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Title	Identification of cheese rancidity-related lipases in Aspergillus oryzae AHU 7139
Author(s)	Chintagavongse, Napaporn; Kumura, Haruto; Hayakawa, Toru; Wakamatsu, Jun-ichi; Tamano, Koichi
Citation	Journal of bioscience and bioengineering, 137(5), 381-387 https://doi.org/10.1016/j.jbiosc.2024.01.016
Issue Date	2024-03-01
Doc URL	http://hdl.handle.net/2115/92618
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Туре	article (author version)
File Information	JBB R2-marked up.pdf



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- 2 AHU 7139
- 3 A short title: Identification of cheese rancidity-related lipases
- 4 Napaporn Chintagavongse¹, Haruto Kumura^{1,*}, Toru Hayakawa¹, Jun-
- 5 ichi Wakamatsu¹, Koichi Tamano²
- 6 ¹Laboratory of Applied Food Science, Graduate School and Research Faculty of
- 7 Agriculture, Hokkaido University, 060-8589, N9, W9, Sapporo, Japan,²Bioproduction
- 8 Research Institute, National Institute of Advanced Industrial Science and Technology
- 9 (AIST), 2-17-2-1 Tsukisamu-Higashi, Toyohira-ku, Sapporo, Hokkaido 062-8517,
- 10 Japan
- 11
- 12
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- 14
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- 17
- 18 * Corresponding author: Haruto Kumura
- 19 Mailing address: Laboratory of Applied Food Science, Graduate School and Research
- 20 Faculty of Agriculture, Hokkaido University, 060-8589, N9, W9, Sapporo, Japan
- 21 Tel: +81-11-706-3642, E-mail: kumura@agr.hokudai.ac.jp
- 22

23 ABSTRACT

24 The adjunct product with enzymatic activity from Aspergillus oryzae is 25 beneficial for flavor enrichment in the ripened cheese. However, an excessive lipolytic 26 reaction leads to the release of volatile free fatty acids. Accordingly, a strong off-flavor 27 (i.e., rancidity) has been detected when A. oryzae AHU 7139 is used. To identify the 28 rancidity-related lipase from this strain, we evaluated the substrate specificity and 29 lipase distribution using five mutants cultured on a whey-based solid medium under 30 different initial pH conditions. The results showed a higher diacylglycerol lipase 31 activity than triacylglycerol lipase activity. Moreover, an initial pH of 6.5 for the 32 culture resulted in higher lipolytic activity than a pH of 4.0, and most of the activity 33 was found in the extracellular fraction. Based on the gene expression analysis by RT-34 PCR and location and substrate specificity, five genes (No. 1, No. 19, mdlB, tglA, and cutL) were selected among 25 annotated lipase genes to identify the respective 35 36 knockout strains. Because $\Delta tglA$ and $\Delta mdlB$ showed an outstanding involvement in 37 the release of free fatty acids, these strains were applied to in vitro cheese curd 38 experiments. In conclusion, we posit that triacylglycerol lipase (TglA) plays a key role 39 as the trigger of rancidity and the resulting diglycerides have to be exposed to diacylglycerol lipase (MdlB) to stimulate rancidity in cheese made with A. 40 41 oryzae AHU 7139. This finding could help screen suitable A.oryzae strains as cheese 42 adjuncts to prevent the generation of the rancid-off flavor.

43 KEYWORDS: Aspergillus oryzae, lipase, substrate, free fatty acids, rancidity, *in vitro*44 curds

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46 **INTRODUCTION**

The filamentous fungus *Aspergillus oryzae* is used in traditional Japanese cuisine (washoku). *A. oryzae* is recognized as safe because it is missing the aflatoxin synthesis gene (1) and is assumed to encode as much as 135 protease genes based on its genomic analysis (2).

51 In contrast to washoku, A. oryzae is rarely used in Western foods. Kumura et 52 al. (3) focused on the high proteolytic potential of A. oryzae and attempted to apply its 53 culture products (CP) as adjuncts for cheese flavor enrichment. As solid-state 54 fermentation is known to be superior to liquid fermentation for abundant enzyme 55 production (4), Kumura et al. (3) prepared solid CP of A. oryzae AHU 7139 as adjuncts 56 for making Gouda-type cheese and confirmed the enhancement of proteolysis during cheese ripening. Moreover, a remarkable content of free fatty acid (FFA) due to 57 58 lipolysis was detected. They reported that the application of CPs whose fermentation 59 had started with a pH of 6.5 resulted in a pronounced increase in FFA compared to the 60 fermentation that started with a pH of 4.0, implying the initial pH of the culture may 61 affect the profile of the lipases produced.

62 Although proper lipolysis provides the characteristic flavor and aroma to 63 cheese, excess FFA accumulation during cheese ripening causes an undesirable off-64 flavor known as rancidity (5) by the release of volatile short-chain fatty acids over an organoleptic threshold. Furthermore, once milk triacylglycerols are degraded by 65 lipases (the first step of lipolysis), the resulting diglycerides are exposed and become 66 67 the subsequent substrates for diacylglycerol lipase (the second step of lipolysis). Thus, 68 the adjunct materials of A. oryzae should be prepared with high proteolytic but limited 69 lipase activity.

70 According to the genomic analysis of a wild-type strain, about 30 genes are 71 annotated to encode lipase genes (FungiDB database, 72 https://fungidb.org/fungidb/app/). Expression of some lipases is likely to be dependent on the culture conditions, and the identification of the rancid-inducible lipase 73 74 molecules could provide valuable information to screen suitable A. oryzae strains for 75 preventing flavor defects in dairy application.

Hence, we compared lipase gene expression under different pH conditions and selected some candidate lipase genes. Then, the influence of mutations on the lipase activity was evaluated. Finally, we identified the most influential lipase molecules on the rancidity, using *in vitro* cheese curd experiments.

80 MATERIALS AND METHODS

81 Strain and culture condition

82 The strain used in this study was A. oryzae AHU 7139 from the culture collection of 83 Hokkaido University and was grown on potato dextrose agar (PDA) (Merck KGaA, 84 Darmstadt, Germany) at 30°C for 7 days. Spore suspensions were prepared by adding 9.0 g L⁻¹ sodium chloride (NaCl) solution into the grown culture on PDA and diluted 85 to the concentration of 2.5×10^5 spores mL⁻¹, which was counted by using 86 haemocytometer (NanoEntek, Seoul, Korea). The spore suspension (150 µL) was 87 88 inoculated on whey solid medium whose pH was adjusted to 4.0 or 6.5, and cultivated 89 at 20° C for 7 days (6).

