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Comparative genomics and genome evolution in yeasts

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Yeasts provide a powerful model system for comparative genomics research. The availability of multiple complete genome sequences from different fungal groups—currently 18 hemiascomycetes, 8 euascomycetes and 4 basidiomycetes—enables us to gain a broad perspective on genome evolution. The sequenced genomes span a continuum of divergence levels ranging from multiple individuals within a species, to species pairs with low levels of protein sequence identity and no conservation of gene order. One of the most interesting emerging areas is the growing number of events such as gene losses, gene displacements and gene relocations that can be attributed to the action of natural selection.

Keywords: Saccharomyces cerevisiae; evolution; bioinformatics; genomics

1. INTRODUCTION

25 The rationale put forward in the late 1980s to justify 26 sequencing the genome of the yeast Saccharomyces 27 cerevisiae was twofold (Dujon 1996). First, yeast is an 28 organism with high economic value in brewing, baking 29 and biotechnology, and one of the primary model 30 organisms for fundamental research into eukaryotic 31 genetics and molecular biology. Second, its genome has 32 many properties that made it an attractive target for 33 early genome sequencing efforts. It has one of the 34 smallest genome sizes among well-studied eukaryotes 35 (only 14 million basepairs), a high gene density (72% of 36 the genome codes for protein), few introns and little 37 repetitive DNA-a combination of factors that made 38 the sequencing relatively easy and cost-effective. The 39 genome project was both doable and worth doing, with 40 the result that S. cerevisiae became the first eukaryote to 41 have a complete chromosome sequenced (Oliver et al. 42 1992), and later to have its whole genome sequenced 43 (Goffeau et al. 1997).

44 These same features have more recently led to yeast 45 species emerging as a powerful eukaryotic model 46 system for comparative genomics and studies of 47 genome evolution. The group of fungi that includes 48 S. cerevisiae—the hemiascomycetes—all have relatively 49 small and non-repetitive genomes, and several are 50 of biotechnological or medical interest. The result is 51 that today we have genome sequences (either complete, 52 or high-quality draft sequences) from 18 species of 53 hemiascomycetes (figure 1). The genome of 54 S. cerevisiae has now been sequenced three times: 55 once from the laboratory strain S288C (Goffeau et al. 56 1997), once from the clinical isolate YJM789 (Gu et al. 57 2005), and once from the vineyard isolate RM11-1a 58 (Broad Institute, unpublished; GenBank accession 59 number AAEG0100000). 60

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A sort of virtuous cycle has developed among the hemiascomycete genomes. The depth of knowledge that now exists about S. cerevisiae genes-such as thousands of microarray transcription experiments, and knockout phenotype information for every genemakes it possible to make quite detailed inferences about the physiology of related yeasts based only on their genome sequences (Hittinger et al. 2004), which makes sequencing those genomes worthwhile. In return, information from the other species can be used to learn more about the S. cerevisiae genome-for example in detecting functional motifs in the regulatory regions of genes based on their evolutionary conservation (Cliften et al. 2001, 2003; Kellis et al. 2003)which further elevates S. cerevisiae as a model organism. Thus, the combined value of the hemiascomycetes as a comparative genomics system is greater than the sum of its parts.

Complementing the recent progress that has been made with hemiascomycete genomes, significant strides have been made in sequencing genomes from the other major fungal clades. As shown in figure 1, we now have complete genome sequences from eight euascomycetes (filamentous ascomycetes such as Neurospora crassa), four basidiomycetes (including the mushroom Coprinopsis cinerea) and the archiascomycete Schizosaccharomyces pombe (Wood et al. 2002; Galagan et al. 2003; Dean et al. 2005; Loftus et al. 2005; and unpublished data available in GenBank). The euascomycete and basidiomycete genomes pose a somewhat greater technical challenge for sequencing than the hemiascomycetes owing to their larger sizes (typically 30-40 Mb) and greater number of introns, but we can see that they already present a rich resource for comparative genomics.

