time that is provided for each compound to find the respective peaks in the **Experiment Total Ion Chromatogram** view (cmp. Fig. 3b). Here, peaks with the same annotation and description are visually connected by **link-and-brush** (Highlighting an annotated peak will cause all other peaks with the same annotation and description to highlight too. This visually links signals that derive from the same original molecule throughout measurements). Peaks that were not annotated automatically (or annotated incorrectly) are easily spotted and may be annotated semi-automatically now. This is possible using the **Annotate** tool that is offered for each peak and allows to fill an annotation form manually, but it also allows to select peaks with similar spectra and similar retention time from other chromatograms to be annotated in the same way (cmp. Fig. 3c). Another tool, **ReQuantify Peaks for Existing Annotations**, provides means to locally run the **Warped Peak Detection** with relaxed thresholds and using existing annotations of a certain compound as samples to be matched. Using these tools, even large experiments of several hundred chromatograms can be annotated with high coverage in a reasonable time frame.

In the following, tools for visualization and statistical analysis are demonstrated. The **Heatmap** tool is a helpful choice (cmp. Fig. 4a). For a quick insight or visual control, all chromatograms and all compounds should be selected. Abundances are taken from peak areas, without additional scaling. In this case, missing values were replaced with zeros to make them easily spottable in the heatmap, as zero values are displayed as black boxes. The **Heatmap** tabularly represents all compound abundances in all chromatograms or their mean abundances in chromatogram groups according to a blue-to-red (low to high abundance) color scale. Rows (compounds) and columns (chromatograms or chromatogram groups) can be rearranged freely. Rows may additionally be sorted alphabetically, by mean abundance or by coefficient of variance. The color scale is either normalized on the whole experiment or on each single compound. The color scale may be shifted by a factor using a slider. The **Heatmap** as currently set can be exported as a PNG file.

When analyzing data of multiple—and as in this case even overlapping classes—it is generally interesting whether and which clusters are formed. Typical workhorses to explore this are **Principal Component Analysis** (pca) and **Hierarchical Cluster Analysis** (hca). Figure 4b shows a screenshot of the interactive scatter plot of the pca, which was calculated on all chromatograms, and all compounds that were detected in all chromatograms (option 'strict' for missing values). Measured peak areas were taken for quantification. Data points with the same color belong to the same chromatogram group. As the pca shows, samples are clearly separated by years along the first principal component (x-axis), which explains 75.24% of the variance in the data. Less clear but still visible is a separation according to the cultivation type, mainly oriented along the second principal component (y-axis), which explains 6.28% of the data. A mouse over on a data point reveals its chromatogram name and coordinates in the plot (cmp. Fig. 4b.1). Similarly, the hca as shown in Figure 4c mainly clusters data according to their years, whereas samples of the same cultivation type group cluster hardly at all. The hca was performed as a Ward clustering with the same basic settings as the pca.

## 4 DISCUSSION AND CONCLUSION

MeltDB 2.0 was developed to comprehensively provide means to complete the entire process from RD to DD within a software platform that supports researchers of diverse scientific backgrounds and fosters collaborations in complex metabolomics research projects. It was a further goal to make the final exploration of produced results and statistical outcomes effective and efficient. This has been achieved by improving the MeltDB tool set throughout all four stages RD, PD, ID and DD. These recent developments leveraged MeltDB to an interactive RIA platform that supports researchers of diverse scientific backgrounds and to support larger and/or long-term projects.

As MeltDB was first published in 2008, a few other tools have been released that take a similar line. The MetaboAnalyst (2.0) web server offers a feature set similar to MeltDB, also supplying means to cover the pipeline from RD to DD. MetaboAnalyst is merely made for a one-time web service-like usage though, whereas MeltDB offers a project and user management that supports collaborative work, allows to manually refine and annotate processing results and stores data for documentation purposes and to support larger and/or long-term projects.

The MetabolomeExpress web server is dedicated to making reviewed datasets publicly available. For that, it offers a fixed pipeline and a set of statistical tools that comprises a clearly smaller set of features, compared with MeltDB or MetaboAnalyst.

The hierarchical data model of MeltDB 2.0 serves single-factorial experiment designs best. It is still possible to address even complex multi-factorial designs, but then a careful organization (sometimes multiple organizations) of chromatogram groups is necessary. The application example shown is a multi-factorial experiment differentiating wheat samples of different years and farming schemes. It is part of a study that additionally considers different cultivars as a third factor, which was also investigated in MeltDB 2.0.

Despite the advantages and opportunities of web platforms, this technology also has its drawbacks. The most critical aspect probably is the lack of tools, which lets users browse the original RD and its raw signals in a smooth interactive way as known from desktop applications. This can be of particular importance especially for *de novo* identification of molecules. Thus,
metabolomics web platforms, including MeltDB 2.0, are mostly useful for experiments with large numbers of chromatograms, which ask for the statistical comparison of sample groups.

Web platforms are also still limited in the consequent analysis of LC-MS data. Here, spectral deconvolution is of critical importance to reveal the interrelationships of mass signals, which may lead to the identification of the original molecules. Although software like the R tool CAMERA (Kuhl et al., 2012) exists to address this, it has not yet been integrated into a larger metabolomics software platform. We are currently working to overcome that limitation.

Further developments also must include support for multistage (MS²) data, another inevitable tool for proper de novo identification.

From a more global and systemic point of view into the future, the potential of integrated analysis with other omics data needs to be explored more intensively. As an example, the MeltDB 2.0 API allows the ProMeTra (Neuweger et al., 2009) software to map relative metabolite abundances to pathway maps, together with either proteome or transcriptome data. Nevertheless, to our knowledge, there is no software available for truly statistical multiomics approaches.

In total, MeltDB has undergone substantial improvements in its capacity as a ‘one-stop-shop’ providing a wide spectrum of necessary tools to answer biological and statistical questions, beginning from chromatographics RD files. The addition of supervised machine-learning tools now allows to directly apply its capacity as a ‘one-stop-shop’ providing a wide spectrum of knowledge, there is no software available for truly statistical multiomics research projects.

ACKNOWLEDGEMENTS

The authors A. Goesmann and Tim W. Nattkemper both want to be considered as senior authors for this manuscript. The authors thank Leonhard J Stutz for his support as a research assistant. The authors appreciate the helpful comments by the anonymous reviewers of this manuscript very much. N.K. is supported by a fellowship from the CLIB Graduate Cluster Industrial Biotechnology. The authors also thank the BRF team for expert technical support.

Funding: German Federal Ministry of Food, Agriculture and Consumer Protection under the Federal Scheme of Organic Farming and other forms of sustainable agriculture [Project 08OE023].

Conflict of Interest: none declared.

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