1	Title

2	Culture-based analysis of fungi in leaves after the primary and secondary fermentation processes
3	during Ishizuchi-kurocha production and lactate assimilation of P. kudriavzevii
4	
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24 Abbreviations¹

¹ CFU	Colony forming unit
ITS	Internal transcribed spacer
LAB	Lactic acid bacteria
NBRP	National Bio-Resource Project
PCR	Polymerase Chain Reaction
PDA	Potato dextrose agar
YNB	Yeast nitrogen base

26 Abstract

27 Ishizuchi-kurocha is a Japanese traditional fermented tea that is produced by primary aerobic and 28 secondary fermentation steps. The secondary fermentation step of Ishizuchi-kurocha is mainly 29 mediated through lactic acid bacteria. Here, we performed quantitative analyses of the culturable 30 fungal communities at each step and identified several morphologically representative fungal isolates. While filamentous fungi (median, 3.2×10^7 CFU/g sample) and yeasts (median, 3.7×10^7 31 CFU/g) were both detected after the primary fermentation step, only yeasts (median, 1.6×10^7 32 CFU/g) were detected in the end of the secondary fermentation step, suggesting that the fungal 33 34 community in tea leaves are dramatically changed between the two steps. Pichia kudriavzevii and 35 Pichia manshurica, the prevalent fungal species at the end of the secondary fermentation step, grew 36 well in exudate from the secondary fermentation step. P. kudriavzevii also grew well in media 37 containing D- or L-lactate as the sole carbon source. The growth of the disruptant of cyb2A encoding a cytochrome b₂ lactate dehydrogenase in P. kudriavzevii was severely impaired on medium 38 39 supplemented with L-lactate, but not D-lactate, suggesting that Cyb2Ap plays a crucial role in the use 40 of L-lactate, and P. kudriavzevii efficiently uses both L- and D-lactate as carbon sources. Thus, lactate 41 assimilation seems to be a key phenotype to become a prevalent species in the secondary 42 fermentation step, and Cyb2Ap has a pivotal role in L-lactate metabolism in P. kudriavzevii. Further 43 understanding and engineering of *P. kudriavzevii* and *P. manshurica* will contribute to the control of

lactic acid bacteria fermentation during the fermented tea production and also to other industrial
uses. **Keywords**: Post-fermented tea, *Aspergillus* section *Nigri, Pichia kudriavzevii, Pichia manshurica, cyb2A cyb2A*

1. Introduction

54	Tea, which is among the most common beverages worldwide, is produced from the leaves
55	of the tea tree, especially two prominent varieties, Camellia sinensis var. sinensis and C. sinensis var.
56	assamica. While the processing of green tea does not involve fermentation, the production of four
57	types of traditional teas (Ishizuchi-kurocha, Goishi-cha, Awa-bancha, and Batabata-cha) in Japan, as
58	well as Pu-erh tea (Yunnan, China), Lahpetso (Myanmar), and Miang (northern Thailand) include
59	post-fermentation processes (Horie et al., 2017).
60	The manufacturing processes among Japanese post-fermented teas have important
61	differences (Horie et al., 2017). For example, the production of Batabata-cha, similar to Pu-erh tea,
62	includes an aerobic fermentation process by fungi. On the other hand, Awa-bancha production
63	includes a fermentation process by lactic acid bacteria (LAB). Different from these post-fermented
64	teas, Ishizuchi-kurocha and Goishi-cha include two fermentation steps. The primary fermentation
65	step is aerobic, in which filamentous fungi play a pivotal role, while the secondary fermentation step
66	is driven by Lactobacilli as the major contributors (Horie et al., 2017).
67	The constituents of the fungal community of Pu-erh tea have been intensely investigated.
68	Zhao et al. (2010) showed that the dominant filamentous fungal genus was Aspergillus, especially
69	Aspergillus niger. More recently, Zhang et al. (2016a) reported that the dominant species in tea aged
70	through a high-temperature pile-fermentation process were A. niger, A. fumigatus, Rhizomucor

