Untargeted Analysis of *Lemna minor* Metabolites: Workflow and Prioritization Strategy Comparing Highly Confident Features between Different Mass Spectrometers

Rofida Wahman 1,2, Stefan Moser 3, Stefan Bieber 4, Catarina Cruzeiro 5, Peter Schröder 5, August Gilg 6, Frank Lesske 6 and Thomas Letzel 1,4,*

Citation: Wahman, R.; Moser, S.; Bieber, S.; Cruzeiro, C.; Schröder, P.; Gilg, A.; Lesske, F.; Letzel, T. Untargeted Analysis of *Lemna minor* Metabolites: Workflow and Prioritization Strategy Comparing Highly Confident Features between Different Mass Spectrometers. *Metabolites* 2021, 11, 832. [https://doi.org/10.3390/metabo11120832](https://doi.org/10.3390/metabo11120832)

Abstract: Metabolomics approaches provide a vast array of analytical datasets, which require a comprehensive analytical, statistical, and biochemical workflow to reveal changes in metabolic profiles. The biological interpretation of mass spectrometric metabolomics results is still obstructed by the reliable identification of the metabolites as well as annotation and/or classification. In this work, the whole *Lemna minor* (common duckweed) was extracted using various solvents and analyzed utilizing polarity-extended liquid chromatography (reversed-phase liquid chromatography (RPLC)-hydrophilic interaction liquid chromatography (HILIC)) connected to two time-of-flight (TOF) mass spectrometer types, individually. This study introduces and discusses three relevant topics for the untargeted workflow: (1) A comparison study of metabolome samples was performed with an untargeted data handling workflow in two different labs with two different mass spectrometers using the same plant material type. (2) A statistical procedure was observed prioritizing significant detected features (dependent and independent of the mass spectrometer using the predictive methodology Orthogonal Partial Least Squares-Discriminant Analysis (OPLS-DA). (3) Relevant features were transferred to a prioritization tool (the FOR-IDENT platform (FI)) and were compared with the implemented compound database PLANT-IDENT (PI). This compound database is filled with relevant compounds of the Lemnaceae, Poaceae, Brassicaceae, and Nymphaceae families according to analytical criteria such as retention time (polarity and LogD (pH 7)) and accurate mass (empirical formula). Thus, an untargeted analysis was performed using the new tool as a prioritization and identification source for a hidden-target screening strategy. Consequently, forty-two compounds (amino acids, vitamins, flavonoids) could be recognized and subsequently validated in *Lemna* metabolic profile using reference standards. The class of flavonoids includes free aglycons and their glycosides. Further, according to our knowledge, the validated flavonoids robinetin and norwogonin were for the first time identified in the *Lemna minor* extracts.

Keywords: FOR-IDENT; metabolomics; OPLS-DA; PLANT-IDENT; polarity-extended chromatography; quadrupole time-of-flight (QTOF); TOF
1. Introduction

Metabolomics is the approach that deals with the investigation of a biological system (cell, tissue, or organism) by determining its overall metabolite profile at a given time point with the specified set of conditions. The modern development of analytical techniques expanded the coverage of metabolomics in investigations of biological systems. This approach permits remarkable insights into regulation mechanisms as well as the responses to different perturbations. Over the centuries, quantitative (targeted) metabolomics analysis has become a routine application in different fields [1]. In untargeted metabolomics studies, the identification of (unknown) metabolites is the fundamental step to transform analytical data into biological knowledge. Unfortunately, this transformation is still considered the major bottleneck. So far, the number of identified metabolites in untargeted metabolomics studies is in general below 50% [1]. Several analytical techniques are available in the untargeted metabolomics analysis such as liquid chromatography-mass spectrometry (LC-MS) and gas chromatography-mass spectrometry (GC-MS) [2] to analyze a large number of different chemical metabolites classes within one single analysis, respectively. However, the thermal stability of the stationary phase, metabolites, and their derivatives, which might introduce artifacts, limit the metabolome coverage derived by GC-MS. Thus, the usage of LC-MS has expanded rapidly over the past ten years in untargeted metabolomics analysis [3].

In approaches based on liquid chromatography–high-resolution mass spectrometer (LC-HRMS), metabolites can mostly be identified if the retention time (RT), the ion mass, and MS/MS fragment spectrum of each compound are successfully matched with an authentic reference standard measured in the same instrument. However, due to the presence of over 200,000 to 1,000,000 different metabolites in the plant kingdom, this can be a challenging task [4,5]. Especially the identification of highly specific metabolites that might have significant roles in protecting plants against predators or microbial infection is difficult due to the unavailability of reference standards [4,5]. To overcome this challenge, several MS/MS databases have been established to facilitate metabolite annotation, such as spectral databases MassBank (https://massbank.eu/MassBank/ accessed on 30 November 2021), METLIN (http://metlin.scripps.edu/index.php accessed on 30 November 2021), LipidBlast (https://fiehnlab.ucdavis.edu/projects/lipidblast accessed on 30 November 2021), and ReSpect (http://spectra.psc.riken.jp accessed on 30 November 2021). Moreover, plant metabolomics databases have been developed for potential identification such as KNAPsACh (http://kanaya.naist.jp/KNAPsACh/ accessed on 30 November 2021), MetaCyc (http://metacyc.org accessed on 30 November 2021), KEGG (http://www.genome.jp/kegg/ accessed on 30 November 2021), and PRIME. Hence, the choice of databases has an essential influence on the interpretation of a scientific study. Small- and medium-sized databases might miss relevant metabolites while large databases contain more molecules so unexpected hits may lead to better information. On the other hand, the small, well-defined databases might suggest more meaningful hits compared to searches in large databases, which may propose compounds that are not related to the respective scientific and biological question [4–7].

The PLANT-IDENT (PI) database (https://water.for-ident.org accessed on 30 November 2021) for example is a specific compound database containing currently 3019 metabolites of the Lemnaceae family in addition to Poaceae, Brassicaceae, and Nymphaeaceae, as well as other families due to the chemotaxonomy filter in addition to analytical characters. This compound database is implemented for rapid search retrieval into the FOR-IDENT (FI) platform (https://water.for-ident.org accessed on 30 November 2021) using batch search for large numbers of compounds. The FI platform ‘translates’ the analytical features with different analytical parameters like retention time (RT), ion mass, (and MS/MS spectrum) via retention time index (RTI) (by calculating the difference between the LogD values calculated with via RTI of the target mixture and the one one is listed in the FI), and accurate mass into logD values, and molecular formula. These parameters are compared with the physicochemical parameters of the metabolites in the compound database PI which
gives a prioritizing hit list (with Look at a function in the FI). Therefore, in general, a large number of molecules from extracts can be annotated and prioritized for the study of these plant families without prior focusing on these compounds. This can enhance the potential coverage of indicators in untargeted metabolomics analysis. A remaining challenge for metabolomic analysis is its weak efficiency and the difficulty of comparing results obtained in different laboratories that use different hardware or data processing. Currently, it may well happen that molecules identified in one laboratory may not be replicable to others [6].

Usually, as it is exemplarily shown in [8], the first step toward metabolites identification is extraction. In the next step, there is an efficient chromatographic separation, providing a reproducible, precise, and wide range of polarity separation. A serial coupling of two LC columns with orthogonal polarities extends the separation of compounds from the complex matrix [8–10].

This analysis provides a large amount of data, requiring an adequate workflow to process and analyze it. Although an overall view would be more relevant using all mass spectrometric ion polarity information, exclusively positive ionization mode was applied in this study. This workflow reflects a subset of the untargeted screening results but leads also to statistical useable datasets. HRMS, such as quadrupole time-of-flight (Q-TOF) mass spectrometry and quadrupole-orbitrap (Q-Orbitrap) mass spectrometry have become the most widely used MS tools in metabolomics [6,11]. The metabolomics data processing continues to be problematic, especially the metabolite identification process. The associated workflow contains numerous steps, which have to be precise, compatible, and with less effort and time-consuming. All steps required quality control procedures to ensure trustworthy metabolomics analysis outcomes. Starting from peak picking, signal-to-noise threshold detection, and background correction to the different datasets were aligned. The workflow ended with the application of the automated metabolite identification algorithms and the biological interpretation of the data [12,13].

The most common proposed workflow applies to food and nutrition sciences, where metabolomic analysis has early been adopted as an analytical tool [14]. In addition, medical research [15] uses metabolomics analysis to gain new insights into the interactions between drugs and the human body that are correlated with other clinical variables [16]. In addition, metabolomic analysis is considered in environmental sciences as a technique to evaluate the physiological responses of organisms to the effect of drugs, toxic substances, or metabolic disorders, from the initial chemical interaction to the final adverse outcome [15].