90 **RNA isolation and RT-PCR**

91 The solid culture product (CP) was recovered and immediately freeze-dried and 92 ground with a mortar and pestle to obtain freeze-dried powder (FDP). On average, 93 0.27 g FDP was obtained from 1 g CP. The FDP (0.5 mg) was transferred into 5 mL of neutralized phenol-saturated water containing 2 mol L⁻¹ NaCl, mixed well, and 94 95 incubated in a waterbath 55°C for 10 min. The sample was centrifuged at $10,000 \times g$, 96 4°C for 15 min and the upper aqueous phase was transferred to a new tube followed 97 by the addition of chloroform: isoamylalcohol (49:1; CIA). After centrifugation at $10,000 \times g$, 4°C for 10 min, the upper phase was transferred to a new tube followed by 98 99 addition of water-saturated acidic phenol/CIA (1:1) and centrifuged again (this step 100 was repeated two times). Then, the upper phase was poured into a new tube and CIA was added into the tube, followed by centrifugation for 5 min. After transferring the 101 upper phase to a new tube, 0.5 volume of 7.5 mol L⁻¹ LiCl (Kanto Chemical, Tokyo, 102

103 Japan) was added to the tube. The mixture was incubated at 4°C overnight, followed 104 by centrifugation at 12,000 ×g, 4°C for 30 min. The supernatant was removed and the pellet was rinsed with 2.5 mol L⁻¹ LiCl and centrifuged twice at 12,000 × g , 4°C for 5 105 106 min. The air-dried pellet was finally dissolved in formamide to obtain total RNA and 107 stored at -80°C. The RNA concentration was estimated by measuring the absorbance 108 at 260 nm with a NanoDrop spectrophotometer (Thermo scientific, US). The recovery 109 of total RNA was evaluated by electrophoresis using a commercial kit (Dynamarker 110 RNA High for Easy Electrophoresis, Biodynamics Laboratory, Tokyo, Japan) according to the manufacturer's instructions (Fig. S1). 111

112 The mRNA was isolated from the total RNA using a commercial kit (Pharmacia 113 Biotech, Stockholm, Sweden) according to the manufacturer's instructions. The mixture containing 50 ng of mRNA sample, random primer (Promega, Wisconsin, US), 114 115 and dNTP was heated at 65°C for 5 min and immediately put on ice. Then, the mixture 116 containing 5x RT buffer, RNase inhibitor (Toyobo, Osaka, Japan), and reverse 117 transcriptase (M-MLV, Nippon Gene, Tokyo, Japan) was added and incubated at 37°C 118 for 60 min according to the manufacturer's instructions. The diluted cDNA sample 119 $(0.5 \ \mu L)$ was used as a template and mixed with the designed primers (1.5 μL), nuclease-free water (3 µL), and 5 µL of the PCR mixture solution (GoTaq[®] DNA 120 121 polymerase and 5x green reaction buffer, Promega, Wisconsin, US). The PCR 122 amplification involved denaturing at 95°C for 2 min, followed by 40 cycles at 95°C for 123 45 s, annealing at 55°C for 30 s for all candidate lipase genes, and an extension at 73 °C for 1 min. A final extension at 73°C for 10 min was followed by cooling to 4°C. All 124 125 the primers used in this study are listed in Table S1. The reliability of all primer sets 126 was checked by PCR using genomic DNA from A. oryzae AHU 7139 and RIB40 as 127 the template (data not shown).

128 Disruption of individual genes in *A. oryzae* AHU 7139

Five candidate genes (AO090012000690, AO090005001319, AO090701000644, AO090003001507, and AO090005000029, named No.1, No. 19, *mdlB*, *tglA*, and *cutL*, respectively) were selected to detect the lipase activity after gene disruption. All the primers used throughout the gene disruption are shown in Table S2, based on the genomic sequence of *A. oryzae* wild-type strain RIB40 (2), used as a reference for designing primers.

135 Prior to the candidate gene disruption, the pyrG gene (AO090011000868) was 136 knocked out in AHU 7139 by homologous recombination under selective pressure by 137 5-fluoroorotic acid (5-FOA). The *pyrG* knockout was aimed at conferring the uracil auxotrophy to AHU 7139 so as to make the pyrG available as a selectable marker. The 138 139 DNA cassette for *pyrG* knockout was prepared by fusion PCR. Briefly, the promoter 140 and terminator regions of *pyrG* were amplified using LUpyrG/LLpyrG and 141 RUpyrG/RLpyrG primer pairs, respectively. The AHU 7139 chromosomal DNA was 142 used as the template, and the KOD-PLUS DNA polymerase was used for DNA 143 amplification (Toyobo Co. Ltd., Osaka, Japan). The amplified DNA fragments were purified by gel extraction using the Wizard SV Gel and PCR Clean-Up System 144 145 (Promega Co., Madison, WI, USA) according to the manufacturer's instructions. Two 146 types of purified DNA fragments were then mixed and joined by fusion PCR using a 147 LUpyrG/RLpyrG primer pair and KOD-PLUS (Fig. S2). The resulting 2035 bp-long 148 DNA fragment was purified by gel extraction and then applied to the transformation 149 of AHU 7139. Protoplasts of AHU 7139 were prepared as reported previously (7) and used for the transformation. To generate transformants, Czapek-Dox (CD) minimal 150 agar medium supplemented with 1.2 mol L^{-1} sorbitol, 1 mg m L^{-1} 5-FOA, 5 mmol L^{-1} 151 uridine, and 10 mmol L⁻¹ uracil was used. Because the non-homologous end-joining 152

153 (NHEJ) activity is originally high in filamentous fungi including *A. oryzae*, about 20 154 single colonies of transformants were screened. After single-spore isolation of 155 transformants was performed three times using CD agar supplemented with 1 mg mL⁻ 156 ¹ 5-FOA, 5 mmol L⁻¹ uridine, and 10 mmol L⁻¹ uracil, the *pyrG* knockout was checked 157 by PCR using a LUpyrG/cLpyrG primer pair. Only one positive homokaryon clone of 158 the *pyrG* knockout mutant was acquired and named AHU 7139 [pyrG⁻] (Fig. S3).

159 Then, five candidate genes were individually disrupted using AHU 7139 [pyrG⁻] as a 160 host. To increase the probability of the locus-specific homologous recombination, the 161 *pvrG* marker-split method was applied to the candidate gene disruption (8). The former 162 and latter parts of pyrG were amplified by PCR using PU/PLsplit and PUsplit/PL 163 primer pairs, respectively. The 548 bp-long DNA region overlapped between them. 164 Approximately 1 kb of promoter and terminator of the candidate genes were also amplified by PCR, using LU/LL and RU/RL primer pairs, respectively. After 165 166 purification of these four types of DNA fragments by gel extraction, both pairs of 167 promoter/pvrG (former part) and pvrG (latter part)/terminator were concatenated by 168 fusion PCR, using LU/PLsplit and PUsplit/RL primer pairs, respectively. As a result, 169 two types of fusion PCR DNA fragments were prepared as the gene disruption 170 cassettes per candidate gene. The triple crossover between the chromosomal DNA and 171 the two cassettes that occurred in the marker-split method is illustrated in Fig. S4A–E. 172 Protoplasts of AHU 7139 [pyrG⁻] were prepared as mentioned above. The protoplasts 173 were applied to co-transformation with the two types of DNA fragments as gene 174 disruption cassettes. In other words, two types of DNA fragments were simultaneously 175 introduced into the protoplasts for each candidate gene disruption. Consequently, the transformants were generated on CD agar supplemented with 1.2 mol L⁻¹ sorbitol, and 176 177 five single colonies were selected per candidate gene. The selected clones were

subjected to single-spore isolation on CD agar at least three times, followed by
evaluating the candidate gene disruption by PCR, using cU/cL primer pairs (data not
shown).