71	pusillus, and Rasamsonia emersonii. Similarly, culture-independent analysis conducted by Zhang et
72	al. (2016b) found that the most abundant fungal species in raw or ripened Pu-erh tea were
73	Aspergillus sp., including A. niger, and Blastobotrys adeninivorans. Although Tamura et al. (1994)
74	characterized the microbial communities in the fermentation steps during the production of
75	Ishizuchi-kurocha, Goshi-cha, and Awa-bancha, the fungi were only identified at the genus level,
76	which included Aspergillus, Penicillium, and Mucor. Following that publication, Okada et al. (1996)
77	isolated the filamentous fungi A. fumigatus, A. niger, Penicillium sp., and Scopulariopsis brevicaulis,
78	and the yeast Debaryomyces hansenii from the primary and secondary fermentation steps,
79	respectively, and identified the isolated molds by morphological observation, but not genetic
80	identification.
81	LAB are major players in the secondary fermentation step (Horie et al., 2019), and
82	changes in culture or media conditions such as metabolite production including lactate and pH
83	changes from the primary fermentation step suggest that the prevalent fungal species in the
84	secondary fermentation step exhibit distinct advantages over the prevalent species in the primary
85	
	fermentation step. Ueno showed that the use of lactate by Candida glabrata was important for its
86	fermentation step. Ueno showed that the use of lactate by <i>Candida glabrata</i> was important for its adaptation in a mouse intestine (Ueno et al., 2011), suggesting this utility to be an important factor
86 87	fermentation step. Ueno showed that the use of lactate by <i>Candida glabrata</i> was important for its adaptation in a mouse intestine (Ueno et al., 2011), suggesting this utility to be an important factor for prevailing. Hence, we focused on lactate utility of a prevalent fungal species in the secondary

89 the conversion from L-lactate to pyruvate and the reverse reaction. Cyb2p is essential for the

90 utiliz	ation of L-lactate	by Sa	ccharomyces	cerevisiae	(Guiard,	1985)).
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- 91 In this study, the culturable fungal communities in the primary and secondary fermentation
- 92 steps of Ishizuchi-kurocha processing were quantitatively analyzed. In parallel, the fungal species
- 93 involved in both fermentation steps were identified. The results showed that *Pichia kudriavzevii*
- 94 (Issatchenkia orientalis, NCBI:txid4909) and P. manshurica (NCBI:txid121235) were the prevalent
- 95 fungal species from tea leaves during the secondary fermentation step. Also, disruptants of *cyb2A*
- 96 and homologues of *P. kudriavzevii* were prepared to assess fungal growth in media with L- or
- 97 D-lactate as the sole carbon source.
- 98

99 2. Materials and methods

100 **2.1. Tea leaf samples**

Tea leaf samples from the end points of the primary fermentation step (26 samples obtained at four to six days after the fermentation) and secondary fermentation step (11 samples obtained at 12 to 25 days after the fermentation) produced between June to October in 2015, 2016, and 2017 were kindly provided from three manufacturers of Ishizuchi-kurocha in Saijo City, Ehime Prefecture, Japan.

2.2. Colony counting and isolation of fungal strains

108	To determine the colony forming units (CFU) of fungi, 10 g of tea leaves from each
109	sample were weighed into filtrate Bag II (Eiken Chemical Co., Ltd., Tokyo, Japan), suspended in a
110	9-fold weight of sterilized water, and homogenized at 230 rpm for 1 min with a Stomacher 400
111	Circulator paddle blender (Seward Ltd., Worthing, UK). Then, 500 μ L of each stomached sample
112	were serially diluted by ten-fold (10^{-1} to 10^{-6}) with sterilized water. Each $100-\mu$ L sample was loaded
113	on plates containing potato dextrose agar (PDA) supplemented with 100 μ g/mL of chloramphenicol,
114	and incubated at room temperature for 2 days. Plates in which 15-200 colonies appeared were used
115	for counting.
116	Several representative colonies determined morphologically from each sample were
117	transferred to fresh PDA slants for preparation of genomic DNA. A. niger YM1 and Meyerozyma
118	caribbica YM9 from tea leaves of the primary fermentation, and P. kudriavzevii (P. kudriavzevii)
119	YM48 and <i>P. manshurica</i> YM63 from tea leaves of the secondary fermentation that were isolated in
120	2015 were used for growth analyses, as described below. Saccharomyces cerevisiae S288C, provided
121	by the National Bio-Resource Project (NBRP) of the Ministry of Education, Culture, Sports, Science
122	and Technology of Japan was also used in this study.
100	

