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#### 1 Extensive splicing across the Saccharomyces cerevisiae genome

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24 Abstract

25 Pre-mRNA splicing is vital for the proper function and regulation of eukaryotic gene expression.

- 26 Saccharomyces cerevisiae has been used as a model organism for studies of RNA splicing
- 27 because of the striking conservation of the spliceosome components and its catalytic activity.
- Nonetheless, there are relatively few annotated alternative splice forms, particularly when
- 29 compared to higher eukaryotes. Here, we describe a method to combine large scale RNA
- 30 sequencing data to accurately discover novel splice isoforms in *Saccharomyces cerevisiae*.
- Using our method, we find extensive evidence for novel splicing of annotated intron-containing
- 32 genes as well as genes without previously annotated introns and splicing of transcripts that are 33 antisense to annotated genes. By incorporating several mutant strains at varied temperatures,
- we find conditions which lead to differences in alternative splice form usage. Despite this, every
- 34 we find conditions which lead to differences in alternative splice form usage. Despite this, e
   35 class and category of alternative splicing we find in our datasets is found, often at lower
- frequency, in wildtype cells under normal growth conditions. Together, these findings show that
- there is widespread splicing in *Saccharomyces cerevisiae*, thus expanding our view of the
- 38 regulatory potential of RNA splicing in yeast.

39

# 40 Author Summary

41 Pre-mRNA splicing is a fundamental step in eukaryotic gene expression. Saccharomyces

42 cerevisiae, also known as brewer's yeast, is a model organism for the study of pre-mRNA

43 splicing in eukaryotes. Through the process of pre-mRNA splicing, a single gene is capable of

44 encoding multiple mature mRNA products, but it is often difficult to identify the splice events that

45 lead to these mRNA products. Here, we describe a method to accurately discover novel splice

46 events in Saccharomyces cerevisiae and find evidence for extensive splicing in

47 Saccharomyces. By utilizing a variety of strains and growth conditions, we are able to

characterize many splice forms and correlate cellular conditions with prevalence of novel spliceevents.

50

# 51 Introduction

52 Eukaryotic genes are composed of coding sequences termed exons interrupted by non-coding

53 sequences called introns. Introns are removed from RNA by the large macromolecular complex

54 known as the spliceosome through the process of RNA splicing. By selectively including

different combinations of exons, a single gene can produce multiple RNA products. Pre-mRNA

56 splicing is crucial for the proper expression of eukaryotic genes, and spliceosome components

are highly conserved from yeast to mammals at the sequence, structure, and functional levels

- 58 [1, 2].
- 59 Despite the high conservation between the spliceosomal components in yeast and mammals,

60 Saccharomyces cerevisiae has a more compact genome with approximately 300 annotated

- 61 intron-containing genes. Even with the relatively streamlined splicing landscape in *S*.
- 62 *cerevisiae*, the prevalence of alternative splicing in this organism has not been well
- 63 characterized. Several studies have found instances of alternative splicing in individual genes
- in *S. cerevisiae* [3, 4, 5], and there have been high-throughput methods to find novel splicing [6,
- 7, 8, 9, 10]. Most of these methods focus primarily or entirely on intron retention or exon
- skipping, while there has been little description of novel 5' or 3' splice site usage.

67 Here we describe a method for discovery of unannotated splice sites in *Saccharomyces* 

- *cerevisiae* by RNA-seq. We tested the method with wildtype strains as well as strains and
- 69 conditions that do not lead to direct changes in the splicing machinery itself, but that impact
- broad cellular conditions, RNA turnover, chromatin remodeling, and histone composition. Many
- of these strains and conditions help us observe changes in splicing that are not or are rarely
- 72 detected in wildtype cells under normal growth conditions. We also analyze several strains that
- <sup>73</sup> include *prp43-1*, a temperature sensitive mutation of the *prp43* gene, an RNA helicase directly
- involved in spliceosome disassembly to determine if modulating disassembly might affect our
- ability to detect aberrant splice site sampling. Interestingly, this mutation of *prp43* leads to a
- decrease in splicing of both annotated introns as well as novel splicing.

