Data Article

Mass spectrometry data of metabolomics analysis of Nepenthes pitchers

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ABSTRACT

Hybridisation plays a significant role in the evolution and diversification of plants. Hybridisation among Nepenthes species is extensive, either naturally or man-made. To investigate the effects of hybridisation on the chemical compositions, we carried out metabolomics study on pitcher tissue of Nepenthes ampullaria, Nepenthes rafflesiana and their hybrid, Nepenthes × hookeriana. Pitcher samples were harvested and extracted in methanol:chloroform:water via sonication-assisted extraction before analysed using LC-TOF-MS. MS data were analysed using XCMS online version 2.2.5. This is the first MS data report towards the profiling, identification and comprehensive comparison of metabolites present in Nepenthes species.

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Specifications

| Subject area       | Biology                          |
|--------------------|----------------------------------|
| More specific subject area | Metabolomics                  |
| Type of data       | Analysed MS data                |

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How data was acquired
High resolution mass spectrometry data were acquired from MicrOTOF-Q III (Bruker Daltonic) using an ESI positive ionisation coupled with UltiMate 3000 UHPLC system (Dionex).

Data format
Raw mzXML files and analysed results in .xlsx format

Experimental factors
Metabolites were extracted from the pitcher of *N. ampullaria, N. rafflesiana* and their hybrid *N. × hookeriana*

Experimental features
Sonication-assisted extracted samples in methanol:chloroform:water (3:1:1) analysed with a LC-TOF-MS and processed using XCMS online version 2.2.5.

Data source location
Bangi, Malaysia (2°55'11.5"N 101°47'01.4"E)

Data accessibility
Supplementary Table 1 and additional files

Value of the data

- LC-MS data allow the profiling of metabolites for the first time in the three species of Nepenthes.
- This enables the identification of highly expressed metabolites or biomarkers through comprehensive comparison between the three species.
- Metabolomics analysis provides further understanding on the metabolite expression in hybrid plant of Nepenthes. This dataset can be combined with previous transcriptomics studies [1–3] to elucidate the biosynthesis pathways of secondary metabolites in Nepenthes species.

1. Data

This dataset comprises the acquired MS raw data (mzXML files), and analysed data in MS Excel (.xlsx) file (Supplementary Table 1) generated from XCMS online analysis of MS data from pitcher extracts of *N. ampullaria, N. rafflesiana* and their hybrid *N. × hookeriana*.

2. Experimental design, materials and methods

2.1. Samples collection

Three lowland *Nepenthes* species, *N. ampullaria, N. rafflesiana* and *N. × hookeriana* were sampled from the experimental terrace of Universiti Kebangsaan Malaysia (2°55'12.7"N, 101°46'59.7"E). Pitcher tissues were harvested 7 days after pitcher opening when the pitchers achieved fully functionality [4]. The pitcher fluids were emptied and rinsed with sterile deionised water before immediately frozen in liquid nitrogen and stored at −80 °C.

2.2. Phytochemical extraction

Each sample was crushed and ground until fine powder before lyophilised for 48 h. Extraction from the dried powder was performed according to [5] with slight modifications. Dried powder samples (10 mg) were extracted with 200 µL of methanol:chloroform:water (3:1:1). Samples were vortexed, sonicated at room temperature for 15 min, vortexed again and then centrifuged at 10,000 g for 10 min. Filtered extracts through a 0.22 µm PTFE membrane were stored at −80 °C.

2.3. Liquid Chromatography-Mass Spectrometry (LC-MS) analysis

The chromatographic separation was performed on Thermo Scientific C18 column (Acclaim™ Polar Advantage II, 3x150 mm, 3 µm particle size) on an UltiMate 3000 UHPLC system (Dionex). The gradient elution was performed at 0.4 mL/min flow rate at 40 °C using (A) water containing 0.1% formic acid, and (B) 100% acetonitrile with 22 min total run time. The gradient started at 5% solvent B
for 3 min (0–3 min), then the gradient increased to 80% solvent B for 7 min (3–10 min) and main-
tained at 80% solvent B for 5 min (10–15 min). Finally, the gradient returned to 5% solvent B in 7 min
(15–22 min).

High resolution MS was carried out using a MicrOTOF-Q III (Bruker Daltonic) using an ESI positive
ionisation with the following settings: capillary voltage at 4500 V, nebuliser pressure at 1.2 bar and
drying gas flow at 8 L/min with the source temperature at 200 °C and m/z range from 50 to 1000 Da.

2.4. Mass spectrometry data handling

The acquired MS raw data were converted to the mzXML file format by Bruker Compass DataAna-
lysisViewer version 4.2. The converted data were processed by XCMS online software package [6] with
Arabidopsis thaliana selected as the bio-source and other default settings to carry out feature detection,
peak alignment, retention time correction, statistical analysis, annotation and identification.

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Transparency document. Supplementary material

Transparency document data associated with this article can be found in the online version at
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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.
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