Evaluation of genes involved in oxidative phosphorylation in yeast by developing a simple and rapid method to measure mitochondrial ATP synthetic activity

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Abstract

Background: Measurement of mitochondrial ATP synthesis is a critical way to compare cellular energetic performance. However, fractionation of mitochondria requires large amounts of cells, lengthy purification procedures, and an extreme caution to avoid damaging intact mitochondria, making it the highest barrier to high-throughput studies of mitochondrial function. To evaluate 45 genes involved in oxidative phosphorylation in Saccharomyces cerevisiae, we aimed to develop a simple and rapid method to measure mitochondrial ATP synthesis.

Results: To obtain functional mitochondria, S. cerevisiae cells were lysed with zymolyase followed by two-step, low- then high-speed centrifugation. Using a firefly luciferin-luciferase assay, the ATP synthetic activity of the mitochondria was determined. Decreasing the ATP synthesis in the presence of mitochondrial inhibitors confirmed functionality of the isolated crude mitochondria. Deletion of genes encoding mitochondrial ATP synthesis-related protein showed their dependency on the oxidative phosphorylation in S. cerevisiae.

Conclusions: Compared with conventional procedures, this measurement method for S. cerevisiae Mitochondrial ATP Synthetic activity in High-throughput (MASH method) is simple and requires a small amount of cells, making it suitable for high-throughput analyses. To our knowledge, this is the first report on a rapid purification process for yeast mitochondria suitable for high-throughput screening.

Keywords: Mitochondria purification, ATP synthesis, High-throughput analysis

Background

Mitochondria are central organelles controlling the life and death of the cell. They participate in key metabolic reactions, synthesize the majority of ATP in a cell, and regulate a number of signaling cascades, including apoptosis [1]. ATP synthesis is vital for various biological reactions. Many studies have measured ATP content or qualitative changes in total cellular ATP production, but few have quantified ATP production from oxidative phosphorylation of isolated mitochondria [2].

Owing to the ease of genetic manipulation and its importance for bio-industry, the budding yeast Saccharomyces cerevisiae is an ideal organism for the study of many basic cellular mechanisms in eukaryotic cells. Their organelles can be rapidly enriched in sufficient quantities for the analysis of specific functions such as metabolite or protein transport. Therefore, S. cerevisiae is a valuable model cell for studying the molecular and cellular mechanisms underlying the essential biological functions of mitochondria. However, mitochondrial proteins have many subunits, the functions of which are still largely unknown because a method for easy mutational analysis and sensitive assay development is still lacking [3]. One of the biggest problems is that the fractionation of mitochondria requires large amounts of cells, long
procedures, and an extreme caution to avoid damaging intact mitochondria [4,5]. In general, to obtain intact mitochondria, the contents of yeast cells are made accessible by a combination of enzymatic digestion of the cell wall and physical disruption of the resulting spheroplasts [6]. To separate the cellular contents by their variable densities, differential centrifugation, which allows for the separation of the constituents of cells based on their different sedimentation properties, is employed to isolate an enriched mitochondrial fraction and is the most common strategy used to obtain crude mitochondria [7]. Crude preparations of mitochondria are contaminated by other organelles such as lysosomes, peroxisomes, tubular Golgi membranes, and, to some extent, small amounts of endoplasmic reticulum. To achieve mitochondria with higher purity, additional time- and labor-consuming purification steps using sucrose density centrifugation are needed. Recently, Frezza et al. described a step-by-step method to isolate mitochondria from mouse liver, muscle, and cultured fibroblasts using modified differential centrifugation steps and a modified sugar concentration for the osmolyte in the isolation buffer [4]. Based on this technique and protocol for purification of mitochondria from yeast cells [5], we modified this general mitochondria extraction method to quickly obtain crude, but functional mitochondria from yeast cells. The method we developed includes only several steps of differential centrifugation and no sucrose density gradient is needed, which is more suitable for high-throughput screening than the conventional method (Figure 1). By using this method, we evaluated 45 genes involved in oxidative phosphorylation for mitochondrial ATP synthesis in *S. cerevisiae*.