*Lemna minor* is a small, free-floating duckweed species. Fast growth, microbial reduction, and high nutrient and metal accumulation potential are the factors that candidate *Lemna* for phytoremediation research [17]. It belongs to the family Lemnaceae, along with four other aquatic genera, containing various chemical constituents such as amino acids, organic acids, sterol, terpenes, and flavonoids [18,19].

This study is divided into three parts according to the following objectives:

(a) A comparison study of metabolome samples was performed with an untargeted data handling workflow in two different labs with two mass spectrometers (TOF and QTOF) using the same plant material type;

(b) Further, the metabolomics data from different mass spectrometers were analyzed and compared with predictive methodology orthogonal partial least squares—discriminant analysis (OPLS-DA). The discrimination method has been adapted to validate the features that were extracted with the workflow. Consequently, the standard statistical methods of metabolomics data investigation were used in the identification of relevant variables (i.e., conditional attributes to each solvent), which related to the discrimination analysis. Three different extracts were systematically analyzed with this workflow;

(c) Furthermore, the plant metabolites identification workflow was described from the theoretical predictions to the final analyses in *Lemna* metabolic profile using reference materials.
The usage of the open-source PI compound database to identify the different metabolites introduces another helpful tool for plant metabolomics. The PI compound database contains plant metabolites that enable and encourages untargeted screening in plant metabolomics.

2. Results and Discussion

In the current work, the data analysis and comparison of *Lemna* metabolic profiles were performed, as well as a potential identification of the metabolites. The analysis strategy (Figure 1) was conducted on the three extracts (100% MeOH, 50% MeOH, and 100% H$_2$O) due to the key findings presented in the recent publication of Wahman et al. [8]. According to the analytical and statistical results, the three solvents could extract *Lemna*’s metabolic profile using the extended chromatographic method. Initially, each extract was analyzed (in triplicate) using a RPLC-HILIC-ESI-TOF-MS (system A) obtaining retention time information and accurate masses for ions. Further, the same extract types were analyzed (each in triplicate) using a RPLC-HILIC-ESI-QTOF-MS/MS (system B). The raw data of both were checked for analytical performance and quality by applying internal standards (details in Material and Methods Section 3.5).

![Figure 1. The non-target workflow flowchart (from precursor ion mass to ‘molecular names’) using a RPLC-HILIC-ESI-TOF-MS (system A) and evaluating the analytical data (including MS/MS) obtained with RPLC-HILIC-ESI-QTOF-MS/MS (system B). The parameters of peak picking and alignments were set according to the internal standards. The features were prioritized using the PLANT-IDENT database in the FOR-IDENT platform after various data filtration steps. The workflow of the data analysis with system A was also mentioned in [8,20].](image-url)

2.1. *Lemna minor* Extracts Untargeted Analysis Using Systems A and B

The obtained data was preprocessed according to the workflow mentioned in Section 3.6 and as shown in Figure 1. A comprehensive strategy was applied to identify metabolites in untargeted *Lemna minor* metabolomics using different labs with different mass spectrometers. Consequently, the mentioned parameters led to 1069, 1111, and 1098 features (in system A in lab 1 (RPLC-HILIC-TOF-MS)) and 1121, 1287, and 1498 features (in system B in lab 2 (RPLC-
HILIC-QTOF-MS/MS)) for 100% MeOH, 50% MeOH, and 100% H2O extracts, respectively. The RT-mass plots of the obtained features in three solvents (100% MeOH, 50% MeOH, and 100% H2O) are plotted in Figure 2 according to their RT (min.) and m/z. The dataset was differentiated at 15 min according to the expected features LogD values. The features of the *Lemna minor* metabolic profile eluting from 5 min to 15 min contained the very polar to moderate polar features with expected LogD < 0 (because eluting from the HILIC column). The features eluting later than 15 min represent the nonpolar features with LogD > 0 (because eluting from endcapped C18-RPLC column).

**Figure 2.** Retention time (RT)—the mass plot of *Lemna minor* metabolic profile extracted with 100% MeOH, 50% MeOH, and 100% H2O; using (a) system A (single TOF-MS) and (b) system B (QTOF-MS/MS). The blue color represents the HILIC eluting metabolites and the red color the RPLC eluting metabolites. The included table reflects the corresponding amount of features.

*Lemna minor* metabolic profile was obtained with the single TOF-MS (system A) in Figure 2a and with the QTOF-MS/MS (system B) in Figure 2b with differentiated values for HILIC and RPLC eluting features (see Table in Figure 2). Interestingly, the data showed significant differences in absolute values between the two MS 100% MeOH, 50% MeOH, and 100% H2O extracts, respectively (Figure 2). This might be explained by the higher detection sensitivity of the newer system B compared to the older system A, the different data handling processes as well as, the lower number of biological replicates of *Lemna minor*. Another hint for the sensitivity differences can be seen in the retention window from 15 to 23 min. The fundamental differences between both systems in the number of features there are obvious. This section is mainly affected by the start of compounds elution from RPLC and minor changes in the gradient of the mobile phase (details were mentioned in Wahman et al., 2020 [8]).

Further, in our previous study [8,20], the amino acids (phenylalanine, proline, tryptophan, alanine, tyrosine, aspartic acid, isoleucine, serine, and valine) were identified in *L. minor* using system A in the HILIC part [8,19]. Despite being also identified in system B in the HILIC part (Table S1) using target analysis, they were not detected with the common metabolic profile between both systems A and B. This is due to differences in the intensities of the amino acids between the two extracts see Section 3.2.

The overlap of common features between system A and B detected in HILIC and RPLC are small (data not shown). The detailed comparison using multivariate analysis is discussed in the next Section 2.2.
2.2. OPLS-DA Analysis of Lemna minor Metabolic Profiles Obtained with Systems A and B

Regarding the untargeted analysis, it is fundamental to assign the common metabolic profile between the different datasets of the same material and/or different materials obtained from the same and/or different laboratories. The untargeted analysis concept depends mainly on prioritizing databases. Thus, the datasets of each system were compared and identification of possible L. minor common metabolites was performed via the FOR-IDENT platform and the PLANT-IDENT compound databases.

The OPLS-DA calculated differences between the three extracted Lemna samples, which colored regarding the analyzed features with single TOF-MS (system A) and QTOF-MS/MS (system B). Hence, coloring the loading score plot (Figure 3a) according to the variables (features) that characterize each extract emphasizes the separation between the mass spectrometer and the dot shape reflects the different extraction solvents measured. Each feature is present in the three extracts but with different intensities (due to their physicochemical conditions and matrix effects in ionization). The OPLS-DA score plot (in Figure 3b) explained 99.2% of the variations in the various extracts ($R^2_Y (cum)$) with a higher predictive value ($Q^2 (cum)$ = 0.877). The first component ($t1$) separated the 100% H$_2$O and 50% MeOH extracts in the negative part (of the score plot) and the 100% MeOH extract in the positive part. The orthogonal component ($t2$) distinguishes significantly between the 100% MeOH and 50% MeOH in the negative and 100% H$_2$O positive parts, respectively. Concluding, from the statistical analysis, the L. minor metabolic profile has common features between the single TOF and QTOF-MS. In both systems, the different extracts could significantly be discriminated, allowing users to apply this analysis for subsequent Lemna metabolite measurements.

![Figure 3. (a) The loading plot displays the relation between the different Lemna minor samples 100% MeOH (triangular), 50% MeOH (stars), and 100% H$_2$O (squares) extracts analyzed with TOF-MS (violet) and QTOF-MS/MS (yellow); (b) OPLS-DA score scatter plot of Lemna minor samples with 95% confidence limit.](image)

2.3. OPLS-DA Analysis of Lemna minor 100% MeOH and 100% H$_2$O Extracts

The 100% MeOH and 100% H$_2$O extracts of Lemna using single TOF-MS (system A) were compared with the same extracts obtained using the QTOF-MS/MS (system B) as an example to validate the information transfer between different mass spectrometric systems and/or laboratories.

