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- Title: The natural diversity and ecology of fission yeast.

  Running head: Natural fission yeast diversity and ecology

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- 11
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15	
16	Abstract
17	While the fission yeast is a powerful model of eukaryote biology, there have been
18	few studies of quantitative genetics, phenotypic or genetic diversity. Here I survey
19	the small collection of fission yeast diversity research. I discuss what we can infer
20	about the ecology and origins of Schizosaccharomyces pombe from microbiology
21	field studies and the few strains that have been collected.
22	
23	Introduction
24	Schizosaccharomyces pombe research began in the 1940s (Fantes and Hoffman
25	2016) and is now a potent model of eukaryote biology, with a well-annotated
26	curated genome (Wood et al. 2002; McDowall et al. 2015), an extensive battery
27	of technical methods and genome-scale tools (Hoffman, Wood and Fantes 2015;
28	Hagan et al. 2016) and regular international meetings devoted to its study. Part of
29	the important utility of fission yeast as a model is that it contains many vertebrate
30	orthologs that are not present in budding yeast (Hoffman, Wood and Fantes
31	2015), so it provides a complement for studies of cell biology.
32	The majority of fission yeast research has used the strains described by
33	Leupold with its three mating types (Leupold 1949), and mutants derived from
34	these strains. Studies of diversity or quantitative genetics have been few and far
35	between. By contrast there is an extensive literature describing diversity and
36	quantitative genetics in the budding yeast Saccharomyces cerevisiae and its wild
37	relative Saccharomyces paradoxus, and a range of related species (Peter and
38	Schacherer 2016). These include QTL studies
39	(Swinnen, Thevelein and Nevoigt 2012; Liti and Louis 2012; Fay 2013;
40	Bloom et al. 2013; Märtens et al. 2016), genome-scale analysis of diversity (Liti
41	et al. 2009; Schacherer et al. 2009) and analysis of diversity and evolution in the
42	natural environment (Robinson, Pinharanda and Bensasson 2016; Leducq et al.
43	2016). In this review, I survey fission yeast diversity research, and I discuss what
44	little is known about the origins and natural ecology of this species.
45	
46	Defining fission yeast species
47	Collections of Schizoaccharomyces strains were classified into three groups based
48	on crossing and protoplast fusion (Sipiczki et al. 1982), phenotypic characters

(Bridge and May 1984), DNA optical reassociation and physiological characters

(Vaughan Martini 1991), simplifying the rather complex list of potential 'species'
 into three (*Schizoaccharomyces pombe*, *S. japonicus*, *S. octosporus*).
 *Schizosaccharomyces cryophilus* was identified much later as a contaminant of a
 *S. octosporus* strain (CBS7191) from Denmark, and the species description was

accompanied by a draft genome (Helston et al. 2010).

The genomes and transcriptomes of *S. japonicus*, *S. octosporus* and an improved *S. cryophilus* genome were described in 2011, showing that the *Schizosaccharomyces* genus is as divergent on the protein level as the human-amphioxus divergence (~55% amino acid identity) (Rhind *et al.* 2011). This analysis described the conservation of orthologous groups, conservation of transcription, the evolution of mating type regions and transposons. It also features the first sequencing of a non-reference strain of *S. pombe*, concluding that the within-species diversity was < 1% (confirmed later with studies of more strains (Fawcett *et al.* 2014; Jeffares *et al.* 2015)). The current clade of only four highly divergent fission yeast species is a limitation for evolutionary studies, since evolutionary constraints can be estimated only inaccurately, and non-coding sites that are in general subject to weaker purifying selection tend to be saturated (Rhind *et al.* 2011). None of the *Schizosaccharomyces* species is sufficiently

closely related to S. pombe to reliably determine ancestral nucleotide states.

# Early (pre-genome sequence) diversity studies

An early field study of this species was conducted by Florenzano *et al.*, who showed that *S. pombe* was frequently present on grapes in Sicilian vineyards (Florenzano, Balloni and Materassi 1977). Phenotypic characterization began with analysis of xerotolerance (resistance to high solute concentrations) in 27 *S. pombe* strains (Ganthala, Marshall and May 1994). One the first genetic analysis of diversity within *S. pombe* described the intron content of mitochondrial genomes in 26 strains, showing presence/absence polymorphisms in group I and group II introns (Zimmer *et al.* 1987). Interestingly, there appears to be no intron presence polymorphisms in the nuclear genomes of sequenced strains (Mourier & Jeffares, unpublished analyses), though on the longer scale fission yeasts have certainly undergone intron gain and loss (Mourier and Jeffares 2003; Jeffares, Mourier and Penny 2006; Rhind *et al.* 2011).

In a prelude to genome-scale analyses, three studies began to explore

genetic and phenotypic diversity on a larger scale. Gomes et al., collected 27

strains from seven Brazilian cachaca distilleries, and characterised osmotolerance. trehalose accumulation and ethanol tolerance, showing that these strains could grow in 50% glucose and 10% ethanol (Gomes et al. 2002). They also explored population structure using RAPD-PCR (random amplified polymorphic DNA PCR), demonstrating local population structure in Brazilian cachaça strains. RAPD-PCR was a useful method to characterise diversity prior to next generation sequencing, but the development of 26 primers for microsatellite PCR now provide a simple method to genotype strain collections (Patch and Aves 2007). Brown et al. assembled 81 natural isolates of S. pombe including samples from all continents (except Antarctica), and measured a large assembly of phenotypic characters, including growth parameters in 42 liquid media and cell length (Brown et al. 2011). This analysis also described diversity at three locations, and estimated that the global effective population size of this species is 10<sup>7</sup> (a figure that remained after genome-wide analysis (Farlow et al. 2015)). Most interestingly, this work described extensive karyotype diversity within this collection, including reciprocal translocations, duplications and inversions, showing that the ribosomal repeats were located on different chromosome ends in different strains.