**Results and discussion**

**Preparation of crude mitochondria by the MASH method**

In the conventional method of mitochondrial purification, yeast cells are subjected to mechanical homogenization or detergent treatment followed by differential centrifugation because the variable density of the organelles will allow separation of the mitochondria from the remaining cellular structures. In the “measurement method for Mitochondrial ATP Synthetic activity in High-throughput” (MASH method) used in the present study, yeast cell walls were lysed with zymolyase (1.2 mg g\(^{-1}\) wet cells) at 37°C for 1 h. Zymolyase is an enzyme prepared from *Arthrobacter luteus* that effectively lyises yeast cell wall. The lysis of the cell wall and the formation of the protoplast were verified.
under a microscope. The crude mitochondria solution was obtained by a two-step, low- (2,500 × g) then high- (20,000 × g) speed centrifugation with 1.2 M and 0.7 M sorbitol, respectively (Figure 1). Although this suspension is enriched in mitochondria, it may also contain other organelles such as the endoplasmic reticulum, Golgi, and vacuoles. To get more pure mitochondria, this crude mitochondrial fraction can be subjected to further fractionation. However, the crude mitochondria solution obtained using the MASH method is sufficient for the analysis of mitochondrial ATP synthetic activity and therefore was used directly in the ATP assay.

ATP synthesis with inhibitors of mitochondrial respiratory chain
To confirm the functionality of the mitochondria, several inhibitors of the respiratory chain were used.

ATP synthesis is inhibited by antimycin A, CCCP, and DCCD [8-11]. Antimycin A, a strong inhibitor of the electron transfer of complex III, functions by binding to the quinone reduction site of the cytochrome bc₁ complex [9]. CCCP is an inhibitor of the proton motive force [10]. DCCD is a specific inhibitor of subunit c of complex V (mitochondria FₐFₐ₁-ATP synthase) [11]. To test the effect of inhibitors on the mitochondria, antimycin A, CCCP, and DCCD were individually added to the reaction mixture. The addition of either antimycin A or CCCP completely abolished ATP production (Figure 2). The addition of DCCD inhibited ATP production by 80%. These results confirmed that this assay could detect ATP synthetic activity of the crude mitochondria solution using the MASH method and be further applied to high-throughput measurement.

High-throughput measurement of mitochondrial ATP synthesis
Mitochondrial inhibitors confirmed the functionality of the crude mitochondria isolated using the MASH method. By using this method, the ATP synthetic activities of mutants from the single-gene deletion library of S. cerevisiae, including 45 ATP synthesis-related mutants were measured (Table 1). The Saccharomyces Genome Deletion Project created a set of isogenic mutant strains with each individual nonessential gene deleted [12]. This mutant collection has facilitated genome-wide studies to identify genes required for resistance to various cellular insults [13,14]. The set of 45 ATP synthesis-related mutants, which are divided into six genes categories including NADH dehydrogenase (Nde1/Nde2; Complex I), Succinate dehydrogenase (Sdh1b, etc.)/Fumarate reductase (Frd1/Osm1; Complex II), Cytochrome c reductase (Cor1, etc.)/Cytochrome bc₁ complex (Cbp4, etc.; Complex III), Cytochrome c oxidase (Cbp4, etc.; Complex IV), FₐFₐ₁-ATP synthase (Atp1, etc.; Complex V), and others including electron transferring-flavoprotein dehydrogenase (Cir2) and ADP/ATP translocator (Aac1/Aac3). The selected mutant strains and the parental strain were inoculated into 5 ml of YPD medium, grown overnight, and examined for their ability to synthesize ATP using MASH method. The results were shown as the relative value (%) of ATP synthetic activity per mg protein, and the mutant strain values were compared with that of the parental strain. We observed that most of gene deletions in this set resulted in partial loss of the ATP synthetic activity (Table 1).

