

Reinvestigation of the *Saccharomyces cerevisiae* genome annotation by comparison to the genome of a related fungus: *Ashbya gossypii*

Sophie Brachat^{*}, Fred S Dietrich^{*†}, Sylvia Voegeli^{*}, Zhihong Zhang[†], Larissa Stuart[†], Anita Lerch^{*}, Krista Gates[‡], Tom Gaffney[‡] and Peter Philippsen^{*}

Addresses: ^{*}Institute of Applied Microbiology, Biozentrum der Universität Basel, Klingelbergstrasse 70, CH-4056 Basel, Switzerland. [†]Department of Molecular Genetics and Microbiology, Duke University Medical Center, Durham, NC 27710-3568, USA. [‡]Syngenta, Research Triangle Park, NC 27709, USA.

Correspondence: Peter Philippsen. E-mail: Peter.philippsen@unibas.ch

Published: 25 June 2003

Genome Biology 2003, 4:R45

The electronic version of this article is the complete one and can be found online at <http://genomebiology.com/2003/4/7/R45>

Received: 13 February 2003

Revised: 7 May 2003

Accepted: 28 May 2003

© 2003 Brachat et al.; licensee BioMed Central Ltd. This is an Open Access article: verbatim copying and redistribution of this article are permitted in all media for any purpose, provided this notice is preserved along with the article's original URL.

Abstract

Background: The recently sequenced genome of the filamentous fungus *Ashbya gossypii* revealed remarkable similarities to that of the budding yeast *Saccharomyces cerevisiae* both at the level of homology and synteny (conservation of gene order). Thus, it became possible to reinvestigate the *S. cerevisiae* genome in the syntenic regions leading to an improved annotation.

Results: We have identified 23 novel *S. cerevisiae* open reading frames (ORFs) as syntenic homologs of *A. gossypii* genes; for all but one, homologs are present in other eukaryotes including humans. Other comparisons identified 13 overlooked introns and suggested 69 potential sequence corrections resulting in ORF extensions or ORF fusions with improved homology to the syntenic *A. gossypii* homologs. Of the proposed corrections, 25 were tested and confirmed by resequencing. In addition, homologs of nearly 1,000 *S. cerevisiae* ORFs, presently annotated as hypothetical, were found in *A. gossypii* at syntenic positions and can therefore be considered as authentic genes. Finally, we suggest that over 400 *S. cerevisiae* ORFs that overlap other ORFs in *S. cerevisiae* and for which no homolog can be detected in *A. gossypii* should be regarded as spurious.

Conclusions: Although, the *S. cerevisiae* genome is rightly considered as one of the most accurately sequenced and annotated eukaryotic genomes, we have shown that it still benefits substantially from comparison to the completed sequence and syntenic gene map of *A. gossypii*, an evolutionarily related fungus. This type of approach will strongly support the annotation of more complex genomes such as the human and murine genomes.

Background

A major breakthrough in the field of genomics came with the publication of the 13 Mb genome of the budding yeast *Saccharomyces cerevisiae* [1], which was the first eukaryotic

genome to be fully sequenced and annotated. Since then, DNA sequencing has developed with an increasing speed, and sequences of much larger genomes, such as those of *Caenorhabditis elegans* [2], *Drosophila melanogaster* [3],

Arabidopsis thaliana [4], *Homo sapiens* [5,6], *Anopheles gambiae* [7] and *Mus musculus* [8] have been published. However, increased sequencing capacity was not matched by a corresponding development in annotation and the gene annotation process is now the rate-limiting step in whole-genome sequencing projects. Despite progress in gene prediction programs, comparisons to expressed sequence tag (EST) databases and to genomic sequences, preferably of related organisms, is still the most favored approach to the annotation of complex genomes.

The original annotation of the *S. cerevisiae* genome was especially challenging because, at the time of its completion [9], only limited genomic sequence information from other eukaryotes was available. Despite the functional characterization of a large number of orphan open reading frames (ORFs) and several efforts to re-evaluate the sequence at the gene level or for an entire chromosome [10,11], a significant number of uncertainties still remain. It is, for example, not known whether all protein-coding genes have been identified and which of the close to 2,000 genes annotated as hypothetical represent real genes. A careful comparison to a related genome should help clarify several of these issues.

The recently completed genome sequence of the filamentous ascomycete *Ashbya gossypii* revealed an unexpected high degree of gene homology and gene order conservation with *S. cerevisiae* (F.S.D., S.V., S.B., A.L., K.G., C. Mohr, S. Steiner, P. Luedi, T.G. and P.P., unpublished work). The two species diverged more than 100 million years ago, and both genomes differ substantially in their GC content (38.3% in *S. cerevisiae* and 51.9% in *A. gossypii*). However, 95% of the 4,700 *A. gossypii* protein-coding genes were found to have a homolog in *S. cerevisiae* and 90% of these homologous genes map at syntenic positions. Despite these striking genomic similarities, the average conservation at the DNA level is 55% in coding regions but drops to 33% in noncoding regions. Thus, significant sequence similarities are restricted to coding regions. Altogether, these findings open up the possibility of a whole-genome reinvestigation of the *S. cerevisiae* annotation.

We carried out an extensive search for homology at the amino-acid level between *A. gossypii* coding regions and *S. cerevisiae* 'annotation-free' regions: stretches of sequence bearing no annotated genomic features such as ORFs, RNA genes, or transposable elements. Focusing on syntenic regions, we identified a total of 95 inconsistencies, suggesting the following four types of changes in the *S. cerevisiae* annotation: novel genes, novel introns, potential ORF extensions, and neighboring ORF fusions. Furthermore, we provide evidence that information from the complete *A. gossypii* genome is also a major resource for recognizing real genes among the numerous *S. cerevisiae* hypothetical ORFs.

Results and discussion

We searched for homology at the amino-acid level between annotated *A. gossypii* coding regions and *S. cerevisiae* 'annotation-free regions'. As a result, we identified 95 regions in the *S. cerevisiae* genome, which had not been annotated as protein coding, that showed both homology and synteny to *A. gossypii* genomic sequences. In this context, synteny refers to a relaxed synteny (loose synteny), which results from several hundred genomic rearrangements in the *A. gossypii* and *S. cerevisiae* lineages and from frequent loss of one of the two gene copies (twin genes) in *S. cerevisiae* after the proposed doubling of the genome [12,13]. As a result, all remaining duplicated genes in *S. cerevisiae* have a single homolog in *A. gossypii* (F.S.D., S.V., S.B., A.L., K.G., C. Mohr, S. Steiner, P. Luedi, T.G. and P.P., unpublished work). On close inspection of these 95 *S. cerevisiae* syntenic loci, we found evidence for novel ORFs, and for substantial boundary changes of annotated ORFs. Figure 1 outlines the categories of changes suggested by this comparative genomics approach.

We first present data supporting novel protein-coding genes and provide detailed analysis of the different types of boundary changes of annotated ORFs due to novel exons, 5'- or 3'-end extension, or even fusion of adjacent ORFs. Second, we will focus on the validation of the approximately 2,000 hypothetical ORFs. We will present evidence that 50% of these hypothetical ORFs are real, and provide arguments to consider several hundred as probably spurious.

Novel ORFs

In 23 annotation-free regions, we discovered homology to syntenic small *A. gossypii* ORFs as outlined in Figure 1a and summarized in Table 1. These presumptive novel *S. cerevisiae* ORFs are 52 to 134 codons long. Twenty have a size below 100 codons, the arbitrary cut-off for small and nonhomologous yeast ORF annotation, several contain an intron, and one contains two introns. The short length and the presence of introns explain why these ORFs remained so far undiscovered. An additional example of a novel *S. cerevisiae* ORF identification by comparison to the *A. gossypii* genome was recently published [14].

We carried out homology searches for all novel ORFs against the available fungal databases. This analysis revealed that all but one of the novel ORFs are present in hemiascomycetes and for 15 ORFs, homologs were found in at least two of the following databases: hemiascomycetes, *Candida albicans*, *Schizosaccharomyces pombe* and *Neurospora crassa* (Table 1). This suggests that they represent conserved fungal proteins. For two genes, YMR194C-B and YNLO24C-A, we identified homologs in higher eukaryotes, including mouse and human. The conservation in other species, and particularly their syntenic positions in *A. gossypii*, strongly support the authenticity of these novel genes.