The OPLS-DA was calculated for both systems, the 100% MeOH and 100% H$_2$O extracts from Lemna samples as shown in Section 2.2. The OPLS-DA loading score in Figure 4a reflects that the number of the QTOF-MS/MS features (on the negative side of the first predictive component (pq1)) is higher than the features number of the single TOF-MS (on the positive side of pq1) the right side of the plot. As stated above it shows the effect of the physicochemical conditions, the matrix effect, and that each dataset was obtained with the different workflow of different software. However, the score plot of OPLS-DA explained 99.8% of the variations in the various extracts ($R^2_Y (cum)$) with a higher predictive value ($Q^2 (cum)$ = 0.891) (Figure 4b). The first components ($t1$) separated...
the different extracts according to the mass spectrometers. The statistical comparison of the data was revealed that the two sets are significantly different with a 95% confidence level. Further, the S-plot was created from the OPLS-DA model. Thereby, common metabolites can be highlighted that could be detected by the two MS in both extracts (Figure 4c). The features were with higher p(corr) values in 100% MeOH extract means that they have higher intensity in 100% MeOH extract compared to 100% H2O extract and vice versa. Hence, the common compounds were chosen according to p(corr) values which were small and approximately equal. The compounds marked in red represent the common metabolic profile of *Lemna minor* in both solvents using systems A and B (Figure 4c) and are listed as features in Table S4.

![Figure 4](image)

**Figure 4.** (a) The loading plot displays the relation between the different *Lemna minor* samples 100% MeOH and 100% H2O extracts analyzed with single TOF-MS (system A) and QTOF-MS/MS (system B); (b) OPLS-DA score scatter plot of *Lemna minor* samples 100% MeOH (triangular) and 100% H2O (squares) extracts analyzed with single TOF-MS (violet) and QTOF-MS/MS (yellow). The confidence limit is 95%; (c) The S-plot of *Lemna minor* 100% MeOH and 100% H2O extracts with red marked molecules that represent the common ones between the two machines. Tryptophan, aspartic acid, alanine, and valine were prioritized by the PLANT-IDENT database and identified using reference standards as shown in Sections 2.4 and 2.5.

### 2.4. The Strategy of *Lemna minor* Metabolites Identification Based on the PLANT-IDENT Database (Using System B)

Data from system B was preferred due to the presence of fragment spectra from MS/MS measurements. Metabolites identification from known databases depends substantially on the following: (1) comprehensive data integration and prioritization of retention time, mass, and MS/MS fragments, and (2) considering or neglecting of MS/MS intensity information [21]. With the PI database, the uploaded analytical data can be compared and transferred via the FI platform into physicochemical parameters such as polarity and empirical formula. RT and mass are applied simultaneously as a prerequisite with MS/MS data for scoring evaluation (details in Section 3.6.2). Then, to reduce the false fragments determination, the intensity threshold is defined. The threshold differentiates the line between the signals and noise level, see Section 3.6.2.
2.4.1. PLANT-IDENT Batch Searching and Scoring

(a) First, the features eluted from the HILIC column with RT < 15 min were uploaded into the FI platform using the PI database (Figure S1a). The search was scored according to the mass screening, and MS/MS, each with a 50% score. Subsequently, 239 candidates were suggested according to a successive elimination approach. First, the matching features with the highest score and labeled as ‘look at’ in the platform, according to mass matching screening, and MS/MS results were considered. Then, the results were filtered and features with LogD > 0 were eliminated. After that, the chemotaxonomic filter was applied. The results were accepted, which were found in the Lemnaceae. In the end, the results with priority were considered and standards reference organized and injected for confirmation (Figure S1a). The filtered parameters decreased the number of the results from 239 to 41 potential candidates. Those metabolites were annotated and classified in the second classification level [22,23]. They could be classified to level one by confirmation with standards reference injections (see Section 2.4.2);

(b) Secondly, the RP part with RT > 15 min, thus the second part of each dataset, was uploaded. Additionally, the retention times of the reference standard mixture were uploaded to normalize the RT (Table S3 and Figure S1b). Here, the scoring of suspected compounds depends on the same parameters in addition to RTI screening [24]. Each is 33% of the total score (mass screening, RTI screening, and MS/MS). One hundred and eighty-eight features were suggested as a matching candidate with the highest score and labeled as ‘look at’ in the platform. After, the adjusted logD > 0 and chemotaxonomic filter exclusively 42 candidates were considered. Those metabolites were annotated and classified in the second classification level. They could be classified to level one (i.e., an identification) by confirmation with standards reference injections (see Section 2.4.2).

2.4.2. Confirmation of Lemna minor Metabolites Using the QTOF-MS/MS

The metabolic profile was interpreted according to the following workflow from ‘precursor ion mass into molecule names’ (using data from system B, Section 2.4.1). The annotated metabolites (in classification level 2) were classified to level 1 via proven with standard reference injection [22,23]. Forty-four compounds (16 from the 41 HILIC candidates in Section 2.4.1 (a) and 28 from the 42 RPLC candidates in Section 2.4.1 (b)) could be identified with the reference standards and classified into identification level 1. These compounds are listed in Table 1 regarding their retention times. The 44 compounds have RT mostly deviations of less than 0.3 min (36 compounds below 0.3 min, five compounds below 0.5 min, one compound at 0.6 min and two compounds above 0.6 min), and a mass deviation of less than 5 ppm between the reference standard mass and the Lemna minor candidate feature mass. Further, MS/MS fragments of the references standards were compared with the literature. Then, the most important fragments (which were mentioned in each corresponding literature (Table 1)) were compared with unknown features fragments.
Table 1. Compounds identified in *Lemna minor* metabolic profile via PLANT-IDENT database. Retention time (RT) of standards (S), measured features (M), and the RT deviation, as well as the mass of standards (S), measured features (M), and the deviation between them, were listed. The mean fragments of standards and measured were compared with the literature and listed with the references; (*) means compounds detected in system A (single TOF-MS).

| Compound Name                          | RT (S) [Min] | RT (M) [Min] | ∆RT [Min] | Mass (S) [Da] | Mass (M) [Da] | ∆ppm | MSMS Fragments | References                        |
|----------------------------------------|--------------|--------------|-----------|--------------|--------------|------|----------------|-----------------------------------|
| Vitexin                                | 7.5          | 7.3          | 0.2       | 433.1133     | 433.1129     | 0.8  | 433;415;397;379;337;313;283 |
| Niacin *                               | 7.6          | 7.8          | −0.2      | 124.0394     | 124.0393     | 0.7  | 124;96;80;78 |
| Nicotinamide                           | 7.8          | 7.6          | 0.1       | 123.0554     | 123.0553     | 0.8  | 123;106;80;78 |
| Phenylalanine *                        | 11.0         | 11.1         | −0.1      | 166.0866     | 166.08627    | 2.0  | 120;103;77    |
| Leucine/Isoleucine *                   | 11.2         | 11.2         | −0.1      | 132.1018     | 132.1020     | 2.0  | 86;69;44;30   |
| Tryptophan *                           | 11.7         | 11.7         | 0.0       | 205.0973     | 205.0970     | 1.8  | 188;146;144   |
| Valine *                               | 12.1         | 11.9         | 0.1       | 118.0863     | 118.0862     | 0.8  | 72;71;55      |
| Tyrosine *                             | 12.3         | 12.2         | 0.1       | 182.0811     | 182.0810     | 1.9  | 136;123;119   |
| Proline *                              | 12.4         | 12.4         | 0.0       | 116.0705     | 116.0707     | 0.3  | 70;68;43      |
| Glutamic acid *                        | 12.5         | 12.6         | −0.1      | 147.0434     | 147.0430     | 0.3  | 130;102;84    |
| Aspartic acid *                        | 12.7         | 12.7         | 0.0       | 134.0447     | 134.0447     | 0.0  | 134;115       |
| Leucine/Isoleucine *                   | 13.4         | 13.1         | 0.3       | 179.0706     | 179.0708     | 0.9  | 147;137       |
| Alanine *                              | 13.3         | 13.2         | 0.2       | 90.0550      | 90.0548      | 2.1  | 44;28         |
| Threonine *                            | 13.6         | 13.4         | 0.2       | 120.0656     | 120.0653     | 2.8  | 73;56         |
| Serine *                               | 14.0         | 13.8         | 0.2       | 106.0500     | 116.0499     | 0.9  | 60;42;43      |
| Apigenin-6,8-di-C-glucopyranoside *    | 15.8         | 15.7         | 0.1       | 595.1659     | 595.1658     | 0.2  | 595;383       |
| Apigenin-6-C-arabopyranoside-8-C-glucopyranose * | 23.4 | 23.3 | 0.2 | 565.1550 | 565.1557 | −1.2 | 565;547;379;337;325;295;121 |
| Luteolin-3′,7-di-O-glucoside            | 23.8         | 23.6         | 0.3       | 611.1640     | 611.1622     | 2.8  | 611;449;287   |
| Saponarin                              | 23.8         | 24.0         | −0.2      | 595.1638     | 595.1663     | −4.2 | 433;415;397;367;337;283;271 |
| Isoorientin                            | 24.7         | 24.4         | 0.2       | 163.0796     | 163.0791     | 0.9  | 135;107       |
| Chlorogenic acid *                     | 24.8         | 24.9         | −0.1      | 303.0549     | 303.0546     | 1.7  | 303;285;257;229;165 |
| Luteolin                               | 24.6         | 24.6         | 0.2       | 287.0562     | 287.0557     | 1.6  | 287;269;241;153 |
| Naringenin-7-O-glucoside               | 25.0         | 24.1         | 0.9       | 435.1298     | 435.1285     | 2.9  | 435;273       |
| Myricetin                              | 25.1         | 25.1         | 0.0       | 319.0440     | 319.0453     | 4.0  | 301;283;265;111 |
| Orientin                               | 25.2         | 25.2         | −0.1      | 449.1123     | 449.1134     | −2.6 | 449;329       |
| Oeninidin                              | 25.6         | 25.2         | 0.4       | 302.0785     | 302.0792     | −2.4 | 302;283;197   |
| Chrysoeriol                            | 26.9         | 26.8         | 0.1       | 301.0731     | 301.0722     | 2.9  | 286;121       |
| Tricin                                 | 26.8         | 26.3         | 0.6       | 331.0811     | 331.0796     | 4.7  | 331;315       |
| Apigenin                               | 26.8         | 26.7         | 0.1       | 271.0603     | 271.0604     | −0.6 | 271;253;153   |
| Acacetin                               | 28.8         | 29.1         | −0.3      | 285.0759     | 285.0760     | −0.4 | 265;242;153   |
| Kaempferol                             | 29.0         | 29.1         | −0.1      | 287.0531     | 287.0540     | −1.1 | 287;269;231;165;153;133 |
| Galangin                               | 29.4         | 29.4         | −0.1      | 271.0602     | 271.0608     | 2.3  | 271;253       |
| 6-Methoxyflavone                       | 29.9         | 29.6         | 0.2       | 223.0756     | 223.0748     | 3.6  | 223;178;152;121 |
| 5-Hydroxy-6-Methoxyflavone             | 30.4         | 30.6         | −0.2      | 253.0879     | 253.0881     | −0.7 | 253;238;210   | NIST                             |
| Flavone (2-Phenylicromone)             | 31.3         | 31.1         | 0.1       | 269.0823     | 269.0819     | 1.3  | 269;254;104   | (MoNA)                           |

