Enhancement of Ethanol Fermentation in \textit{Saccharomyces cerevisiae}
Sake Yeast by Disrupting Mitophagy Function

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\textit{Saccharomyces cerevisiae} sake yeast strain Kyokai no. 7 has one of the highest fermentation rates among brewery yeasts used worldwide; therefore, it is assumed that it is not possible to enhance its fermentation rate. However, in this study, we found that fermentation by sake yeast can be enhanced by inhibiting mitophagy. We observed mitophagy in wild-type sake yeast during the brewing of Ginjo sake, but not when the mitophagy gene (ATG32) was disrupted. During sake brewing, the maximum rate of CO\textsubscript{2} production and final ethanol concentration generated by the \textit{atg32}Δ laboratory yeast mutant were 7.50% and 2.12% higher than those of the parent strain, respectively. This mutant exhibited an improved fermentation profile when cultured under limiting nutrient concentrations such as those used during Ginjo sake brewing as well as in minimal synthetic medium. The mutant produced ethanol at a concentration that was 2.76% higher than the parent strain, which has significant implications for industrial bioethanol production. The ethanol yield of the \textit{atg32}Δ mutant was increased, and its biomass yield was decreased relative to the parent sake yeast strain, indicating that the \textit{atg32}Δ mutant has acquired a high fermentation capability at the cost of decreasing biomass. Because natural biomass resources often lack sufficient nutrient levels for optimal fermentation, mitophagy may serve as an important target for improving the fermentative capacity of brewery yeasts.

\textit{S}ake is a traditional Japanese alcoholic beverage produced from steamed rice and koji. During the manufacturing process, glucose is produced (saccharification) from the starch present in rice by the actions of enzymes produced by the koji fungus \textit{Aspergillus oryzae}. Glucose is fermented to ethanol by \textit{Saccharomyces cerevisiae} sake yeast strains (1). Sake contains the highest ethanol concentration of all the brewed alcoholic beverages worldwide. This high ethanol concentration is generated by technologies that include successive addition of enzymes and nutrients derived from koji during sake brewing (2, 3), a 3-step pitching process, brewing in winter, and the historical selection of high-ethanol-producing sake yeast strains (1). Sake yeast strains have been selected through a long history of cultivation, ranging from 100 to 400 years. The most frequently used sake yeast at present is Kyokai no. 7 (K7), which was isolated from sake mash in 1946 (4, 5). This strain produces a high concentration of ethanol, because it lacks functions of proteins encoded by \textit{MSN4}, \textit{PPT1}, and \textit{RIM15}, which are required to mount a stress response (6–8). For this reason, researchers in this field believe that it is difficult to further augment the fermentation rate of this sake yeast.

Although rice is used as a raw material to brew sake, the surface of rice contains many constituents such as amino acids that impart a heavy and complex taste to sake. Because Japanese consumers tend to prefer a light and clear taste, rice with a polished surface is used for sake brewing. Sake is categorized into two representative types depending on the extent of rice polishing; these types have been specified by the official guidelines of the Japanese government (http://www.nta.go.jp/shiraberu/sennonjoho/sake/hyoji/seisshu/gaiyo/02.htm). When the weight of the removed surface is less than 30% or more than 40% of the total weight of rice, sake is categorized as either normal sake or premium Ginjo sake, respectively. Because sake yeast strains are cultured in the presence of low nutrient concentrations during Ginjo sake brewing (30% lower amino acid concentration than normal sake brewing) (9), sake yeast produces flavors imparted by ethyl caproate and isoamyl acetate (10).

Autophagy is a bulk degradative and recycling process involving the transport of cytoplasmic components and organelles to the vacuole (plant and fungal cells) or lysosome (mammalian cells); it is required for homeostasis and is induced under conditions of nutrient starvation (11). Mitophagy is a selective form of autophagy that specifically degrades mitochondria (12, 13) and plays critical roles in the pathogenesis of Parkinson’s disease. Mitophagy is mediated by the activity of the serine/threonine protein kinase PTEN-induced putative kinase 1 (PINK1) and the ubiquitin ligase Parkin (PARK2) (14, 15). PINK1 phosphorylates mitofusin 2 (MFN2), which functions as a Parkin receptor for culling damaged mitochondria in response to mitochondrial depolarization (16). Although yeasts undergo autophagy during wine fermentation (17–19), selective modes of autophagy, such as mitophagy, have not been reported. Moreover, there are no published studies on the relationship between mitophagy and the fermentation characteristics of yeast.