2.3. Genomic DNA preparation and Sanger sequencing

125	Genomic DNA preparation, PCR amplification, and Sanger sequencing were performed as
126	described previously (Onishi et al., 2017; Toyotome et al., 2019).
127	
128	
129	2.4. Preparation of yeast nitrogen base (YNB) liquid media supplemented with various carbon
130	sources and filtered exudate from a secondary fermentation sample
131	YNB with ammonium sulfate (MP Biomedicals, LLC., Santa Ana, CA, USA) liquid media
132	containing 2% (wt/vol) glucose (YNBG), 2% (vol/vol) L-lactate (YNBLL), or D-lactate (YNBDL)
133	were prepared. YNBLL and YNBDL media were adjusted to pH 7.0 or pH 4.0 with 4 M NaOH
134	before autoclaving.
135	Air was removed from leaves with weight stone during the secondary fermentation step.
136	Exudate (acidity around 10%) was recovered from a bucket after the secondary fermentation and
137	filtered with a 0.45-µm filter. The filtered exudate was stored at -25° C until use.
138	
139	2.5. Growth analysis in liquid media and exudate
140	The spores of A. niger were collected in 1 mL of 0.9% NaCl solution. P. kudriavzevii, P.
141	manshurica, and M. caribbica were cultured in YNBG at 37°C overnight, then the cells were

142	collected and re-suspended in 0.9% NaCl solution. The spore or yeast concentration in each
143	suspension was determined by cell-counting with an improved Neubauer hemocytometer.
144	The concentration of each suspension was adjusted to 4×10^5 cells/mL in each liquid
145	medium. A 100- μ L aliquot of each cell suspension was added to the wells of Round- and
146	Flat-Bottomed 96-Well Test Plates with lid (AS ONE Co., Osaka, Japan) for endpoint analysis and
147	for growth curve analysis, respectively. In YNBG, YNBLL, or YNBDL, the fungal cells were
148	cultured at 37°C. For endpoint analysis, the AnaeroPack-MicroAero cultivation system (Mitsubishi
149	Gas Chemical Co., Inc., Tokyo, Japan) was used, and after culture for 48 h, the optical density (OD)
150	in each well was determined with a GENios Pro Multifunction Microplate Reader (Tecan Group Ltd.,
151	Männedorf, Switzerland). For the growth curve analysis, plates covered with a clear plate seal was
152	incubated at 37°C in the microplate reader, and OD at 620 nm was determined every hour until 72
153	hours after inoculation. Wells without fungal cells were used as blank samples.
154	2.6. Gene disruption in <i>P. kudriavzevii</i>
155	Gene disruption in <i>P. kudriavzevii</i> was performed as described below using the primers
156	shown in Table A.1. Plasmids pHM874-8 (NBRP ID: BYP8929) and pHM785-1 (NBRP ID:
157	BYP8919) provided by NBRP were used as templates of the hygromycin B and G418/kanamycin
158	resistance genes, respectively. The upstream and downstream flanking genomic regions at 1 kb from
159	the target gene cyb2A (JL09_g341 in P. kudriavzevii SD108) and the homologue cyb2B (JL09_g1432

160	in <i>P. kudriavzevii</i> SD108) and <i>cyb2C</i> (JL_09_g3241) were fused by the double-joint PCR method
161	(Yu et al., 2004) after amplification of the resistance genes with each specific primer pair.
162	The fused constructs were transformed into P. kudriavzevii YM48 using an electroporation
163	method (Zemanova et al., 2004) with slight modifications. Yeast cells were pretreated with 10 mM
164	dithiothreitol before electroporation and electric pulses were applied using a Gene Pulser (Bio-Rad
165	Laboratories, Hercules, CA, USA) and a 2-mm cuvette (Nepa Gene Co., Ltd., Chiba, Japan) at 2.5
166	kV, 200 $\Omega,$ and 25 $\mu F.$ After electroporation, transformants were selected with 400 $\mu g/mL$
167	hygromycin B (FUJIFILM Wako Pure Chemical Co., Osaka, Japan) or 400 μ g/mL hygromycin B
168	and 400 μ g/mL G418 (FUJIFILM Wako Pure Chemical Co.). The colonies were picked-up and the
169	remaining gene copies were determined using a relative quantification by real-time PCR. The
170	primers listed in Table A.1, THUNDERBIRD SYBR qPCR Mix (Toyobo Co., Ltd., Osaka, Japan),
171	and LightCycler 480 Instrument II (Roche Diagnostics, Rotkreuz, Switzerland) were used for
172	real-time PCR. Briefly, the P. kudriavzevii YM48 genome was serially diluted and standard curves
173	for a housekeeping gene <i>act1</i> and each <i>cyb2</i> gene were determined from cycle threshold (Ct) values.
174	Ct values of disruptants based on the determined standard curves were used for relative
175	quantification. As shown in Fig. A.1, quantification of gene copies indicated that each cyb2 gene in
176	single disruptants was half of that of act1 gene copies, indicating that a single copy of each gene was

disrupted. Since no gene copy was present in each double disruptant, each *cyb2* gene was not

178 detected (Fig. A.1).

