Optimization of Extraction Method for GC-MS based Metabolomics for Filamentous Fungi

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Abstract

Metabolite extraction is considered as one of the most important for quality controlling steps in metabolomic research. In the present study, optimal extraction protocol for fungal intracellular metabolites was investigated. Two extraction protocols with different solvent systems, methanol extraction and biphasic extraction under different pH conditions were compared. Number of detected peaks, sample-to-sample variation, and throughput of whole process were taken into account as criteria. Extracted metabolites were analyzed by gas chromatography-mass spectrometry (GC-MS) and principal component analysis (PCA). The extraction protocol using methanol as extraction solvent provided the highest number of detected peaks (~ 300 peaks) with the ease and speed, suggesting that the methanol-extraction method was suitable for intracellular metabolites extraction from P. chrysosporium. However, depending on the nature of metabolites of interest, the biphasic extraction method under different pH conditions is suitable for a targeted analysis.

Keywords: Basidiomycetes; Extraction protocol; Optimization; Metabolomics

Introduction

Metabolome is complex because of the highly chemically and physically diverse nature of metabolites and of often occurring the multiplication of metabolic pathways for some particular metabolites. To obtain as much information about the metabolome as possible, using of productive sample preparation, merged with selective extractions associated with different analytical techniques and combined with the efficient mathematical interpretation of the data are crucial to examine [1,2]. Furthermore, the method should be optimized for each type of sample organisms of interest. Thus, the suitable selection for extraction protocols is considered as one of the most important quality controlling steps in microbial metabolite studies.

Mass spectrometry coupled with chromatography has become the preferred technique available for the quantification of metabolome. These analytical techniques possess high sensitivity, selectivity and simultaneous quantification of many different metabolites and require only a small sample volume for analysis [3,4]. In metabolomics, the number and amounts of metabolites extracted must be maximized, while the degradation of metabolites during sample preparation process and the analytical variation should be minimized [5]. Gas chromatography-mass spectrometry (GC-MS) is one of the most commonly used analytical techniques with a high liability and a capability of high-throughput and automated analysis.

Recently, the interest of using filamentous fungi such as Ascomycetes and Basidiomycetes as microbial cell factories to produce high-valued metabolites has been increased. Several researches about extraction strategy for intracellular metabolite of fungi were reported, especially in yeast [6-10], but only little is known about metabolites and their dynamics in filamentous fungi. Hence, the development of reliable method for metabolites determination in the organism has been prompted [11,12].

The white-rot Basidiomycete, Phanerochaete chrysosporium, has served as a model organism for studies of lignin biodegradation [13]. One possible target function is a fungal degradation of lignin which is well known to retard effective bio-energy production from woody materials. Effective and comprehensive extraction of intracellular metabolites from P. chrysosporium is difficult because this fungus has robust cell wall and very thick extracellular glucan layer [14].

In this study, different extraction methods were examined for its suitability for GC-MS based metabolomics in P. chrysosporium. Aiming to develop an extraction protocol for filamentous fungi with maximizing the number and amount of metabolites, whereas minimizing the analytical variation. GC-MS was utilized in the present work as an analytical apparatus because of its advantages for quick identification of low-molecular-weight metabolites based on retention times and fragmentation patterns. Optimized extraction protocol for P. chrysosporium would be applicable for other filamentous fungi.

Materials and Methods

Chemicals

All chemicals were of analytical grade and purchased from Wako Pure Chemicals Industries, Ltd, except N-methyl-N-trifluoroacetamide (MSTFA) from Thermo Fisher Scientific Inc. Deionized water was obtained with a Milli Q system (Millipore).

Microorganism and culture conditions

The culture medium (HCLN) containing 1% glucose, 1.2 mM

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ammonium tartrate, 20 mM dimethylsucinate and 10% (v/v) Kirk salts [15]. P. chrysosporium was grown on PDA slant for 7 days at 37°C, then, 1% conidial suspension was inoculated to HCLN medium (20 mL in a 200-mL Erlenmeyer flask) and grown at 37°C under static conditions. After a 2-day preincubation, mycelium mat was homogenized and transferred into 1 L production medium in a 2 L Erlenmeyer flask and incubated at 37°C, 140 rpm for 4 days under air.

Quenching of cellular metabolism

One volume of culture broth was quickly harvested and filtered through a membrane filtration (0.45 μm) and washed twice with deionized water. Wet mycelium was then mixed with 1 volume of 50% cold methanol (-30°C). After quenching, wet mycelium was centrifuged (5 min, 11900 × g) at -20°C and ground to powder by using a liquid N₂-cooled pestle and mortar prior to freeze dry. Freeze-dried sample was stored at -80°C.

Intracellular metabolites extraction

Dried mycelium was mixed with small amount of sea sand (20-35 mesh), ground to powder by using a liquid N₂-cooled pestle and mortar. A brown-glass centrifuge tube containing 30 mg of powdered mycelium was mixed with internal standards; 50 μL of a ribitol stock solution (0.2 mg/mL) and 50 μL of a nonadecanoic acid methyl ester stock solution (2 mg/mL). Four different extraction protocols were used and experiment was replicated five times in each extraction protocol. The relative concentration of intracellular metabolites were determined and compared.