181 The candidate gene disruption was further confirmed by Southern hybridization (Fig. 182 S5A–E). Aliquots (8 µg) of genomic DNA were digested with each restriction enzyme 183 (HincII, PstI, SphI, and PvuII), fractionated on 0.6% agarose gel, and transferred onto 184 a Nytran SuPerCharge membrane (GE Healthcare Co., Piscataway, NJ, USA). 185 Hybridization and signal detection were performed using a digoxigenin (DIG) system 186 according to the manufacturer's instructions (GE Healthcare Co.). Briefly, DIG-187 labeled probes (300-800 bases long) were prepared using a PCR DIG Probe Synthesis 188 Kit (Roche Applied Science AG, Mannheim, Germany) with SBU/SBL primer pairs, 189 followed by agarose gel extraction. The probes were hybridized to the DNA fragments 190 bound to the membrane, and the signals of the probes were detected via 191 chemiluminescence arising from the anti-DIG Fab fragment alkaline phosphatase 192 conjugate (Roche Applied Science AG), using the CDP-Star Detection Reagent (GE 193 Healthcare Co.). A WSE-6100H LuminoGraph I gel imager (ATTO Co. Ltd., Tokyo, 194 Japan) was used for the signal detection of Southern hybridization. The acquired 195 disruptants of the candidate genes ($\triangle 1$, $\triangle 19$, $\triangle mdlB$, $\triangle tglA$, and $\triangle cutL$) and the 196 parent strain were used to measure lipase activities and FFA levels in the in vitro 197 cheese model.

198 Biomass measurement by glucosamine assay

The glucosamine content was determined to estimate the fungal biomass in the solid
medium, using the method of Fuji et al. (9) and Kasuga (10) with some modifications.
The CP (0.5 g) was washed with 50 mmol L⁻¹ sodium phosphate buffer pH 7.0 and

shake vigorously by hand to obtain a homogeneous dispersion. Then, the samples were centrifuged at 1,070 ×g for 10 min at room temperature, and the supernatant was discarded. After washing three times, 15 mL of the same buffer was added to disperse the solid culture, followed by the addition of 20 mg of Yatalase (Takara Bio, Tokyo, Japan). Then, the tube was shaken at 60 rpm, 37°C for 1.5 h, followed by centrifugation. The supernatant (10 mL) was filtered through a 0.45 μ m filter and collected in a 15 mL conical tube. The samples were kept at -20°C until use.

209 One hundred microliters of the sample or N-acetylglucosamine (GlcNAc) standard 210 solution were transferred to a 1.5 mL microtube and then mixed with 40 µL of 0.4 mol 211 L^{-1} potassium tetraborate tetrahydrate solution. The sample tubes were heated at 100°C for 3 min and were then cooled down at room temperature. Then, 600 µL of p-212 213 dimethylaminobenzaldehyde (DMAB) solution (0.1 g of DMAB powder dissolved in a mixture of 17.3 mol L⁻¹ acetic acid and 0.37 mol L⁻¹ HCl) was added to the tube and 214 215 incubated at 37°C for 20 min. Finally, 150 µL of each sample solution was transferred 216 in duplicate into a 96-well microplate to measure absorbance at 595 nm, using a 217 microplate reader (Infinite F200 PRO, Tecan, Switzerland). One milligram of dried 218 koji mycelium contains 0.628 µmol of GlcNAc according to Fujii et al. (9).

219 Preparation of the extracellular and intracellular enzymes

The CPs were mixed with an equal weight of deionized water and treated by a stomacher for 5 min. The materials were then transferred to a centrifugal tube and centrifuged at $21,130 \times g$, $4^{\circ}C$, 10 min. The supernatant was recovered and the precipitate was re-extracted with deionized water another two times. The pooled supernatants were used as the extracellular enzyme. The precipitate was washed once more with deionized water and freeze-dried. The freeze-dried sample was ground into a fine powder with a mortar and pestle and then dispersed in a simulated milk
ultrafiltrate (SMUF) at pH 5.5 (11), which was used as the intracellular enzyme.

228 Measurement of fractionated lipase activity

229 The extracellular enzyme (70 µL) was mixed with 230 µL of SMUF buffer pH 5.5, 230 while the suspension comprising 0.05 g of intracellular FDP dispersed in 300 μ L of 231 SMUF pH 5.5 was used as the intracellular enzyme. The lipase activity was determined 232 as previously described, with some modifications (12). In brief, one gram of purified 233 triolein or diolein emulsified with 100 mL of 2% polyvinyl alcohol (degree of 234 polymerization 2000, Kishida Chemical, Osaka, Japan) was used as the substrate. 235 Purification of triolein was carried out by loading commercial triolein (Kanto 236 Chemical, Tokto, Japan) on the column packed with silica gel 60 N (Kanto Chemical, 237 Tokyo, Japan) pre-equilibrated with an organic solvent mixture of hexane and 238 chloroform (5:1). Triolein was recovered by the isocratic elution of the same solvent. 239 Purification of diolein was carried out by loading commercial diolein (Kanto Chemical, 240 Tokyo, Japan) on the same column pre-equilibrated with the organic solvent mixture 241 of hexane and chloroform (5:1). Then, diolein was recovered by the isocratic elution 242 of another solvent mixture of hexane and chloroform (1:1). The purity of these lipids 243 was confirmed by conventional thin layer chromatography (TLC) (Fig. S6).

The substrate was divided into 2 mL for each test tube and subjected to pre-incubation at 30°C for 5 min, followed by the addition of the enzyme to be tested (300 μ L). The reaction was carried out at 30°C for 30 min. The reaction was terminated by adding 7.5 mL of extract solution (heptane : isopropanol : 0.5 mol L⁻¹ sulfuric acid = 48:48:4). FFA measurement was performed using a spectrophotometer at 570 nm, and the values were corrected by subtracting the value obtained from the blank, which was subjected to the same treatment except for the incorporation of the enzyme after the addition of the extract solution. Enzyme activity was expressed as μ mol of released oleic acid from the substrate per 1 h at the defined temperature from one gram of the CPs (1 μ mol oleic acid/h/g-CP = 1 lipase unit: LU). The specific lipase activity was calculated based on the biomass and expressed as the total lipase unit per mg of dried koji biomass (LU/ mg koji).

256 Study of the FFA levels of mutants in a cheese curd model

The *in vitro* cheese curds used in this study were prepared as previously described (6) with the addition of 2% NaCl. After producing the curds, 0.2 g of FDP prepared using the culture with an initial pH of 4.0 or 6.5 was mixed with 200 g of curds, and 20 g was divided into nine bags, and then vacuum-sealed. The culture with the parent strain (P4, P6.5) or mutants, including $\Delta mdlB$ (M4, M6.5) and $\Delta tglA$ (T4, T6.5), were used. The control cheese (C) had no addition of any adjunct material. The curd packs were ripened at 12°C and sampled at 4 weeks.

For FFA determination, the triplicate curd sample (0.2 g) was transferred to 2 mL of 7.7 mol L⁻¹ HCl in a test tube and placed into boiling water to dissolve the sample. FFAs, extracted by the same extraction reagent as was used in the lipase activity measurement were determined by the phenol-red method (13) and its content was expressed as millimoles per 100 g of curds using oleic acid as the standard.

269 Statistical analysis

270 FFA content in the curds was analyzed using Tukey-Kramer's multiple comparison

test. The data were analyzed by JMP software (Pro 17; SAS Institute, Inc., Tokyo,

Japan). Differences were considered to be statistically significant at p < 0.05.

273 RESULTS AND DISCUSSION

Qualitative gene expression analysis by RT-PCR to pre-screen of candidate lipase genes for mutation

The results of RT-PCR performed with twenty-five candidate primers revealed that initial cultures with a pH of either 4.0 or 6.5 resulted in the expression of eight and six genes respectively (Fig. 1). gDNA contamination in cDNA was unlikely because no amplification was found when the mRNA preparation was used as a template (Fig. S7). Annotated lipase genes Nos. 1, 20, 22, *mdlB*, *tglA*, and *cutL* were expressed in *A. oryzae* AHU 7139 with either an initial pH 4.0 or 6.5, whereas Nos. 19 and 21 showed limited expression in the culture with an initial pH of 4.0.