179

180 **3. Results**

3.1. Fungal community shifts from molds and yeasts in the primary fermentation step to LAB
 and yeasts in the secondary fermentation step

183 Culturable fungal cells were quantified at the end of each fermentation step (Fig. 1A). The

184 median of fungal cell number was 8.8×10^7 (range, 2.2×10^6 to 7.9×10^8) CFU/g sample in tea leaf

- samples from the primary fermentation. In the primary fermentation, yeasts and molds were
- observed as culturable cells (Fig. 1B). Molds were detected in every sample. On the other hand,
- 187 yeasts were not detected in 3 (11.5%) of 26 samples, suggesting that molds are essential to the
- aerobic fermentation step, while yeasts are auxiliary. In contrast to the primary fermentation samples,
- only yeasts were detected in all secondary fermentation samples (median number, 1.6×10^7 CFU/g
- sample; the range was from 8.5×10^4 to 4.8×10^8 CFU/g sample), suggesting that yeasts support the
- 191 process with LAB in the secondary fermentation.
- 192 The culturable fungal species in the primary fermentation step were identified
- 193 morphologically to elucidate the abundance (Fig. 1C). Aspergillus Section Nigri was the major mold
- in 23 (88.5%) of 26 primary fermentation samples. Internal transcribed spacer (ITS) sequencing of

195	the picked-up colonies confirmed the presence of A. niger sensu lato, but not A. tubingensis. In 2
196	(7.7%) of 26 samples, A. fumigatus was the major mold species. Zygomycetes (e.g., Mucor and
197	Rhizopus) were found in some tea leaf samples. Overall, 43 yeast strains were obtained from
198	fermented leaf samples after the primary fermentation including Candida intermedia (2 isolates),
199	Clavispora lusitaniae (2), Cutaneotrichosporon dermatis (12), Cyberlindnera saturnus (1),
200	Debaryomyces hansenii (2), Dipodascus geotrichum (2), Meyerozyma caribbica (4),
201	Rhodosporidiobolus ruineniae (5), Pichia kudriavzevii (2), and Pichia manshurica (11). On the other
202	hand, yeasts (43 isolates) isolated from the secondary fermentation samples except for one isolate
203	(Cutaneotrichosporon dermatis) were P. kudriavzevii (16) and P. manshurica (26). These data
204	suggest that A. niger s.l. and the yeasts P. kudriavzevii and P. manshurica play important roles in the
205	primary and the secondary fermentation steps during Ishizuchi-kurocha production, respectively.
206	
207	3.2. P. kudriavzevii have advantages in lactate-containing media
208	To clarify the reason why mycological abundance was shifted from the primary
209	fermentation to secondary fermentation, we examined the growth of five fungal species in filtered
210	exudate of the secondary fermentation. A. niger YM1 and Meyerozyma caribbica YM9 isolated from
211	the primary fermentation samples, P. kudriavzevii YM48 and P. manshurica YM63 from the
212	secondary fermentation samples, and Saccharomyces cerevisiae S288C were used in this experiment.

213	As shown in Fig. 2 and Fig. A.2, <i>P. kudriavzevii</i> and <i>P. manshurica</i> grew better in the exudate from
214	leaves in the secondary fermentation than the three other species analyzed here, indicating that P.
215	kudriavzevii and P. manshurica were the prevalent species and have advantages in the environment
216	of the secondary fermentation.
217	Because the exudate and fermented leaves contained large amounts of lactate produced by
218	LAB, we assumed that efficient lactate assimilation by <i>P. kudriavzevii</i> and <i>P. manshurica</i> may be an
219	advantage in the secondary fermentation. LAB produce large quantities of L-lactate and D-lactate,
220	and lower the pH in the secondary fermentation. To examine the utilization of lactate, P. kudriavzevii
221	and <i>P. manshurica</i> were cultured in YNB supplemented with L- or D-lactate (pH 7.0 or 4.0) and
222	YNB supplemented with glucose as a control. As shown in Fig. 3 and Fig. A.3, P. kudriavzevii
223	rapidly propagated in liquid media supplemented with lactate. Although P. manshurica propagated in
224	L-lactate containing media (Fig. A3), its growth was inefficient as compared with <i>P. kudriavzevii</i> .
225	Further, the propagation of <i>M. caribbica</i> , <i>A. niger</i> , or <i>S. cerevisiae</i> in lactate containing media was
226	slow (Fig. A3).
227	
228	3.3 Cyb2Ap of <i>P. kudriavzevii</i> has a pivotal role in L-lactate assimilation
229	We focused on the ability of the <i>cyb2</i> genes to assimilate lactate in <i>P. kudriavzevii</i> . CYB2
230	of S. cerevisiae (Guiard, 1985) and Cyb2p of C. glabrata (Ueno et al., 2011) are known as L-lactate