The hemiascomycete species whose genomes have 123 been sequenced span a very broad evolutionary range 124 (figure 1). It is difficult to put an absolute time-scale 125 onto this phylogenetic tree because of the lack of fossil 126 data for yeasts (Berbee & Taylor 1993). However, an 127 intuitive feeling for the depth of the divergences among 128

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 ⁶³ One contribution of 15 to a Discussion Meeting Issue 'Bioinformatics:
 64 from molecules to systems'.



Figure 1. The sequenced fungal genomes (June 2005). All these genomes have been sequenced to at least 3× coverage and the
data are in the public domain (available through GenBank or from the Sanger Institute website). Thickened branches indicate
the lineages that show whole-genome duplication. The phylogenetic tree is not drawn to scale and may contain errors. It is a
composite drawn from several sources (Berbee & Taylor 1993; Cai *et al.* 1996; Fungal Genome Initiative Steering Committee
2003; Kurtzman & Robnett 2003; Rokas *et al.* 2003) and D. R. Scannell and K. H. W., unpublished results.

yeast species can be gained by comparing the average level of protein sequence difference between S. cerevisiae and other yeasts, to that between human and other animals (Dujon et al. 2004). By this measure, the divergence between S. cerevisiae and Candida glabrata is slightly greater than that between humans and fish, while the divergence between S. cerevisiae and Yarrowia lipolytica (the deepest-branching hemiasco-mycete whose genome has been sequenced) is about the same as that between human and the sea squirt Ciona (Dujon et al. 2004). Similarly, the divergences among the Saccharomyces sensu stricto species are comparable to those among different orders of mammal. These levels of divergence are surprisingly high and demonstrate the evolutionary depth that is encompassed by the available fungal genome sequences. They also suggest that we should be cautious about inferring that the function of a gene in one species will be exactly the same as that of its

ortholog in *S. cerevisiae* (for an example of an exception, see Kadosh & Johnson 2001).

2. THE YEAST GENOME BROWSER

Nested within the phylogenetic tree of hemiascomy-cetes is a whole genome duplication (WGD) event (figure 1). This event was first detected by analysis of the S. cerevisiae genome sequence alone, which showed that in many places in the genome a series of genes on one chromosome had a series of paralogs on another chromosome, usually in the same order along the chromosome and with conserved transcriptional orien-tation (Wolfe & Shields 1997). We identified 55 such duplicated regions in the S. cerevisiae genome, and found that the centromere-to-telomere orientation was almost always conserved in each pair of regions. We interpreted this as evidence that entire duplicated chromosomes had existed at some stage during yeast's evolutionary past, and that the mosaic of duplicated



271 Figure 2. Illustration of our model of gene order evolution 272 following whole-genome duplication (WGD). The box at the top shows a hypothetical region of chromosome containing 273 ten genes numbered 1-10. After WGD, the whole region is 274 briefly present in two copies. However, many genes 275 subsequently return to single-copy state because there is no 276 evolutionary advantage to maintaining both copies. In this 277 example, only genes 1, 6 and 10 remain duplicated. However, 278 the arrangement of these three homolog pairs in the post-279 WGD species (bottom) would be sufficient to allow the sister 280 regions to be detected using that genome sequence alone. 281 Also, the order of genes in sister regions in post-WGD species 282 have well-defined relationships to the gene order that existed 283 in the pre-WGD genome (top), which will also be similar to the gene order seen in any species that diverged from the 284 WGD lineage before the WGD occurred. Modified from 285 Montcalm & Wolfe (in press). 286

288 regions currently identifiable were the result of 289 subsequent interchromosomal rearrangements, princi-290 pally reciprocal translocations (Wolfe & Shields 1997; 291 Keogh et al. 1998; Seoighe & Wolfe 1998). Within any 292 pair of duplicated regions, the paralogous genes were 293 interspersed with many single-copy genes (figure 2), 294 which indicated that large numbers of genes had been 295 lost (deleted) from the genome after WGD. The 296 limited amount of gene order information available at 297 that time from other yeast species showed that some 298 species, such as Kluyveromyces lactis and Saccharomyces 299 kluyveri, had gene orders consistent with them having 300 diverged from the S. cerevisiae lineage before WGD 301 occurred in the latter, as shown in figure 1 (Keogh et al. 302 1998; Seoighe & Wolfe 1999). These findings were 303 comprehensively confirmed in 2004 when the complete 304 genome sequences of Ashbya gossypii (Dietrich et al. 305 2004), Kluyveromyces waltii (Kellis et al. 2004) and 306 K. lactis (Dujon et al. 2004) were published. Each of 307 these species shows a gene order similar to that inferred 308 to have existed in an ancestor of S. cerevisiae 309 immediately before the WGD happened. 310