# Genome-wide sequence analyses

The creation and analysis of the only fission yeast recombinant strain library was published in 2014 (Clément-Ziza *et al.* 2014). This study used a two-parent segregant panel and described expression QTLS (eQTLs) from both protein-coding and non-coding transcripts, during growth and stress conditions. Interestingly this study discovered a larger proportion of associations between genetic variants and non-coding transcripts than coding transcripts. The most significant variant, that affected 44% of eQTL associations and growth rate, was a frameshift in the *swc5* gene - part of a complex that affects histone deposition. Detailed analysis showed that this frameshift caused increased antisense transcription and decreased sense transcription, providing an example of the molecular events that influenced a complex trait such as growth. Further analyses of segregant panels are in progress, describing positive selection and the genetic control of RNA and protein levels (Clément-Ziza, pers. comm.).

An analysis of segregant pool based mapping (bulk segregant analysis) from

a two-parent cross showed that this method was feasible in fission yeast (Hu, Suo

120 and Du 2015). Hu et al. localised the probable causal allele of maltose deficiency 121 by sequencing pools grown with and without maltose. The analysis was 122 complicated by an inversion in the reference strain, but few other wild strains 123 (Jeffares et al. 2017), which reduces the local recombination rate (Clément-Ziza et 124 al. 2014). 125 Two genome-wide analyses of genetic diversity in S. pombe were published 126 soon afterwards (Fawcett et al. 2014; Jeffares et al. 2015). Both analyses 127 described recombination rate and population structure, and showed that exons, 128 UTRs and introns were the main targets of purifying selection. Estimates of diversity ( $\pi$ ) were ~3 × 10<sup>-3</sup> (pairwise comparison have an average of 3 SNPs/kb), 129 slightly higher than the budding yeast Saccharomyces cerevisiae (1x 10<sup>-3</sup>) (Liti et 130 131 al. 2009). From the genetic diversity and mutation rates, the effective population 132 size of S. pombe has been estimated to be 12 million, on a similar scale to budding 133 yeast (3 million) (Farlow et al. 2015). 134 The analysis of Fawcett et al. (32 strains) described some unusual patterns 135 of diversity that were likely due to soft selective sweeps, and either balancing 136 selection or introgression from some unknown fission yeast outgroup (Fawcett et 137 al. 2014). Jeffares et al. (161 strains) described transposon insertions and included 138 analysis of quantitative traits, their heritability and quantitative genetics using the 139 genome-wide association study (GWAS) approach (Jeffares et al. 2015). This 140 study located 1.400 variants that were significantly associated with traits despite 141 the very small sample size, showing that the combination of simple tractable 142 genetics with the capability to measure traits accurately with abundant repeat 143 measurements in well-controlled environments, is a powerful combination. 144 Further analysis with the same strain collection described structural variants 145 showing that they are both transient and contribute considerably to quantitative traits and reproductive isolation (Jeffares et al. 2017). Interestingly the variance in 146 147 wine-making traits, such as malic acid accumulation and glucose/fructose 148 ultilisation (Benito et al. 2016), appeared to be caused entirely by structural 149 variants. Two genome-scale analyses of the mutation rate estimated the point 150 mutation rate to be 1.7 x 10<sup>-10</sup> (or 2.0 x 10<sup>-10</sup>) per base per generation (Farlow et 151 152 al. 2015; Behringer and Hall 2015), very similar to estimates for the budding yeast Saccharomyces cerevisiae (estimates at 3 and 1.67 x 10<sup>-10</sup>) (Lynch et al. 153 154 2008; Zhu et al. 2014). Both studies noted a strong bias towards small insertions,

over deletions, which occur primarily in the non-protein regions of the genome, a pattern that is retained in natural genetic diversity (Jeffares et al. 2015).

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# Reproductive isolation

158 159 One topic that has received particular attention is the study of mating types 160 and reproductive isolation. Since the outset of fission yeast research, it was clear homothallic strains could mutate to more or less stable heterothallic genotypes (h<sup>+</sup> or h<sup>-</sup>) (Leupold 1949). Natural isolates also vary genetically at mating type regions and in their mating behavior, with some strains mutating more frequently from h<sup>+</sup> to h<sup>-</sup> and vice versa (Schlake and Gutz 1993). In an interesting demonstration that reproductive isolation could evolve via pre-zygotic mechanisms, Sieke et al. created three novel reproductive groups with different pheromone-receptor pairs (Seike, Nakamura and Shimoda 2015). Given these changes it is likely that pre-zygotic reproductive isolation occurs within some populations. 170 Several studies described the low spore viability that results from many inter-strain matings (Kondrat'eva and Naumov 2001; Teresa Avelar et al. 2013; Zanders et al. 2014; Naumov and Kondratieva 2015; Jeffares et al. 2015). Viability ranges from pairs showing < 1% viable offspring to strains with 90% viable, similar a range observed for *species* of budding yeast with that have much higher genetic divergence than fission yeast strains (Liti, Barton and Louis 2006), consistent with S. pombe strains being 'on the verge of speciation' (Naumov and Kondratieva 2015) (Figure 1A). Some homothallic strains are also ineffective at mating with their own genotype (Kondrat'eva and Naumov 2001; Jeffares et al. 2015). Since most crosses do produce mating bodies and asci (Xavi Marsellach, pers. comm.), the isolation is generally post-zygotic (intrinsic reproductive isolation). The accumulation of genetic factors that reduce mating success withinin these relatively closely related strains is probably due to the low 184 frequency of outbreeding in fission yeast. Based on the decay in linkage between wild isolates Farlow et al. estimated that S. pombe mate with a genetically dissimilar individual on average every 800,000 generations (Farlow et al. 2015), far less frequently than the estimates 50,000 generation for S. cerevisiae (Ruderfer et al. 2006). Given this frequency of, it is not surprising that the existing strains

have accumulated genetic factors that preclude interbreeding in the ~2300 years since these strains have drifted apart (Jeffares *et al.* 2015).