231	dehydrogenases that convert L-lactate to pyruvate. A Protein BLAST search identified three
232	homologues of S. cerevisiae CYB2 in P. kudriavzevii strain SD108, namely JL09_g341 (Cyb2Ap),
233	JL09_g1432 (Cyb2Bp), and JL09_g3241 (Cyb2Cp), with Cyb2Ap having the highest homology.
234	Next, disruptants of <i>P. kudriavzevii</i> YM48 were prepared. <i>P. kudriavzevii</i> was a diploid and harbored
235	two copies of each cyb2 gene. Elimination of both copies were confirmed by real-time PCR (Fig.
236	A.1). As shown in Fig. 4, <i>P. kudriavzevii</i> $\Delta \Delta cyb2A$, but not $\Delta \Delta cyb2B$ or $\Delta \Delta cyb2C$, strongly impaired
237	cell growth in YNBLL medium, suggesting that Cyb2Ap has a pivotal role in L-lactate assimilation.
238	In contrast, all disruptants grew in YNBDL medium, suggesting that the three Cyb2 homologues are
239	dispensable for D-lactate assimilation.
240	
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 241 242 243 244 245 246 	 4. Discussion In this study, we quantified and qualified the fungal communities in the primary and secondary fermentation steps of Ishizuchi-kurocha production. Culturable filamentous fungi (median, 3.2 × 10⁷ CFU/g sample) and yeasts (median, 3.7 × 10⁷ CFU/g) were observed in the primary fermentation step, indicating that the numbers of filamentous fungi and yeasts were comparable (Fig. 1). In contrast, only yeasts were isolated from leaf samples of the secondary fermentation step.
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249	species can be identified, and molds may underestimate because of the tubular structure of hyphae.
250	Nevertheless, more than 100 strains were obtained using the culture-based analysis from the primary
251	and secondary fermentation samples and the genome data of <i>P. manshurica</i> YM63 have recently
252	been released. Although it seems that other weaknesses may be a bias of the primer pair and
253	remaining DNA derived from the primary fermentation step in the secondary fermentation step.
254	However, the mycobiome analyzes by amplicon sequencing with next-generation sequencers are
255	useful to obtain an overview of fungal flora that should be conducted in near future. The information
256	on prevalent fungal species both fermentation steps serve as a good pilot guide of further
257	mycobiome analyzes. A. niger s.l. was prevalent in most samples of the primary fermentation step.
258	In an earlier study, <i>Rhizomucor variabilis</i> was isolated from Ishizuchi-kurocha (Yanai et al., 2008).
259	In the present study, R. variabilis was not detected, although some zygomycetes species, including
260	Rhizopus sp. and Mucor sp., were isolated from the primary fermentation samples. A. niger was
261	reported as the major fungal species during the fermentation process of other teas, such as Pu-erh tea
262	(Abe et al., 2008; Xu et al., 2005) and Goishi-cha (Okada et al., 1996), suggesting that the fungal
263	community has changed over the period of several years from Rhizomucor-type to A. niger-type,
264	similar to that of Pu-erh tea and Goishi-cha. A. fumigatus was dominant in two primary fermentation
265	samples. Since fungal communities in the primary fermentation are unstable, as shown in Figure 1C,
266	the surveillance of fungal communities or the use of A. niger strains as a starter might prove useful

267	for producing more stable products. Additionally, the opportunistic pathogen A. fumigatus was also
268	found in few samples. A preliminary experiment using A. oryzae as a secure starter showed A. oryzae
269	to be a prevalent species at the end of the primary fermentation step. Currently, we possess and
270	identified 71 A. niger isolates and believe that the data and isolates obtained in this study will
271	contribute to production of more stable Ishizuchi-kurocha end samples.
272	In contrast to the primary fermentation step, P. kudriavzevii and P. manshurica were
273	isolated from the secondary fermentation step. Of the LAB, Lactobacillus plantarum was the most
274	prevalent strain and a major player in the fermentation step (Horie et al., 2018). As reported in a
275	review article by (Dalié et al., 2010), LAB and associated products can potentially control
276	filamentous fungi, as some L. plantarum strains show antifungal activity (Russo et al., 2017), due to
277	the antifungal potency of lactic acid. As shown in Fig. 2, the growth of A. niger was not detected in
278	the cell-free exudate from the secondary fermentation, which is consistent with the findings of an
279	earlier study (Russo et al., 2017). Propagation of the yeasts S. cerevisiae and M. caribbica was also
280	not detected in the exudate for 48 hours after inoculation. In contrast, P. kudriavzevii and P.
281	manshurica were growable in the exudate. We hypothesized that this difference was due to the
282	ability to assimilate lactic acid, rather than an antifungal effect. As shown in Fig. 3, examination of
283	these yeast species and A. niger using synthetic media supplemented with different carbon sources
284	indicated that <i>P. kudriavzevii</i> rapidly propagated in media supplemented with lactic acid. Notably,