In order to visualize the synteny relationship among 311 the sequenced yeast genomes, our laboratory recently 312 developed a bioinformatics tool, the Yeast Gene Order 313 314 Browser (YGOB; http://wolfe.gen.tcd.ie/ygob; Byrne & 315 Wolfe in press). YGOB was designed in particular 316 to handle the 1:2 genomic relationship between 317 'pre-WGD' species and 'post-WGD' species (figure 3). 318 In this shorthand notation, post-WGD means the 319 group of species whose common ancestor underwent 320 WGD, and pre-WGD means any species that diverged

from the lineage leading to S. cerevisiae before the 321 WGD happened. Under the WGD hypothesis, any 322 genomic region in a pre-WGD species should have two 323 counterparts in each post-WGD species; we refer to the 324 paired regions in the post-WGD species as 'sister 325 regions'. At present YGOB includes genome sequence 326 data from three post-WGD species (S. cerevisiae, 327 S. castellii and C. glabrata), and four pre-WGD species 328 (K. waltii, K. lactis, A. gossypii and S. kluyveri). The 329 database underlying YGOB consists of sets of hom-330 ologous genes from each species, corresponding to the 331 columns of genes shown in figure 3. Each set (which we 332 refer to as a 'pillar') can maximally consist of one gene 333 from each pre-WGD species (i.e. orthologs) and two 334 genes (a pair of paralogs) from each post-WGD 335 species. The assignments of orthology in YGOB were 336 based initially on BLAST results and on the annota-337 tions provided by the sequencers of each genome, but 338 the pillars have been refined subsequently by extensive 339 manual editing and curation. The strong conservation 340 of gene order among this group of yeast species has the 341 consequence that it is generally straightforward to 342 identify the orthologs of any locus in each species, or to 343 identify sites from which genes have been deleted 344 (figure 3). The number of singleton genes (i.e. genes 345 without a syntenic ortholog in at least one other 346 species) in each species in YGOB is less than 11% 347 (Byrne & Wolfe in press), and many of these are in 348 genomic regions close to telomeres (see below). 349

For on-screen presentation of gene order information, the user selects one gene (from any species) to focus on. This gene is shown in the central pillar of the display, and other aspects of the display (such as gaps caused by gene deletions/insertions, and interchromosomal rearrangements in any species) are calculated relative to it. The user can walk along any chromosome in any species by sequentially focusing on neighbouring genes. Details of the data and visualization algorithms in YGOB are given in Byrne & Wolfe (in press).

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One unanticipated outcome of developing YGOB 361 was the discovery of several loci that are evolving very 362 rapidly. One example is YKL023W, an S. cerevisiae gene 363 whose function is unknown but which encodes a 364 protein that co-purifies with the variant histone Htz1 365 (Krogan et al. 2003). None of the other published yeast 366 genomes, except for the S. sensu stricto species, included 367 a gene annotated as an ortholog of YKL023W. When 368 we curated the original set of genomes in YGOB, we 369 370 noticed that A. gossypii, K. lactis, K. waltii and 371 C. glabrata were each annotated as having singleton 372 genes in the interval between their orthologs of URA6 and CDC16, which corresponds to the position of 373 374 YKL023W in S. cerevisiae (figure 3). None of these singletons hits any of the others in a BLASTP search 375 (i.e. the BLASTP E-value between any of them is 376 >10), but their coincident locations and identical 377 transcriptional orientations suggested that they were in 378 fact orthologs whose sequence had diverged too much 379 to be detectable by BLAST. After noticing this we 380 381 examined the S. castellii genome sequence and found Q1 that it too contains a large ORF in the same region. 382 383 The S. castellii gene does hit some of the others in 384 BLASTP searches, but only very weakly. Multiple