Our laboratory focuses on the effects of mitochondrial activities and the metabolic engineering of sake yeast strains that influence their ability to ferment substrates (1, 20–26). Mitochondria depolarize during anaerobiosis, which corresponds to the conditions used for industrial alcoholic fermentation (27–29), and mitophagy occurs when mitochondrial electron potential decreases...
with the flanking regions of the strains were disrupted using a PCR-based method. To disrupt the first
ATG32
 promoter, primers atg32nat1fw (5’-TGAATTCCTAATACAAAAAGAAGAAAAAAAAATCTGCCAGGAAC GTCACATACCATACGTTTAGGACAC-3’) and atg32nat1rv (5’- TAGTAAAAGTGAGTAAAGCCTCTGTTTGTATGTATGGGAAAAGGATAGCTACTCATTAGGAG-3’) were amplified as a template.

(30). Moreover, mitophagy is induced in yeast cells when the availability of nutrients is limited (11). Indeed, sake yeasts are cultured under such conditions during sake brewing, because the nutrient-rich surface of rice substrate is polished and removed. Therefore, we hypothesized that mitophagy plays a role in the fermentation characteristics of sake yeast.

In the present study, we demonstrate that mitophagy occurs during sake brewing and that the fermentative capacity is improved by inhibiting mitophagy. This novel approach will be valuable for improving the fermentative capacity of other brewery yeasts.

MATERIALS AND METHODS

Yeast strains. The S. cerevisiae strains used in this study are listed in Table 1. BY4743 (MATa α his3Δ1α/Δ1 leu2Δ1Δ/Δ1 met15Δ1Δ/Δ1 MET15 lys2Δ1/SYS2/lys2Δ1Δ/ura3Δ1/Δ1 and the atg8A, atg11Δ, and atg32Δ mutants (BY4743 background) were purchased from Life Technologies. The sake yeast strain K7 was purchased from the Brewing Society of Japan. The K7 (BY4743 background) were purchased from Life Technologies. The sake yeast strain RAK1536 was obtained from Life Technologies.

To propagate yeast cells, yeast extract-peptone-dextrose (YPD) medium containing a 0.67% (wt/vol) yeast nitrogen base without amino acids (Beckton Dickinson), 800 mgliter (YPD) medium containing 2% (wt/vol) Bacto peptone, 1% (wt/vol) Bacto yeast extract, 2% (wt/vol) glucose was used. To propagate cells harboring mitochondrion-targeted green fluorescent protein (GFP), minimal synthetic medium containing a 0.67% (wt/vol) yeast nitrogen base without amino acids (Beckton Dickinson), 800 mg liter⁻¹ complete supplement mixture Drop-out-HIS + 40 ADE (Formedium), and 2% (wt/vol) glucose was used. For fermentation tests, minimal synthetic medium containing a 0.67% (wt/vol) yeast nitrogen base without amino acids (Beckton Dickinson), 790 mg liter⁻¹ complete supplement mixture Drop-out: Complete (Formedium), and 15% (wt/vol) glucose was used.

Construction of yeast mutants. The two copies of ATG32 in the K7 strains were disrupted using a PCR-based method. To disrupt the first copy of ATG32, a DNA fragment containing kanMX was amplified using the forward primer atg32kanMXf (5’-AGATCACTGTCCTCCTC GAC-3’) and the reverse primer atg32kanMXr (5’-TGCACTACACATT TACAGCGA-3’) and atg32Δ DNA (BY4743 background) as a template.
the manufacturer’s protocol (Thermo Scientific). The extracted protein solution was concentrated by extraction using chloroform-methanol-water (4:1:3), and the precipitated protein was solubilized in sterile water. The protein concentrations of the disrupted cells were determined using the Bradford method. Equal amounts of protein samples (20 μg) were applied to a 12.5% SDS-polyacrylamide gel and electrophoresed at 40 mA for 50 min. Proteins were transferred to a polyvinylidene difluoride (PVDF) membrane (100 mA, 2 h). The membrane was blocked with skim milk (0.3% [wt/vol] in Tris-buffered saline-Tween 20 [TBST]). Incubation with the first antibody was performed overnight using an antibody against GFP (diluted 1:1,000) (mFX73; Wako Chemicals), and immune complexes were detected using an alkaline phosphatase-conjugated goat anti-mouse IgG (diluted 1:30,000) (Sigma-Aldrich). The membrane was visualized using the BCIP/NBT (5-bromo-4-chloro-3-indolyl-phosphate with nitro blue tetrazolium) liquid substrate system (Sigma-Aldrich).