285	pH had no effect on growth. Conversely, P. manshurica propagated inefficiently on media
286	supplemented with lactic acid compared with P. kudriavzevii. It is noteworthy that other
287	compositions derived from tea leaves and metabolites produced by LAB might inhibit other fungi
288	found in the primary fermentation and promote <i>P. manshurica</i> and <i>P. kudriavzevii</i> . The growth of <i>P</i> .
289	manshurica was promoted particularly in the exudate of the secondary fermentation. In a further
290	study, we will examine those components that promote this growth.
291	The ability of lactate assimilation was examined with the use of $cyb2$ gene disruptants. A
292	homologue of CYB2, a cyb2A disruptant of P. kudriavzevii, severely impaired growth on YNBLL,
293	but not YNBDL. These data indicate that <i>cyb2A</i> has a pivotal role in L-lactate assimilation. DLD1 is
294	the major mitochondrial D-lactate dehydrogenase in S. cerevisiae, which oxidizes D-lactate to
295	pyruvate. A homologue was also found in <i>P. kudriavzevii</i> , suggesting that the enzyme has a role in
296	D-lactate assimilation. Although we attempted the disruption of the gene, but no <i>dld1</i> disruptants
297	were obtained. Therefore, further studies are warranted to determine whether <i>dld1</i> in <i>P. kudriavzevii</i>
298	is important for assimilating D-lactate. P. manshurica is also a major fungus in the secondary
299	fermentation and common in fermentations and food spoilage. P. kudriavzevii and P. manshurica are
300	broadly used for industrial applications, especially food production(Douglass et al., 2018). The P.
301	kudriavzevii and P. manshurica strains isolated in this study may provide further useful resources as
302	starters for the secondary fermentation. We recently sequenced and released the genome data of P.

303	manshurica YM63 isolated from the secondary fermentation step. Further understanding and
304	engineering of lactate metabolism in <i>P. kudriavzevii</i> and <i>P. manshurica</i> will contribute to not only
305	the control of LAB fermentation during the fermented tea production, but also other industrial uses.
306	In summary, we elucidated the culturable fungal communities and the shift between two
307	fermentation steps during Ishizuchi-kurocha production. These results are useful reference data to
308	produce Ishizuchi-kurocha stably and securely as well as to extend current knowledge. Our
309	identification of the efficient propagation of <i>P. manshurica</i> and <i>P. kudriavzevii</i> in the secondary
310	fermentation environment presents advantageous information about the fermentation steps. We also
311	determined the gene cyb2A to play a pivotal role in L-lactate assimilation in P. kudriavzevii.
312	
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390	Fig. 1. The amount of culturable fungi in fermented tea leaf samples. CFUs per gram of total
391	fungi (A) and molds and yeasts (B) in fermented leaf samples are shown in the box-plot graphs (C).
392	The percentages of A. niger, A. fumigatus, zygomycetes, and yeasts in leaf samples from each
393	primary fermentation step. The numbers shown below the graph indicate the manufacturing years of
394	each sample.
395	
396	Fig. 2. Fungal growth in exudates from a secondary fermentation sample. The bars show the

- 397 optical density (OD) at 620 nm after culturing for 96 hour under microaerobic conditions. Pk, *Pichia*
- 398 kudriavzevii; Pm, Pichia manshurica; Mc, Meyerozyma caribbica; Sc, Saccharomyces cerevisiae;
- 399 An, Aspergillus niger.
- 400

401 Fig. 3. Fungal growth in media supplemented with glucose or lactate. The bars show the OD at
402 620 nm after culturing for 48 hours under microaerobic conditions. Filled bars, YNBG; gray bars,
403 YNBLL (pH 7); open bars, YNBLL (pH 4); hatched bars, YNBDL (pH 7); dotted bars, YNBDL (pH
404 4). Pk, *Pichia kudriavzevii*; Pm, *Pichia manshurica*; Mc, *Meyerozyma caribbica*; Sc, *Saccharomyces*405 *cerevisiae*; An, *Aspergillus niger*.