283 To narrow down the candidates for mutation, we evaluated the substrate specificity and distribution of the lipase in the culture products (CPs) 284 285 of A. oryzae AHU 7139 to compare with the gene expression profile. Introducing 286 purified substrates revealed that the parent strain of A. oryzae AHU 7139 produced 287 DG lipolytic activity approximately twenty times higher than TG lipase activity at both 288 initial pHs (Fig. 2). The activity was predominantly found in the extracellular fraction 289 although the CPs prepared with an initial pH of 4.0 showed a higher ratio of 290 intracellular DG lipase than those prepared at pH 6.5. It should be noted that the 291 number of genes expressed at pH 4.0 was greater than at pH 6.5 (Fig. 1) despite the 292 lower total lipase activity of the initial culture at pH 4.0 compared to pH 6.5 (Fig. 2). 293 The gene No.1 (AO090012000690) annotated as a triglyceride lipase and the gene No. 294 19 (AO090005001319), annotated as a lipase and carboxyl ester hydrolase, were the 295 focus of further study because of the limited information available regarding substrate 296 specificity. On the other hand, Nos. 20 (AO090005001602), 21 (AO090023000717)

and 22 (AO090003000839) were screened out because they were reported to be intracellular TG lipases (14) while we found limited intracellular lipase activity toward triolein under either initial pH (Fig. 2). The expression of the *cutL* (15), *mdlB* (16), and *tglA* (17) genes has been demonstrated in liquid culture and this study demonstrated their expression in a solid-state medium as well. Taking the lipase activity and the gene expression into account, two annotated lipase genes (Nos. 1 and 19), *mdlB*, *tglA*, and *cutL* were selected to obtain mutants.

304 Lipase production and distribution in the mutant strains of *A. oryzae* AHU

305 **7139**

306 We introduced a single-gene knockout of A. oryzae AHU 7139 to confirm the 307 lipase productivity. Considering the growth rate difference of the mutants, the specific 308 lipase activity that represent the activity based on the biomass was compared (Table 309 1). Reduced specific TG lipase activity was observed in $\Delta 1$ (694.1 LU/mg koji), $\Delta 19$ 310 (430.3 LU/mg koji) and $\Delta cutL$ (1079.8 LU/mg koji) when cultivation was initiated at 311 pH 4.0, but the activity was elevated when cultivation was initiated at pH 6.5 (9165.5, 312 20209.0, and 12047.9 LU/mg koji, respectively) compared to the parent strain (1934.7 313 and 5570.3 LU/mg koji at pH 4.0 and 6.5, respectively). CutL from A. oryzae was 314 assumed to be one of the factors responsible for rancidity in dairy products since it 315 showed higher activity on esters of short-chain fatty acids (15). However, CutL as well 316 as Nos. 1 and 19, were unlikely to be related to rancidity because the rancidity was 317 more pronounced in cheese prepared using the adjuncts produced at an initial culture 318 pH of 6.5 rather than pH 4.0. In contrast, $\Delta m dl B$ showed the opposite response because 319 specific TG lipase activity at pH 6.5 was reduced to 61.9% (3450 LU/mg koji) 320 compared with that of the parent strain with slight increase of 2153.7 LU/mg koji at

321 pH 4.0. Although no substrate specificity of MdlB toward triacylglyceride has been 322 reported (16). Lan et al. (18, 19) reported that the single point mutation of mono- and 323 diacylglyceride lipase increased the lipolytic activity toward TG and DG making it 324 function as a TG lipase. Thus, we cannot rule out that certain modifications of *mdlB* 325 in A. oryzae AHU 7139 conferred hydrolytic capability toward triacylglycerides. On 326 the other hand, an 83% reduction of TG lipase activity was detected in $\Delta tglA$ (968.1 327 LU/mg koji) when the cultivation was initiated at pH 6.5 whereas only a 25% reduction 328 of TG lipase activity was found in $\Delta tglA$ when the cultivation was initiated at pH 4.0 329 (1451 LU/mg koji). Regarding the DG lipase, Nos. 1, 19, and CutL were proven to be 330 active toward DG as well as TG since the specific lipase activity of the culture initiated 331 at pH 4.0 was reduced by 59%, 77% and 34.6% in the $\Delta 1$ (19889.9 LU/mg koji), $\Delta 19$ 332 (11003.0 LU/mg koji) and $\Delta cutL$ (31720.8 LU/mg koji) mutants, respectively. We 333 initially speculated that the biological significance of these lipases under acidic 334 circumstance was related to the transition into the sporulation phase of A. oryzae (14) 335 because the spore-forming ability was recognized in the CP initiated at pH 4.0 but not 336 in that initiated at pH 6.5 (Fig. S8). In fact, it has been suggested that the metabolism 337 of storage lipids (20) and two intracellular lipases in A. oryzae BCC7051 (14) are 338 involved in fungal spore production. However, $\Delta 1$, $\Delta 19$ and $\Delta cutL$ exhibited similar 339 spore formation at the end of the cultivation (data not shown), which indicated that 340 these three lipases are independent on spore formation. It is unclear that an increase in 341 the specific DG and TG lipase activity in $\Delta 19$ in the culture initiated at pH 6.5, despite 342 No. 19 was unexpressed under the initial culture condition at pH 6.5 (Fig.1). ΔmdlB 343 showed the greatest lipase reduction (84%) compared to the parent strain under either 344 pH condition (from 48519.8 LU/mg koji to 7844.1 LU/ mg koji at pH 4.0 and from 345 104114.6 LU/ mg koji to 16393.0 LU/ mg koji at pH 6.5). Thus, major DG lipase

346 activity could be ascribed to MdlB. Furthermore, major intracellular DG lipase found 347 in the culture at 4.0 was regarded as MdlB because $\Delta mdlB$ showed limited intracellular 348 lipase accumulation (Fig. 2). The MdlB found in the intracellular fraction was 349 probably due to the limited secretion period, followed by the rapid life cycle transition 350 into conidiation. As Toida et al. (17) reported the hydrolytic ability of TglA toward 351 DG, partial reduction of specific DG lipase activities of $\Delta tglA$ was confirmed in both 352 initial pHs. Overall, in the culture initiated at pH 4.0, the predominant TG lipase 353 activity was derived from Nos. 1, 19 and CutL, whereas that was due to TglA in the 354 culture initiated at pH 6.5 (Table 1). These results demonstrate that the production of 355 triglyceride lipase molecules depends on the initial culture pH. On the other hand, 356 MdlB was the major DG lipase irrespective of the initial culture pH. Considering the 357 higher level of total lipase activity in the culture initiated at pH 6.5, TglA and MdlB 358 were more likely to be related to rancidity. Therefore, $\Delta tglA$ and $\Delta mdlB$ were selected 359 for further study.

360 In vitro studies of FFA profiles in the cheese curd model

361 Finally, we conducted an in vitro cheese curd experiment to evaluate the effects 362 of triacylglycerol lipase (TglA) and mono-, diacylglycerol lipase (MdlB) on the free 363 fatty acid (FFA) levels in the curds. Table 2 shows the whole lipase activity of freeze-364 dried powder (FDP), used for mixing with curds. The lipase activity was higher in all 365 CPs with an initial pH of 6.5 compared to those with an initial pH of 4.0, except for 366 the activity of mutant $\Delta tglA$ toward TG. The triacylglycerol lipase activity in all 367 mutants was lower than that in the parent strain for both initial culture pHs. Moreover, 368 lipase activity toward DG was not detected in $\Delta mdlB$ initially cultured at pH 4.0, and 369 low activity was observed with an initial pH of 6.5 (302.3 LU). After 4 weeks of 370 ripening, the addition of the CPs from the mutant $\Delta tglA$ resulted in a significantly 371 lower amount of FFA than that from the parent strain (p < 0.05) and retained a 372 comparable FFA level to that of the control sample (Fig. 3). Moreover, the addition of 373 CPs from the mutant $\Delta mdlB$ initially cultured at pH 6.5 resulted in a significantly lower 374 FFA level than that of the parent, although higher than that of the control cheese (Fig. 375 3).