Figure 3. Screenshot from the Yeast Gene Order Browser (YGOB; Byrne & Wolfe in press). The image shows how gene order is related between two sister genomic regions in each of the post-WGD species *S. cerevisiae* (chromosome XI genes in dark blue and chromosome IX genes in light blue), *S. castellii* and *C. glabrata*, and the single homologous genomic region in the pre-WGD species *A. gossypii*, *K. lactis* and *K. waltii*. In this representation, each rectangle represents a gene and homologs are arranged as vertical columns. Arrows below the rectangles show transcriptional orientation. Gray bars connect genes that are adjacent but do not indicate the actual gene spacing on the chromosome. The large arrow indicates the rapidly evolving locus *YKL023W* discussed in the text.

alignment of the proteins shows that only a short region
at the N-terminus of this large protein is conserved
among species, but this is sufficient to recognize that
the genes are orthologs, which is why we show them in
the same column in figure 3. Other examples of rapidly
evolving loci are discussed in Wolfe (2004) and Byrne &
Wolfe (in press).

427 3. PAIRS OF CENTROMERES

We have also been able to use YGOB to investigate the evolutionary changes in chromosome number that occurred before and after the WGD (Montcalm & Wolfe in press). Genome duplication can occur either by autopolyploidization (doubling of the chromosome set of a species) or by allopolyploidization (hybridiz-ation between two species). In the former case the number of chromosomes should be doubled, and in the latter the new species should have the sum of the numbers of chromosomes in its parents. The process of gene loss sketched in figure 2 should not alter the chromosome number. The number of chromosomes in post-WGD species is approximately twice that in pre-WGD species (Keogh et al. 1998) but the arithmetic is not precise. In the pre-WGD species, there are 8 (K. waltii and S. kluyveri), 7 (A. gossypii) or 6 (K. lactis) chromosomes (Neuveglise et al. 2000; Dietrich et al. 2004; Dujon et al. 2004; Kellis et al. 2004). In the post-WGD species there are 16 (S. cerevisiae and the other sensu stricto species), or 13 (C. glabrata; Dujon et al. 2004). We suspect that

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the estimate of nine chromosomes in the post-WGD species S. castellii (Petersen et al. 1999) is an underestimate because it is based on pulsed-field electrophoresis (a technique that often tends to under-count); the genome sequence for this species (Cliften et al. 2003) is a $4 \times$ draft consisting of numerous contigs rather than complete chromosome sequences.

The number of chromosomes in a species is determined by the number of centromeres, so we used YGOB to examine the history of yeast centromeres and the protein-coding genes near them (figure 4; Montcalm & Wolfe in press). The 16 centromeres of S. cerevisiae can be arranged into eight pairs based on the sister relationships resulting from the WGD (Wong et al. 2002), and these are in conserved gene order relationships with the eight centromeres of K. waltii (Kellis et al. 2004). This observation shows that the WGD involved an 8-chromosome ancestral genome doubling (or two 8-chromosome species hybridizing) to form a 16-chromosome descendant. The other changes in chromosome number (reduction from 8 to 7 in A. gossypii; reduction from 8 to 6 in K. lactis; reduction from 16 to 13 in C. glabrata) can all be attributed to chromosome fusions. Each of these fusions seems to have been accompanied by the inactivation of one of the two centromeres in the newly formed dicentric chromosome (figure 4), without other changes in the local gene order