407 Fig. 4. Fungal growth of cyb2A, cyb2B, or cyb2C disruptant in media supplemented with

- 408 glucose or lactate. The bars show the OD at 620 nm after culturing for 48 hours under microaerobic
- 409 conditions. Filled bars, YNBG; gray bars, YNBLL (pH 7); open bars, YNBLL (pH 4); hatched bars,
- 410 YNBDL (pH 7); dotted bars, YNBDL (pH 4).
- 411

412 Fig. A.1. Quantification of each cyb2 gene to confirm disruption. Each single disruptant ($\Delta cyb2$)

413 possesses a half copies of each *cyb2* compared with the number of *act1* copies. Each *cyb2* gene was

- 414 not detected in each double disruptant. ND, not detected.
- 415

416 Fig. A.2. Growth curves of Pichia kudriavzevii (Pk), Pichia manshurica (Pm), Saccharomyces

417 cerevisiae (Sc), Meyerozyma caribbica (Mc), and Aspergillus niger (An) in exudate from the

- 418 secondary fermentation step. Those curves were drawn by averaging the data at each point of time
- 419 across the two (Pk) or three (others) replicates.

420

421 Fig. A.3. Growth curves of Pichia kudriavzevii (Pk), Pichia manshurica (Pm), Saccharomyces

422 cerevisiae (Sc), Meyerozyma caribbica (Mc), and Aspergillus niger (An) cultured in YNBG, YNBLL

423 (pH 7), YNBLL (pH 4), YNBDL (pH 7), or YNBDL (pH 4). Those curves were drawn by averaging

424 the data at each point of time across the three replicates.