FFA accumulation during cheese ripening is mainly attributed to the action of 376 377 exogenous lipases because the endogenous milk lipase is heat-labile and can be 378 inactivated by pasteurization (21). We found that the released FFA in the *in vitro* curds 379 seemed to correlate with the TG lipase activity levels in the adjuncts (Fig.3, Table 2). 380 Since the FFA accumulation in $\Delta tglA$ was significantly reduced in the ripening curds 381 regardless of the initial pH, we suggested that TglA triggers hydrolysis that results in 382 rancidity. As described above, lipases No. 1, No. 19, and CutL should still be active in 383 either CPs from $\Delta tglA$ and $\Delta mdlB$. However, the CPs from $\Delta tglA$ showed comparable 384 FFA levels to the control, which demonstrated that those lipases were not as important 385 as TglA under the cheese ripening circumstance. This is probably due to the reduced 386 catalytic activity of other lipases at the temperature of cheese ripening (12°C) and/ or 387 due to the degradation by concomitant rennet, which is supplied as the milk 388 coagulation enzyme. The reduction of the FFA levels of $\Delta mdlB$ indicates that MdlB 389 lipase should not be overlooked when considering rancidity. Okumura et al. (22) 390 reported that mono- and diacylglycerol lipases, from Penicillium cyclopium M1, more 391 easily hydrolyze acylglycerols than triacylglycerol lipase, and the synergistic action of 392 mono- and diacylglycerol lipase with triacylglycerol lipase could accelerate lipolysis. 393 Wong et al. (23) found elevated FFA productivity in A. oryzae RIB 40 when they

applied the overexpressed strain AO090701000644 (*mdlB*). Based on these results, we
postulated a rancidity process for ripened cheese caused by *Aspergillus* lipolysis (Fig.
4). Extracellular triacylglycerol lipase (TglA) is crucial for the degradation of
triglycerides in milk and the release of FFA and DG as the first step of rancidity.
Subsequently, the resulting DGs are provided as the substrate for MdlB lipase. The
dual effect of TglA, followed by MdlB, is likely to cause an excess amount of volatile
fatty acids and rancidity in cheese products made with *A. oryzae*.

401 It might be possible to distinguish MdlB from TglA only if highly purified 402 triolein is used as the substrate. Suzuki et al. (24) found a poor correlation between 403 lipase activity and butyric acid levels in cheese products made with A. oryzae when 404 they used a synthetic substrate of *p*-nitrophenol butyrate, which might represent whole 405 esterase activity. Even though butter oil was used as the substrate for lipase activity 406 measurement, FFA levels in the resulting cheese products were not correlated with the 407 lipase activities in the adjuncts prepared using Aspergillus CPs (6). As butter oil 408 included TG as the major component, with DG and MG as minor components, it could 409 be as the substrate not only for TG lipase but also for the abundant DG lipase. Thus, 410 when suitable strains of A. oryzae are screened with respect to lipase activity for the 411 preparation of adjuncts for cheesemaking, a strict evaluation of TG activity should be 412 performed.

In conclusion, *A. oryzae* is expected to generate adjuncts to enrich the flavor of ripened cheese if the excessive accumulation of volatile fatty acids due to lipase activity is reduced. In this study, we demonstrated that *tglA* is the most pivotal lipase gene encoded in *A. oryzae*. Therefore, a TglA-inactive strain might be a candidate for use as an adjunct for cheese ripening to prevent the production of rancid-off flavor. We are currently investigating the reduction of TglA catalytic activity by substituting amino acid(s) and inserting or deleting nucleotides, leading to a frameshift of *tglA* transcripts. Furthermore, genome editing might be an alternative to obtain desirable strains.

422 ACKNOWLEDGEMENTS

This research was partially supported by grants from The Tojuro Iijima Foundation for Food Science and Technology and Sapporo Bioscience Foundation. The authors are very grateful to Yu Toba and Hiroki Matsumo, Field Science Center for Northern Biosphere, Hokkaido University, for their assistance with cheese curds preparation. We also thank Professor Satoshi Koike for the experimental support.

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498 Figure Legends

- 499 Fig. 1. Agarose gel (1%) electrophoresis of RT-PCR products with cDNA from CP
- 500 initial pH 4.0 (A) and pH 6.5 (B). M = DNA ladder marker, lane no.1-22 = annotated
- 501 lipase no. 1-22, m = mdlB, t = tglA, and c = cutL gene. Red box shows the expected
- 502 amplified band while others are non-specific bands.
- 503 Fig. 2. Lipase activities of *A. oryzae* AHU 7139 (parent) and five selected mutants
- 504 ($\Delta mdlB$, $\Delta tglA$, $\Delta cutL$, $\Delta 1$, $\Delta 19$) toward purified triolein (TG) and diolein (DG) from
- 505 the solid culture products (CP) initiated with pH 4.0 and pH 6.5. Values are expressed
- 506 in average value (n = 2).
- 507 **Fig. 3.** Free fatty acid (FFA) profiles in cheese model at 4 weeks.
- 508 C = curds only (control), P4 = parent AHU 7139 with initial culture pH 4, M4 = mutant

509 AHU 7139 $\Delta mdlB$ with initial pH 4, T4 = mutant AHU 7139 $\Delta tglA$ with initial pH 4,

- 510 P6.5 = parent AHU 7139 with initial pH 6.5, M6.5 = mutant AHU 7139 $\Delta mdlB$ with
- 511 initial pH 6.5, T6.5 = mutant AHU 7139 $\Delta tglA$ with initial culture pH 6.5. Results of
- 512 FFA level are expressed as the mean \pm SE (n = 3). The different letters indicate a
- significant difference between the samples using Tukey-Kramer test (p < 0.05)

514 Fig. 4. The proposed rancidity process in the ripened cheese caused by Aspergillus 515 lipolysis. As triacylglycerol (TG) is the major lipid in milk, triacylglycerol lipase 516 (TglA) hydrolyzes TG at any fatty acid position ((A) sn-1, 3 or (B) sn-2) and generates 517 diacylglycerol (DG) and free fatty acids (FAs) during ripening. Subsequently, 518 diacylglycerol lipase (MdlB and, to some extent, TglA) hydrolyzes DG and produces 519 FFA. The predominant lipases (TglA or MdlB) at each hydrolytic step are shown in 520 bold, although other genes might be involved. The excess amounts of free volatile fatty 521 acids released by fungal lipolysis could cause rancidity.

522 Supplementary Figure Legends

Fig. S1. RNA integrity check by non-denaturing agarose gel electrophoresis. Total RNA was isolated from *A. oryzae* AHU 7139 inoculated on whey solid culture adjusted to pH 4.0 and pH 6.5. +con = positive control of 28S and 18S RNA bands from bovine endometrial stromal cells.

527 Fig. S2. Construction of the DNA fragment for *pyrG* (AO090011000868) knockout

528 in A. oryzae AHU 7139. The 2035 bp-long DNA fragment was constructed for the

529 knockout. Primers used for the construction and clone check are shown as colored

530 and black arrows, respectively.