https://mona.fiehnlab.ucdavis.edu accessed on 30 November 2021.
Sixteen compounds were eluted from the HILIC column and could be identified with the following reference compounds: vitexin, niacin, nicotinamide, phenylalanine, Leucine/isoleucine, tryptophan, valine, tyrosine, proline, glutamic acid, aspartic acid, di-L-alanine, 4-methoxy cinnamic acid, alanine, threonine, and serine. They have LogD < 0 at (pH = 7). In a previous study, the amino acids (phenylalanine, proline, tryptophan, alanine, tyrosine, aspartic acid, isoleucine, serine, and valine) were identified in L. minor using system A. The RT and mass deviations were also less than 0.3 min and 5 ppm, respectively [8,20,41]. The RT and mass deviations of both systems were less than 0.5 min. and 10 ppm. As an example for HILIC-eluting compounds the extracted ion chromatogram, m/z spectrum, and MS/MS spectrum of niacin (from Table 1) are compared in Figure S2 with the feature eluting at 7.8 min.

Twenty-six compounds were eluted from the RPLC column and could be identified with the following reference compounds: apigenin-6,8-di-C-glucopyranoside, apigenin-6-C-arabinopyranoside-8-C-glucopyranose, apigenin, robinetin, quercetin, luteolin, kaempferol, acacetin, orientin, isoorientin, peonidin, 6-methoxy-flavone, flavone, naringenin-7-O-glucoside, quercetin-3-O-glucoside, saponarin, 5-hydroxy-6-methoxy-flavone, luteolin-3',7-di-O-glucoside, apiin, chrysoeriol, umbelliferone, norwogonin, isovitexin, tricin, galangin, and myricetin. Further, the MSMS fragments of the reference standard and the identified peaks in the Lemna metabolic profile were compared with the literature, as shown in (Table 1).

As an example for RPLC-eluting compounds the extracted ion chromatogram, mass spectrum, and MS/MS spectrum of apiin are compared in (Figure S3) with the feature eluting at 23.8 min.

According to chemotaxonomy criteria results, 42 compounds had previously been separated from Lemnaceae or Araceae family. According to the available literature, robinetin and norwogonin were not reported from Lemnaceae or Araceae family so far. Those two compounds could not pass the last filter of the chemotaxonomy because they are a pentahydroxy-flavone and 5,7,8-trihydroxy-flavone, respectively.

Besides, robinetin is known as 5-deoxy-myricetin. Therefore, robinetin and norwogonin reference standards were injected. In addition, luteolin, quercetin, kaempferol, apigenin, and myricetin (hydroxylated-flavone and flavanol) and their glucosides have been identified in the family before [31,42]. According to our knowledge, robinetin and norwogonin were identified for the first time in the Lemna minor extracts.

2.5. The Strategy of Lemna minor Metabolites Identification Based on the PLANT-IDENT Database (Using Common Data from Both MS Systems)

The common compounds in the metabolic profile of Lemna minor (systems A and B) were chosen according to p(corr) values which were marked in red (Figure 4c and Table S4). The obtained features were classified according to RT, and, prioritized using the PI database on the FI platform. The search was scored according to the mass screening (and MS/MS in system B). The PLANT-IDENT database suggested pegamine, tryptophan, aspartic acid, alanine, valine, and betaine (trimethylglycine) candidates matching features in both 100% MeOH, and 100% H₂O extracts. They have LogD < 0 at pH = 7 (except pegamine, which was eliminated). They also have priority ‘look at’ and standards reference injected for validation except for betaine (Figure S1a). Subsequently, the filtration parameters decreased the number of the results to 5 metabolites, listed in Table S5. Those metabolites were annotated and classified in the second identification level. They could be classified into level one by confirmation with standards reference injections.

Regarding the RP part with RT > 15 min, the second part of each dataset was uploaded into the PI database in the FI platform, individually. Two features were suggested as a matching candidate in 100% MeOH, and 100% H₂O extracts. They did not have any matches in the PI database. Those metabolites were annotated and classified in the third level.
Thus, the identification process using physicochemical parameters and the knowledge from mass spectrometric results can easily be transferred into further knowledge and to other mass spectrometers with the continuous updating of the PI database.

3. Materials and Methods

3.1. Reagents and Chemicals

LC-MS grade methanol (MeOH) and water (H$_2$O; LC-MS grade) were obtained from VWR, Darmstadt, Germany. Acetonitrile (LC-MS grade) was purchased from Riedel-de-Haen (Honeywell, Seelze, Germany). 6-Amino-1,3-dimethyl-5-(formylamino)uracil, chlorotoluron, famotidine, etilefrine, and 2,4-diamino-6-(hydroxymethyl)pteridine hydrochloride, and apigenin, monuron, chloridazon, carbetamide, metaboluron, sotalol, metconazol, vitexin, flavone (2-phenylchromone), DL-Ala-DL-Ala, nicotinic acid, nicotinamide, galangin, and acacetin were obtained from Merk, Darmstadt, Germany. Chlorbromuron was obtained from Dr. Ehrenstorfer, Augsburg, Germany. Apigenin-6-C-arabinoside-8-C-glucoside, Apigenin-6,8-di-C-glucoside, apiin, orientin, kaempferol, peonidin, norwogonin, luteolin-3,7-O-di-glucoside, 6-methoxy flavone, 4-methoxy cinnamic acid, naringenin-7-O-glucoside, isovitexin, tricin, quercetin-3-O-glucoside, saponarin, myricetin, 5-hydroxy-6-methoxy flavone, chrysoeriol, isovitexin, robinetin, and umbelliferone were kindly provided by Center of Life and Food Science Weihenstephan, Biotechnology of Natural Products, Technical University of Munich, Germany.

3.2. Plant Samples

*Lemna minor* L. was cultivated according to Obermeier et al. \[43\] with small modifications. Briefly, plants were grown at 23 °C with a photoperiod of 16–8 h and an average light intensity of 43 µmol m$^{-2}$ s$^{-1}$. *Lemna* fronds were subcultured every two weeks in 24 L of Steinberg medium. After harvesting, plants were shortly rinsed with distilled water, dried with lint-free tissue paper, and frozen in liquid nitrogen. Samples were kept at −80 °C until further processing. *Lemna* was kindly provided by the German Research Center for Environmental Health, Plant–Microbe Interactions, Helmholtz Centrum of Munich, Germany.