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513	VCI VCD		577
514	$\operatorname{Sc} 7 \longrightarrow \operatorname{FGL} O \longrightarrow \operatorname{FGR}$	Sc 4 <u>TDL</u> <u>TDR</u>	578
515	Cg 1 - A00627	Cg 13 M10463 M10439	579
516	A00781	VBI VBR	580
517	Sc 1 $ -$	$\operatorname{Sc} 2 \longrightarrow \operatorname{CO4267} \longrightarrow \operatorname{CO4222}$	581
518	Cg 7	Cg 3 - C04307 - C04323	582
519	s47 17596 s47 17598	s27 10006 s27 10001	583
520	Kw 4	$Kw 2 - \frac{S27_{10990}}{1000} + \frac{S27_{10991}}{1000}$	584
521	Ag 6 <u>AFL</u> <u>AFR</u>	Ag 5 <u>AEL AER</u>	585
522	KI 3 C18513	$E17369 \times E17325$	586
523			587
524			588
525	Sc 14 YNL YNR	Sco YIL YIR	589
526	$H_{13949} = H_{13949}$	50.9 - 102156 - 102200	590
527	$Cg 8 \longrightarrow 100000$	Cg 10 O 0	591
528	Sc 3 YCL O YCR	Sc 5 YEL YER	592
529	B04675	Se 5 H09768	593
530	Cg 2		505
532	Kw 8 s0_281 s0_284	Kw 6 s55_20901 s55_20903	595 506
533	A 1 AAL AAR	A ADL ADR	590
534	Ag 1	Ag 4	598
535	Kl 1	KI 4 DI 3810 DI 3772	599
536			600
537			601
538	Sc 16 $\underline{\qquad YPL \qquad \bigcirc \qquad YPR \qquad }$	Sc 11 $\underline{\qquad}$ YKL $\underline{\qquad}$ YKR	602
539	Cg 12 L09042 L09086	Cg 12 L02321 L02299	603
540			604
541	Sc 6 \underline{YFL} $$ \underline{YFR}	Sc 8 YHL YHR	605
542	Cg 6 $-\frac{F02343}{O} = \frac{F02299}{F02299}$	Cg 11 $- \frac{K02849}{0} - \frac{K02893}{0}$	606
543		-	607
544	Kw 1 825_5150 825_5158	Kw 3 $\underline{\hspace{0.5mm}}^{s26}\underline{\hspace{0.5mm}}^{/923}\underline{\hspace{0.5mm}}^{s26}\underline{\hspace{0.5mm}}^{/924}\underline{\hspace{0.5mm}}^{/924}$	608
545	Ag 7 <u>AGL</u> AGR	Ag 2 <u>ABL</u> <u>ABR</u>	609
546	E14344 E14300	KI 2 B13277 B13321	610
547			611
548			612
549	Sella YLL YLR	So 15 YOR YOL	613
551	D05808 D05874 ??	5015 - 602585 = 602541	615
552	Cg4 — O — $D03000$ — $D03074$	Cg 5 -	616
553	Sc 10 YJL YJR	Sc 13 YML YMR	617
554	$D03058 \qquad D03036$		618
555	Cg 4 — X — — — — — — — — — — — — — — — — —	Cg 9	619
556	Kw 5 s33_15200 s33_15203	Kw 7 s56_22443 s56_22447	620
557	< AAL174C $>$ ACR029C	A 2 ACR ACL	621
558	Ag El2802	Ag 5 D01200	622
559	Kl 6	Kl 4 \longrightarrow D01267 \longrightarrow D01309	623

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Figure 4. Relationships among centromeric regions of S. cerevisiae (Sc), C. glabrata (Cg), K. waltii (Kw), A. gossypii (Ag) and K. lactis (Kl) chromosomes. Each of the eight panels shows a group of genomic regions that are related by virtue of their gene 562 contents close to a centromere (or a similar noncentromeric region). Black circles represent centromeres in pre-WGD species, 563 and white circles represent centromeres of post-WGD species. X symbols indicate the absence of a centromere. Names indicate 564 chromosome arms, or genes close to the centromere on each arm. Hooked lines indicate loss of relatedness. The assignment of 565 the centromere of C. glabrata chromosome 9 to the group on the bottom-right is less certain than for the other centromeres, and 566 is based on the linkage of CgCEN9 to the gene CAGL0108107g, which is an ortholog of the genes KLLA0D01243g (K. lactis) and s56_22439 (K. waltii) which are close to the corresponding regions in those species. Gene names in K. lactis and C. glabrata have 567 been shortened by writing A00627 instead of CAGL0A00627g, etc. Reproduced with permission from Montcalm & Wolfe 568 (in press). 569

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(Montcalm & Wolfe in press). It is noteworthy that, 572 despite the growing number of hemiascomycete 573 574 genomes that have been sequenced, no examples of 575 a gain of a centromere by a mechanism other than 576 WGD have yet been observed.