531 Fig. S3. Clone check of the *pyrG* knockout mutant isolated after transformation.

532 Transformant clone Nos. 1-3 are shown. Lane M: 1 kb DNA ladder marker [1-10 kb],

lane R: RIB40 strain as a negative control, lane A: AHU 7139 strain as a negative
control. Amplified DNA size: Positive clone, 1.0 kb; Negative clone, 2.1 kb. The clone

535 No. 3 in red was considered a positive homokaryon, which was named AHU 7139 536 [pyrG⁻].

Fig. S4A. Construction of the DNA fragment for disrupting AO090003001507 in *A*.

oryzae AHU 7139 [pyrG⁻]. The 2048 bp-long and 2375 bp-long DNA fragments were
constructed for the disruption. Primers used for the construction and clone check are
shown as colored and black arrows, respectively.

541 **Fig. S4B.** Construction of the DNA fragment for disrupting AO090701000644 in A.

542 oryzae AHU 7139 [pyrG⁻]. The 2021 bp-long and 2377 bp-long DNA fragments were

543 constructed for the disruption. Primers used for the construction and clone check are

shown as colored and black arrows, respectively.

545 Fig. S4C. Construction of the DNA fragment for disrupting AO090005000029 in A.

546 *oryzae* AHU 7139 [pyrG⁻]. The 2035 bp-long and 2392 bp-long DNA fragments were

- 547 constructed for the disruption. Primers used for the construction and clone check are548 shown as colored and black arrows, respectively.
- 549 Fig. S4D. Construction of the DNA fragment for disrupting AO090012000690 in A.
- 550 *oryzae* AHU 7139 [pyrG⁻]. The 2031 bp-long and 2403 bp-long DNA fragments were
- 551 constructed for the disruption. Primers used for the construction and clone check are
- shown as colored and black arrows, respectively.
- 553 Fig. S4E. Construction of the DNA fragment for disrupting AO090005001319 in A.
- 554 *oryzae* AHU 7139 [pyrG⁻]. The 2045 bp-long and 2408 bp-long DNA fragments were
- 555 constructed for the disruption. Primers used for the construction and clone check are
- shown as colored and black arrows, respectively.
- 557 Fig. S5A. Southern hybridization analysis of AO090003001507 disruptant constructed
- 558 from AHU 7139. 1: AHU 7139; 2: AO090003001507 disruptant.
- 559 Fig. S5B. Southern hybridization analysis of AO090701000644 disruptant constructed
- 560 from AHU 7139. 1: AHU 7139; 2: AO090701000644 disruptant.
- 561 Fig. S5C. Southern hybridization analysis of AO090005000029 disruptant constructed
- 562 from AHU 7139. 1: AHU 7139; 2: AO090005000029 disruptant.
- 563 Fig. S5D. Southern hybridization analysis of AO090012000690 disruptant constructed
- 564 from AHU 7139. 1: AHU 7139; 2: AO090012000690 disruptant.
- 565 Fig. S5E. Southern hybridization analysis of AO090005001319 disruptant constructed
- 566 from AHU 7139. 1: AHU 7139; 2: AO090005001319 disruptant.
- 567 **Fig. S6.** Thin layer chromatography of the purified lipids. M = control monoglyceride,
- 568 D = control diglyceride, T = control triglyceride (tributyrin), PD = purified commercial
- 569 diolein using silica gel chromatography, PT = purified commercial triolein using silica
- 570 gel chromatography, CD = commercial diolein, CT = commercial triolein. Migration
- 571 of the sample on the silica gel plate was carried out using chloroform : methanol :

- 572 formic acid : DI water (45:20:2.5:1), stained with 0.05% primulin dissolved in
- 573 acetone : water (4:1 v/v), and observed under UV light.
- 574 Fig. S7. Agarose gel (1%) electrophoresis of PCR using mRNA from CP with initial
- 575 pH 4.0 (A) and pH 6.5 (B) as a template with all candidate lipase gene primers. M =
- 576 DNA ladder marker, lane no.1-22 = annotated lipase no. 1-22, m = mdlB, t = tglA, and
- 577 c = cutL gene. Confirming no gDNA in the mRNA sample.
- 578 Fig. S8. Growth and appearance of *A. oryzae* AHU 7139 on the WPC media incubated
- 579 at 20°C for 7 days. The initial culture pH were of (A) pH 4.0 and (B) pH 6.

Samula	dried biomass	TG	DG
Sample	(mg)	LU ⁽¹⁾ / mg koji	LU ⁽¹⁾ / mg koji
CP initial pH 4.0			
parent	0.008	1934.7	48519.8
Δ1	0.014	694.1	19889.9
Δ19	0.027	430.3	11003.0
$\Delta m dl B$	0.004	2153.7	7844.1
$\Delta tglA$	0.013	1451.0	32850.0
ΔcutL	0.015	1079.8	31720.8
CP initial pH 6.5			
parent	0.016	5570.3	104114.6
Δ1	0.018	9165.5	99601.9
Δ19	0.005	20209.0	330880.8
$\Delta m dl B$	0.013	3450.7	16393.0
$\Delta tglA$	0.017	968.1	88648.6
ΔcutL	0.017	12047.9	108810.5

	Whole	lipase activity
Sample	TG	DG
P4	25.6	1632.4
M4	15.4	0.0
T4	8.9	808.9
P6.5	86.6	2433.3
M6.5	66.8	302.3
T6.5	4.8	2226.6
arent strain of A. oryzad	e AHU 7139, M = $\Delta mdlB$, T = Δ	<i>tglA</i> . FDP recovered from cultu



Fig. 1. Agarose gel (1%) electrophoresis of RT-PCR products with cDNA from CP initial pH 4.0 (A) and pH 6.5 (B). M = DNA ladder marker, lane no.1-22 = annotated lipase no. 1-22, m = mdlB, t = tglA, and c = cutL gene. Red box shows the expected amplified size band while others are non-specific bands.





Fig. 2. Lipase activities of *A. oryzae* AHU 7139 (parent) and five selected mutants $(\Delta mdlB, \Delta tglA, \Delta cutL, \Delta 1, \Delta 19)$ toward purified triolein (TG) and diolein (DG) from the solid culture products (CP) initiated with pH 4.0 and pH 6.5. Values are expressed in average value (n = 2).



Fig. 3. Free fatty acid (FFA) profiles in cheese model at 4 weeks.

C = curds only (control), P4 = parent AHU 7139 with initial culture pH 4, M4 = mutant AHU 7139 $\Delta mdlB$ with initial pH 4, T4 = mutant AHU 7139 $\Delta tglA$ with initial pH 4, P6.5 = parent AHU 7139 with initial pH 6.5, M6.5 = mutant AHU 7139 $\Delta mdlB$ with initial pH 6.5, T6.5 = mutant AHU 7139 $\Delta tglA$ with initial culture pH 6.5. Results of FFA level are expressed as the mean \pm SE (n = 3). The different letters indicate a significant difference between the samples using Tukey-Kramer test (p < 0.05)



Fig. 4. The proposed rancidity process in the ripened cheese caused by *Aspergillus* lipolysis. As triacylglycerol (TG) is the major lipid in milk, triacylglycerol lipase (TglA) hydrolyzes TG at any fatty acid position ((A) sn-1, 3 or (B) sn-2) and generates diacylglycerol (DG) and free fatty acids (FAs) during ripening. Subsequently, diacylglycerol lipase (MdlB and, to some extent, TglA) hydrolyzes DG and produces FFA. The predominant lipases (TglA or MdlB) at each hydrolytic step are shown in bold, although other genes might be involved. The excess amounts of free volatile fatty acids released by fungal lipolysis could cause rancidity.