3.3. Sample Preparation

Five hundred milligrams of freeze-dried and milled whole *Lemna* powder were extracted with (a) 100% MeOH, (b) 50% MeOH, and (c) 100% H$_2$O, respectively. The dissolved plant powders were sonicated (35 kHz, Sonorex super RK 106, Bandelin, Berlin, Germany) for 10 min at 4 °C with a 35 kHz frequency. Then, samples were centrifuged (NuWind Multi-Application Bench Top Refrigerated Centrifuge 2, NuAire, Plymouth, MN 55447, USA) at 1500 rpm/261.6 × g for 20 min at 4 °C and the supernatants were transferred into clean glass test tubes. The extraction process was triplicated in identical experimental conditions. Finally, the extracts were evaporated to dryness (using a SpeedVac, Fischer Scientific, Göteborg, Sweden) and dissolved in 1 mL of 50% MeOH \[8,41\]. The extracts were diluted at 1:100 before the analysis.

For this work, separate batches of *Lemna* were used in each system analysis (A and B). In system B, two different biologicals were used, which were injected separately with a time interval of one month.

3.4. Instruments

3.4.1. Chromatographic System for Polarity Extended Separation

*Lemna* metabolomics and reference standards were separated with LC (Agilent 1260 Infinity, 5301 Stevens Creek Blvd., Santa Clara, CA, USA) consisting of an autosampler, two binary pumps, an online degasser, a mixing chamber, a UV detector, and an isotropic pump. The LC-system was used to perform separations in a serial coupling of reversed-phase and zwitterion hydrophilic interaction liquid chromatography (HILIC). The reversed-phase separation column was a Poroshell 120 EC-C18 (50.0 × 3.0 mm, 2.7 µm; Agilent Technologies,
The HILIC column was a ZIC-HILIC column (150 × 2.1 mm, 5 µm, 200 Å; Merck Sequant, Umeå, Sweden). Columns were coupled through a T-piece (Upchurch, IDEX Europe GmbH, Erlangen, Germany). The third port of the T-piece was connected to the HILIC flow pump. The injection volume was 10 µL.

The mobile phase of the serial coupling (RPLC-HILIC-TOF-MS) was employed as follows: Solvent A: 10 mM of ammonium acetate in 90:10 (v/v) water–acetonitrile; solvent B: 10 mM of ammonium acetate in 10:90 (v/v) water–acetonitrile (RPLC); solvent C: acetonitrile; solvent D: water (HILIC). Further mobile phase conditions are summarized in Table 2. Further details, like the RPLC-HILIC serial coupling settings, are described in [8,44,45]. The effluent of the HILIC column was connected with another T-piece (Upchurch, IDEX Europe GmbH, Erlangen, Germany). An isocratic pump was connected to the second port of the T-piece. It provided a continuous flow of 50 µL/min acetonitrile-water (90:10, v:v) with 125 nM of purine and 6.25 nM of HP-921 MS tuning mix (Agilent Technologies, Waldbronn, Germany) for constant MS recalibration. The third port of the T-piece was connected to the ion source of the mass spectrometer.

| Time (min) | Flow Rate (mL/min) | A% | B% | Time (min) | Flow Rate (mL/min) | C% | D% |
|------------|--------------------|----|----|------------|--------------------|----|----|
| 1          | 0.05               | 100| 0  | 0          | 0.4                | 100| 0  |
| 7          | 0.05               | 100| 0  | 6          | 0.4                | 100| 0  |
| 12         | 0.05               | 50 | 50 | 13         | 0.4                | 60 | 40 |
| 13         | 0.1                | 50 | 50 | 32         | 0.4                | 60 | 40 |
| 22         | 0.1                | 0  | 100| 33         | 0.8                | 100| 0  |
| 32         | 0.1                | 0  | 100| 53         | 0.8                | 100| 0  |
| 33         | 0.1                | 100| 0  | 54         | 0.4                | 100| 0  |
| 53         | 0.1                | 100| 0  | 58         | 0.4                | 100| 0  |
| 54         | 0.05               | 100| 0  |
| 58         | 0.05               | 100| 0  |

3.4.2. Mass Spectrometric Detection System A in Lab 1 (Single TOF-MS)

Samples were analyzed and detected with a ‘time-of-flight’ mass spectrometer (G6230A; Agilent Technologies, Waldbronn, Germany), equipped with the Jet Stream II ESI interface (system A). Ions were detected in positive ionization mode with a mass range of 50–2100 Dalton. The parameters were as follows: 325 °C gas temperature, 10 L/min drying gas flow, 325 °C sheath gas temperature, 7.5 L/min sheath gas flow, 45-psi nebulizer operating pressure, and 100 V fragmentor voltage.

3.4.3. Mass Spectrometric Detection System B in Lab 2 (QTOF-MS/MS)

The other utilized MS system was a Triple TOF 5600 (AB SCIEX, Darmstadt, Germany) (system B) with a Duospray ion source and a TurbolonSpray ESI probe in positive ion mode. A mass range of m/z 65 to 2000 was scanned in full range. MS/MS was acquired in independent data acquisition mode with a mass tolerance of 50 mDa. The intensity threshold was 50 cycles per second (cps). The MS/MS data were collected in the eight parallel experiments at a collision energy of 40 ± 20 eV. The ionization parameters were set as the following: 2000 kV ion spray voltage floating (ISVF) and 650 °C turbo spray temperature. The declustering potential (DP) and collision energy (CE) were set to 46 and 40 V, respectively. The nebulizer gas (gas 1), the heater gas (gas 2), and the curtain gas were set to 44, 50, and 29 psi, respectively. All gas flows were nitrogen.
3.5. Internal Standards

Each sample and blank were spiked with a standards mixture of 12 substances. The mixture consists of 6-amino-1,3-dimethyl-5-(formylamino) uracil, chlortoluron, famotidine, vidarabine, etilefrine, monuron, carbetamide, metobromuron, sotalol, chlorbromuron, metconazol, and 2,4-diamino-6-(hydroxymethyl) pteridine hydrochloride to obtain a final concentration of 5 µM each. These internal standards were used to estimate the mass deviation as an absolute variation between theoretical monoisotopic mass and the mean of measured isotopic masses were calculated as (Δppm) as well as the retention time stability across all experiments. The results are summarized in (Table S2). These parameters were used to correlate the unknown features in different samples.

3.6. Data Collection and Preprocessing
3.6.1. Single TOF-MS (System A)

Data were acquired with MassHunter Workstation LC/MS Data Acquisition software B 05.00 (Agilent Technologies, Waldbronn, Germany), and was subsequently analyzed with Profinder B.06.00 (Agilent Technologies) extracting the so-called ‘features’ by their RT, molecular mass, and their peak intensity in various extracts. This was performed in a combination of the three injections of each sample after removing the features found in the corresponding blank samples. The parameters are set to a peak filter of 1000 counts peak height, ion species to ‘positive ions’ with H⁺, Na⁺, K⁺, and NH₄⁺, ‘charge state’ to ±1, the ‘expected RT’ to ±3.00 min, and the mass to ±10 ppm. The extracted ion chromatograms (EICs) were smoothed with a Gaussian function using 9 points function width and 5000 points Gaussian width. This limits the result finally to 2000 compound groups. The detailed workflow was discussed in Wahman et al., 2022 [20].

3.6.2. QTOF-MS/MS (System B)

The data were obtained with Analyst Software (version TF 1.7.1). The data evaluation consisted of peak picking, alignment, and filtering, which was done with MarkerView Software (AB SCIEX, version 1.3.1). The metabolites identification was performed using the vendor software package (Sciex OS).

Peak picking: The peak picking parameters were optimized according to the mass and retention deviations of the internal standards in all samples and blanks. The minimum and maximum retention times were 5 and 34 min, respectively. The subtraction offset was 15 scans. The noise threshold and subtraction multiplication factors were 50 and 1, respectively. Further, the minimum spectral peak width was 2 ppm (Figure 1). The noise threshold and subtraction multiplication factors have a fundamental impact on peak picking especially for the low abundance metabolites due to the matrix effect of unfractionated extracts: This was achieved using internal standards. Transformation and/or normalization of the data was also acquired in this step, which is required for univariate and multivariate statistical analysis.