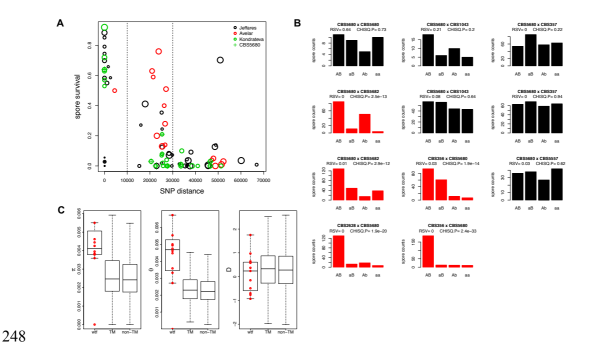
There are at least three (non-exclusive) genetic causes for the reproductive isolation of fission yeasts. Spore killing (meiotic drive), has been proposed to be a mechanism (Kondrat'eva and Naumov 2001; Zanders *et al.* 2014; Naumov and Kondratieva 2015). Many of the crosses analysed by Kondratieva *et al.* from genetically divergent strains and produced strong deviations from expected Mendelian ratios (Kondrat'eva and Naumov 2001; Naumov, Kondratieva and Naumova 2015) (Figure 1B), while the analyses of Zanders *et al.* concluded that there were meiotic drive elements on all three chromosomes (Zanders *et al.* 2014).

Two recent analyses have demonstrated that members of the *wtf* gene family mediate drive with a spore killer-antidote system (Hu *et al.* 2017; Nuckolls *et al.* 2017). Hu *et al.* demonstrate that *wtf9* and *wtf27* genes from the non-reference strain (CBS5557/JB4) drive segregation distortion in when mated to the reference strain, that this drive is independent of genomic location. Nuckolls *et al.* show that *wtf4* promotes distortion in crosses between the reference strain and the kombucha strain (SPK1820/YFS276/JB1180, as initially sequenced by the Broad Institute (Rhind *et al.* 2011)). Other strains analysed by Kondratieva *et al.* also show very biased segregation (Figure 1B).

Collectively, these analyses show that the spore killer (or poison) and antidote functions can be separated by mutations. In the natural state, there are two transcripts that mediate killer/antidote functions (Nuckolls *et al.* 2017). While the killer protein variant is distributed in all four spores of the asci, the antidote remains only within cells with the relevant *wtf* genotype. Since *wtf* genes encode membrane-spanning domains they may travel between asci. The genetics of the poison-antidote systems are complex, in that there are multiple *wtf* genes in different strains that have degenerated to contain the poison and antidote functions, antidote only, or no function. Both analyses show that *wtf* genes are particularly genetically diverse (Figure 1C). However, they do not show an excess of high Tajima's D values (Tajima 1989)(Figure 1C), a genetic diversity parameter which is one of the expected signatures of balancing selection.

Reproductive isolation may also be the result of the aneuploidy that occurs when parents differ in chromosomal inversions and translocations. For example, engineered inversions and translocations reduce spore viability by ~40% (Teresa Avelar *et al.* 2013). *S. pombe* strains do have extensive karyotype differences

224 (Brown et al. 2011; Naumov, Kondratieva and Naumova 2015; Jeffares et al. 225 2017), including a strain that maintains four (rather than the usual three) chromosomes (Brown et al. 2014). There is a significant association between 226 227 viability and the SV-distance between parents (Jeffares et al. 2017), though 228 viability declines at less than 40% viability per variant. This is probably because 229 natural structural variants are biased to chromosome ends that do not contain 230 essential genes (Jeffares et al. 2015), due to selection for those that do not cause 231 lethal aneuploidies. Structural variants may also contribute to drive (Zanders et al. 232 2014). 233 Formally, reproductive isolation may also be due to Bateson-Dobzhansky-234 Muller interactions (BDMIs) or any of the other genetic mechanisms of negative 235 epistasis (Nei and Nozawa 2011). However segregation data from random spores 236 (Kondrat'eva and Naumov 2001; Naumov and Kondratieva 2015) and dissected 237 tetrads is inconsistent with simple two-locus BDMIs, which are expected to 238 produce small deviations from expected segregation patterns (even when the 239 affected alleles were strongly linked to markers) (Hou and Schacherer 2016). 240 Ultimately meiotic drive, epistasis and structural variants may have interacting 241 effects on viability, since locally adapted haplotypes are predicted to develop 242 within areas of reduced recombination (Kirkpatrick and Barton 2006). 243 With all these studies of population genetics (reproductive isolation, 244 divergence dating, diversity measures, population size etc.) the analyses are based 245 on a small collection of strains that are a worldwide sample of mostly human 246 commensals (see below), so conclusions may not represent natural populations.



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Figure 1. Intrinsic reproductive isolation in S. pombe.