Table S1. Primers used for gene expression analysis.				
Cono	Drimor	Second (52, 22)	Predicted amplified size (bp)	
Gene	sequence (5 - 5)		genomic DNA	cDNA
Triacylglycerol lipase (tglA)	Forward	CTTTCACGGAGCCCTTCCAC	057	765
(AO090003001507)	Reverse	CGTAAATTAGTTCGCAGCCGC	937	705
Mono-, di-acylglycerol lipase (mdlB)	Forward	TGAGTAGACCCTGCGAAGCAC	1024	021
(AO090701000644)R	Reverse	CGAATTAGCGCAATGGCAATCCAG	1024	921
Cutinase 1 (<i>cutL</i>)	Forward	GCTTTGCCCCAGGAAGAAT	087	924
(AO090005000029)	Reverse	ACTGACCTCCATCAAATGGGAG	987	034
A Q000012000600 (1)	Forward	GAGATTTGATCGCACTGATGGC	1002	964
A0090012000090 (1)	Reverse	GCCAAGCTTTTGGCTTTCAC	1005	804
A 0000701000602 (2)	Forward	CTACCTTAGCCCGCAAGTGC	1245	1038
A0090701000092 (2)	Reverse	CCACTAACAAACCATCCCTTCAAC		
A Q 0 0 0 0 2 0 0 1 4 2 2 (2)	Forward	GCCGTATGTCCTCTGCCTCTG	1362	1362
A0090003001432 (3)	Reverse	CCACAATCAAGGAATTAGCCGACG		
A Q000002001215 (4)	Forward	TGAGCTTTTCACCGTCTCCC	12(0	1269
A0090003001313 (4)	Reverse	ACACATTCGTCCAAACATGGC	1209	
A Q000001000142 (5)	Forward	GCTGACTTCGCAAGGGATATTC	1056	888
A0090001000143 (3)	Reverse	TCAAGGCTGGCAGAAGTCTC	1036	
A Q000010000610 (6)	Forward	CGGCACTGAAGTGTCTGCATAATC	1505	1368
A0090010000019 (0)	Reverse	TAGATGGTGGGGGTCTCTGGGAG		
A Q000012000102 (7)	Forward	AGTGCGATCAATTCCCTACCC	1420	1353
A0090012000105 (7)	Reverse	CCCACATAACTCCCGTTTGAG	1420	
A Q000124000015 (8)	Forward	CCTTCTGTCCCTTGGACCTG	1102	804
A0090124000015 (8)	Reverse	CAGTACTGTAGGTCAGCTTTCC	1102	894

, , , , , , , , , , , , , , , , , , ,			Predicted amplified	
Gene	Primer	Sequence (5' - 3')	size () genomic DNA	cDNA
A 0 0 0 0 1 0 0 0 0 1 7 0 (0)	Forward	CGGGCAGTTCCCATGTCATC	1244	1244
A0090103000172 (9)	Reverse	GGCACCACAAAGAAGCGAAAG	1544	1344
A Q0001200001(7 (10)	Forward	TCCCGACCTCACCAGGATAG	1000	004
A0090138000107 (10)	Reverse	CGATCTTTCTCAGCTATAATTCGGC	1009	894
A Q000028000214 (11)	Forward	CTGTTGCTTTGTATAGTGTCACG	1152	1065
A0090038000214 (11)	Reverse	TCTGTCACCCAGAGCAGATG	1155	1005
	Forward	AAGAGATACCCAACACCCAC	1121	057
A0090020000171 (12)	Reverse	CCCAGTGACCCGATGTATC	1151	937
A Q00002000600 (12)	Forward	GCACGCCAAAGACGCTTAATG	1500	1356
A0090020000009 (15)	Reverse	CACAATCTGACCACATCATTCGG	1388	
A Q000002000022 (14)	Forward	ACTGCGCTTTCCCAACAAAC	1327	1263
A0090005000022 (14)	Reverse	TCTATCACCCCGTCATTGCG		
A O 0 0 0 0 2 3 0 0 0 1 2 5 (1 5)	Forward	AACCACGCCACACCCATATC	1781	1644
A0090023000123 (13)	Reverse	GGGAAAGTGTTGGTTTAGACTGG		
10000138000014 (16)	Forward	CCCATATAACGCTCGAAACATC	1054	915
A0090138000014 (10)	Reverse	ACGGATAAGGGTTGTAACTTAGG		
A O 0 0 0 0 2 0 0 0 8 5 1 (17)	Forward	TCGACCAGCATTACTCCGC	1033	933
A0070003000031 (17)	Reverse	TAAACCCATGTGGTAGCCCC		
A O 090005000930 (18)	Forward	TGACCCAAGTTCACATCCGC	1190	1125
A0070003000730 (18)	Reverse	CATCCAAGACATTGATCCCGC		
A O 0 0 0 0 5 0 0 1 3 1 0 (1 0)	Forward	CAGCCCACCCCTTCAATAG	918	918
A0090003001319 (19)	Reverse	ACGGAACGATTCTTGACGAAAAC		
A O 0 9 0 0 5 0 0 1 6 0 2 (20)	Forward	GGTTGCTCGTGGTAGGAATG	425	284
A0090003001002 (20)	Reverse	CAGTCGGCCCAGAATGAGTC		
A O 0 9 0 0 2 3 0 0 7 1 7 (21)	Forward	GGTGCTGTATCTGAAAATGCG	- 449	361
A0070023000717 (21)	Reverse	ATCACAACGACCTTTCGCTG		
$\Delta \cap 0 0 0 0 0 3 0 0 8 3 0 (22)$	Forward	GCGCACCGATATCCCTATTAG	837	780
A0070003000037 (22)	Reverse	CACTCGACCTTCTTGTCATCC	057	

Table S2. Primers	s used for can	didate gen	e disruption.	
Use application		Name	Sequence (5' to 3')	
Knockout of	Front arm	LUpyrG	CGACTAAGCCACGATCTCGATCAT	
AO090011000868		LLpyrG	aaaggagtacgtatccaccactacCAATTGCCGCGAAAAATTAA	
(pyrG)	Rear arm	RUpyrG	GTAGTGGTGGATACGTACTC	
		RLpyrG	CTCCGTTGCGGATCTTGCTGCTTG	
	Clone check	cLpyrG	GAGTACGTATCCACCACTAC	
<i>pyrG</i> _forme	er part	PU	GTCCATATATCGAGGCAGGT	
		PLsplit	ATTGACCTACAGCGCACGC	
pyrG_latter	· part	PUsplit	CCGGTAGCCAAAGATCCCTT	
		PL	TCCTCATTTACTCCCGAGAT	
Disruption of	Front arm	LU1	CGACACGGAGAATTTCCCGATGTA	
AO090003001507		LL1	agacacctgcctcgatatatggacAAGTGGAAGGGCTCCGTGAA	
	Rear arm Clone check	RU1	gcagatctcgggagtaaatgaggaTTTACGATAAGGGCTCCATG	
		RL1	ACGCTTTTGTAAGAGCCAGCCCAC	
		cU1	TCCGAAGTCAGTAGATCGAC	
		cL1	AGCCACTTATTTCCGTGACC	
Disruption of	Front arm	LU2	GAGGGAAATTCTGGACGATTCTCG	
AO090701000644		LL2	agacacctgcctcgatatatggacCTTTGCCAGTGTGCTTCGCA	
	Rear arm	RU2	gcagatctcgggagtaaatgaggaCTTACATGATTTGGACGGAC	
		RL2	GCATCACTCGGCAATCCTACCTAA	
	Clone check	cU2	CTCGACTACGTCAGAGGAGC	
		cL2	CAACCACAGAATCTATCACC	
Disruption of	Front arm	LU3	CTGTCTTCTTTTATTCCGCTCACCAC	
AO090005000029		LL3	agacacctgcctcgatatatggacGAGGAAAGTGTTTTAGAAAGCG	
	Rear arm	RU3	gcagatctcgggagtaaatgaggaTCTGCTGCCTTGCGTCGGAT	
		RL3	TTGCTGAACCATGCCCTGCTATCG	
	Clone check	cU3	CAAGCTACTATGGTGTGGAT	
		cL3	GCAACCTCCACATCATCTCC	