Alignment and filtering: The three injections of each extract were compared and aligned. The RT variations between different injections were expected to be as good as those, observed with the internal standards. Further, mass tolerance was set to 5 ppm based on QTOF specification [46], and observations from internal standards. Features were accepted if they were found in all three injections of a sample. Then, the background was deleted (i.e., the features found in the blank corresponding to each solvent were deleted from the same extract) (Figure 1). *Lemna minor* metabolic profile investigation demands alignments of features, which were found in different extracts and the different injections of similar extracts. Further, the features were deleted, which were found in the corresponding blank.

In addition, the RTI and mass tolerance were determined according to the results of the internal standard, which led to a decrease in the number of false-positive, as well as, negative peaks (i.e., reduction of the number of features from the same metabolite). The
minimum variation in the noise threshold was affected the peak extraction by increasing
the number of false-positive peaks or subtracting real peaks apart.

PLANT-IDENT batch searching and scoring: After alignment and filtering, data
were uploaded into the FI platform. The molecular masses of detected compounds were
compared with compounds stored in the PI database, resulting in suggestions for their
identification. The compound database PI containing various plant metabolites was created
and implemented in the FI platform. It contains up to over 3000 plant metabolites from
Lemnaceae, Poaceae, Brassicaceae, Nymphaceae families, in addition to, flavonoids, and
nitrogen-containing plant metabolites. The compound’s name and plant source were col-
clected from the available literature (articles and/or books) which was cited in the database.
The chemical identifiers (smiles, LogD, InChI Key) and physicochemical properties were
gathered from the PubChem and Chemsipider databases. The LogD values (obtained by
ChemAxon) were used to support the identification of compounds through hydrophilicity
for the HILIC eluted compounds. Further, the RTI was used to support the identification
of RPLC eluted compounds via hydrophobicity. The retention time of the standards mix-
ture (Table S2) was also uploaded to normalize the RT. The normalization was performed
according to the calibration curve (correlation between the RT and Log D at pH = 7) of
the target analysis of the standard mixture, for more details readers referred to [24]. The
mixture consists of compounds, which have affixed logD at different pHs (i.e., compounds
with stable logP values) (Table S2).

The features with the highest score (the sum of different prioritization criteria e.g.,
mass deviation, RT, and MSMS fragments with equal weighting factors) were marked in
the ‘look at’ column in the FOR-IDENT platform, which was calculated via the platform.
Consequently, the marked hits fragments were compared through in-silico fragmenta-
tion MetFrag [47]. Moreover, chemotaxonomy was applied because the PLANT-IDENT
contained the plant name and/or family name, which was reported to contain the corre-
sponding metabolites.

Those features were considered and standards reference injected for validation. After
injection of references standards, the important peak of each has been characterized using
the available literature (Table 1). The fragments were compared with the different features
using the vendor software package (Sciex OS).

3.6.3. PI Batch Searching and Scoring of System A Data

The processed data from systems A were also uploaded into the FI platform. The
search process was performed according to the following parameters: pH = 7, precursor
ion mass deviation 5 ppm, and ion species was positive. The metabolites were considered
according to their score, which is the same as system B data except for the MS/MS data.
Then, the results were considered when they have negative LogD at pH = 7. In contrast, the
metabolites were suspected in the RP column according to the score of the same parameter
in addition to the RTI [24]. The metabolites were considered when they have positive LogD.
The last step was the chemotaxonomy criteria. Each metabolite has a plant and/or family
name reference according to the available literature (Figure 1).

3.6.4. Classification Scheme

The different extracts were analyzed according to the previously mentioned workflow
(Figure 1). Then, the plant metabolites classification was performed adopted to the scheme
of Letzel et al., 2014, and Schymanski et al., 2014 and Letzel et al., 2014 [22,23], which
complies five levels:

1. The identification by the reference standard;
2. The identification was performed by various criteria such as (retention time behav-
ior, accurate mass (i.e., empirical formula), fragmentation, and chemotaxonomical
criteria);
3. The identification was performed by comparison of accurate mass and fragments
from different laboratories;
4. The identification is done by molecular formula or fragments comparison;
5. Mass recognition without further information; this classification scheme enhances the identification of plant metabolites in untargeted metabolomics analysis.

3.6.5. Orthogonal Partial Least Square—Discriminant Analysis

The quality of features was statistically investigated from both systems (Figures 3 and 4). In each experiment, *Lemna* was grown in the same condition and extracted with 100% MeOH, 50% MeOH, and 100% H$_2$O extracts. The data matrix was built in the Microsoft Access Database file according to the aligned qualitative data (i.e., feature annotations/abundance as columns and sample annotations as rows), along with related metadata (i.e., plant name, extraction solvent, and mass spectrometer type). The RTs, masses, and abundances were connected to the corresponding plant, extraction solvent, and mass spectrometer type. Once the matrix was built, comprehensive statistical analyses could be performed by using the vast range of functions provided by the SIMCA software. Additionally, the mass spectrometric system was included as a secondary observation. The dataset organization and different statistical parameters were mentioned in detail in [9].

OPLS is the orthogonal modification of PLS regression analysis methods, which both are supervised statistical models on the contrary of PCA [48]. The OPLS fits well for metabolomics analysis because it can analyze a large number of variables for a small sample size. The OPLS separates the variables (X) (i.e., 8940 features) into two directions linear and orthogonal to Y. The lower number of biological replicates in this study might affect the statistical conclusion. Thus, in further studies, a community-harmonized number of more biological replicates will be analyzed to reach a power analysis > 0.8. The OPLS model can be visualized with the score plot and the loading plot, which describe the contribution of the variables. The OPLS quality is described with the cumulative variation in the matrix or cumulative variation ($R^2_X$; cum), the cumulative variation in the Y matrix (extracts) or $R^2_Y$(cum), and the cross-validated predictive ability or $Q^2$(cum) values. $R^2$ is defined as a fraction of the variance explained by a component. Cross-validation of $R^2$ gives $Q^2$, which represents the proportion of total variation predicted by a component. Thus, the $R^2$ indicates how well the variation of a variable is explained, and $Q^2$ how well a variable could be predicted and estimated by cross-validation [49].

4. Conclusions

*Lemna*’s metabolic profile was investigated using an extended polarity liquid chromatographic system.

The applied workflow(s) was (were) investigated with statistical analysis to validate the reliability and information transfer between different mass spectrometric systems and/or laboratories. The identification strategy of untargeted data was applied using an open access PLANT-IDENT database. The identification of *Lemna* metabolites proceeded according to different filters: LogD, mass deviation, MSMS fragment comparison, and chemotaxonomy filter. After prioritization, compounds were identified using reference standards leading to compounds of identification level 1. Further, from the statistical analysis, the *L. minor* metabolic profile has common features between the single TOF and QTOF-MS. The different extracts could be significantly discriminated in both MS, which could be used for further generations of *Lemna* metabolite measurements.

Thus, the untargeted plant metabolomics research will be enhanced via utilizing the workflow combined with the PLANT-IDENT database and with continuous updating of the database.

**Supplementary Materials:** The following are available online at [https://www.mdpi.com/article/10.3390/metabo11120832/s1](https://www.mdpi.com/article/10.3390/metabo11120832/s1), Figure S1: (a) Search for the feature masses, which were separated with the HILIC column from the 100% MeOH extract; (b) RTI handling according to the pH of the compounds as the prioritization tool in PLANT-IDENT database in the FOR-IDENT platform for more information readers referred to the FOR-IDENT manual ([https://water.for-ident.org/#!home](https://water.for-ident.org/#!home)), Figure S2: (a) Extracted ion chromatogram (EIC) of niacin shows intensity in counts per second (cps),
appears on the y-axis, while RT appears on the x-axis; (b) mass spectrum of niacin represents the masses and intensity of the ions with particular mass-to-charge (m/z) values in Daltons; (c) MS/MS spectrum of niacin shows the fragments and relative intensity (%), Figure S3: (a) Extracted ion chromatogram (EIC) of apiin shows intensity in counts per second (cps), appears on the y-axis, while RT appears on the x-axis; (b) mass spectrum of apiin represents the masses and intensity of the ions with particular mass-to-charge (m/z) values in Daltons; (c) MS/MS spectrum of apiin shows the fragments and relative intensity (%), Table S1: The compounds of *Lemna minor* metabolic profile identified in the different samples which have been measured with Agilent and Sciex machines, Table S2: The internal standards mean monoisotopic mass in Daltons, mean mass standard deviation (SD), mean RT in minutes, mean RT standard deviation (SD), and relative standard deviation (RSD) were listed, Table S3: The RTI calibration compounds mean monoisotopic mass of in Daltons, mean mass standard deviation (SD), mean RT in minutes, mean RT standard deviation (SD), and relative standard deviation (RSD) were listed, Table S4: The list of marked compounds in the S-plot, Table S5: The proposed compounds in the HILIC part of *Lemna minor* metabolomics by the PLANT-IDENT database.