Among the mutant strains examined, deletion of genes directly related to the mitochondria respiratory chain (NADH dehydrogenase, succinate dehydrogenase, electron transferring-flavoprotein dehydrogenase, cytochrome c reductase, cytochrome bc₁ complex, cytochrome c oxidase, and FₐFₐ₁-ATP synthase) showed lower ATP synthetic activity compared with that of the parental strain, indicating that these components of the mitochondria respiratory chain were indispensable for ATP synthetic activity.
In the case of F_F1-ATP synthase (complex V), 12 single gene deletion strains were used to measure their ATP synthetic activity by the MASH method. Deletion of ATP4 had almost no effect on the ATP synthetic activity because ATP4 encodes b subunit which affects on the stability of oligomeric F_F1-ATP synthases, not ATP synthetic activity [15]. As the result, especially in both the ΔATP2 strain and ΔATP20 strain, the ATP synthetic activities were drastically decreased compared to the other mutant strains lacking gene encoding a subunit of the F_F1-ATP synthase. Their ATP synthetic activities were 20% of that of the parental strain. ATP2 and ATP20 encode β subunit and γ subunit of the F_F1-ATP synthase, respectively. The β subunit is the catalytic subunit of F_F1-ATP synthase and the γ subunit constructs a stalk structure connecting the proton-motive force generated in F_o-part and ATP synthesis in F_F1-part of F_F1-ATP synthase [16]. Thus, the deletions of these functionally important genes, ATP2 and ATP20, indicate completely loss of ATP synthetic ability of F_F1-ATP synthase. These reasonable results guarantee the validity of this assay. The remaining 20% activities of ATP synthesis compared to the parental strain in the ΔATP2 strain and ΔATP4 strain indicate the ATP synthesis by mitochondrial adenylate kinase (2ADP→AMP + ATP) encoded by ADK2 [17]. Some deletion mutants (QCR2, QCR7, CYT1, CBP4 and COX7) showed lower ATP synthetic activities rather than 20% of the parental strain. This result indicates that the deletion of these genes enhance the hydrolysis of ATP resulted from adenylate kinase. The ATP hydrolysis would be catalyzed through reversible reaction of ATP synthesis by F_F1-ATP synthase because of the lower H^+-gradient formed between inside and outside of mitochondrial inner membrane.

Aside from the respiratory chain mutants, the ADP/ATP translocator, which is mainly responsible for transferring ADP/ATP in or out of the mitochondria [18], was also tested. Notably, deletion of genes AAC1 and AAC3 encoding the ADP/ATP translocator, resulted in a 50% - 60% loss of ATP synthetic activity compared with
that of the parental strain. Previous studies demonstrated that disruption of AAC1 or AAC3 did not influence the content of the ADP/ATP translocator, and both AAC1 and AAC3 genes did not substantially participate in mitochondrial ADP/ATP transport under normal growth conditions [19]. On the contrary, our result demonstrated that AAC1 and AAC3 caused some loss of ATP synthetic activity, indicating that although these genes may not be directly involved in mitochondrial ADP/ATP transport, they are potentially responsible for mitochondrial ATP synthesis. Further elucidation of the underlying mechanism is needed.

When deleted, three genes (QCR2, QCR7, and Cyt1) encoding subunits of cytochrome c reductase (complex III) have been shown to have undetectable ATP synthetic activity (Table 1). Cytochrome c reductase is essential to the energy-generating process of oxidative phosphorylation [20]. Qcr2 is one of the core subunits of complex III, and its mutation has been demonstrated to cause either a severe decrease or a total block in complex III activity and respiratory growth [21]. Cyt1 (cytochrome c1) is one of the catalytic subunits of the cytochrome bc1 complex and is essential for electron transfer and for the respiratory growth [22]. Therefore, the deletion of Qcr2 and Cyt1 led to dramatically decreased ATP synthetic activity. Together with the result that the addition of antimycin A completely suppressed ATP synthetic activity (Figure 1), this result further indicated that cytochrome c reductase played an important role in mitochondrial ATP synthesis.