Table A.1. Primers used for this study

Name	Sequence (5' to 3')	Purposes and reference		
Primers for species	identification			
ITS4	TCCTCCGCTTATTGATATGC	Primer to amplify and sequence ITS region. (White et al., 1990)		
ITS5	GGAAGTAAAAGTCGTAACAAGG	Primer to amplify and sequence ITS region. (White et al., 1990)		
NL1	GCATATCAATAAGCGGAGGAAAAG	Primer to amplify and sequence partial region of 28S rDNA. (Kurtzman and Robnett, 1997)		
NL4	GGTCCGTGTTTCAAGACGG	Primer to amplify and sequence partial region of 28S rDNA. (Kurtzman and Robnett, 1997)		
Primers for cyb2A d	isruption			
IoCyb2a-upF	ACTTGTTTTCCTCTTTCCCCTAGAG	Forward primer to amplify cyb2A upstream region in P. kudriavzevii.		
IoCyb2a-upR	GGAGGGTAFTCTGGGCCTCCATGTCTGTGTGCAACTAGGTTTATGTGGAG	Reverse primer to amplify cyb2A upstream region in <i>P. kudriavzevii</i> with 25-bp 5'-terminal region of hygromycin B resistance gene amplicon or G418/kanamycin resistance gene amplicon.		
IoCyb2a-hygroRdownF	CTCGTCCGAGGGCAAAGGAATAGGTTTGTTGGAAATATATTATTCATAAAGGC	Forward primer to amplify cyb2A downstream region in <i>P. kudriavzevii</i> with 25-bp 3'-terminal region of hygromycin B resistance gene amplicon.		
IoCyb2a-kanRdownF	CGTATGTGAATGCTGGTCGCTATACTGTTGTTGGAAATATATTATTCATAAAGGC	Forward primer to amplify <i>cyb2A</i> downstream region in <i>P. kudriavzevii</i> with 25-bp 32 terminal region of G418/congressing center amplicon		
IoCvb2a-downR	TTAAACAACCCAACACAGACCATTC	Reverse primer to amplify <i>cvb2A</i> downstream region in <i>P kudriavzevii</i>		
Primers for cvb2B d	isruntion	te onse primer to uniprir y cyto21 downstream region in 1. martur22/m.		
InCyb2b-unF	CGTGCGCCTGGTCGATETICACCTTCTTTG	Forward primer to amplify wh? Bunctream region in P kudriavravii		
IoCyb2b-upR	GGAGGGTAFTCTGGGGCCTCCATGTCTTTGAAGGTAFTAFCACTGCTGTTG	Reverse primer to amplify <i>cyb2b</i> upstream region in <i>P. kudriavzevii</i> with 25-bp 5'-terminal region of hygromycii B resistance gene amplicon or G418/kanamycin revietnos enna emplicon		
IoCyb2b-hygroRdownF	CACTCGTCCGAGGGCAAAGGAATAGGTGTCTGATATTTGCTAAATTGAAATGAAC	Forward primer to amplify cyb2B downstream region in P. kudriavzevii with 25-bp 3'-terminal region of hygromycin B resistance gene amplicon.		
IoCyb2b-kanRdownF	CGTATGTGAATGCTGGTCGCTATACTGGTCTGATATTTGCTAAATTGAAATGAAC	Forward primer to amplify cyb2A downstream region in P. kudriavzevii with 25-bp 3'-terminal region of G418/kanamycin resistance gene amplicon.		
IoCyb2b-downR	TCCTGATCTCACAGCACATTTCTAC	Reverse primer to amplify cyb2A downstream region in P. kudriavzevii.		
Primers to amplify	selection marker gene cassette			
TEFp-F	GACATGGAGGCCCAGAATACC	Forward primer to amplify hygromycin B resistance gene in pHM874-8 or G418/kanamycin resistance gene in pHM785-1.		
Hygro-R	ACCTATTCCTTTGCCCTCGGACGA	Reverse primer to amplify hygromycin B resistance gene in pHM874-8.		
KanR-R	CAGTATAGCGACCAGCATTCACATACG	Reverse primer to amplify G418/kanamycin resistance gene in pHM785-1.		
Primers for cyb2C d	isruption			
IoCyb2c-upF	TCTTCAAGTCTCCCGATTCC	Forward primer to amplify the upstream region of cyb2C in P. kudriavzevii.		
IoCyb2c-upR	TGGTCTCGTCAAAGTATTTGCAAGGAC	Reverse primer to amplify the upstream region of cyb2C in P. kudriavzevii.		
IoCyb2c-downF	TTTACGAATCCATGTCTGATTCAAGTTGGG	Forward primer to amplify the downstream region of cyb2C in P. kudriavzevii.		
IoCyb2c-downR	TCTCAAATGTTTTCTTTTGATTTACTTAAATACTAA	Reverse primer to amplify the downstream region of cyb2C in P. kudriavzevii.		
DIoCyb2c-F	CAATAGTTATTTCTGAGTTTCTTATGGCACATTGTCCTTGCAAATACTTTGACGAGACCAGACATGGAGGCCCAG	Forward primer to amplify hygromycin B resistance gene in pHM874-8 or G418/kanamycin resistance gene in pHM785-1 with 60-bp upstream region adjacent to cyb2C gene in P. kudriavzevii.		
DIoCyb2c-R	AAGGGAGAAATGTGGAATACATTTTCACATCCCAACTTGAATCAGACATGGATTCGTAAACTTCGAGCGTCCCAA	Reverse primer to amplify hygromycin B resistance gene in pHM874-8 with 60-bp downstream region adjacent to cyb2C gene in P. kudriavzevii.		
DIoCyb2cKanR-R	CCCAACTTGAATCAGACATGGATTCGTAAACAGTATAGCGACCAGCATTCACATACG	Reverse primer to amplify G418/kanamycin resistance gene in pHM785-1 with 25-bp		
Primers for real-tim	e PCR	downstream region adjacent to cyo2c gene in F. kuartavzevit.		
IoCvb2a-F	TCGATTTGGGTGATTCCGCA	Forward primer to check the disruption of cvb2A in P. kudriavzevii by real-time PCR.		
IoCyb2a-R	TTAGTCCACGACTTGACCGC	Reverse primer to check the disruption of cyb2A in P. kudriavzevii by real-time PCR.		
IoCyb2b-F	TCTTTGTCACTGTGGACGCT	Forward primer to check the disruption of cyb2B in P. kudriavzevii by real-time PCR.		
IoCyb2b-R	TGGCAATGCACCAGAGGTAG	Reverse primer to check the disruption of cyb2B in P. kudriavzevii by real-time PCR.		
IoCyb2c-F	AACATTTTGAGGGCATGCGG	Forward primer to check the disruption of cyb2C in P. kudriavzevii by real-time PCR.		
IoCyb2c-R	GGCGAGGCACTCTCCATAAA	Reverse primer to check the disruption of cyb2C in P. kudriavzevii by real-time PCR.		
IoACT1-F	CCGGTTTTGCCGGTGATGAT	Forward primer to amplify partial act1 in P. kudriavzevii by real-time PCR.		
IoACT1-F	CCTTTTGGCCCATACCAACCA	Reverse primer to amplify partial act1 in P. kudriavzevii by real-time PCR.		



Figure 1 C



■ A. niger ■ A. fumigatus ■ Zygomycetes □ Yeast

Figure 2



Figure 3



Figure 4





Figure A.2



Figure A.3



OD₆₂₀