**Author Contributions:** R.W. and T.L. designed the metabolomics study. R.W. prepared the samples, performed the untargeted metabolomics analysis, and ran the analyses in the metabolomics platform. S.M. created the statistical design for data evaluation and data interpretation and performed the realization together with R.W. and T.L., S.B. supported the study with analysis concepts on the QTOF system and proofing the analytical realization with R.W. R.W. and T.L. conceived and drafted the manuscript. S.M., S.B., C.C., P.S., A.G. and F.L. proof-read and corrected the manuscript. T.L., A.G., and F.L. concerted the FOR-IDENT platform and included compound databases. R.W. created the PLANT-IDENT compound database. All the authors contributed with critical intellectual input. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research was partially funded by the Bavarian State Ministry of the Environment and Consumer Protection, by the German Federal Ministry of Education and Research (funding ‘FOR-IDENT-Nachhaltigkeit’ (02WRS1456)) as well as the Cultural Affairs and Mission Sector of the Egyptian Ministry of Higher Education. C.C. was funded by the Water Joint Programming Initiative (WATER 21015 JPI) through the European research project ‘IDOUM - Innovative Decentralized and low-cost treatment systems for Optimal Urban wastewater Management’.

**Institutional Review Board Statement:** Not applicable.

**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** The data presented in this study are available on request from the corresponding author. The data are not publicly available due to continuous work on it.

**Acknowledgments:** We thank Axel Besa and Sciex for providing the TripleTOF mass spectrometer and valuable support.

**Conflicts of Interest:** The authors declare that they have no conflict of interest.

**References**

1. Piovesana, S.; Cavaliere, C.; Cerrato, A.; Montone, C.M.; Laganà, A.; Capriotti, A.L. Developments and pitfalls in the characterization of phenolic compounds in food: From targeted analysis to metabolomics-based approaches. *TrAC Trends Anal. Chem.* 2020, 133, 116083. [CrossRef]
2. Ikram, M.M.M.; Ridwani, S.; Putri, S.P.; Fukusaki, E. GC-MS based metabolite profiling to monitor ripen-ing-specific metabolites in pineapple (Ananas comosus). *Metabolites* 2020, 10, 134. [CrossRef] [PubMed]
3. Aretz, I.; Meierhofer, D. Advantages, and pitfalls of mass spectrometry-based metabolome profiling in systems biology. *Int. J. Mol. Sci.* 2016, 17, 632. [CrossRef] [PubMed]
4. Doppler, M.; Bueschl, C.; Kluger, B.; Koutnik, A.; Lemmens, M.; Buerstmayr, H.; Rechthaler, J.; Krska, R.; Adam, G.; Schuhmacher, R. Stable Isotope-assisted plant metabolomics: Combination of global and trac-er-based labeling for enhanced untargeted profiling and compound annotation. *Front. Plant Sci.* 2019, 10, 1366. [CrossRef]
5. Yang, Z.; Nakabayashi, R.; Okazaki, Y.; Mori, T.; Takamatsu, S.; Kitamura, S.; Kikuchi, J.; Saito, K. Toward better annotation in plant metabolomics: Isolation and structure elucidation of 36 specialized metabolites from Oryza sativa (rice) by using MS/MS and NMR analyses. *Metabolomics* 2014, 10, 543–555. [CrossRef]
6. Gika, H.G.; Theodoridis, G.A.; Earl, M.; Snyder, R.W.; Sumner, S.J.; Wilson, I.D. Does the mass spectrometer define the marker? A comparison of global metabolite profiling data generated simultaneously via UPLC-MS on two different mass spectrometers. *Anal. Chem.* 2010, 82, 8226–8234. [CrossRef] [PubMed]

7. Sorokina, M.; Steinbeck, C. Review on natural products databases: Where to find data in 2020. *J. Cheminform.* 2020, 12, 1–51. [CrossRef]

8. Wahnman, R.; Graßmann, J.; Sauvêtre, A.; Schröder, P.; Letzel, T. Lemna minor studies under various storage periods using extended-polarity extraction and metabolite non-target screening analysis. *J. Pharm. Biomed. Anal.* 2020, 188, 113362. [CrossRef] [PubMed]

9. Chen, L.; Zhong, F.; Zhu, J. Bridging targeted and untargeted mass spectrometry-based metabolomics via hybrid approaches. *Metabolites* 2020, 11, 2. [CrossRef]

10. De Vos, R.C.; Schipper, B.; Hall, R.D. High-performance liquid chromatography-mass spectrometry analysis of plant metabolites in Brassicaceae. In *Plant Metabolomics*; Humana Press: Clifton, NJ, USA, 2012; pp. 111–128.

11. Chen, L.; Zhong, F.; Zhu, J. Bridging targeted and untargeted mass spectrometry-based metabolomics via hybrid approaches. *Metabolites* 2020, 10, 348. [CrossRef] [PubMed]

12. Martins, M.C.M.; Caldana, C.; Wolf, L.D.; de Abreu, L.G.F. The importance of experimental design, quality assurance, and control in plant metabolomics experiments. In *Plant Metabolomics*; Humana Press: New York, NY, USA, 2018; pp. 3–17.

13. Want, E.J.; Masson, P.; Michopoulos, F.; Wilson, I.D.; Theodoridis, G.; Plumb, R.S.; Shockcor, J.; Loftus, N.; Holmes, E.; Nicholson, J. Global metabolic profiling of animal and human tissues via UPLC-MS. *Nat. Protoc.* 2013, 8, 17–32. [CrossRef]

14. Salem, M.A.; De Souza, L.P.; Serag, A.; Fernie, A.R.; Farag, M.A.; Ezzat, S.M.; Alseekh, S. Metabolomics in the context of plant natural products research: From sample preparation to metabolite analysis. *Metabolites* 2020, 10, 37. [CrossRef]

15. Zhang, X.W.; Li, Q.H.; Xu, Z.D.; Dou, J.J. Mass spectrometry-based metabolomics in health and medical science: A systematic review. *RSC Adv.* 2020, 10, 3092–3104. [CrossRef]

16. Bedía, C.; Cardoso, P.; Dalmau, N.; Garreta-Lara, E.; Gómez-Canela, C.; Gorrochategui, E.; Navarro-Reig, M.; Ortiz-Villanueva, E.; Puig-Castellví, F.; Tauler, R. Chapter nineteen—applications of metabolomics analysis in environmental research. In *Comprehensive Analytical Chemistry*; Jaumot, J., Bedía, C., Tauler, R., Eds.; Elsevier B.V.: Amsterdam, The Netherlands; Oxford, UK; Cambridge, MA, USA, 2012; Volume 82, pp. 533–582.

17. Al-Snafi, A. Lemna minor: Traditional uses, chemical constituents, and pharmacological effects—A review. *IOSR J. Pharm.* 2019, 9, 6–11.

18. Jensen, S.R.; Nielsen, B.J. Chemical characters. In *The Families of the Monocotyledons: Structure, Evolution, and Taxonomy*; Dahlgren, R.M.T., Clifford, H.T., Yeo, P.F., Eds.; Springer: Berlin/Heidelberg, Germany, 1985; pp. 17–22.

19. Kotowska, U.; Piotrowska, A.; Isidorova, A.G.; Bajguz, A.; Isidorov, V.A. Gas chromatographic-mass spectrometric investigation of the chemical composition of the aquatic plant Wolffia arrhiza (Lemnaceae). *Oceanol. Hydrobiol. Stud.* 2013, 42, 181–187. [CrossRef]

20. Wahnman, R.; Cruzeiro, C.; Graßmann, J.; Schröder, P.; Letzel, T. The changes in Lemna minor metabolomic profile: A response to diclofenac incubation. *Chemosphere* 2022, 287, 132078. [CrossRef]

21. Zhao, X.; Zeng, Z.; Chen, A.; Lu, X.; Zhao, C.; Hu, C.; Zhou, L.; Liu, X.; Wang, X.; Hou, X.; et al. Com-prehensive strategy to construct an in-house database for accurate and batch identification of small molecule metabolites. *Anal. Chem.* 2018, 90, 7635–7643. [CrossRef]

22. Letzel, T.; Lucke, T.; Schulz, W.; Sengl, M.; Letzel, M. In a class of its own—OMI (Organic Molecule Identification) in water using LC-MS/MS: Steps from “unknown” to “identified”: A contribution to the discussion. *Lab More Int.* 2014, 4, 24–28.