A) Random spore viability from three studies shows a decline in spore survival with genetic distance (SNP distance) between parents. The size of circles indicates the lowest self-mating viability of parents. Data from (Kondrat'eva and Naumov 2001; Teresa Avelar et al. 2013; Jeffares et al. 2015). Crosses involving the strain CBS5680 (as in part B) are indicated with cross hairs. The range of genetic differences that have highly variable effects on viability (10,000 - 30,000 SNPs)is indicated with vertical dashed lines. The outlier at top right is JB848/CBS10475 (Brazil) x JB870/CBS10499 (South Africa), which appears to be real (Xavier Marsellach, pers. comm.). B) segregation of control markers in random spore analysis show strong deviations from the expected 1:1:1:1 ratio, data from (Kondrat'eva and Naumov 2001). For one strain (CBS5680/JB873, from Poland) we show the counts of control markers (aB and Ab are parental types, AB, ab are recombinants, see Kondrateva et al. for details). Segregation counts whose  $\chi^2$  test P-values were < 0.05 are plotted with red bars. Plot text shows the parents of the cross, the random spore viability (RSV) and the  $\gamma^2$  test P-value (CHISQ.P). C) wtf genes have high pairwise diversity within strains compared to all other transmembrane domain containing and non-TM genes ( $\pi$ , left panel), high numbers of segregating sites ( $\theta$ , middle panel), but are not outliers for Tajima's D (which is calculated from the ratio of the two, D, right panel). Plots show diversity estimators from 57 strains, red circle indicate individual values for wtf genes.

Predicted transmembrane proteins were collected from a query of Pombase (www.pombase.org), diversity data from (Jeffares *et al.* 2015).

# Genetics and the reference strain

The fission yeast community has worked almost exclusively with one reference strain, and spontaneous mutants generated from this strain (Fantes and Hoffman 2016). This laboratory strain is a natural isolate, and is not an unusual strain phenotypically. It does not appear to be adapted to the standard rich or minimal media, since it does not grow particularly rapidly in these media compared to wild strains. There are several important discoveries that are relevant to the fission yeast researcher. Firstly, Wild strains can differ from the reference by up to 68,000 SNPs and up to 24 structural variations, which contribute to phenotypic variation between strains (Clément-Ziza et al. 2014; Jeffares et al. 2015; Hu, Suo and Du 2015; Jeffares et al. 2017). I summarise the structural differences between strains in Supplementary Figure 1. Secondly, the structural differences and meiotic drive elements that wild strains contain complicate crosses between strains, by reducing spore viability and skewing the proportions of alleles that are produced in the offspring (Kondrat'eva and Naumov 2001; Kondrateva and Naumov 2011; Clément-Ziza et al. 2014; Hu, Suo and Du 2015; Nuckolls et al. 2017; Hu et al. 2017).

# The ecology of fission yeast

There have been few published attempts to systematically collect fission yeast strains (Gomes *et al.* 2002; Benito *et al.* 2013; Hellberg 2013). However, fission yeasts have been serendipitously discovered in a variety of microbiological studies (Table 1, Figure 2). Sources have generally been traditional non-industrialised fermentations, produced without any intentional inoculation from substrates that contain high concentrations of sugars. When quantitative estimates of species abundances are included *Schizosaccharomyces* yeasts were generally minor components of these fermentations, with the exceptions of kombucha, some cachaça fermentations and baijiu (from tea, sugar cane and sorghum respectively) (Pataro, Guerra and Peixoto 2000; Teoh, Heard and Cox 2004; Wu, Xu and Chen 2012).

Perhaps more informative for fission yeast ecology, are the cases where

fission yeasts have been discovered in natural substrates such as palm wine (a fermentation of palm sap) (Theivendirarajah and Chrystopher 1987; Amanchukwu, Obafemi and Okpokwasili 1989; Ouoba *et al.* 2012). Fission yeast are also present in natural fermentations of fruits such as *Coffea arabica* and *Theobroma cacao* (from which coffee and cocoa beans are harvested respectively) (Silv *et al.* 2000; Schwan and Wheals 2004). Collectively, the field studies show that fission yeasts are a component of natural microbial communities that ferment botanical sugars in several geographic regions.

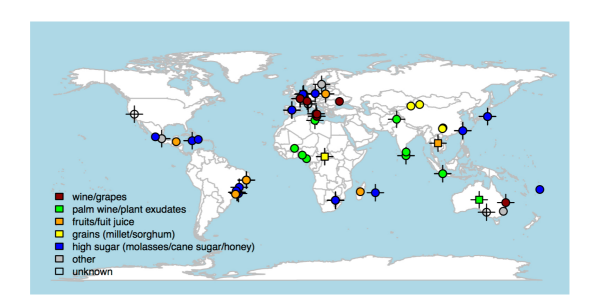
Including the strains present in stock collections and in field studies the most common substrates for fission yeast have been palm wine, grape wine, high-sugar substrates (molasses, cane sugar, honey) and fruits (Figure 2). Three selective media to have been described to enrich for fission yeast (Florenzano, Balloni and Materassi 1977; Hellberg 2013; Benito *et al.* 2013), so further systematic collections from similar locations and substrates should be possible in the future.

Table 1. Schizosaccharomyces in field microbiology

Substrate	Location	Reference
Grape must	Sicily	(Florenzano, Balloni and
		Materassi 1977)
Grapes	Ukraine	(Bayraktar 2014)
Palm wine	Sri Lanka	(Atputharajah,
		Widanapathirana and
		Samarajeewa 1986;
		Theivendirarajah and
		Chrystopher 1987)
Palm wine	Nigeria	(Sanni and Lönner 1993;
		Amanchukwu, Obafemi
		and Okpokwasili 2006)
Palm wine	Burkina Faso	(Ouoba et al. 2012)
Rum	Haiti	(Fahrasmane, Ganou-
		Parfait and Parfait 1988)
Molasses, raisin	Japan/Thailand/Taiwan	(Ishitane 1985)

Tequila	Mexico	(Lachance 1995)
Coffee cherries	Brazil	(Silv et al. 2000)
	Madagascar	(Ravelomanana et al.
		1984)
Cachaça	Brazil	(Pataro, Guerra and
(from sugar cane)		Peixoto 2000; Gomes et
		al. 2002)
Kombucha	Australia**	(Teoh, Heard and
(fermented tea)		Cox 2004)
Cocoa pulp	Belize	(Schwan and Wheals
		2004)
Baijiu	China	(Wu, Xu and Chen 2012)
(distillate of fermented		
sorghum)		
Traditional breweries	China	Fen-Yang Bai,
		pers. comm.
Honey	Fiji	(Ponici and Wimmer
		1986)
Honey	Spain	(Benito et al. 2014)

<sup>\*</sup> Not microbiological study itself, refers to earlier work.