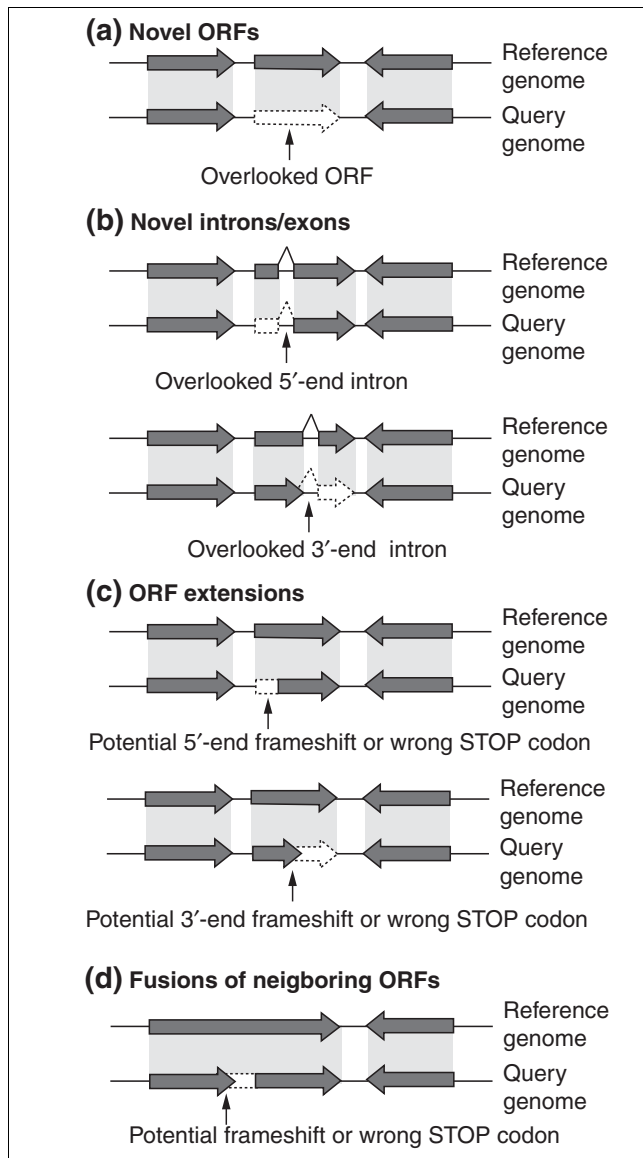


Figure 1
Genome reinvestigation using comparative genomics. Translated DNA comparison allows the detection of homology outside annotated features of a query genome. This can lead to the detection of (a) novel ORFs, (b) novel introns/exons, (c) ORF extensions, and (d) ORF fusions (fusion of adjacent ORFs or merging of overlapping ORFs with same transcription direction). Gray areas illustrate regions of homology at the protein level and dashed lines depict suggested modifications in the query genome. Cases of changed splice sites may also be detected but are not drawn here.

We screened the 23 protein sequences for the presence of known domains but did not find any significant hits. These novel *S. cerevisiae* ORFs were not deleted by the yeast gene deletion consortium [15] but the *A. gossypii* homologs of YIL156W-B, YJL127C-B, YMR194C-B and YNL138W-A have been deleted (K.G. and T.G., unpublished data). One deletion is lethal; the others did not exhibit any apparent phenotype under normal growth conditions. Recently, two of the novel

ORFs - YPL096C-A (*ERI1*, ER-associated Ras Inhibitor 1) and YKL138C-A (*HSK3*, Helper of *Ask1*) - were added to the *Saccharomyces* Genome Database (SGD) as reserved gene names, indicating unpublished functional data, and one novel ORF, YPR036W-A, was shown to be expressed in response to drug treatment [16].

A similar approach based on the so-called Génolevures project, a partial shotgun sequencing of 13 hemiascomycete genomes [17], suggested the presence of 50 overlooked ORFs in *S. cerevisiae*, distinct from the set of 23 described in this paper. These 50 ORFs were recently incorporated in the SGD. Arguing that the species under consideration were too closely related, Wood *et al.* [18] recommended further investigations before considering these 50 novel ORFs as real. Having the *A. gossypii* genome to hand allowed us to evaluate the authenticity of these proposed ORFs. Indeed, we found 20 of the suggested novel ORFs at syntenic positions (see Additional data files). Similarly, the comparison between the *S. pombe* and the *S. cerevisiae* genome annotations identified three additional novel *S. cerevisiae* ORFs [18,19], distinct from the 23 ORFs discussed above. All three correspond to syntenic homologs in *A. gossypii*, which confirms the assumption that they are real ORFs. More recently, 84 *S. cerevisiae* small ORFs, called smORFs, were identified on the basis of homology to a larger fungal database and experimental evidence for transcription products [20]. Upon re-evaluation, we found that five smORFs correspond to novel ORFs described here and five others match sequences of ORF extensions as discussed below. Several smORFs correspond to RNA genes or match the opposite strand of previously annotated ORFs in both *S. cerevisiae* and *A. gossypii* and thus do not represent protein-coding genes. For the remaining smORFs, there were no homologs found in *A. gossypii*.

Novel introns and exons

Splicing rules and intron positions are generally conserved in *A. gossypii* and *S. cerevisiae*. On this basis we were able to identify 13 cases of probably overlooked introns in *S. cerevisiae*, as schematically represented in Figure 1b. Splicing of the novel introns and fusion of the novel exon extend the *S. cerevisiae* ORFs up to 236 codons and lead in most cases to substantially increased similarity between homologs of the two species (data not shown). The ORFs under consideration, the overall size increases, and other supporting evidence are shown in Table 2, which summarizes all 72 ORF extensions. Perfect splice consensus sequences were found for only three genes, which explains the difficulty in recognizing these introns. Finally, for one gene, *SEF1* (YBLO66C), we propose a base-pair change in addition to an intron. We tested the authenticity of the proposed introns for YKRO04C (*ECM9*), YML017W (*PSP2*) and YOLO48C using 5' rapid amplification of cDNA ends (5' RACE). In all three cases, the intron could be confirmed by sequencing the cDNA obtained (AY245791, AY245792, and AY245793). cDNA and genomic sequence alignment confirms that the intron of YKRO04C is spliced at

comment
reviews
reports
deposited research
referred research
interactions
information

Table 1**Novel *S. cerevisiae* ORFs identified by homology and synteny to *A. gossypii* ORFs**

<i>A. gossypii</i> ORF		Novel <i>S. cerevisiae</i> ORF(s)		% Similarity	Homologs*			
Name	Size	Name(s)	Size(s)		Hemiascomycetes	<i>C. albicans</i>	<i>S. pombe</i>	<i>N. crassa</i>
AGL322W	57	YBL039W-A	59	53.70	x			
ADL343C	99	YBR111W-A	98	65.28	x	x		
AAL005W	72	YCL005W-A	73	79.17	x	x		x
AFR743C-A	132	YCL058W-A	132	45.00	x			
AER271W	81	YCR075W-A/YNR034W-A†	75/99	44.00/46.15	x			
ACL158W	95	YDL160C-A	80	55.70	x			
AGR097W-A	66	YER180C-A	85	83.70	x			
AFR298C	73	YIL156W-B	73	71.23	x	x	x	
AFL216C	49	YJL127C-B	52	61.22	x	x		
AAL130W	93	YJR005C-A/YGR169C-A†	92/93	74.44/50.82	x	x		
ABR148C-A	117	YJR112W-A	109	56.44	x	x		x
ABR099W-A	73	YKL068W-A	78	33.33	x			
AFR204W	68	YKL138C-A	68	57.35	x			
AFR289W	81	YKR095W-A	83	58.75	x	x	x	
ADL036W-A	75	YLR307C-A	87	50.00				
AFL165W	94	YMR194C-B	73	51.47	x	x	x	x
ADL210W	72	YNL024C-A	72	81.94	x	x	x	x
AFR059W	84	YNL138W-A	85	52.38	x			
AAR108W-A	70	YOR020W-A	90	35.71	x	x		
ABR192C-A	71	YPL096C-A	68	45.59	x			
AFL069C	58	YPL189C-A	68	69.64	x	x		
ACR178C	65	YPR036W-A	67	51.06	x	x		
AEL262C-A	86	YPR170W-B	85	85.88	x	x		x

*x indicates that a homolog to the novel ORF was found in the hemiascomycete, *C. albicans*, *S. pombe* or *N. crassa* databases. †Novel cases of gene duplication with homology to a single *A. gossypii* gene, remnant from the postulated genome doubling in the *S. cerevisiae* lineage [12,13].

perfect consensus splice sequences and that both YML017W and YOLO48C bear non-consensus splice sites with the respective acceptor/donor sites GTATGT--CACTAAC--CAG and GTAAGT--CACTAAC--TAG. In three cases of novel introns - YBL091C-A, YHR079C-A and YOLO48C - splicing has already been proposed by either Blandin *et al.* [17] or Wood *et al.* [18].