23. Schymanski, E.L.; Jeon, J.; Gulde, R.; Fenner, K.; Ruff, M.; Singer, H.P.; Hollender, J. Identifying small molecules via high-resolution mass spectrometry: A case study with pyridine derivatives. *Rapid Commun. Mass Spectrom.* 2014, 48, 2097–2098. [CrossRef] [PubMed]

24. Grosse, S.; Letzel, T. *User Manual for FOR-IDENT Database*; TUM: Munich, Germany, 2017; pp. 1–35.

25. Wardel, P.; Wolfender, J.L.; Ndjoko, K.; Hobby, K.R.; Major, H.J.; Hostettmann, K. Evaluation of quadrupole-time-of-flight tandem mass spectrometry and ion-trap multiple-stage mass spectrometry for the differentiation of C-glycosidic flavonoid isomers. *J. Chromatogr. A* 2001, 926, 29–41. [CrossRef]

26. Ndolo, V.U.; Fulcher, R.G.; Beta, T. Application of LC-MS-MS to identify niacin in aleurone layers of yel-low corn, barley, and wheat kernels. *J. Cereal Sci.* 2015, 65, 88–95. [CrossRef]

27. Stadler, R.; Jenny, T.A.; Fay, L.B. Tandem mass spectrometric accurate mass performance of time-of-flight and Fourier transform ion cyclotron resonance mass spectrometry: A case study with pyridine derivatives. *Rapid Commun. Mass Spectrom.* 2001, 15, 1840–1848. [CrossRef]

28. Wang, X.; Zhong, X.J.; Zhou, N.; Cai, N.; Xu, J.H.; Wang, Q.B.; Li, J.J.; Liu, Q.; Lin, P.C.; Shang, X.Y. Rapid characterization of chemical constituents of the tubers of Gymnadenia conopsea by UPLC-Orbitrap-MS/MS analysis. *Molecules* 2020, 25, 898. [CrossRef]

29. Sakalem, M.E.; Negri, G.; Tabach, R. Chemical composition of hydroethanolic extracts from five species of the Passiflora genus. *Rev. Bras. Farm.* 2012, 22, 1219–1232. [CrossRef]

30. Simirgiotis, M.J.; Schmeda-Hirschmann, G.; Börquez, J.; Kennelly, E.J. The Passiflora tripartita (Banana Passion) Fruit: A source of bioactive flavonoid C-Glycosides isolated by HSCCC and characterized by HPLC–DAD–ESI/MS/MS. *Molecules* 2013, 18, 1672–1692. [CrossRef]
31. Akhtar, T.A.; Lees, H.A.; Lampi, M.A.; Enstone, D.; Brain, R.A.; Greenberg, B.M. Photosynthetic redox imbalance influences flavonoid biosynthesis in Lemma gibba. _Plant Cell Environ._ 2010, 33, 1205–1219. [CrossRef] [PubMed]

32. Pereira, C.; Yariwake, J.; McCullagh, M. Distinction of the C-glycosylflavone isomer pairs orientin/isoorientin and vitexin/isovitexin using HPLC-MS exact mass measurement and in-source CID. _Phytochem. Anal. Int. J. Plant Chem. Biochem. Tech._ 2005, 16, 295–301. [CrossRef] [PubMed]

33. Gao, X.; Sun, W.; Fu, Q.; Niu, X. Ultra-performance liquid chromatography coupled with electrospray ionization/quadrupole time-of-flight mass spectrometry for the rapid analysis of constituents in the traditional Chinese medical formula Danggui San. _J. Sep. Sci._ 2014, 37, 53–60. [CrossRef]

34. Jang, G.H.; Kim, H.W.; Lee, M.K.; Jeong, S.Y.; Bak, A.R.; Lee, D.J.; Kim, J.B. Characterization and quantification of flavonoid glycosides in the Prunus genus by UPLC-DAD-QTOF/MS. _Saudi J. Biol. Sci._ 2018, 25, 1622–1631. [CrossRef]

35. Crow, F.W.; Tomer, K.B.; Looker, J.H.; Gross, M.L. Fast atom bombardment and tandem mass spectrometry for structure determination of steroid and flavonoid glycosides. _Anal. Biochem._ 1986, 155, 286–307. [CrossRef]

36. Tsimogiannis, D.; Samiotaki, M.; Panayotou, G.; Oreopoulou, V. Characterization of flavonoid subgroups and hydroxy substitution by HPLC-MS/MS. _Molecules_ 2007, 12, 593–606. [CrossRef]

37. Le Gall, G.; DuPont, M.S.; Mellon, F.A.; Davis, A.L.; Collins, G.J.; Verhoeyen, M.E.; Colquhoun, I.J. Characterization and content of flavonoid glycosides in genetically modified tomato (Lycopersicon esculentum) fruits. _J. Agric. Food Chem._ 2003, 51, 2438–2446. [CrossRef] [PubMed]

38. Ma, Y.L.; Li, Q.M.; Van den Heuvel, H.; Claeyts, M. Characterization of flavone and flavonol aglycones by collision-induced dissociation tandem mass spectrometry. _Rapid Commun. Mass Spectrom._ 1997, 11, 1357–1364. [CrossRef]

39. Xu, L.L.; Xu, J.J.; Zhong, K.R.; Shang, Z.P.; Wang, F.; Wang, R.F.; Zhang, L.; Zhang, J.Y.; Liu, B. Analysis of non-volatile chemical constituents of Menthae haplocalycis herba by ultra-high performance liquid chromatography-high resolution mass spectrometry. _Molecules_ 2017, 22, 1756. [CrossRef] [PubMed]

40. Hughes, R.J.; Croley, T.R.; Metcalfe, C.D.; March, E.R. A tandem mass spectrometric study of selected characteristic flavonoids. _Int. J. Mass Spectrom._ 2001, 210, 371–385. [CrossRef]

41. Wahman, R.; Grassmann, J.; Schröder, C.A.; Helmreich, B.; Schröder, P. The enzymatic and antioxidative stress response of Lemna minor to copper and a chloroacetamide herbicide. _Environ. Sci. Pollut. Res._ 2015, 22, 18495–18507. [CrossRef] [PubMed]

42. Tsolmon, B.; Fang, Y.; Yang, T.; Guo, L.; He, K.; Li, G.-Y.; Zhao, H. Structural identification and UPLC-ESI-QTOF-MS2 analysis of flavonoids in the aquatic plant Landoltia punctata and their in vitro and in vivo antioxidant activities. _Food Chem._ 2020, 343, 128392. [CrossRef]

43. Obermeier, M.; Schröder, C.A.; Helmreich, B.; Schröder, P. The enzymatic and antioxidative stress response of Lemma minor to copper and a chloroacetamide herbicide. _Environ. Sci. Pollut. Res._ 2015, 22, 18495–18507. [CrossRef] [PubMed]

44. Biber, S.; Greco, G.; Grosse, S.; Letzel, T. Plant metabolomic workflows using reversed-phase LC and HILIC with ESI-TOF-MS. _LCGC N. Am._ 2019, 37, 8–15.

45. Tsolmon, B.; Fang, Y.; Yang, T.; Guo, L.; He, K.; Li, G.-Y.; Zhao, H. Structural identification and UPLC-ESI-QTOF-MS2 analysis of flavonoids in the aquatic plant Landoltia punctata and their in vitro and in vivo antioxidant activities. _Food Chem._ 2020, 343, 128392. [CrossRef] [PubMed]

46. Ruttkies, C.; Schymanski, E.L.; Wolf, S.; Hollender, J.; Neumann, S. MetFrag relaunched: Incorporating strategies beyond in silico fragmentation. _J. Cheminform._ 2016, 8, 3. [CrossRef]

47. Blasco, H.; Błaszczyński, J.; Billaut, J.; Nadal-Desbarats, L.; Pradat, P.; Devos, D.; Moreau, C.; Andres, C.; Emond, P.; Corcia, P.; et al. Comparative analysis of targeted metabolomics: Dominance-based rough set approach versus orthogonal partial least square-discriminant analysis. _J. Biomed. Inform._ 2015, 53, 291–299. [CrossRef] [PubMed]

48. Löffstedt, T.; Trygg, J. OnPfLS—a novel multiblock method for the modelling of predictive and orthogonal variation. _J. Chemom._ 2011, 25, 441–455. [CrossRef]