<sup>\*\*</sup> From commercial kombucha brewers.

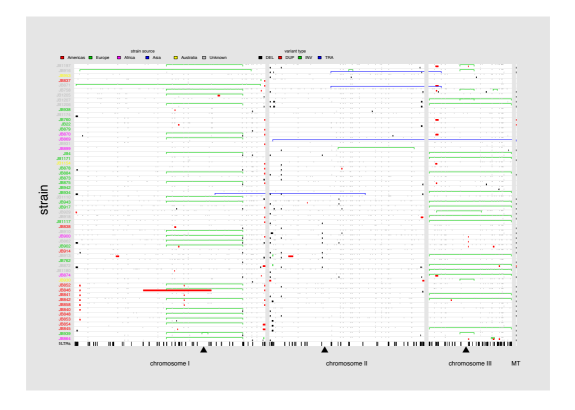
328 Figure 2. Fission yeast locations and substrates. The locations and 329 substrates where fission yeast have been discovered, including all strains that have 330 been sequenced from stock centers (Fawcett et al. 2014; Jeffares et al. 2015), and 331 reports from field studies (Table 1). Sequenced strains are marked with cross-332 hairs, and strains isolated from uncertain locations are marked with a square. 333 334 The origin of fission yeast 335 S. pombe is now globally distributed (Figure 2), but we know little about its 336 origin and dispersal. We have estimated that these strains began to spread globally 337 in from ~340 BCE (95% confidence interval 1875 BCE-1088 CE), and that the 338 current collection of strains from Brazilian cachaça originated from the remainder 339 in about ~1620 CE (confidence interval 1422–1752 CE) (Jeffares et al. 2015), a 340 hint that like budding yeast and *C. elegans*, this model has probably been 341 dispersed as a commensal (most likely in fermented beverages). 342 The reference strain originated from French grapes (Osterwalder 1924). The 343 common belief is that S. pombe originated from Africa, perhaps because the initial 344 species description was from an African millet beer isolate (Lindner 1893; 345 Vorderman 1894). While genetic analysis is consistent with exchange between 346 African and European stocks (Jeffares et al. 2015), and some strains have been 347 collected from traditional African fermentations, there is no scientific evidence for 348 an African origin of this species. There are very few studies of the microbial 349 constituents of millet beer from Africa (I could fine none than specifically 350 mentioned S. pombe, and one description of sorghum beer that did not mention S. 351 pombe (Kayode et al. 2011)). Since fission yeasts can be major components of 352 kombucha, which has been traditionally produced in China (Sreeramulu, Zhu and 353 Knol 2000; Teoh, Heard and Cox 2004), palm wine which is widely produced in 354 Asia (Table 1, Figure 2), and in traditional Chinese breweries (Fen-Yang Bai, 355 pers. comm.), China is an equally good candidate for the initial origin of S. 356 pombe. 357 358 Why study diversity in fission yeast? 359 The small genomes of budding yeasts enabled the early development of 360 population genomics methods (Liti et al. 2009; Schacherer et al. 2009), and now

large scale accurate quantitative genetics analyses (Bloom et al. 2013; Märtens et

al. 2016). The continuing advance of sequence throughput, analysis software and

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363	laboratory methods (eg: RAD-seq) have now made population genomics
364	approaches available to any species. However, the abundance of genome-scale
365	data and technical tools and the small non-redundant genomes of yeasts make
366	them attractive models for systems biology, including approaches to
367	understanding genetic diversity and traits (Parts 2014). Fission yeast has the
368	benefit of being haploid (so that F1 generations need not be intercrossed). As with
369	budding yeast, fission yeast has abundant heritable phenotypic diversity in
370	growth, stress responses, cell morphology, and cellular biochemistry that is yet to
371	be explored with powerful quantitative genetics (Brown et al. 2011; Clément-Ziza
372	et al. 2014; Jeffares et al. 2015; 2017). Yeasts are also powerful tools for detailed
373	study of evolutionary processes using pooled time-series sequencing and other
374	high-throughput approaches that would be expensive or unfeasible in other
375	species (Cubillos et al. 2011; Hou et al. 2015). Finally, studies by Benito et al.
376	show that some non-reference <i>S. pombe</i> strains have potential in the winemaking
377	industry (Benito et al. 2014; 2016), so diverse strains could well have potential
378	elsewhere in biotechnology.
379	
380	Acknowledgements
381	I thank Mathieu Clément-Ziza for commentary about unpublished work and
382	Xavier Marsellach for discussions.
383	
384	Supplementary data
385	All used for plots is available at figshare at:
386	https://figshare.com/projects/The natural diversity and ecology of fission yeas
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# **Supplementary Figure 1. Structural variants present in wild fission yeast strains**. Using predictions from short read data (Jeffares *et al.* 2017), I show the genomic location of structural variants (SVs) in wild strains contain that differ from the standard laboratory isolate (Leupold's 972). I show deletions (black), duplications (red), inversions (green) and translocations (blue). SVs present in each of the 57 non-clonal strains are shown within the white horizontal bars, with strain names coloured according to their continent of origin. Tf1-type retrotransposon insertions that are present in some, but not all strains are shown at grey ticks at the tops of bars. The positions of fixed Tf1-type retrotransposon insertions are indicated on the last row (f/LTRs). Centromeres are indicated with black triangles.

# References

Amanchukwu SC, Obafemi A, Okpokwasili GC. Single-cell-protein production by Schizosaccharomyces pombe isolated from palmwine using hydrocarbon feedstocks. *Folia Microbiol* 1989;**34**:112–9.