Measurement of dry cell weight. After fermentation, cells were collected by centrifugation, washed twice with sterile water, suspended in 1 ml sterile water, and added to 250-ml aluminum bottles. The bottles were weighed before adding the cells. The bottles were heated in an oven at 180°C overnight and then weighed. The differences between the weight of bottles before and after addition of cells have been presented as the dry cell weight. The dry cell weight was expressed as biomass.

Measurement of signal intensity of mitochondria. Cells were photographed under a fluorescence microscope (Keyence BZ8000) at a magni-
FIG 1 continued

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C

D

E

Minimal synthetic medium

Ginjo-sake mash

FIG 1 continued
Signal intensities of GFP images were determined using Dynamic Cell Count software (Keyence).

Measurement of cell area. Cells were photographed under a microscope (Olympus BX53) at a magnification of 1,000. The photos were processed using Image J software (National Institutes of Health). One hundred cells were outlined, and areas of the outlined cells were measured using the software.

Quantitative reverse transcriptase real-time PCR. Yeast cells were collected from minimal synthetic medium or Ginjo sake mash by centrifugation. Total RNA was extracted from the cells using a hot phenol method (36). Total RNA was purified using the RNeasy minikit (Qiagen). Real-time PCR was performed using primers 5′-TGGTGTTCAATGCTT TCCAAG-3′ and 5′-CAAGGGTATCACCTTCAAAC-3′, the TaKaRa PrimeScript RT Master mix (Perfect Real Time) (TaKaRa Bio, Inc.), and the Light Cycler 480 system (Roche Diagnostics).

Statistical analysis. The statistical significance of differences between the averages of two data groups with fewer than 30 samples was judged using an unpaired one-tailed Student’s t test without known deviations.
The statistical significance of differences between the averages of two data groups with more than 30 samples was judged using the two-sample \( z \)-test (37).

RESULTS

Mitophagy occurs dependent on Atg32 in sake yeast during sake brewing. To determine whether mitophagy occurs in yeast cells during sake brewing, we generated a sake yeast strain disrupted in \( ATG32 \) and sake yeast strains that express GFP targeted to the mitochondria. \( ATG32 \) encodes a mitochondrial outer membrane protein, which recruits the autophagy adaptor protein Atg11p and the ubiquitin-like protein Atg8p to the mitochondrial surface to initiate mitophagy (11). A study reported that a mutant defective in \( Atg32 \) does not cause mitophagy (11). Sake was brewed using these strains, and mitochondrial and vacuolar structures were observed using a fluorescence microscope. Highly fragmented mitochondria and large swollen vacuoles were observed during the later phase of sake brewing, as reported previously (22, 38). In wild-type sake yeast cells analyzed during the brewing of normal sake, some portion of the mitochondria (green) fused with the vacuolar membrane (red) to generate yellow signals in z-stack images (Fig. 1A, yellow arrow). In contrast, in the \( atg32 \)/H9004 sake yeast, mitochondria (green) were located distant from the vacuolar membrane (red), and virtually no yellow signal was detected that would indicate fusion of mitochondria with the vacuolar membrane (Fig. 1B). We hypothesized that mitophagy would be more evident in cultures with limited nutrient concentration. Mitophagy was clearly observed in sake yeast during the brewing of Ginjo sake (a refined sake), likely because the medium contained low nutrient concentrations. In wild-type sake yeast, a significant portion of mitochondria (green) fused with the vacuolar membrane (red) to generate yellow signals (Fig. 1C), but this was not observed in \( atg32 \)/H9004 sake yeast (Fig. 1D). The \( atg32 \) cells exhibited weaker mitochondrial signals than wild-type sake yeast during the brewing of Ginjo sake (for the wild type, average signal intensity = 75, standard deviation [SD] = 20, \( n = 172 \); for the \( atg32 \) strain, average signal intensity = 41, SD = 8, \( n = 201 \), two-sample \( z \)-test).
In contrast, there was no significant difference between the atg32Δ strain and its parent strain with respect to the amount of GFP (Fig. 1E) or in their levels of expression of GFP mRNA (data not shown) when cells were cultured under the conditions of Ginjo sake brewing. These results indicate that mitochondrial GFP, although expressed normally in the atg32Δ strain, cannot translocate to the mitochondria in the atg32Δ strain because of low mitochondrial electron potential, which is consistent with the report that this mutant accumulates dysfunctional mitochondria (39). Together, these results indicate that mitophagy occurs dependent on Atg32 and this mitophagy is required to maintain mitochondrial quantity in sake yeast during sake brewing.