4. EVOLUTION OF GENE CONTENT

'Use it or lose it' is one of the truisms of genome 637 evolution. If a gene's function is not beneficial to the 638 organism, there will be no natural selection against 639 mutations that damage that gene. It will become a 640

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Figure 5. Gene displacement during evolution of the uracil biosynthesis pathway. The hemiascomycete species are indicated by the blue background. Red and green rectangles represent the URA9 and URA1 genes, respectively, and red and green X symbols on the phylogenetic tree represent losses of those genes. The inferred point of gain of URA1 is shown by the arrow. All three possible outcomes of this horizontal gene transfer have occurred in different lineages descended from it: loss of URA9 (gene displacement, e.g. in S. cerevisiae); loss of URA1 (return to the original state, e.g. in C. glabrata); or retention of both genes (e.g. in S. kluyveri). Based on Gojkovic et al. (2004) and Hall et al. (2005).

pseudogene and eventually disappear from the gen-ome. For example, the plastid genomes of non-photosynthetic plants do not contain photosynthesis genes (Wolfe et al. 1992). Within fungi, the hemi-ascomycetes have lost several genes with functions related to splicing (Aravind et al. 2000) and to RNA interference and heterochromatin formation (Alexan-dersson & Sunnerhagen in press), as compared to euascomycetes and basidiomycetes. These losses are presumably attributable to the relative lack of introns and the simple centromeres in hemiascomycetes. Although the great majority of genes in any hemi-ascomycete species have homologs in each other species, a few examples of recent gene losses (i.e. losses within the hemiascomycetes) have come to light in the past year. Comparative genomics enables us to detect these losses, and they are particularly interesting because in some cases we can infer (or at least guess at) the physiological or ecological changes that rendered the genes unnecessary.

Most hemiascomycetes can grow on galactose as a sole carbon source, and in S. cerevisiae seven genes (the GAL genes) function exclusively in this pathway. Hittinger et al. (2004) showed that, although orthologs of the GAL genes are present in most of the sequenced

yeast genomes, they have been lost independently in three or four lineages. In S. kudriavzevii all seven GAL loci are pseudogenes indicating very recent gene losses and hence a recent change in the metabolic capacity of this species, uniquely among the S. sensu stricto species. Likewise, the GAL genes are completely absent from three other yeasts (C. glabrata, A. gossypii and K. waltii). We do not really know what ecological changes permitted the loss of this pathway in each case, but it is perhaps most readily understood in the case of C. glabrata which (unlike most other hemiascomycetes) is a pathogen of mammals and is unlikely to encounter galactose in this environment.

C. glabrata has also lost the pathway of de novo synthesis of nicotinic acid (the kynurenine pathway; BNA genes), a loss that is directly linked to the pathogenesis of this species (Domergue et al. 2005). In both C. glabrata and S. cerevisiae, the histone deacetylase Sir2 is regulated by the availability of NAD⁺ because Sir2 consumes NAD⁺ when it modifies histones. Nicotinic acid is a precursor of NAD^+ , so S. cerevisiae can replenish its pool of NAD^+ via de novo nicotinic acid synthesis, but C. glabrata cannot. Consequently C. glabrata Sir2 activity depends on the availability of external nicotinic acid. The



Figure 6. Formation of the *DAL* gene cluster during hemiascomycete evolution. The six genes are unlinked in the species shown at the bottom of the phylogenetic tree (from *S. kluyveri* to *Y. lipolytica*). In *S. castellii* and the *S. sensu stricto* species, the six genes are adjacent and located close to a telomere. There is a rearrangement of gene order in the cluster in *S. castellii* relative to the *sensu stricto* species. Formation of the cluster involved apparent transposition of the genes *DAL1*, *DAL2*, *DAL3* and *DCG1*, and duplication of *FUR4* (to form *DAL4*) and *MLS1* (to form *DAL7*). None of the *DAL* cluster genes are present in *C. glabrata*. Modified from Wong & Wolfe (2005).