Table S2. (contin	andidate gene disruption.				
Use application		Name	Sequence (5' to 3')		
Disruption of	Front arm	LU4	CTCTCTCGTGGAAGTATGTAAGCG		
AO090012000690		LL4	agacacctgcctcgatatatggacCAGTGCGATCAAATCTCAAAC		
	Rear arm	RU4	gcagatctcgggagtaaatgaggaATTCTCATCGCCATCACTA		
		RL4	CCAGGGCTCACTATTGAGGTATTG		
	Clone	cU4	GGGTTTAGCGAGATCTTATC		
	check	cL4	TATCATCAGGGTCTGGTAGA		
Disruption of	Front arm	LU5	TACCAAAGTGCCCGTCACCTCATT		
AO090005001319		LL5	agacacctgcctcgatatatggacTGTGATATCGATCACGGTTTC		
	Rear arm	RU5	gcagatctcgggagtaaatgaggaCAAATTCAAGTCAAGGCTATCG		
		RL5	GTGCTGAACAGTAGCCTCAATCCA		
	Clone	cU5	GCATTCAACAATGGCGATGC		
	check	cL5	TTAGCAGCACCATCTAGTCG		
Southern hybridization	Probe 1	003-1507 SBU	CCAACATACTGTTGTCACAC		
		003-1507 _SBL	TCAACAGTACGGTTTCACTC		
	Probe 2	701-0644 _SBU	CTTACATGATTTGGACGGAC		
		701-0644 _SBL	GTGTAGTGTGCTTGGCCGAC		
	Probe 3	005-0029 _SBU	TCTCCTCTCTCTGTGTCT		
		005-0029 _SBL	CTCGCTTCGGATTGTATGAT		
	Probe 4	012-0690 SBU	CGTGAGCGACATGACTAAGT		
		012-0690 SBL	CCGTTGTTGAGCTGGAATGT		
	Probe 5	005-1319 SBU	GTGGATCAAATCGGGATGAA		
		005-1319 SBL	GAATCTTGGACAGGAATCGT		
Tails of primers used for overlapping in fusion PCR are shown in lower case.					



Fig. S1. RNA integrity check by non-denaturing agarose gel electrophoresis. Total RNA was isolated from *A. oryzae* AHU 7139 inoculated on whey solid culture adjusted to pH 4.0 and pH 6.5. +con = positive control of 28S and 18S RNA bands from bovine endometrial stromal cells.



Fig. S2. Construction of the DNA fragment for pyrG (AO090011000868) knockout in *A. oryzae* AHU 7139. The 2035 bp-long DNA fragment was constructed for the knockout. Primers used for the construction and clone check are shown as colored and black arrows, respectively.

M R A 1 2 3 M



Fig. S3. Clone check of the *pyrG* knockout mutant isolated after transformation. Transformant clone Nos. 1-3 are shown. Lane M: 1 kb DNA ladder marker [1-10 kb], lane R: RIB40 strain as a negative control, lane A: AHU 7139 strain as a negative control. Amplified DNA size: Positive clone, 1.0 kb; Negative clone, 2.1 kb. The clone No. 3 in red was considered a positive homokaryon, which was named AHU 7139 [pyrG⁻].



Fig. S4A. Construction of the DNA fragment for disrupting AO090003001507 in *A. oryzae* AHU 7139 [pyrG⁻]. The 2048 bp-long and 2375 bp-long DNA fragments were constructed for the disruption. Primers used for the construction and clone check are shown as colored and black arrows, respectively.



Fig. S4B. Construction of the DNA fragment for disrupting AO090701000644 in *A. oryzae* AHU 7139 [pyrG⁻]. The 2021 bp-long and 2377 bp-long DNA fragments were constructed for the disruption. Primers used for the construction and clone check are shown as colored and black arrows, respectively.



Fig. S4C. Construction of the DNA fragment for disrupting AO090005000029 in *A. oryzae* AHU 7139 [pyrG⁻]. The 2035 bp-long and 2392 bp-long DNA fragments were constructed for the disruption. Primers used for the construction and clone check are shown as colored and black arrows, respectively.



Fig. S4D. Construction of the DNA fragment for disrupting AO090012000690 in *A. oryzae* AHU 7139 [pyrG⁻]. The 2031 bp-long and 2403 bp-long DNA fragments were constructed for the disruption. Primers used for the construction and clone check are shown as colored and black arrows, respectively.



Fig. S4E. Construction of the DNA fragment for disrupting AO090005001319 in *A. oryzae* AHU 7139 [pyrG⁻]. The 2045 bp-long and 2408 bp-long DNA fragments were constructed for the disruption. Primers used for the construction and clone check are shown as colored and black arrows, respectively.



Fig. S5A. Southern hybridization analysis of AO090003001507 disruptant constructed from AHU 7139. 1: AHU 7139; 2: AO090003001507 disruptant.



Α



Fig. S5B. Southern hybridization analysis of AO090701000644 disruptant constructed from AHU 7139. 1: AHU 7139; 2: AO090701000644 disruptant.



С

Fig. S5C. Southern hybridization analysis of AO090005000029 disruptant constructed from AHU 7139. 1: AHU 7139; 2: AO090005000029 disruptant.



Fig. S5D. Southern hybridization analysis of AO090012000690 disruptant constructed from AHU 7139. 1: AHU 7139; 2: AO090012000690 disruptant.



Fig. S5E. Southern hybridization analysis of AO090005001319 disruptant constructed from AHU 7139. 1: AHU 7139; 2: AO090005001319 disruptant.



Ε

Fig. S6. Thin layer chromatography of the purified lipids. M = control monoglyceride, D = control diglyceride, T = control triglyceride (tributyrin), PD = purified commercial diolein using silica gel chromatography, PT = purified commercial triolein using silica gel chromatography, CD = commercial diolein, CT = commercial triolein. Migration of the sample on the silica gel plate was carried out using chloroform : methanol : formic acid : DI water (45:20:2.5:1), stained with 0.05% primulin dissolved in acetone : water (4:1 v/v), and observed under UV light.



Fig. S7. Agarose gel (1%) electrophoresis of PCR using mRNA from CP with initial pH 4.0 (A) and pH 6.5 (B) as a template with all candidate lipase gene primers. M = DNA ladder marker, lane no.1-22 = annotated lipase no. 1-22, m = mdlB, t = tglA, and c = cutL gene. Confirming no gDNA in the mRNA sample.



Fig. S8. Growth and appearance of *A. oryzae* AHU 7139 on the WPC media incubated at 20°C for 7 days. The initial culture pH were of (A) pH 4.0 and (B) pH 6.5.