420	Amanchukwu SC, Obafemi A, Okpokwasili GC. Hydrocarbon degradation and
421	utilization by a palm-wine yeast isolate. FEMS microbiology letters
422	2006; <b>57</b> :151–4.
423	Atputharajah JD, Widanapathirana S, Samarajeewa U. Microbiology and
424	biochemistry of natural fermentation of coconut palm sap - ScienceDirect.
425	Food Microbiol 1986.
426	Bayraktar VN. Biodiversity of yeast cultures isolated from littoral areas of the
427	Regional Landscape Park «Tiligulskiy». J. Optimization and Protection of
428	Ecosystems 2014.
429	Behringer MG, Hall DW. Genome-Wide Estimates of Mutation Rates and
430	Spectrum in Schizosaccharomyces pombe Indicate CpG Sites are Highly
431	Mutagenic Despite the Absence of DNA Methylation. G3 2015;6:149–60.
432	Benito Á, Jeffares DC, Palomero F et al. Selected Schizosaccharomyces pombe
433	Strains Have Characteristics That Are Beneficial for Winemaking. PLoS ONE
434	2016; <b>11</b> :e0151102.
435	Benito S, Gálvez L, Palmero F et al. Schizosaccharomyces selective differential
436	media. African Journal of Microbiology Research 2013;7:3026–36.
437	Benito S, Palomero F, Calderón F et al. Selection of appropriate
438	Schizosaccharomyces strains for winemaking. Food Microbiol 2014;42:218-
439	24.
440	Bloom JS, Ehrenreich IM, Loo WT et al. Finding the sources of missing
441	heritability in a yeast cross. <i>Nature</i> 2013; <b>494</b> :234–7.
442	Bridge PD, May JW. A numerical classification of fission yeasts of the genus
443	Schizosaccharomyces Lindner. <i>Microbiology</i> 1984; <b>130</b> :1921–32.
444	Brown WRA, Liti G, Rosa C et al. A geographically diverse collection of
445	Schizosaccharomyces pombe isolates shows limited phenotypic variation but
446	extensive karyotypic diversity. Fay JC (ed.). G3 2011;1:615–26.
447	Brown WRA, Thomas G, Lee NCO et al. Kinetochore assembly and
148	heterochromatin formation occur autonomously in Schizosaccharomyces

149	pombe. Proc Natl Acad Sci USA 2014;111:1903–8.
450	Clément-Ziza M, Marsellach FX, Codlin S et al. Natural genetic variation impacts
451	expression levels of coding, non-coding, and antisense transcripts in fission
452	yeast. Mol Syst Biol 2014; <b>10</b> :764–4.
453	Cubillos FA, Billi E, Zörgö E et al. Assessing the complex architecture of
454	polygenic traits in diverged yeast populations. <i>Mol Ecol</i> 2011; <b>20</b> :1401–13.
455	Fahrasmane L, Ganou-Parfait B, Parfait A. Research note: Yeast flora of Haitian
456	rum distilleries. Mircen Journal 1988;4:239–41.
457	Fantes PA, Hoffman CS. A Brief History of Schizosaccharomyces pombe
458	Research: A Perspective Over the Past 70 Years. <i>Genetics</i> 2016; <b>203</b> :621–9.
459	Farlow A, Long H, Arnoux S et al. The Spontaneous Mutation Rate in the Fission
460	Yeast Schizosaccharomyces pombe. <i>Genetics</i> 2015; <b>201</b> :737–44.
461	Fawcett JA, Iida T, Takuno S et al. Population Genomics of the Fission Yeast
462	Schizosaccharomyces pombe. <i>PLoS ONE</i> 2014; <b>9</b> :e104241.
463	Fay JC. The molecular basis of phenotypic variation in yeast. Current Opinion in
464	Genetics and Development 2013; <b>23</b> :672–7.
465	Florenzano G, Balloni W, Materassi R. Contributo alla ecologia dei lieviti
466	Schizosaccharomyces sulle uve. Vitis 1977;16:38–44.
467	Ganthala BP, Marshall JH, May JW. Xerotolerance in fission yeasts and the role
468	of glycerol as compatible solute. <i>Archives of microbiology</i> 1994; <b>162</b> :108–13.
469	Gomes FCO, Pataro C, Guerra JB et al. Physiological diversity and trehalose
470	accumulation in Schizosaccharomyces pombe strains isolated from
471	spontaneous fermentations during the production of the artisanal Brazilian
472	cachaça. <i>Can J Microbiol</i> 2002; <b>48</b> :399–406.
473	Hagan IM, Grallert A, Carr AM et al. eds. Fission Yeast: a Laboratory Manual.
474	Cold Spring Harbour Laboratory Press, 2016.
475	Hellberg JEEU. Finding wild fission yeast: Where are they? Genetics Society
476	News 2013: <b>68</b> :59–60.