Methanol extraction

Sample was extracted with 1.5 mL cold methanol (-30°C), vigorously mixed, sonicated and centrifuged (20 min, 100 x g) at room temperature. The supernatant was transferred to a new vial and the precipitated pellet was re-extracted with 1.5 mL of cold methanol. All extracts were pooled, frozen in liquid N₂, concentrated by freeze-dryer, and stored at -80°C.

Biphasic extraction under neutral conditions

The final ratio of methanol-chloroform-water extraction method was 1:1:1:9 [16]. Sample was homogenized twice in 1.5 mL cold methanol (-30°C), 0.75 mL cold chloroform (-30°C) and 0.6 mL cold water (4°C) to reach the first ratio of 1:0.5:0.4. Then, another 0.75 mL cold chloroform was added and mixed to form the second ratio of 1:1:9. Finally, 0.75 mL cold water was added to get the final ratio of 1:1:9, and samples were vortexed and centrifuged (10 min, 1000 x g) at room temperature. The upper phase (methanol and water) was separated from the lower phase (chloroform) using a glass syringe, transferred to a new vial independently. Both fractions were frozen in liquid N₂, freeze-dried, and stored at -80°C until further analysis.

Biphasic extraction under acidic conditions

The extraction protocol was the same as previously described in methanol-chloroform-water extraction under neutral conditions. The alkaline extraction was carried out with 0.1 M KOH. The upper phase compose of methanol and potassium hydroxide aqueous solution (pH ~ 12) was neutralized by the addition of 0.1 M formic acid checking with pH paper and removed precipitated salts by centrifugation. The supernatant was frozen in liquid N₂, freeze-dried, and stored at -80°C until further analysis.

Biphasic extraction under basic conditions

The extraction protocol was the same as previously described in methanol-chloroform-water extraction under neutral conditions. The alkaline extraction was carried out with 0.1 M KOH. The upper phase (methanol and potassium hydroxide aqueous solution (pH ~ 12) was neutralized by the addition of 0.1 M formic acid checking with pH paper and removed precipitated salts by centrifugation. The supernatant was frozen in liquid N₂, freeze-dried, and stored at -80°C until further analysis.

GC-MS analysis

O-Methylhydroxylamine Hydrochloride (100 μL of 30 mg/mL in dry-pyridine) was added to dried metabolites, incubated at 30°C, 200 rpm for 120 minutes to increase the degree of methoximation, and then dried under N₂ [17]. Then 100 μL M STFA and 50 μl pyridine were added, incubated at 37°C for 90 minutes [18]. The derivatized samples were analyzed by GC-MS coupled with quadrapole detector (GCMS-QP2010 plus, Shimadzu, Japan). A fused silica column of Rtx-5MS (30 m x 0.25 mm i.d., 0.25 μm film thickness, RESTEK) was used and temperature was programmed from 80 to 320°C at 8°C/min, and hold for 5 minutes. The injection temperature was 280°C. Mass spectra were obtained with electron impact ionization at 70 eV with the split ratio of 25:1.

Data processing and statistical analysis

The raw data files obtained from GC-MS runs were converted to vendor-independent NetCDF (network Common Data Format format. MZmine software version 0.60 (http://mzmine.sourceforge.net/) was applied for data processing. MZmine processing consists of several stages, which are input file manipulation, spectral filtering, peak detection, chromatographic alignment, normalization, visualization and data export, each of which is controlled by several parameters. The preprocessed GC-MS data was then imported into SIMCA-P 11.5 software (Umetrics Inc., Umeå, Sweden). Principal component analysis (PCA) was applied to identify the degree of similarity between the data.

Results and Discussion

Effect of biomass to methanol ratio

Our previous data indicated that fungal biomass in the culture system causes minimal deviations during the early stage [19]. Effect of organic solvents on the amount of intracellular metabolites extracted from mycelium of P. chrysosporium was investigated. Among solvents tested, methanol was found to be the best based on the highest amount of extracts. Methanol has been reported to be a good organic solvent for extraction with many advantages; easy to remove after extraction, good recovery of metabolites, good reproducibility, and broad range of metabolites extractable [2]. To determine the best protocol, four biomass-to-methanol ratios; 2.5, 5, 10 and 20 mg/mL were examined. As the ratios of biomass to methanol increased, the number of detected peaks was increased until it reached the maximum at the ratio of 10 mg/mL (Table 1). Since one important goal in this study was to develop the extraction method which enables to extract the metabolites as many as possible. Therefore, the ratio of biomass to methanol at 10 mg/mL was selected for further studies. The balance between extraction efficiency and occurrence of metabolite fragmentation should be considered for optimization of extraction protocols; however, we put
priority to a number of metabolite-derived peaks at this stage, since a later analysis can point out the true metabolic data and fragment data but any analytical methods cannot discuss on a metabolite with no peak appearance.