In addition to overlooked introns, we identified one case of a potentially wrongly assigned 5' splice site in *CPT1* (YNL130C), which codes for the sn-1,2-diaclyglycerol cholinephosphotransferase [21]. The current annotation proposes that *CPT1* would be spliced at a mismatched splice acceptor sequence. However, comparison with the *A. gossypii* homolog strongly suggests an intron of 92 base-pairs (bp), instead of 441 bp, with perfect consensus splice sequences. This would result in a protein of 407 amino acids with increased similarity to its *A. gossypii* homolog. This suggestion is supported by comparison with other fungal species, for example *C. albicans* and *S. pombe* (Table 2). Finally, a size of 407 amino acids for this

enzyme was already proposed in the first publication describing it [21].

A special case, not listed in Table 2, is the intron in *STO1* (YMR125W), a gene that encodes the large subunit of the nuclear cap-binding protein complex, a transcriptional activator of glycolytic genes. The comparison with *A. gossypii* cannot distinguish between two alternatives: presence or absence of an intron, as shown in Figure 2. The *S. cerevisiae* sequence currently available at SGD is annotated with an intron. Although we noticed the presence of an equivalent intron in *A. gossypii*, homology is conserved between the two non-spliced forms of these genes in the two organisms as well. Therefore, it may be possible that the *STO1* locus in both organisms encodes two proteins with differently charged amino ends.

5' and 3'ORF extensions

In 35 cases, it was possible to extend the boundaries of ORFs into annotation-free regions by artificially introducing single

Table 2**Summary of different types of ORF extensions proposed for annotated *S. cerevisiae* ORFs**

<i>S. cerevisiae</i>	Type*	Proposed reason†	Extension size (codons)	Supporting evidence‡				
				Hemiascomycetes	<i>C. albicans</i>	<i>S. pombe</i>	<i>N. crassa</i>	<i>S. cerevisiae</i> §
YAL013W	3' extension	FS	57	++	++		++	Resequenced
YAR044W/YAR042W	Fusion	FS		++	++	++	++	Resequenced; duplication
YBL066C	3' intron	Intron + FS	3	+	+			
YBL091C-A	5' intron	Intron	76	++	++	++		
YBL104C	5' and 3' extensions	FS	128	++	++	++		Resequenced
YBR041W	3' extension	FS	46	++	++		++	Resequenced
YBR074W/YBR075W	Fusion	2FS		+	+	+	+	Resequenced
YBR098W/YBR100W	Fusion	FS		++	++		+	Resequenced
YBR157C	3' extension	1FS	149	++				
YCL001W-A/YCL001W-B	Fusion	2FS						Duplication
YCL008C	3' extension	1FS	89					Resequenced [45]
YCL025C	3' extension	FS	38	++	++	+	++	gb: P25376; duplication
YCL069W	5' extension	PS	124	++	++	++	++	
YDL115C	5' intron	Intron	113	++	++			
YDR179W-A	5' extension	FS	138	++	++			gb: CAA86685
YDR474C/YDR475C	Fusion	2FS		++	++			Duplication
YDR494W	3' extension	FS	78	++	++			
YER039C/YER039C-A	Fusion	PS		++	++	++	++	Duplication
YER066W	5' extension	PS	201	++	++	++	++	
YFL007W/YFL006W	Fusion	FS		++	++		++	[46]
YFR038W	3' extension	FS	75	++	++	++		
YFR040W	5' extension	FS	97	++				Duplication
YFR045W	5' extension	FS + PS	106	++	++	++	++	[47]
YGL046W/YGL045W	Fusion	3FS		++	++		+	
YGL059W	3' extension	FS	46	++	++		++	
YGL183C	5' intron	Intron	45	++	++			
YGL211W	3' extension	3FS	164	++	++	++	++	
YGR006W	3' extension	FS	32	++	++	++		[48]
YGR225W	3' extension	FS	31	++	++	++	++	[49]
YGR272C/YGR271C-A	Fusion	FS		++	++			
YHR056C	5' extension	FS	51					gb: P38781; duplication
YHR079C-A	3' intron	Intron	45	++	++	++	++	
YHR176W	3' extension	FS	67	++	++		++	
YJL012C/YJL012C-A	Fusion	FS		++	++	++		Resequenced
YJL017W/YJL016W	Fusion	FS			++			Resequenced
YJL019W/YJL018W	Fusion	FS						Resequenced
YJL020C/YJL021C	Fusion	FS						Resequenced
YJL031C	5' intron	Intron	37	++	++	++	++	
YJL108C/YJL107C	Fusion	FS		++	++	++	++	Resequenced
YJL159W	3' extension	FS	4	++	++			Resequenced; duplication

Table 2 (Continued)**Summary of different types of ORF extensions proposed for annotated *S. cerevisiae* ORFs**

YJL160C	3' extension	FS	107	++	++			Resequenced; duplication
YJL178C	5' extension	FS	75	++	++			Resequenced
YJR013W	5' extension	FS	98	++	++	++	++	Resequenced
YKL033W-A	3' extension	FS	176	++		++	++	Resequenced
YKL199C/YKL198C	Fusion	FS		++	++	++	++	Resequenced; duplication
YKL207W	5' extension	FS	22	++	++	++		Resequenced
YKR004C	5' intron	Intron	85					5' RACE verified
YKR056W	5' extension	FS	21	++	++	++	++	Resequenced
YKR058W	5' extension	2FS	138	++	++			Resequenced [50]; duplication
YKR100C	3' extension	FS	114	++				Duplication
YKR103W/YKR104W	Fusion	PS		++	++	++	++	
YLL017W/YLL016W	Fusion	FS			+			Duplication
YLL052C/YLL053C	Fusion	FS		++	++	++		
YLR054C	5' intron	Intron	212	++	++			
YLR205C	5' extension	2FS	44	++	++			Resequenced
YLR389C	3' extension	FS	40	++		++		Resequenced
YLR401C	3' extension	FS	59	++	++	++	++	
YLR445W	3' intron	Intron	54					
YML002W/YML003W	Fusion	FS		++	++		++	
YML017W	5' intron	Intron	15					5' RACE verified
YMR084W/YMR085W	Fusion	FS		++	++	++	++	Duplication
YMR207C	5' extension	FS	59		++	++		Duplication
YMR269W	5' extension	FS	69	++	++			Resequenced
YNL083W	3' extension	FS	51	++	++	++	++	[47]
YNL130C	Other intron	Intron	21	++	++		++	[21]; duplication
YOL048C	5' intron	Intron	236	++	++			5' RACE verified; duplication
YOL163W/YOL162W	Fusion	PS+FS		++	++	++	++	
YOR069W	5' extension	Annotation	364	++	++	++		gb: Q92331; duplication
YOR298C-A	5' extension	FS	92	++	++	++	++	Resequenced; gb: BAA33217
YPL109C	3' intron	Intron	47	++	++	++	++	
YPR090W/YPR				++	++	++		
YPR090W/YPR089W	Fusion	FS						Resequenced
YPR098C	5' intron	Intron	53	++	++			

*Extension, ORF extension. †FS, frameshift; PS, premature STOP. ‡Supporting evidence from other fungal sequences: ++ indicates that homology supports the corrected *S. cerevisiae* ORF; + indicates that weak homology supports the corrected *S. cerevisiae* ORF. §Supporting evidence from other *S. cerevisiae* sequences: from published work (reference given); from sequence databases (GenBank accession number); from this study (resequenced or 5'RACE); from duplicated *S. cerevisiae* sequences matching the proposed sequence (duplication).