177 178	Helston RM, Box JA, Tang W <i>et al.</i> Schizosaccharomyces cryophilus sp nov., a new species of fission yeast. <i>Fems Yeast Res</i> 2010; <b>10</b> :779–86.
179 180 181	Hoffman CS, Wood V, Fantes PA. An Ancient Yeast for Young Geneticists: A Primer on the Schizosaccharomyces pombe Model System. <i>Genetics</i> 2015; <b>201</b> :403–23.
182 183 184	Hou J, Friedrich A, Gounot J-S <i>et al.</i> Comprehensive survey of condition-specific reproductive isolation reveals genetic incompatibility in yeast. <i>Nat Commun</i> 2015; <b>6</b> :7214.
185 186 187	Hou J, Schacherer J. On the Mapping of Epistatic Genetic Interactions in Natural Isolates: Combining Classical Genetics and Genomics. <i>Methods Mol Biol</i> 2016; <b>1361</b> :345–60.
188 189	Hu W, Jiang Z-D, Suo F <i>et al.</i> A large gene family in fission yeast encodes spore killers that subvert Mendel's law. <i>eLife</i> 2017; <b>6</b> :3025.
190 191	Hu W, Suo F, Du L-L. Bulk segregant analysis reveals the genetic basis of a natural trait variation in fission yeast. <i>Genome Biol Evol</i> 2015;7:3496–510.
192 193	Ishitane. Identification of yeasts isolated from high-sugar foods. Tokuoka K, Ishitane T, Goto S, et al. (eds.). <i>J Gen Appl Microbiol</i> 1985; <b>31</b> :411–27.
194 195 196	Jeffares DC, Jolly C, Hoti M <i>et al.</i> Transient structural variations have strong effects on quantitative traits and reproductive isolation in fission yeast. <i>Nat Commun</i> 2017; <b>8</b> :14061.
197 198 199	Jeffares DC, Jolly C, Hoti M <i>et al</i> . Transient structural variations have strong effects on quantitative traits and reproductive isolation in fission yeast. <i>Multiple values selected</i> <b>8</b> :14061EP–.
500 501	Jeffares DC, Mourier T, Penny D. The biology of intron gain and loss. <i>Trends Genet</i> 2006; <b>22</b> :16–22.
502 503	Jeffares DC, Rallis C, Rieux A <i>et al.</i> The genomic and phenotypic diversity of Schizosaccharomyces pombe. <i>Nature Genetics</i> 2015; <b>47</b> :235–41.
504 505	Kayode APP, Vieira-Dalode G, Linnemann AR <i>et al.</i> Diversity of yeasts involved in the fermentation of tchoukoutou, an opaque sorghum beer from Benin.

506	African Journal of Microbiology Research 2011; <b>5</b> :2737–42.
507	Kirkpatrick M, Barton N. Chromosome inversions, local adaptation and
508	speciation. <i>Genetics</i> 2006; <b>173</b> :419–34.
509	Kondrat'eva VI, Naumov GI. The Phenomenon of Spore Killing in
510	Schizosaccharomyces pombeHybrids. Doklady Biological Sciences 2001.
511	Kondrateva VI, Naumov GI. POPULATION ANTAGONISM IN THE YEASTS
512	SCHIZOSACCHAROMYCES POMBE. Ecological genetics 2011;9:21–6.
513	Lachance MA. Yeast communities in a natural tequila fermentation. Antonie Van
514	Leeuwenhoek 1995; <b>68</b> :151–60.
515	Leducq J-B, Nielly-Thibault L, Charron G et al. Speciation driven by
516	hybridization and chromosomal plasticity in a wild yeast. Nat Microbiol
517	2016; <b>1</b> :15003.
518	Leupold U. Die Vererbung von Homothallie und Heterothallie bei
519	Schizosaccharomyces pombe. Compt Rend Trav Lab Carlsberg Ser
520	1949; <b>24</b> :381–480.
521	Lindner P. Schizosaccharomyces pombe n. sp., ein neuer Gährungserreger.
522	Wochenschrift für Brauerei 1893; <b>10</b> :1298–300.
523	Liti G, Barton DBH, Louis EJ. Sequence diversity, reproductive isolation and
524	species concepts in Saccharomyces. <i>Genetics</i> 2006; <b>174</b> :839–50.
525	Liti G, Carter DM, Moses AM et al. Population genomics of domestic and wild
526	yeasts. <i>Nature</i> 2009; <b>458</b> :337–41.
527	Liti G, Louis EJ. Advances in quantitative trait analysis in yeast. Fay JC (ed.).
528	PLoS Genet 2012;8:e1002912.
529	Lynch M, Sung W, Morris K et al. A genome-wide view of the spectrum of
530	spontaneous mutations in yeast. <i>Proc Natl Acad Sci USA</i> 2008; <b>105</b> :9272–7.
531	Märtens K, Hallin J, Warringer J et al. Predicting quantitative traits from genome
532	and phenome with near perfect accuracy. Nat Commun 2016;7:11512.

533	McDowall MD, Harris MA, Lock A <i>et al.</i> PomBase 2015: updates to the fission yeast database. <i>Nucleic Acids Res</i> 2015; <b>43</b> :D656–61.
535	Mourier T, Jeffares DC. Eukaryotic intron loss. Science (New York, NY)
536	2003; <b>300</b> :1393–3.
537	Naumov G, Kondratieva V. Hybrid sterility of the yeast Schizosaccharomyces
538	pombe: Genetic genus and many species in statu nascendi? Microbiology
539	(00262617) 2015.
540	Naumov GI, Kondratieva VI, Naumova ES. [Hybrid Sterility of the Yeast
541	Schizosaccharomyces pombe: Genetic Genus and Many Species in statu
542	nascendi?]. Mikrobiologiia 2015; <b>84</b> :192–203.
543	Nei M, Nozawa M. Roles of mutation and selection in speciation: from Hugo de
544	Vries to the modern genomic era. Genome Biol Evol 2011;3:812–29.
545	Nuckolls NL, Bravo Núñez MA, Eickbush MT et al. wtf genes are prolific dual
546	poison-antidote meiotic drivers. <i>eLife</i> 2017; <b>6</b> :2235.
547	Osterwalder A. Schizosaccharomyces liquefaciens n.sp., eine gegen freie
548	schweflige Säure widerstandsfähige Gärhefe. Mitt Geb Lebensmittelunters
549	<i>Hyg</i> 1924; <b>15</b> :5–28.
550	Ouoba LII, Kando C, Parkouda C et al. The microbiology of Bandji, palm wine of
551	Borassus akeassii from Burkina Faso: identification and genotypic diversity
552	of yeasts, lactic acid and acetic acid bacteria. J Appl Microbiol 2012, DOI:
553	10.1111/jam.12014.
554	Parts L. Genome-wide mapping of cellular traits using yeast. <i>Yeast</i> 2014; <b>31</b> :197–
555	205.
556	Pataro C, Guerra JB, Peixoto MP. Yeast communities and genetic polymorphism
557	of Saccharomyces cerevisiae strains associated with artisanal fermentation in
558	Brazil. Journal of Applied Microbiology 2000;89:24–31.
559	Patch A-M, Aves SJ. Fingerprinting fission yeast: polymorphic markers for
560	molecular genetic analysis of Schizosaccharomyces pombe strains.
561	Microbiology (Reading, Engl) 2007;153:887–97.