**Enhanced ethanol fermentation by the atg32Δ laboratory strain.** Because the above results indicated clearly that mitophagy occurred in sake yeast during sake brewing, we hypothesized that mitophagy may affect the characteristics of brewed sake. Therefore, sake was first brewed with the atg32Δ and its parent laboratory yeast strains. Analysis of the ability of the atg32Δ laboratory strain to produce CO2 indicated that it has an improved fermentation (P < 0.05; Fig. 2A). This result indicated that mitophagy plays a role in ethanol fermentation. We then asked whether inhibiting general autophagy would have the same effect. To address this question, we brewed sake with mutants defective in genes required for general autophagy (atg11Δ and atg8Δ) laboratory yeast strains). However, in contrast to what was seen for the atg32Δ mutant, fermentation by these mutants was significantly reduced (P < 0.05; Fig. 2B), which is consistent with the results of previous studies of wine fermentation (40). These results indicate that disrupting mitophagy, but not general autophagy, enhances fermentation. To determine the basis for this effect, we assessed the fermentation profiles of atg32Δ mutants and found that the CO2 production rate (Fig. 2C), the final ethanol concentration (Fig. 2E), and CFU (Fig. 2F) were significantly higher than those of the parental strain (P < 0.05). The maximum CO2 production rate of the parent laboratory strain was 10.3 ± 0.167 g liter⁻¹ day⁻¹, and that of the atg32Δ mutant was 11.0 ± 0.146 g liter⁻¹ day⁻¹, which represents an increase of 7.50% relative to that of the parent strain (P < 0.05). These results indicate that the atg32Δ mutant has an enhanced ethanol fermentation rate. The normalized CO2 production rate of the atg32Δ strain per cell was significantly lower than that of the parent strain (P < 0.05) (Fig. 2D), suggesting that the metabolic competence of the atg32Δ mutant per cell did not mediate its high fermentation rate.
Enhanced ethanol fermentation by the \textit{atg32} sake yeast mutant when cultured under Ginjo sake (a refined sake) brewing conditions and also in minimal synthetic medium. In order to elucidate if \textit{atg32} sake yeast also shows improved fermentation profile, the fermentation profile of \textit{atg32} sake yeast during the brewing of normal sake was investigated. In contrast to the \textit{atg32} laboratory strain, the \textit{atg32} sake yeast strain (K7H868, a haploid strain) showed a significantly ($P < 0.05$) lower ability to produce CO$_2$ (Fig. 3A), CO$_2$ production rate (Fig. 3B), final ethanol concentration (Fig. 3D), and CFU (Fig. 3E) than its parent sake yeast strain. The maximum CO$_2$ production rates by the parent and \textit{atg32} sake yeast strains were 16.8 $\pm$ 0.470 g liter$^{-1}$ day$^{-1}$ and 13.6 $\pm$ 0.293 g liter$^{-1}$ day$^{-1}$, respectively, representing a 19.1% decrease relative to that of the parent strain ($P < 0.05$). Intriguingly, the normalized CO$_2$ production rate per cell of \textit{atg32} sake yeast was higher than that of its parent strain under this condition (Fig. 3C), suggesting the metabolic impact of the \textit{atg32} mutation. Increase of fermentation and final ethanol concentration during the brewing of normal sake were not observed in \textit{atg32} diploid sake yeast strains either (see Fig. S1 in the supplemental material). These data led us to conclude that laboratory and sake yeasts may respond differently to nutrient concentrations, as described previously (41), because laboratory yeasts have long been cultured in media very rich in nutrients such as YPD, and sake yeasts have long been cultured in nutrient-poor media that contain polished rice. Therefore, sake yeast should exhibit the same response as laboratory yeast in the presence of low nutrient concentrations similar to those used for Ginjo sake brewing. Consistent with this hypothesis, under the conditions used to produce Ginjo sake, the \textit{atg32} sake yeast strain showed significantly ($P < 0.05$) increased CO$_2$ production (Fig. 4A), fermentation rate (Fig. 4B), final ethanol concentration (Fig. 4D), and CFU (Fig. 4E) compared to the parent strain. The maximum CO$_2$ production rate of the parent sake strain was 13.2 $\pm$ 0.179 g liter$^{-1}$ day$^{-1}$, and that of the \textit{atg32} mutant was 14.1 $\pm$ 0.126 g liter$^{-1}$ day$^{-1}$, representing an increase of 6.28% relative to that of the parent strain ($P < 0.05$). The \textit{atg32} sake yeast strain showed a normalized CO$_2$ production rate that was not significantly different from that of its