base-pair changes in the *S. cerevisiae* genomic sequence. These changes eliminated presumptive frameshifts or premature stop codons, as outlined in Figure 1c. The ORFs affected, the increase in ORF size (more than 70 amino acids for 50%

of the ORFs), and other supporting evidence are listed as part of Table 2. In 15 cases, we resequenced the region of the proposed change and confirmed the sequencing error. Finally, homology searches also supported the proposed sequence

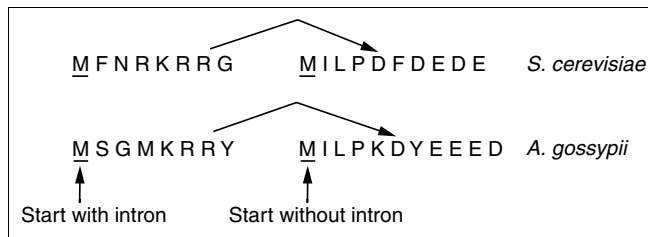


Figure 2
S. cerevisiae *STO1* (YMR125W) and its *A. gossypii* syntenic homolog show two possible amino termini. The *STO1* mRNA was proposed to be spliced in *S. cerevisiae*. However, both the spliced and non-spliced versions show homology to the *A. gossypii* genome, suggesting two alternative variants.

corrections. All regions of suggested change were inspected using BLAST searches against the Génolevures, *C. albicans*, *S. pombe* and *N. crassa* sequence data. In more than 70% of cases, we found homologous sequences in two or more databases that matched the *A. gossypii* annotation (Table 2).

A special case of ORF extension concerns *VPS5* (YOR069W), for which we propose an annotation rather than a sequence correction. The *A. gossypii* homolog is much longer and consideration of a further upstream start codon for *VPS5* would result in an 5' extension 364 codons long with strongly enhanced homology to the *A. gossypii* homolog.

ORF fusions

Another 22 proposed modifications resulted in the fusion of two previously distinct *S. cerevisiae* ORFs, as outlined in Figure 1d. A compilation of the *A. gossypii* ORFs, the fused *S. cerevisiae* ORFs, and their sizes is given in Table 2 and Table 3. As for the ORF extensions, we obtained supporting evidence for the validity of these fusions from database searches. For 17 of the proposed fusions, we found homologs of similar sizes in two or more fungal databases. Moreover, 10 of the ORF fusions had already been reported but not yet been included in databases, and seven are supported by a much better alignment to a duplicated copy in *S. cerevisiae*.

It should be pointed out that *S. cerevisiae* carries pseudogenes [22] and that confirmed pseudogenes may have homology over their entire length to single *A. gossypii* ORFs; see examples in Table 3 for three pseudogenes annotated as YER039C/YER039C-A, YLL017W/YLL016W, and YOL163W/YOL162W. Consequently, discrepancies observed between ORFs of the two species may either result from sequencing errors or may represent real pseudogenes. Therefore, we experimentally investigated nine of the proposed ORF fusions by resequencing the respective genomic regions in *S. cerevisiae* strain S288C, the reference strain of the yeast genome sequencing project [1]. In eight cases, a sequencing error was found, confirming the fusion of eight pairs of neighboring genes (Table 2 and Table 3). On the other hand,

resequencing also revealed that YJL107W/YJL108W is a novel pseudogene that bears a single point mutation in S288C.

In addition, programmed ribosomal frameshifting has been demonstrated in *S. cerevisiae* [23,24] and this might explain some of the observed differences between *S. cerevisiae* and *A. gossypii* genomic sequences. Therefore, resequencing all the questioned regions in *S. cerevisiae* would be needed to be able to discriminate between sequencing errors, pseudogenes and functional frameshifts.

Gene extensions revealing additional functional domains

We analyzed the presence of known functional domains in *S. cerevisiae* proteins with or without the proposed changes. Most of these extensions did not generate additional domains in the proteins. YKLO33W-A, however, could be extended at the 3' end by 176 codons, adding a HAD (haloacid dehalogenase) domain (InterPro: PF00702) and suggesting that this protein of previously unknown function may have a role in the assimilation of halogenated compounds. Similarly, YMR269W was described as a hypothetical protein of 164 amino acids. We propose an amino-terminal extension of 69 amino acids, which would generate a protein of 211 amino acids with 11% greater similarity to its *A. gossypii* homolog (Figure 3a,b). This proposal was confirmed by resequencing. Domain analysis revealed the presence of a putative RNA-binding domain (D111: PS50174) in both the extended *S. cerevisiae* and the annotated *A. gossypii* proteins (Figure 3c). YMR269W might therefore have a role in RNA-mediated cellular processes such as splicing, transcription, or translation. Indeed, YMR269W was recently found to interact with the translation initiation factor GCN3 (YKRO26C) in a whole-genome two-hybrid screen [25], which also points to a role in translation. Finally, evaluation of expression data from over 100 genome-scale experiments showed that YMR269W is regulated in a very similar manner as genes involved in protein synthesis [26]. It thus appears very likely that the 'extended' version of YMR269W is involved in protein synthesis.

Confirmation of hypothetical ORFs as real ORFs

In 1996 the yeast genome sequencing consortium faced the difficulty of annotating the first eukaryotic genome. Many potential ORFs in the newly sequenced genome did not have homology with entries in the existing databases, and discrimination between 'real' and 'chance' ORFs was often not possible. Novel *S. cerevisiae* genes lacking homology to any database were annotated if they were at least 100 codons long. The use of this arbitrary cut-off permitted the annotation of most of the 'real' genes but also led to the annotation of many questionable ORFs. Since then, a substantial number of these ORFs have been functionally characterized. However, many are still annotated as hypothetical ORFs because of a lack of functional data. The identification of homologs of

Table 3**Size comparison of proposed fused ORF in *S. cerevisiae* with *A. gossypii* homolog**

<i>S. cerevisiae</i> ORF1			<i>S. cerevisiae</i> ORF2			Fused ORF		<i>A. gossypii</i>		Supporting evidence
Systematic name	Common name	Size (amino acids)	Systematic name	Common name	Size (amino acids)	Size (amino acids)	Reason*	Name	Size (amino acids)	
YAR044W		859	YAR042W		256	1,188	FS	AER225W	1,300	Resequenced
YBR074W		103	YBR075W		460	976	2FS	AGL209W	1,012	Resequenced
YBR098W	MMS4	108	YBR100W		112	691	FS	ADL318C	715	
YCL001W-A†		84	YCL001W-B		153	313	2FS	AAL001W	386	
YDR474C†		155	YDR475C	JIP4	555	876	2FS	AEL233C	794	
YER039C†	HVG1	249	YER039C-A		72	341	PS	AFR236C	330	Pseudogene [22]
YFL007W†	BLM3	1,804	YFL006W		254	2,143	FS	AGL022W	2,146	[46]
YGL046W		262	YGL045W		229	540	3FS	AFR108W	549	
YGR272C		152	YGR271C-A‡		63	233	FS	ABR143C	214	
YJL012C	VTC4	648	YJL012C-A‡		65	721	FS	AGR316C	714	Resequenced
YJL017W		325	YJL016W		171	561	FS	AGR313W	471	Resequenced
YJL019W	MPS3	620	YJL018W		102	682	FS	AGR312W	617	Resequenced
YJL020C	BBC1	446	YJL021C		710	1,156	FS	AGR306C	924	Resequenced
YJL108C†	PRM10	383	YJL107C		387	770	FS	AFR075C	741	Resequenced
YKL199C†	YKT9	279	YKL198C	PTKI	399	649	FS	AFR372W	775	Resequenced
YKR103W		1,218	YKR104W		306	1,558	PS	NB§	NB§	
YLL017W†	SDC25		YLL016W	SDC25		1,252	FS	ADL038W	1,510	Pseudogene (SGD)
YLL052C†	AQY2	149	YLL053C		152	286	FS	AGL266C	451	
YML002W		737	YML003W		290	1,090	FS	ACR006C	1,072	
YMR084W†		262	YMR085W		432	719	FS	ABL036C	707	
YOL163W†	169	YOL162W			215	553	PS	ACL203C	538	Pseudogene [22]
YPR090W	736	YPR089W			155	888	FS	ABR111C	790	Resequenced

*PS, premature STOP codon; FS, frameshift. †Agreement with proposal of Blandin et al. [17] and Wood et al. [18]. ‡Novel ORFs identified by Blandin et al. [17]. §No homolog was found in *A. gossypii* but YKR103W/YKR104W is a member of a gene family including YLL048C and YHL035C for which *A. gossypii* homologs were found.

these ORFs in related organisms can be taken as strong evidence for their biological significance. Because *A. gossypii* shares as much as 95% of its genes with *S. cerevisiae* (90% being in synteny) it is an excellent organism to evaluate the authenticity of yeast hypothetical ORFs.