562 563	Peter J, Schacherer J. Population genomics of yeasts: towards a comprehensive view across a broad evolutionary scale. <i>Yeast</i> 2016; <b>33</b> :73–81.
564	Ponici L, Wimmer FL. Characterization of the yeasts (Blastomycetes) in some
565	Fijian honeys. Acta alimentaria polonica 1986.
566	Ravelomanana R, Guiraud JP, Vincent JC et al. Study of the yeast flora from the
567	traditional cacao fermentation in Madagascar. Revue des Fermentations et des
568	Industries Alimentaires (Belgium) 1984.
569	Rhind N, Chen Z, Yassour M et al. Comparative functional genomics of the
570	fission yeasts. Science (New York, NY) 2011;332:930-6.
571	Robinson HA, Pinharanda A, Bensasson D. Summer temperature can predict the
572	distribution of wild yeast populations. Ecol Evol 2016; <b>6</b> :1236–50.
573	Sanni AI, Lönner C. Identification of yeasts isolated from Nigerian traditional
574	alcoholic beverages. Food Microbiol 1993;10:517–23.
575	Schacherer J, Shapiro JA, Ruderfer DM et al. Comprehensive polymorphism
576	survey elucidates population structure of Saccharomyces cerevisiae. Nature
577	2009; <b>458</b> :342–5.
578	Schlake T, Gutz H. Mating configurations in Schizosaccharomyces pombe strains
579	of different geographical origins. Curr Genet 1993.
580	Schwan RF, Wheals AE. The microbiology of cocoa fermentation and its role in
581	chocolate quality. Crit Rev Food Sci Nutr 2004;44:205–21.
582	Seike T, Nakamura T, Shimoda C. Molecular coevolution of a sex pheromone and
583	its receptor triggers reproductive isolation in Schizosaccharomyces pombe.
584	Proc Natl Acad Sci USA 2015; <b>112</b> :4405–10.
585	Silv CF, Schwan RF, Sousa Dias ES et al. Microbial diversity during maturation
586	and natural processing of coffee cherries of Coffea arabica in Brazil. Int $J$
587	Food Microbiol 2000; <b>60</b> :251–60.
588	Sipiczki M, Kucsera J, Ulaszewski S et al. Hybridization Studies by Crossing and
589	Protoplast Fusion Within the Genus Schizosaccharomyces Lindner. J Gen
590	Microbiol 1982: <b>128</b> :1989–2000

591	Sreeramulu G, Zhu Y, Knol W. Kombucha fermentation and its antimicrobial
592	activity. J Agric Food Chem 2000;48:2589–94.
593	Swinnen S, Thevelein JM, Nevoigt E. Genetic mapping of quantitative phenotypic
594	traits in Saccharomyces cerevisiae. Fems Yeast Res 2012;12:215–27.
595	Tajima F. Statistical method for testing the neutral mutation hypothesis by DNA
596	polymorphism. <i>Genetics</i> 1989; <b>123</b> :585–95.
597	Teoh AL, Heard G, Cox J. Yeast ecology of Kombucha fermentation. Int J Food
598	<i>Microbiol</i> 2004; <b>95</b> :119–26.
599	Teresa Avelar A, Perfeito L, Gordo I et al. Genome architecture is a selectable
600	trait that can be maintained by antagonistic pleiotropy. Nat Commun
601	2013; <b>4</b> :2235.
602	Theivendirarajah K, Chrystopher RK. Microflora and microbial activity in
603	palmyrah (Borassus flabellifer) palm wine in Sri Lanka. Mircen Journal
604	1987; <b>3</b> :23–31.
605	Vaughan Martini A. Evaluation of phylogenetic relationships among fission yeast
606	by nDNA/nDNA reassociation and conventional taxonomic criteria. Yeast
607	1991; <b>7</b> :73–8.
608	Vorderman AG. Analecta Op Bromatologisch Gebied: I II. Geneeskundig
609	Tijdschrift voor Nederlandsch-Indië, 1894:343–97.
610	Wood V, Gwilliam R, Rajandream M-A et al. The genome sequence of
611	Schizosaccharomyces pombe. <i>Nature</i> 2002; <b>415</b> :871–80.
612	Wu Q, Xu Y, Chen L. Diversity of yeast species during fermentative process
613	contributing to Chinese Maotai-flavour liquor making. Letters in Applied
614	Microbiology 2012; <b>55</b> :301–7.
615	Zanders SE, Eickbush MT, Yu JS et al. Genome rearrangements and pervasive
616	meiotic drive cause hybrid infertility in fission yeast. eLife
617	2014; <b>2014</b> :e02630.
618	Zhu YO, Siegal ML, Hall DW et al. Precise estimates of mutation rate and
619	spectrum in yeast. Proc Natl Acad Sci USA 2014;111:E2310-8.

620	Zimmer M, Welser F, Oraler G et al. Distribution of mitochondrial introns in the
621	species Schizosaccharomyces pombe and the origin of the group II intron in
622	the gene encoding apocytochrome b. Curr Genet 1987;12:329-36.
623	
624	The author declares that there is no conflict of interest.