![FIG 5](http://aem.asm.org/)

**FIG 5** Fermentation profiles of \textit{atg32} sake yeast strain and its parent sake yeast during incubation in minimal synthetic medium. Sake yeast wild-type and \textit{atg32} strains were incubated in minimal synthetic medium containing 15% glucose, and their fermentation profiles were analyzed. (A) CO$_2$ evolution during fermentation. The results are expressed as the weight loss of the culture, which represents the weight of CO$_2$. (B) Fermentation rate (g CO$_2$ liter$^{-1}$ day$^{-1}$). (C) Ethanol concentration on day 11 (% vol/vol). Closed black and open symbols or boxes represent the results for sake yeast strains K7RAK and K7RAK \textit{atg32}, respectively. The results are expressed as the means $\pm$ SEM of eight independent fermentation experiments initiated with respective starter cultures ($n = 8$; $^*$, $P < 0.05$; unpaired one-tailed Student’s $t$ test).
parent strain (Fig. 4C). Furthermore, during fermentation in minimal synthetic medium, the atg32Δ sake yeast strain exhibited significantly ($P < 0.05$) increased CO$_2$ production (Fig. 5A), fermentation rate (Fig. 5B), and final ethanol concentration (Fig. 5C) relative to the parent strain. The maximum CO$_2$ production rate of the parent sake was 9.14 ± 0.526 g liter$^{-1}$ day$^{-1}$, while that of atg32Δ strain was 9.83 ± 0.104 g liter$^{-1}$ day$^{-1}$, representing an increase of 7.44% ($P < 0.05$). The final ethanol concentration, CO$_2$ production, cell/biomass, specific CO$_2$ production rate, specific ethanol production rate, and ethanol yield of the atg32Δ strain were significantly higher than those of the parent sake yeast strain ($P < 0.05$), while the final biomass, OD$_{600}$ and biomass yield of the atg32Δ mutant were significantly lower than those of the parent sake yeast strain ($P < 0.05$) (Table 2).

**DISCUSSION**

In the present study, we have shown that mitophagy occurs dependent on Atg32 in sake yeast during the brewing of Ginjo sake. The atg32Δ laboratory yeast strain showed improvement in fermentation characteristics such as the final ethanol concentration and fermentation rate. Fermentation was also enhanced in sake yeast under the conditions used for Ginjo sake as well as in minimal synthetic medium. Our present study demonstrates that the ethanol fermentation rate of sake yeast, the most active sake fermenter among brewery yeasts, can further be enhanced. The final ethanol concentration of minimal synthetic medium incubated with the atg32Δ strain was 2.76% increased relative to its parent sake yeast strain (Fig. 5C). These findings have significant implications for the bioethanol industry. Because natural biomass resources often lack sufficient nutrient levels required for optimal fermentation, this approach will be valuable for the production of bioethanol from natural biomass resources.

Although several mutations that augment the fermentation rate of sake yeast strains relative to those of other yeast strains have been reported, including MSN4, PPT1, and RLM1 (6–8), we are not aware of any report of a mutation that further augments ethanol fermentation. We have shown here that the atg32Δ mutant enhanced the ability of laboratory and sake yeast strains to produce ethanol during sake brewing and fermentation in minimal synthetic medium. A mutation that further enhances ethanol fermentation has not been reported for sake yeast, suggesting that deletion of ATG32 provides a novel approach for improving the fermentative capacity of sake yeast and potentially those of other strains. Moreover, ethanol tolerance did not differ between the atg32Δ mutant and its parent sake yeast strain ($P > 0.05$; see Fig. S2 in the supplemental material), suggesting that this strain would be a practical strain for industrial application. Enhancing fermentation output is the subject of intensive research on brewery yeasts (42, 43). An obvious benefit of these efforts includes reduced production costs, for example, of bioethanol and alcoholic beverages. Therefore, the present study should stimulate efforts to genetically manipulate brewery yeasts to increase ethanol productivity and final ethanol concentration.