Currently, 1,885 *S. cerevisiae* ORFs are classified as hypothetical ORFs in the *Saccharomyces* Genome Database (SGD). We compared these genes to the *A. gossypii* genome annotation and identified a homolog in *A. gossypii* for 1,041 of them. Most important, 999 of these (96%) share both

Figure 3 (see following page)

Proposed changes in the hypothetical protein YMR269W. (a) Multiple alignment of the translated +2 and +3 frames of the *S. cerevisiae* YMR269W region and the syntenic *A. gossypii* protein. The boxed sequence indicates the region of the potential shift from frame +3 to +2 as suggested by the multiple alignment. (b) *S. cerevisiae* YMR269W region. Light gray depicts the current annotation as available at SGD. Dark gray shows the proposed elongation of the frame +2 translation on the 5' end of the yeast YMR269W gene. (c) Domain organization of YMR269W proteins in *S. cerevisiae* and *A. gossypii*. While the current YMR269W protein sequence of *S. cerevisiae* carries only a nuclear localization signal, both the *A. gossypii* homolog and the proposed extended YMR269W protein have an additional G-patch domain, which has been described as a putative RNA-binding domain.

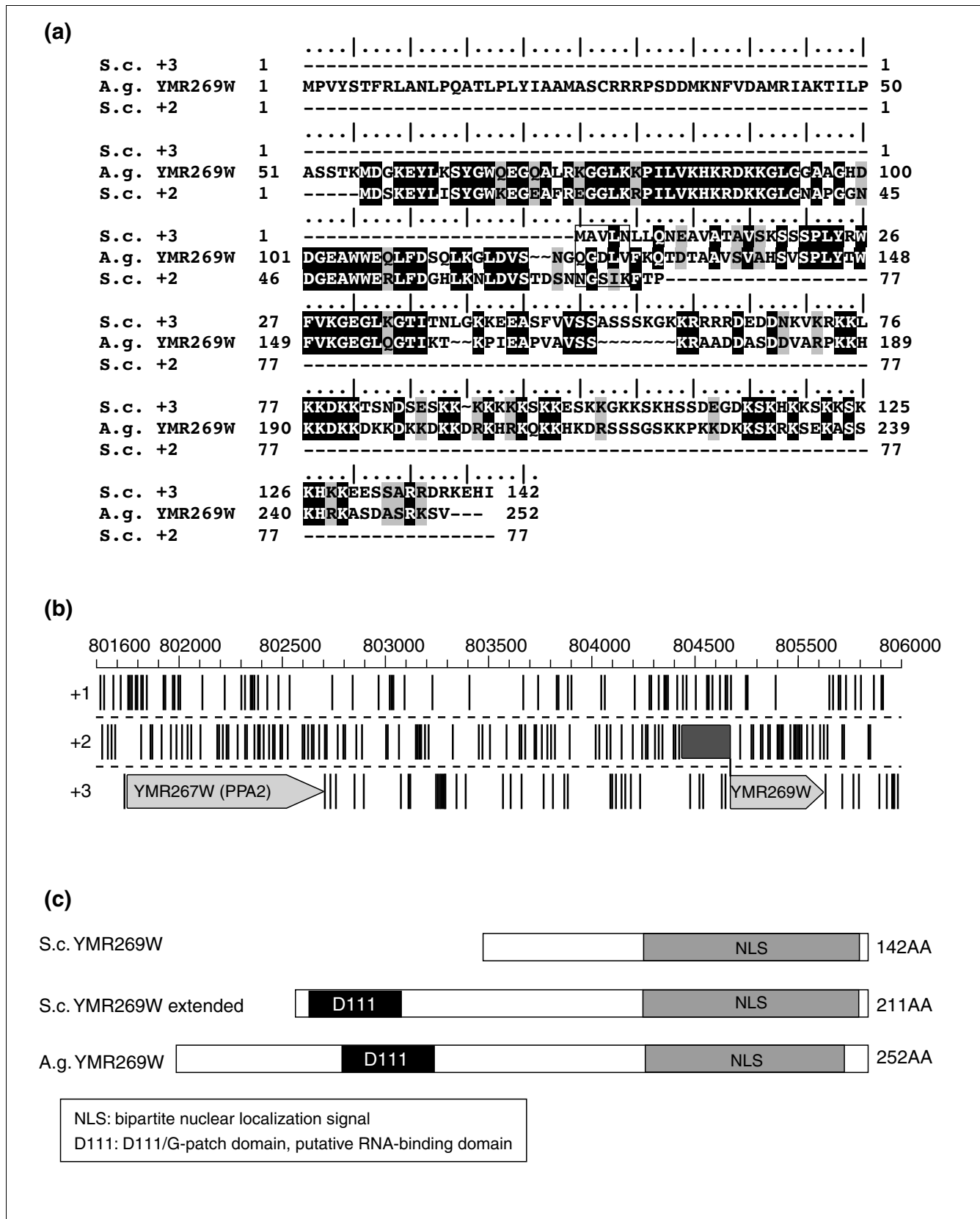


Figure 3 (see legend on previous page)

homology and synteny and can therefore be considered to be orthologs. The full list of hypothetical ORFs that should be regarded as real ORFs because of their homology and synteny with *A. gossypii* is available as an Additional data file. The Munich Information Center for Protein Sequences (MIPS), the other publicly accessible *S. cerevisiae* genome database, lists 988 of the *S. cerevisiae* ORFs classified as hypothetical or questionable (with questionable referring to hypothetical ORFs overlapping functionally characterized ORFs). We found homologs for 279 of these ORFs at syntenic positions in the genome of *A. gossypii* and all belong to the group of ORFs identified above as real among the 1,885 hypothetical ORFs at SGD. This comparison therefore provides strong evidence for the authenticity of a substantial part of the ORFs annotated as hypothetical in both SGD and MIPS.

Spurious ORFs among *S. cerevisiae* hypothetical ORFs

We assume it unlikely that all the remaining 844 hypothetical *S. cerevisiae* ORFs in SGD encode proteins, as only 10% of the known functional *S. cerevisiae* genes have no homolog in *A. gossypii* (F.S.D., S.V., S.B., A.L., K.G., C. Mohr, S. Steiner, P. Luedi, T.G. and P.P., unpublished work). They cannot, however, be directly investigated using this comparative approach owing to the absence of homologous genes in *A. gossypii*. Nevertheless, indirect evidence of the dubiousness of a subgroup of the ORFs absent in *A. gossypii* can be obtained by taking into consideration that they overlap other ORFs. The inspection of all overlapping pairs among annotated *S. cerevisiae* ORFs (based on our revised version of the *S. cerevisiae* ORF boundaries) reveals that, in the vast majority of the cases, one of the two ORFs belongs to the group of hypothetical ORFs lacking a syntenic homolog in *A. gossypii*. Furthermore, although there is some experimental evidence that, in rare cases, functional fungal genes overlap at the 3' ends of their ORFs; in other words, the opposite strand of one ORF acts as transcription terminator for the other ORF and vice versa [27], there is no experimental demonstration that a sequence within a functional ORF can act as promoter for another gene. We used these rules, in addition to the absence or presence of homologs in *A. gossypii*, to validate *S. cerevisiae* overlapping ORFs. We found three different categories among the 419 pairs of overlaps as schematically shown in Figure 4. For only seven pairs, *A. gossypii* carries homologs of both ORFs. Two pairs of homologs overlap in *A. gossypii*; the other five do not. Two cases of 5' end overlapping ORF pairs are probably explained by the assignment of the wrong start codon, and the remaining cases relate to overlapping 3' ends, which supports the hypothesis that ORF overlaps are rare in *S. cerevisiae* and that they involve 3' ends of ORFs (see Figure 4 legend).