**Evaluation of extractable metabolites**

The optimized extraction protocol should be developed for each microorganism of interest. Based on the concept that polar solvents extract polar metabolites and non-polar solvents extract non-polar metabolites; likewise, acid stable metabolites should be extracted with acids whereas alkali stable metabolites should be extracted with alkali [3]. Thus, the effect of solvents and pH conditions on metabolite extraction was investigated using *P. chrysosporium*.

Based on the number of detected peaks, polar phases contained more peaks than non-polar phases (Table 2). Biphasic extraction solvent system composed of methanol with pH-controlled water to extract polar metabolites and chloroform for non-polar metabolites showing the possibility to separate polar from non-polar metabolites. These two phases can be extracted simultaneously and each fraction can be analyzed separately, which may cause better separation because of peak numbers and of column selection. For this reason, methanol/chloroform/water was previously reported as the preferred extraction method for lipid-rich tissues such as liver and brain [20-22]. Phospholipids, a class of lipids and a major component of all cell membranes, can also cause ion suppression in electrospray ionization apparatus, which further prevents ionization and detection of other metabolites [21]. Hence, minimizing lipids contamination is necessary to prevent ion suppression and subsequent poor detection of other metabolites [22]. Moreover, chloroform presented in biphasic extraction solvent mixtures is able to denature of enzymes avoiding further reactions [2]. Therefore, considering the possibility to remove polar out of non-polar metabolites, which is mainly referred to lipid compounds, the biphasic extraction under acidic conditions was preferable.

**Evaluation of different extraction protocols using PCA**

**Polar phase**: Principal components analysis (PCA) was used to compare the sample-to-sample variation of metabolic fingerprints derived from 5 replicates in each different extraction protocols. Representative GC-MS spectra showing the metabolic fingerprints of mycelium extracts in polar phases including methanol presented in (Figure 1). From PCA scores plot (Figure 1A), tetracosanoic acid are metabolites that contribute to the separation of methanol extraction data from biphasic clusters. As shown by the PCA loadings plot (Figure 1B), tetracosanoic acid, hexadecanoic acid and pentadecanoic acid are metabolites that contribute to the separation of methanol extraction data from biphasic clusters.

Furthermore, each peak in the chromatogram was putatively identified by using the data libraries and Venn diagram plot was applied to distinguish the unique metabolites presented in each extraction method. The result revealed that the sample via methanol extraction protocol contained a series of free fatty acids (ca. 30% of total unique metabolites) which were not found in biphasic extraction methods (Figure 2).

**Non-polar phase**: Three sets of data from non-polar phase derived from biphasic extraction methods were further investigated...
for sample-to-sample variation of each condition by PCA. According to PCA scores plot (Figure 3A), three different pH conditions were distinct from each other, although some data points from neutral and acidic conditions were overlapped. Not like Fig. 2, PC2 was not the axis explaining the sample-to-sample variation. On the other hand, the PCA loading plot (Figure 3B) indicated that oleic acid, propanoic acid, ergosterol and 9, 12-octadecadienoic acid (Z, Z) were compounds conducing to the difference among three different extraction pHs.

As shown in Table 3, methanol solvent system presented the highest number of detected peaks with a small sample-to-sample variation, representing the high reproducibility of the method. Furthermore, the advantages of this method were simple and fast. Biphasic extraction method under acidic and basic conditions also gave the high number of detected peaks in both polar and non-polar phases. The biphasic extraction showed the possibility to separate polar from non-polar metabolites, but the weak point of this method was the time consuming.

However, for metabolic fingerprinting studies, an acid and an alkaline extraction may contain advantages depending on the metabolite(s) of interest, but a number of compounds are unstable at high and low pHs (dihydroxyacetone phosphate (DHAP) and phospho(enol)pyruvate (PEP) are well-know examples) and are degraded (NAD+ in acid solution and NADH in alkaline solution) as previously described on the study of glycolytic intermediates levels in Aspergillus niger mycelium [9,12,23]. It was found that the extraction method using chloroform at neutral conditions appeared to be superior to both extractions with acidic and basic conditions, since larger amounts of glycolytic intermediates were extracted. They assumed that it was probably due to instability of these metabolites at high and low pHs [11]. Moreover, acid extraction, which is widely used in protein precipitation and hydrophilic metabolites extraction, is not only caused the damage of metabolite structures, but also make problematics for many analytical methods [20].

It would be noteworthy to mention that the fragmentation of several metabolites could be pointed out from the PCA loading plot seen as several points derived from one metabolite (Figures 1 and 2). However, this type of multiplication was quite similar among 4 protocols in the present study (data not shown), suggesting that the extent of fragmentation during extraction processes were similar. Furthermore, the thermal fragmentation may also occur at the injection port of GC.

### Conclusion

Based on number of detected peaks, sample-to-sample variation, the ease and speed of the experimental efficiency, methanol extraction method is preferable and selected as the method for the extraction of
intracellular metabolites from *P. chrysosporium*. However, depending on the nature of metabolites of interest, the biphasic extraction method under different pH conditions is an optional. Furthermore, pre-separation based on characteristics of metabolites may provide a better strategy for analytical optimization such as column selection for chromatography and ionization selection for MS analysis.

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