To further test the feasibility of the MASH method for measuring ATP synthetic activity, we compared the distribution of protein concentration and ATP synthetic activity in crude mitochondrial solutions from 45 single-gene deletion strains related to mitochondrial ATP synthesis. The relative protein concentration (%) versus relative ATP synthetic activity (%) of each strain compared to the parental strain was plotted (Figure 3). This distribution map indicated that the relative protein concentration and the ATP synthetic activity were not correlated. All tested strains were distributed in the 40-50% range for relative protein concentration except the parental strain. Previous studies demonstrated that disruption of QCR2, QCR7, and Cyt1 showed almost 55% of the relative ATP synthetic activity compared to the parental strain. Therefore, the deletion of QCR2 and Cyt1 led to dramatically decreased ATP synthetic activity. Together with the result that the addition of antimycin A completely suppressed ATP synthetic activity (Figure 1), this result further indicated that cytochrome c reductase played an important role in mitochondrial ATP synthesis.

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This result demonstrates the wide applicability of the MASH method. As shown in this study, the MASH method can propose new areas of study to resolve the cellular ATP synthesis mechanism.

**Potential applications of the MASH method**

The MASH method is a simple and rapid way to obtain a crude mitochondria solution and determine respiratory ATP synthesis in yeast cells. Mitochondria have been isolated from yeast using the combination of zymolyase and Dounce homogenization for many years at least since 1982 from the Schatz’s laboratory [7]. Thus, in the conventional methods, to prepare intact mitochondria from yeast, cells are collected, then disrupted by mechanical homogenization or detergent treatment. Next, the suspension is separated using differential centrifugation, and the fraction containing mitochondria is subjected to several steps of differential gradient centrifugation, which takes 4–5 hours. In contrast, the MASH method can be finished within two hours and is free of contamination that affects the ATP activity assay, simplifying the purification procedure. Moreover, because the method needs only a small amount of cells and has no requirement for
retrieving the band containing mitochondrial fraction from the centrifuged gradient, it is suitable for high-throughput (e.g., 96-well format) analysis of mutants and drugs.

It is worth noting that any components whose defects result in loss of mitochondrial ATP synthesis can be measured by the MASH method. By using a combination of specific mitochondrial inhibitors and single gene deletion mutant strains, the point of defect could be determined. Furthermore, if the original or mutated target genes are added back into the knockout cells by transformation, their function can be investigated in more detail. One application of this method would be a functional test for activities of the respiratory chain complexes I, II, III, IV, V, the ADP/ATP translocator, and other ATP synthesis-related proteins in mitochondria.

**Conclusion**

This method is designated as measurement method for *S. cerevisiae* Mitochondrial ATP Synthetic activity in High-throughput (MASH method). To obtain crude, yet functional mitochondria, only three steps are included (Figure 1). ATP production from isolated mitochondria could be determined by a bioluminescence assay. By using this MASH method, systematic analysis of gene deletion mutants related to the mitochondrial ATP synthesis was carried out to identify genes required for ATP synthesis, providing a global view of these genes in maintaining ATP activity. The MASH method described in this study introduces a quick and reproducible methodology for measuring ATP synthetic activity in isolated mitochondria.

**Methods**

**Yeast strains and growth conditions**

The parental strain *S. cerevisiae* BY4741 (MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0) was cultured in 5 ml of YPD medium containing 10 g L⁻¹ yeast extract, 20 g L⁻¹ bacto-peptone, and 20 g L⁻¹ glucose overnight at 30°C. The collection of yeast knockouts was purchased from Invitrogen. Zymolyase-20T was purchased from Seikagaku Kogyo Co. (Tokyo, Japan). Protease inhibitor cocktail and D-luciferin was purchased from Roche (Basel, Switzerland). Firefly luciferase was obtained from Promega (Madison, WI, USA). Other chemicals were purchased from Nacalai Tesque (Kyoto, Japan) or Wako Chemicals (Osaka, Japan).