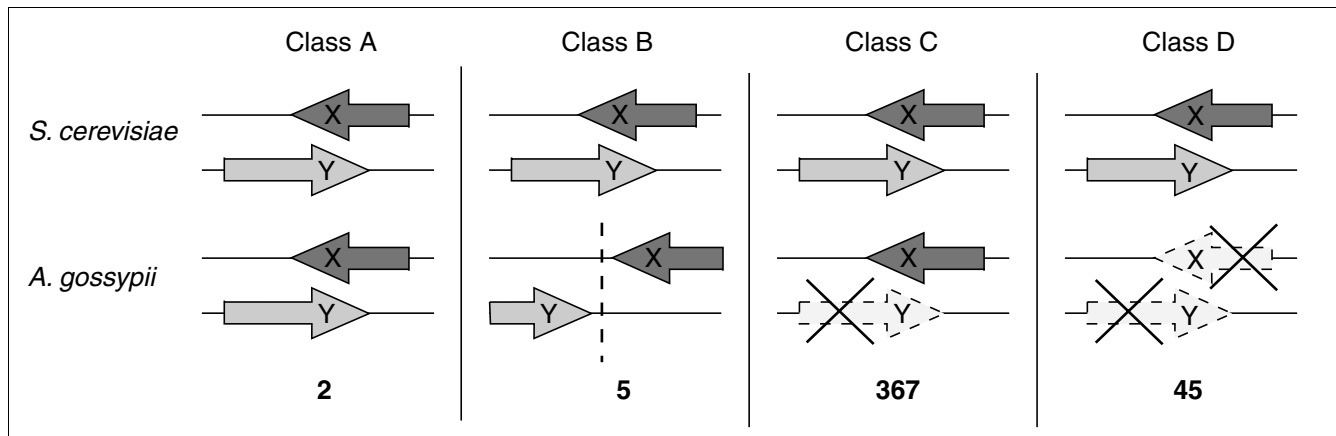
For 367 pairs, one ORF homolog was found in *A. gossypii*, the other not and we propose that the later ones are very likely to be spurious (see Figure 4, class B). These ORFs are listed in the additional data files with information about their present functional annotation, their sizes, and the type of overlap. For

66% of the pairs in this class, one or both presumptive promoter regions overlap an ORF sequence. In the remaining 34%, both terminator regions overlap ORF sequences. These latter cases should be viewed with caution as a very small percentage of them might turn out to be real. Indeed, in two cases marked in the additional data files, published data confirm the authenticity of a suggested spurious ORF. It should be noted that some of the proposed spurious ORFs are reported in very close relative of *S. cerevisiae* such as *S. bayanus*. However, the similarity is often restricted to the overlapping regions and is likely to result from the transfer of conservation from the real coding region to the other frames or strand. As *A. gossypii* is more distantly related, such homology can be found, though seldom, but does not match ORFs owing to the presence of STOP codons. Finally, in the remaining 45 pairs, for two ORFs we could find no homolog in *A. gossypii* and the criteria applied above cannot be used here. However, 36 ORFs could be considered as likely to be spurious as they overlap ORFs with described function or with a size of at least 500 codons (Figure 4, class D, and see Additional data files).

In summary, comparison of pairs of overlapping *S. cerevisiae* ORFs with the *A. gossypii* genome suggests that probably 403 of the remaining 844 hypothetical ORFs should be considered with care as they are likely to be spurious. Wood *et al.* [18] used ORF overlaps as one criterion to disregard 371 *S. cerevisiae* genes annotated as hypothetical, 289 of which are spurious according to the criteria applied above. Our analysis leaves about 450 hypothetical ORFs for which no information is currently available to categorize them as likely to be real or spurious. Additional evidence from similar analyses with other yeast species will be necessary to resolve these problems. Therefore, projects such as the current *Saccharomyces* Genome Project [28] will hopefully allow for a final *S. cerevisiae* genome annotation, seven years after completion of the genome sequence.

Conclusions

We have demonstrated the power of comparative genomics for the annotation of two completely sequenced fungal genomes. Whole-genome comparison guided the identification of novel ORFs, improved gene annotations, revealed sequencing errors, and helped to distinguish between real and spurious ORFs, thereby enhancing our view of the *S. cerevisiae* genome. As a consequence, these results will also contribute to the validity of genome-scale experiments, such as gene-expression profiling, an area where accurate gene annotation is crucial. The forthcoming availability of more yeast species genomes will, in an analogous way, drastically improve speed and accuracy of genome annotation of *S. cerevisiae* and the newly sequenced species. The method described here is straightforward for lower eukaryotes and should be applicable to any two closely related genomes of any complexity.

**Figure 4**

Classes of overlapping annotated ORFs in *S. cerevisiae* derived from comparison with the *A. gossypii* genome. Class A, homologs for both overlapping *S. cerevisiae* ORFs are found at syntenic positions in *A. gossypii* and also overlap. Class B, homologs for both overlapping *S. cerevisiae* ORFs are found in *A. gossypii* but do not overlap (see comments at end of legend). Class C, gene X but not gene Y has a syntenic homolog in *A. gossypii*. Class D, both overlapping *S. cerevisiae* ORFs lack a homolog in *A. gossypii*. Numbers refer to the frequency of the four types of overlapping pairs. Although all three possible directions of overlaps were observed, for convenience only 3'/3'-end overlaps are depicted here. YPL166W/YPL165C and YLR360W/YLR361C are the only two cases for which overlap was observed both in *S. cerevisiae* and *A. gossypii* (class A). The two overlaps are short (24 and 35 nucleotides, respectively, in *S. cerevisiae* and 24 and 14 nucleotides in *A. gossypii*, and involve only terminator-ORF sequence overlap. Class B overlaps comprise three syntenic ORF pairs (YJR012C/YJR013W, YML095C/YML096W, and YGR074W/YGR075C) and two non-syntenic (YPL018W/YPL017C and YBR262C/YBR263W). The lack of synteny reflects chromosomal rearrangements in one or the other species which resulted in either the separation of the two ORFs in *A. gossypii* or in their joining in *S. cerevisiae*. Two of the five class B ORF pairs refer to YML096W/YML095C and YGR074W/YGR075C, both with 6-nucleotide 3-end overlaps. The syntenic *A. gossypii* homologs are separated at their 3' ends by 2 and 51 nucleotides respectively, implying some overlap of terminator and ORF sequences very similar to their syntenic *S. cerevisiae* homologs. For two pairs of ORFs overlapping at their 5' ends in *S. cerevisiae* (YBR262C/YBR263W and YJR012C/YJR013W) both ORFs have a homolog in *A. gossypii*. The alignments of YBR263W and YJR012C with their respective *A. gossypii* homologs strongly suggest an error in selection of their start codons. Both *S. cerevisiae* ORFs are very likely to be 75 codons shorter, thus eliminating the presumptive promoter-ORF overlaps. Directions are provided for classes C and D in the additional data files.

Materials and methods

Yeast strain

S. cerevisiae AB972 (S288C) strain was used for resequencing and 5' RACE-based intron verification.

5' RACE-based verification of proposed introns in *S. cerevisiae*

Reverse transcription and 5' RACE was done using SMART RACE cDNA amplification system (BD Bioscience Clontech). Gene-specific primer (GSP) sequences were selected approximately 200 bp downstream of the putative introns for YKR004C, YML017W, and YOL048C. The amplified cDNA fragments were purified from the gel, cloned in the TOPO-TA cloning vector (Invitrogen) and sequenced on an ABI Prism 310 sequencer (Applied Biosystems). The sequences are available in GenBank (accession numbers AY245791, AY245792, and AY245793).

Resequencing of *S. cerevisiae* genomic regions

The following 25 *S. cerevisiae* genomic regions were resequenced: YAL013W (AY260888), YAR044W/YAR042W (AY260892), YBL104C (AY260889), YBR074W/YBR075W (AY260891), YBR157W (AY260879), YCLO08C (AY260880), YJL012C/YJL012C-A (AY227894), YJL016W/YJL017W

(AY260898), YJL019W/YJL018W, YJL020C/YJL021C, YJL108C/YJL107C (AY227895), YJL159W (AY260881), YJL160C (AY260893), YJL178C (AY260894), YJR013W (AY260895), YKLO33W-A (AY260896), YKL199C/YKL198C, YKL207W (AY260882), YKR056W (AY260897), YKR100C (AY260883), YLR205C (AY260884), YLR389C (AY260885), YMR269W (AY22789), YOR298C-A (AY260886), and YPR089W/YPR090W (AY260887). Primers flanking the region of the putative frameshift or premature stop mutation were selected to be unique within the yeast genome, and to have similar melting points. PCR was carried out using standard protocols on the AB972 (S288C) genomic DNA. PCR products were confirmed by agarose gel electrophoresis and sequencing was carried out on an ABI 310 sequencer using big dye chemistry and protocols from ABI. Sequence assembly was performed using the phred/phrap/consed analysis package [29-31]. Sequence corrections for YJL019W/YJL018W, YJL020C/YJL021C and YKL199C/YKL198C were recently corrected in SGD and were, therefore, not submitted to GenBank.