pathogen appears to exploit this dependence as a way of activating a cell surface adhesion program when it enters the urinary tract of a mammal. Urine is a poor source of nicotinic acid, so C. glabrata cells located in the urinary tract have low Sir2 activity. Consequently the adhesin (EPA) genes, whose transcription is normally repressed owing to Sir2-mediated histone modification, become activated and enable C. glabrata to adhere (Domergue et al. 2005). The loss of BNA genes in C. glabrata may be an example of a pathway loss that was advantageous rather than neutral.

Like redundant factory workers, genes can also be shed from a genome if their function is outsourced instead of being shut down completely. This has occurred in the uracil synthesis pathway in some hemiascomycetes (Gojkovic et al. 2004; Zameitat et al. 2004; Hall et al. 2005). In the deeper-branching hemiascomycetes (figure 5), and in all other eukar-yotes, the enzyme dihydroorotate dehydrogenase (DHODase; product of the URA9 gene) is located in the mitochondrion and its activity is coupled to the mitochondrial respiratory chain because a quinone is the terminal electron acceptor. At one point during hemiascomycete evolution, however, an alternative DHODase enzyme was gained by means of horizontal gene transfer from a bacterium similar to Lactococcus lactis (Gojkovic et al. 2004; Hall et al. 2005). The new DHODase (product of the URA1 gene) is located in the cytosol. Because it uses fumarate as an electron acceptor, uracil biosynthesis can proceed even when the cell is not respiring. In some other yeasts such as *S. kluyveri*, the two enzymes coexist (figure 5). In *S. cerevisiae*, only *URA1* is present and the ortholog of the ancestral eukaryotic gene *URA9* has been completely lost.

Piskur & Langkjaer (2004) proposed that the resulting decoupling of uracil biosynthesis from respiration was an important step in allowing *S. cerevisiae* and its close relatives to develop the ability to grow almost completely without oxygen. This theme was expanded by Hall *et al.* (2005) who carried out a systematic search for genes horizontally transferred from bacteria into the *S. cerevisiae* lineage. They identified 10 *S. cerevisiae* genes that are candidates for having been gained by horizontal transmission, including *URA1* as described above. Notably, for 7 of the 10 genes the most closely related prokaryotic sequences are in anaerobic bacteria. It is also interesting that 9 of the 10 putatively transferred genes are located near telomeres in *S. cerevisiae*.

A similar displacement, but over a longer evolution-ary period, has occurred at the mating-type (MAT) loci of ascomycetes. Comparison of the genes flanking the loci that have been demonstrated genetically to be the MAT loci of Yarrowia lipolytica (a hemiascomycete), Neurospora crassa and Gibberella zeae (two euascomy-cetes) shows that these loci are positionally as well as functionally homologous. That is, the locus that specifies mating type has not moved in these genomes during all the time that has elapsed since

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897 hemiascomycetes and euascomycetes diverged from 898 their common ancestor. The MAT genes in these three species are flanked by the gene APN2 on one side and 899 900 SLA2 on the other (Butler et al. 2004). APN2 and 901 SLA2 code for a DNA repair enzyme and a component 902 of the cytoskeleton, respectively, neither of which appear to have any functional connection to mating 903 904 type determination. Within the hemiascomycetes, 905 several rearrangements of the genes flanking the MAT locus have occurred but a continuous line of descent 906 can still be traced (Butler et al. 2004). The positional 907 908 continuity of the MAT locus is remarkable because the 909 actual contents of this locus have changed considerably. 910 A gene for an alpha-domain protein (homologous to S. 911 cerevisiae $MAT\alpha 1$) is the only gene invariably present at 912 the MAT locus in all ascomycetes. The euascomycetes 913 have in addition genes for two HMG-domain proteins 914 (mat-a1 and mat-A3 in Neurospora) and a coiled-coil 915 protein (mat-A2), whereas S. cerevisiae encodes two 916 homeodomain proteins (MAT α 2 and MATa1). How 917 one type of DNA-binding protein displaced the other 918 during the evolution of this locus remains a mystery.