**Measurement of ATP synthetic activity**

All solutions were stored at 4°C and freshly prepared prior to use. Cultures of *S. cerevisiae* cells were grown aerobically in 5 ml of YPD medium at 30°C for 24 h. The optical density at 600 nm (OD₆₀₀) was measured using a UVmini-1240 spectrophotometer (Shimadzu, Kyoto, Japan). The cells were collected, washed with 0.5 ml of 10 mM EDTA, and centrifuged (400 x g, 5 min, 4°C). The supernatant was discarded, and the pellets were resuspended in 50 mM Tris–HCl (pH 7.5), 1.2 M sorbitol, 10 mM EDTA, 0.3% (v/v) 2-mercaptoethanol, and 1.2 mg g⁻¹ wet cells of the zymolyase solution (4 mg ml⁻¹). After incubation at 37°C for 1 h with rotary agitation, the lysis of the cell wall was verified under a microscope. The supernatant was resuspended in 50 mM Tris–HCl (pH 7.5), 0.7 M sorbitol, 10 mM EDTA, 1 mM PMSE, protease inhibitor cocktail, and 20 mM triethanolamine, and then subjected to the low-speed centrifugation step (2,500 x g, 15 min, 4°C). The debris was discarded and the supernatant was then subjected to the high-speed centrifugation step (20,000 x g, 15 min, 4°C). The resulting pellets obtained were dissolved in 50 mM Tris–HCl buffer (pH 7.5), and stored at 4°C before use.

The ATP assay was conducted as previously described [23,24]. The reaction buffer containing 50 mM Tris–HCl (pH 7.5), 1.3 µg ml⁻¹ luciferase, 0.05 mM D-luciferin, 1 mM DTT, 5 mM MgCl₂, and 0.1 mM EDTA was added to the crude mitochondria solution. The reaction was initiated by addition of 0.1 mM ADP, and the luminescence was measured using EnVision Multilabel Reader 2104 (PerkinElmer, Waltham, MA, USA). The luminescence of each well was measured at 1 s intervals. The ATP synthetic activities of the crude mitochondria solutions were calculated by taking away the background luciferase activity in the presence of ADP. The relative ATP synthetic activity was normalized each protein concentration determined using the Bradford method [25] of the mutant strains. The values are expressed as a percentage of its activity of the parental strain. To test the functionality of mitochondria, the inhibitors (0.1 mM antimycin A, 0.5 mM CCCP, and 0.1 mM DCCD) were incubated with the crude mitochondria solution for 5 min before being subjected to the ATP assay.

**High-throughput measurement for mitochondrial ATP synthesis**

The glycerol stock of yeast gene-deletion mutants was inoculated with a sterilized toothpick to 5 ml of YPD medium. Cells were cultured overnight at 30°C with shaking. The cells were harvested by centrifugation (3,000 x g, 10 min, 4°C) and washed twice with distilled water. The purification of mitochondria was conducted as described above. Protein concentration was measured with the Bio-Rad assay system (Bio-Rad, Hercules, CA, USA) using bovine serum albumin as the standard. Relative specific activity (%) was calculated from the ratio of total activity divided by total protein concentration of the mutant strain to that of the parental strain. Z' of this assay can be calculated as 0.58.
Adenosine 5’-diphosphate; CTP: Cytidine triphosphate; ADP: Adenosine 5’-diphosphate; NAD: Nicotinamide adenine dinucleotide; ATP: Adenosine 5’-triphosphate; G3P: Glycerol 3-phosphate; CCCP: Carbonylcyanide; DCCD: N,N’-dicyclohexylcarbodiimide; pmf: Proton motive force; N,N’-dicyclohexylcarbodiimide; pmf: Proton motive force; NADH: Nicotinamide adenine dinucleotide hydrogen; ADP: Adenosine 5’-diphosphate; EDTA: Ethylenediaminetetraacetic acid; Tris-2-Amino-2-hydroxymethyl-propane-1,3-diol; (hydroxymethyl) aminonitrile; PMF: Phenylmethylsulfonyl fluoride; DTT: Dithiothreitol.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
XY designed the study, performed the experiments, and wrote the manuscript. KM performed additional experiments to revise the manuscript. SHH assisted in performing the experiments. MA and KN were involved in the discussions of the research results and contributed to finalize the manuscript. TH supervised the research. KYH designed the study, edited the manuscript, and coordinated the project. AK coordinated the project. All the authors read and approved the manuscript.

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