Sequence databases and sequence analysis

S. cerevisiae, *S. pombe* and hemiascomycete genomic sequence information was retrieved from GenBank at the

National Center for Biotechnology Information (NCBI) [32,33]. The *S. cerevisiae* genome was used as available at NCBI on 27 October, 2002. This release was submitted by the SGD [34,35]. Sequence data from *C. albicans* and *N. crassa* were obtained from the Stanford Genome Technology Center [36] and from the Neurospora Sequencing Project, Whitehead Institute/MIT Center for Genome Research [37], respectively. Sequence analysis was carried out using the GCG Wisconsin Package (Accelrys), BLAST [38] and FASTA tools [39,40]. Domain analysis was carried out using the InterProScan.pl algorithm from the European Institute of Bioinformatics [41,42]. Functional classifications of *S. cerevisiae* ORFs were taken from the SGD [34] and MIPS [43].

Annotation of *A. gossypii* chromosomes

The 9 Mb genome was sequenced by combining three strategies: end-sequencing of chromosome-sorted plasmid and BAC clones, shotgun sequencing of sheared genomic DNA fragments, and extensive gap filling by primer walking, which resulted in an average accuracy of 99.8% (F.S.D., S.V., S.B., A.L., K.G., C. Mohr, S. Steiner, P. Luedi, T.G. and P.P., unpublished work). In the first round of annotation, all ORFs longer than 50 codons were searched using BLAST against the set of *S. cerevisiae* ORF translations available from SGD [34]. *A. gossypii* ORFs with a hit lower or equal to $E = 1e-2$ were automatically annotated as *S. cerevisiae* homologs. This first draft of the *A. gossypii* genome annotation together with the BLAST results were re-evaluated case by case in a non-automatic procedure. *A. gossypii* ORFs sharing low or high homology with syntenic *S. cerevisiae* ORFs were kept. The synteny was independently assessed by two people. Inter-ORF regions were then compared with the six translation frames of the *S. cerevisiae* genome sequence, leading to the identification of potentially overlooked ORFs in *S. cerevisiae* and *A. gossypii*. In a final step, the remaining *A. gossypii* potential ORFs were searched against other databases and led to the annotation of *A. gossypii* genes with no homolog in *S. cerevisiae*.

Homology screening of all *S. cerevisiae* inter-ORF regions

A. gossypii ORF translations were searched against a locally built yeast inter-ORF sequence database using BLAST 2.0. A cut-off threshold E-value of $1e-2$ was used to filter the results. RNA genes were automatically filtered out and the remaining hits were manually checked for synteny. Regions of discrepancy were carefully checked in *A. gossypii* and in all cases matched good-quality consensus sequence. The current *S. cerevisiae* genome annotation release was re-annotated with proposed changes or novel sequence using the Artemis annotation tool [44] and the Sequin submission tool [32,33].

Experimentally verified sequence corrections are available in GenBank under the accession numbers: AY260888, AY260892, AY260889, AY260891, AY260879, AY260880, AY227894, AY260898, AY227895, AY260881, AY260893,

AY260894, AY260895, AY260896, AY260882, AY260897, AY260883, AY260884, AY260885, AY22789, AY260886, AY260887, AY245791, AY245792 and AY245793.

Additional data files

The following files are available with the online version of this article: a list of the novel *S. cerevisiae* ORFs proposed by Blandin *et al.* and Wood *et al.* [17,18] for which a syntenic homolog was found in the *A. gossypii* genome (Additional data file 1); a list of all *S. cerevisiae* hypothetical ORFs for which a homolog was found in the *A. gossypii* genome (Additional data file 2); a list of all *S. cerevisiae* hypothetical ORFs for which no homolog was found in the *A. gossypii* genome (Additional data file 3). A list of all class C overlaps together with gene sizes, gene functions and overlap directions can be found in Additional data file 4; these genes are suggested to be spurious based on our criteria. A list of class D overlaps: none of the two overlapping genes in *S. cerevisiae* has a homolog in *A. gossypii* can be found in Additional data file 5; the size and function classifications were used to predict spurious genes in some of these cases. A graphical display of the proposed *S. cerevisiae* annotation changes based on comparison with *A. gossypii* can be found in Additional data file 6. GenBank files for each of the *S. cerevisiae* chromosomes that take account of the proposed modifications (prior to confirmation of the sequence) can be found in Additional data file 7. GenBank files for each *A. gossypii* genomic locus used to infer annotation corrections in *S. cerevisiae* can be found in Additional data file 8.

Acknowledgements

We thank Philippe Luedi and Amy Gladfelter for supporting discussions, and Arndt Brachat and Mike Primig for critical reading of the manuscript. This work was supported by grants from the University of Basel and Duke University.

References

- Goffeau A, Barrell BG, Bussey H, Davis RW, Dujon B, Feldmann H, Galibert F, Hoheisel JD, Jacq C, Johnston M, et al.: **Life with 6000 genes.** *Science* 1996, **274**:563-567.
- The *C. elegans* Sequencing Consortium: **Genome sequence of the nematode *C. elegans*: a platform for investigating biology.** *Science* 1998, **282**:2012-2018.
- Adams MD, Celniker SE, Holt RA, Evans CA, Gocayne JD, Amanatides PG, Scherer SE, Li PW, Hoskins RA, Galle RF, et al.: **The genome sequence of *Drosophila melanogaster*.** *Science* 2000, **287**:2185-2195.
- The *Arabidopsis* Genome Initiative: **Analysis of the genome sequence of the flowering plant *Arabidopsis thaliana*.** *Nature* 2000, **408**:796-815.
- Lander ES, Linton LM, Birren B, Nusbaum C, Zody MC, Baldwin J, Devon K, Dewar K, Doyle M, FitzHugh W, et al.: **Initial sequencing and analysis of the human genome.** *Nature* 2001, **409**:860-921.
- Venter JC, Adams MD, Myers EW, Li PW, Mural RJ, Sutton GG, Smith HO, Yandell M, Evans CA, Holt RA, et al.: **The sequence of the human genome.** *Science* 2001, **291**:1304-1351.
- Holt RA, Subramanian GM, Halpern A, Sutton GG, Charlab R, Nusser DR, Wincker P, Clark AG, Ribeiro JM, Wides R, et al.: **The genome sequence of the malaria mosquito *Anopheles gambiae*.** *Science* 2002, **298**:129-149.
- Waterston RH, Lindblad-Toh K, Birney E, Rogers J, Abril JF, Agarwal