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921 5. EVOLUTION OF GENE ORDER

922 As implied in figure 2, and shown for one small region 923 of the genome in figure 3, gene order is generally well 924 conserved among the hemiascomycetes included in 925 YGOB, once the effects of the WGD are taken into 926 account (i.e. formation of sister regions and numerous 927 gene deletions within each sister). In addition, synteny 928 is interrupted by species-specific interchromosomal 929 translocations. In designing YGOB we chose not to 930 include species that are more distant from S. cerevisiae 931 (for example, Candida albicans) because previous 932 analyses have shown that gene order is noticeably 933 more poorly conserved at this depth (Keogh et al. 1998; 934 Llorente et al. 2000; Dujon et al. 2004).

935 In comparing gene order among species, we found a 936 striking example of relocation of a set of genes during 937 recent hemiascomycete evolution (figure 6; Wong & 938 Wolfe 2005). The movement of these genes contrasts 939 starkly with the syntenic stasis seen at most other loci. 940 In S. cerevisiae, six of the eight genes involved in 941 allantoin degradation form a physical gene cluster 942 (called the DAL cluster). Allantoin is a degradation 943 product of purines and can be used by S. cerevisiae as a 944 non-preferred nitrogen source. The DAL genes are also 945 clustered in the other S. sensu stricto species, and in 946 S. castellii. Homologs of the six genes are present in the 947 other hemiascomycetes but they are found at six 948 separate chromosomal locations. The cluster origi-949 nated at one particular point in the phylogenetic tree 950 (on the branch leading to the common ancestor of 951 S. cerevisiae and S. castellii) and is located in a 952 subtelomeric region of the genome. Two of the 953 clustered genes (DAL4 and DAL7) are duplicates of other genes located elsewhere in the S. cerevisiae Q3 954 955 genome. The other four DAL genes appear to have 956 simply transposed to the cluster site, but we suspect 957 that they originated by gene duplications (forming 958 subtelomeric copies), followed by deletion of the 959 original genes. The set of genomic rearrangements 960 that produced the cluster seems so improbable

(figure 6) that one can only conclude that random rearrangements that by chance moved the genes to a subtelomeric location were strongly favoured by natural selection, and that each incremental addition of one gene to the cluster was individually advantageous. We have found that the DAL cluster was formed approximately simultaneously with a reorganization of the first steps in the allantoin degradation pathway: species that have the DAL cluster obtain their allantoin by importing it from outside the cell (using the newly formed allantoin permease, Dal4) instead of by oxidation of urate (as is done in species without the cluster). This finding has suggested to us that, like the displacement of the DHODase gene (figure 5), the evolutionary genomic changes were the result of selection to minimize oxygen consumption during the evolution of the 'petite-positive' subset of hemiascomycetes (Piskur 2001; Wong & Wolfe 2005). This demonstration that natural selection can occasionally flex its muscles and reshape part of the genome suggests that the arrangement of genes elsewhere in fungal genomes might be less random than at first appears.

6. CONCLUSIONS AND PROSPECTS

Comparative genomics enables us to identify the parts of genomes that have changed during recent evolution, which gives us an indication of the evolutionary processes that are currently moulding the genome. Although the overall impression in most parts of the genome is one of strings of genes whose order is stable but that are occasionally disrupted by translocations or other types of rearrangement, the dynamic history glimpsed at some other loci shows that the genome is not a passive letter-rack of genes but an organelle whose form is shaped by its function. Increasing attention is being focused on subtelomeric regions because they seem to be particularly receptive to picking up genes that are newly transferred from bacteria or newly relocated from elsewhere in the genome. Whether this is owing to a mutational cause (e.g. a higher rate of illegitimate DNA recombination in subtelomeres) or a selective cause (e.g. preferential retention of newly integrated DNA if it is located at subtelomeres because it can be regulated effectively by means of histone modification) remains to be seen.

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