The fermentation characteristics of the atg32Δ mutant suggest that the increased fermentative capability of this strain is caused by the increased carbon flux from glucose to ethanol per cell at the cost of decreasing the carbon flux from glucose to biomass. Consistent with this hypothesis, the average cell area of the atg32Δ mutant was 916 ± 20.4 arbitrary units of Image J software ($n = 100$), which was significantly smaller than that of its parent sake yeast strain (Fig. 4C). Furthermore, during fermentation in minimal synthetic medium, the atg32Δ sake yeast strain exhibited significantly ($P < 0.05$) increased CO$_2$ production (Fig. 5A), fermentation rate (Fig. 5B), and final ethanol concentration (Fig. 5C) relative to the parent strain. The maximum CO$_2$ production rate of the parent sake was 9.14 ± 0.526 g liter$^{-1}$ day$^{-1}$, while that of atg32Δ strain was 9.83 ± 0.104 g liter$^{-1}$ day$^{-1}$, representing an increase of 7.44% ($P < 0.05$). The final ethanol concentration, CO$_2$ production, cell/biomass, specific CO$_2$ production rate, specific ethanol production rate, and ethanol yield of the atg32Δ strain were significantly higher than those of the parent sake yeast strain ($P < 0.05$), while the final biomass, OD$_{600}$ and biomass yield of the atg32Δ mutant were significantly lower than those of the parent sake yeast strain ($P < 0.05$) (Table 2).

**TABLE 2** Fermentation characteristics of wild-type and atg32Δ sake yeast strains cultured in minimal synthetic medium

<table>
<thead>
<tr>
<th>Strain</th>
<th>Ethanol (g liter$^{-1}$)</th>
<th>CO$_2$ (mmol g$^{-1}$)</th>
<th>Cell (g liter$^{-1}$)</th>
<th>Biomass (g liter$^{-1}$)</th>
<th>Ethanol yield (g liter$^{-1}$)</th>
<th>Specific ethanol production rate (mmol g$^{-1}$ h$^{-1}$)</th>
<th>Specific CO$_2$ production rate (mmol g$^{-1}$ h$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>0.0580</td>
<td>1.67</td>
<td>0.00091</td>
<td>0.309</td>
<td>0.058</td>
<td>0.064</td>
<td>0.00174</td>
</tr>
<tr>
<td>atg32Δ mutant</td>
<td>0.00093</td>
<td>7.21</td>
<td>0.00069</td>
<td>0.468</td>
<td>0.001</td>
<td>0.041</td>
<td>0.00009</td>
</tr>
</tbody>
</table>

Values are means ± SEM of eight independent experiments initiated with respective starter cultures. Abbreviations and symbols: Yi/j, yield of production of constituent i on constituent j; qCO$_2$, specific CO$_2$ production rate (mmol g$^{-1}$ h$^{-1}$); qEthanol, specific ethanol production rate (mmol ethanol g$^{-1}$ biomass h$^{-1}$); ND, not detectable; *, $P < 0.05$; **, $P < 0.01$ as judged by unpaired one-sided Student’s t-test.
yeast (985 ± 30.8 arbitrary units of Image J software, n = 100; P < 0.01 as judged by the two-sample z-test). This finding may be explained by the difference in cell response of wild-type and \textit{atg32A} cells exposed to stressful conditions induced by accumulation of fermented products and limited nutrient concentration. While wild-type cells use carbon flux to construct cell components by degrading mitochondria and recovering amino acids upon exposure to limited nutrient concentrations, \textit{atg32A} cells lack this response, do not invest in carbon flux directed to biomass, and instead use the carbon flux toward ethanol. 

Although autophagy has been observed under various wine-making conditions (17–19, 44), to our knowledge, no study has defined the role of mitophagy in alcoholic fermentation. The present study demonstrates that mitophagy occurs during alcoholic fermentation and its disruption enhances fermentation. Indeed, very recently, it has been shown that mitophagy occurs during fermentation, independent of respiration (13). Together, these studies indicate that mitophagy during alcoholic fermentation will be a new target in the development of fermentation technologies.

In summary, we have shown here that mitophagy occurs in sake yeast and that deleting the mitophagy gene augments this organism’s capacity to produce ethanol. Considering that natural biomass resources often contain low concentrations of nutrients, this method will provide benefits in the production of bioethanol from biomass resources.

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REFERENCES