- P, Agarwala R, Ainscough R, Alexandersson M, An P, et al.: **Initial sequencing and comparative analysis of the mouse genome.** *Nature* 2002, **420**:520-562.
9. Mewes HW, Albermann K, Bahr M, Frishman D, Gleissner A, Hani J, Heumann K, Kleine K, Maierl A, Oliver SG, et al.: **Overview of the yeast genome.** *Nature* 1997, **387**:7-65.
 10. **Sequence updates at SGD** [<http://genome-www.stanford.edu/Saccharomyces/sequenceupdates.shtml>]
 11. **Chromosome III resequencing information at MIPS** [<http://mips.gsf.de/cgi-bin/proj/yeast/THREE>]
 12. Philippsen P, Kleine K, Pohlmann R, Dusterhoft A, Hamberg K, Hege-mann JH, Obermaier B, Urrestarazu LA, Aert R, Albermann K, et al.: **The nucleotide sequence of *Saccharomyces cerevisiae* chromosome XIV and its evolutionary implications.** *Nature* 1997, **387**:93-98.
 13. Wolfe KH, Shields DC: **Molecular evidence for an ancient duplication of the entire yeast genome.** *Nature* 1997, **387**:708-713.
 14. Zhang Z, Dietrich FS: **Verification of a new gene on *Saccharomyces cerevisiae* Chromosome III.** *Yeast* 2003, **20**:731-738.
 15. Giaever G, Chu AM, Ni L, Connelly C, Riles L, Veronneau S, Dow S, Lucanu-Danila A, Anderson K, Andre B, et al.: **Functional profiling of the *Saccharomyces cerevisiae* genome.** *Nature* 2002, **418**:387-391.
 16. Miura F, Yada T, Nakai K, Sakaki Y, Ito T: **Differential display analysis of mutants for the transcription factor Pdr1p regulating multidrug resistance in the budding yeast.** *FEBS Lett* 2001, **505**:103-108.
 17. Blandin G, Durrens P, Tekaija F, Aigle M, Bolotin-Fukuhara M, Bon E, Casaregola S, de Montigny J, Gaillardin C, Lepingle A, et al.: **Genomic exploration of the hemiascomycetous yeasts: 4. The genome of *Saccharomyces cerevisiae* revisited.** *FEBS Lett* 2000, **487**:31-36.
 18. Wood V, Rutherford KM, Ivens A, Rajandream MA, Barrell B: **A reannotation of the *Saccharomyces cerevisiae* genome.** *Comp Funct Genomics* 2001, **2**:143-154.
 19. Wood V, Gwilliam R, Rajandream MA, Lyne M, Lyne R, Stewart A, Sgouros J, Peat N, Hayles J, Baker S, et al.: **The genome sequence of *Schizosaccharomyces pombe*.** *Nature* 2002, **415**:871-880.
 20. Kessler MM, Zeng Q, Hogan S, Cook R, Morales AJ, Cottarel G: **Systematic discovery of new genes in the *Saccharomyces cerevisiae* genome.** *Genome Res* 2003, **13**:264-271.
 21. Hjelmstad RH, Bell RM: **The sn-1,2-diacylglycerol cholinephosphotransferase of *Saccharomyces cerevisiae*. Nucleotide sequence, transcriptional mapping, and gene product analysis of the CPT1 gene.** *J Biol Chem* 1990, **265**:1755-1764.
 22. Harrison P, Kumar A, Lan N, Echols N, Snyder M, Gerstein M: **A small reservoir of disabled ORFs in the yeast genome and its implications for the dynamics of proteome evolution.** *J Mol Biol* 2002, **316**:409-419.
 23. Morris DK, Lundblad V: **Programmed translational frameshifting in a gene required for yeast telomere replication.** *Curr Biol* 1997, **7**:969-976.
 24. Asakura T, Sasaki T, Nagano F, Satoh A, Obaishi H, Nishioka H, Ima-mura H, Hotta K, Tanaka K, Nakanishi H, et al.: **Isolation and characterization of a novel actin filament-binding protein from *Saccharomyces cerevisiae*.** *Oncogene* 1998, **16**:121-130.
 25. Uetz P, Giot L, Cagney G, Mansfield T, Judson RS, Knight JR, Lockshon D, Narayan VA, Srinivasan M, Pochart P, et al.: **A comprehensive analysis of protein-protein interactions in *Saccharomyces cerevisiae*.** *Nature* 2000, **403**:623-627.
 26. Hughes TR, Marton MJ, Jones AR, Roberts CJ, Stoughton R, Armour CD, Bennett HA, Coffey E, Dai H, He YD, et al.: **Functional discovery via a compendium of expression profiles.** *Cell* 2000, **102**:109-126.
 27. Gerads M, Ernst JF: **Overlapping coding regions and transcriptional units of two essential chromosomal genes (*CCT8*, *TRP1*) in the fungal pathogen *Candida albicans*.** *Nucleic Acids Res* 1998, **26**:5061-5066.
 28. **Saccharomyces Genome Sequencing** [<http://genome.wustl.edu/projects/yeast/>]
 29. Gordon D, Abajian C, Green P: **Consed: a graphical tool for sequence finishing.** *Genome Res* 1998, **8**:195-202.
 30. Ewing B, Green P: **Base-calling of automated sequencer traces using phred. II. Error probabilities.** *Genome Res* 1998, **8**:186-194.
 31. Ewing B, Hillier L, Wendl MC, Green P: **Base-calling of automated sequencer traces using phred. I. Accuracy assessment.** *Genome Res* 1998, **8**:175-185.
 32. **National Center for Biotechnology Information** [<http://www.ncbi.nlm.nih.gov>]
 33. Benson DA, Karsch-Mizrachi I, Lipman DJ, Ostell J, Wheeler DL: **GenBank.** *Nucleic Acids Res* 2003, **31**:23-27.
 34. Cherry JM, Adler C, Ball C, Chervitz SA, Dwight SS, Hester ET, Jia Y, Juvik G, Roe T, Schroeder M, et al.: **SGD: *Saccharomyces Genome Database*.** *Nucleic Acids Res* 1998, **26**:73-79.
 35. **Saccharomyces Genome Database** [<http://www.yeastgenome.org>]
 36. **Stanford Genome Technology Center, *Candida albicans* sequence** [<http://www-sequence.stanford.edu/group/candida/download.html>]
 37. **Whitehead Institute/MIT Center for Genome Research** [<http://www-genome.wi.mit.edu>]
 38. Altschul SF, Gish W, Miller W, Myers EV, Lipman DJ: **Basic local alignment search tool.** *J Mol Biol* 1990, **215**:403-410.
 39. Pearson WR, Lipman DJ: **Improved tools for biological sequence comparison.** *Proc Natl Acad Sci USA* 1988, **85**:2444-2448.
 40. Pearson WR: **Using the FASTA program to search protein and DNA sequence databases.** *Methods Mol Biol* 1994, **25**:365-389.
 41. Apweiler R, Attwood TK, Bairoch A, Bateman A, Birney E, Biswas M, Bucher P, Cerutti L, Corpet F, Croning MD, et al.: **The InterPro database, an integrated documentation resource for protein families, domains and functional sites.** *Nucleic Acids Res* 2001, **29**:37-40.
 42. Zdobnov EM, Apweiler R: **InterProScan - an integration platform for the signature-recognition methods in InterPro.** *Bioinformatics* 2001, **17**:847-848.
 43. Mewes HW, Frishman D, Guldener U, Mannhaupt G, Mayer K, Mokrejs M, Morgenstern B, Munsterkotter M, Rudd S, Weil B: **MIPS: a database for genomes and protein sequences.** *Nucleic Acids Res* 2002, **30**:31-34.
 44. Rutherford K, Parkhill J, Crook J, Horsnell T, Rice P, Rajandream MA, Barrell B: **Artemis: sequence visualization and annotation.** *Bioinformatics* 2000, **16**:944-945.
 45. Li Y, Kane T, Tipper C, Spatrick P, Jenness DD: **Yeast mutants affecting possible quality control of plasma membrane proteins.** *Mol Cell Biol* 1999, **19**:3588-3599.
 46. Robben J, Hertveldt K, Volckaert G: **Revisiting the yeast chromosome VI DNA sequence reveals a correction merging YFL007w and YFL006w to a single ORF.** *Yeast* 2002, **19**:699-702.
 47. Belenkiy R, Haefele A, Eisen MB, Wohlrab H: **The yeast mitochondrial transport proteins: new sequences and consensus residues, lack of direct relation between consensus residues and transmembrane helices, expression patterns of the transport protein genes, and protein-protein interactions with other proteins.** *Biochim Biophys Acta* 2000, **1467**:207-218.
 48. Horowitz DS, Abelson J: **A U5 small nuclear ribonucleoprotein particle protein involved only in the second step of pre-mRNA splicing in *Saccharomyces cerevisiae*.** *Mol Cell Biol* 1993, **13**:2959-2970.
 49. Cooper KF, Mallory MJ, Egeland DB, Jarnik M, Strich R: **Amal1p is a meiosis-specific regulator of the anaphase promoting complex/cyclosome in yeast.** *Proc Natl Acad Sci USA* 2000, **97**:14548-14553.
 50. Cheng C, Mu J, Farkas I, Huang D, Goebel MG, Roach PJ: **Requirement of the self-glucosylating initiator proteins Glg1p and Glg2p for glycogen accumulation in *Saccharomyces cerevisiae*.** *Mol Cell Biol* 1995, **15**:6632-6640.