77 We compare our results with those of earlier studies and find good agreement, however over 78 two thirds of the splice forms our method predicts, 676 total novel splice forms, have not been 79 previously described. Within known intron-containing genes (ICGs), we find that novel isoforms generate longer introns in samples grown at higher temperature. We also find that deletion of 80 81 xrn1, an evolutionarily-conserved 5' to 3' exonuclease, leads to an accumulation of novel splice 82 isoforms that do not use either an annotated 5' nor an annotated 3' splice site, both within known ICGs and unannotated ICGs. While most of our novel splice forms use a known 5' splice 83 84 site and a novel 3' splice site, we find that deletion of ume6, a component of a histone 85 deacetylase complex that regulates early meiosis and that has been recently shown to have a regulatory role during mitosis [11], causes a dramatic increase in the use of novel 5' splice sites 86 87 within ICGs. We also find evidence for splicing of RNAs that are antisense to annotated 88 transcripts. Together, our results indicate that there is significantly more splicing than previously

- 89 thought. This suggests that the opportunity for splicing in the form of "latent" introns is a key
- 90 feature of the yeast genome. Furthermore, changes in the activity of the gene expression
- 91 machinery or the cells' environment can significantly alter this rich splicing landscape.
- 92

# 93 **Results**

94

### 95 Alternative splicing is widespread in Saccharomyces cerevisiae

In order to explore the extent of splicing across the yeast transcriptome in a high confidence 96 97 manner, we implemented a novel approach to allow us to leverage a large amount of RNA sequence data while imposing stringent filters (figure 1). Briefly, a large number of RNA-seq 98 99 datasets are combined such that each novel splice form needs at least a single read that aligns 100 without mismatches to the novel junction to pass the initial filter. This read cannot also align to known transcripts or the Saccharomyces cerevisiae genome, even with a large number of 101 mismatches. Once a splice junction has been identified in this way, all reads are realigned to 102 the newly discovered splice forms, annotated transcripts, and the genome to find the optimum 103 alignment. Each novel splice form is then scored based on how well its 5' splice site, 3' splice 104 105 site, and predicted branch point fit the consensus sequence for Saccharomyces cerevisiae 106 splice signals. These scores are then used to compute p-values that represent how likely a 107 splice product score of this strength is to occur by chance (figure S1).

108 To capture as many novel splice forms as possible, we incorporated datasets from a variety of 109 strains and experimental conditions that are known or suspected to have an impact of splicing,

some accessed from previous studies [12, 13] and some novel (Table S1). In addition to 4 110 wildtype samples grown at three different temperatures, these samples include 2 biological 111 112 replicates each of two deletions in genes involved in decay, including xrn1, a 5'-3' exonuclease, 113 and upf1, required for efficient nonsense-mediated decay. We also include  $htz1\Delta$  which encodes the histone variant H2A.Z and  $swr1\Delta$  which is required to exchange H2A for H2A.Z in 114 115 chromatin as well as double mutants  $swr1\Delta xm1\Delta$ ,  $swr1\Delta upf1\Delta$ ,  $htz1\Delta xm1\Delta$ , and  $htz1\Delta$ upf1*Δ*. Cells that lack H2A.Z are found to have impaired splicing of intron-containing genes 116 117 (ICGs), particularly genes that have suboptimal splice sites [13, 14]. We also analyze RNA-seq 118 data from snf2A, which leads to an increase in use of non-canonical branch point and 5' splice 119 site sequences in annotated ICGs and  $ume6\Delta$ , which derepresses genes implicated in meiosis 120 in Saccharomyces cerevisiae, and the double mutants  $upf1\Delta snf2\Delta$ ,  $xrn1\Delta snf2\Delta$ , and  $ume6\Delta$ 121 snf2 $\Delta$ . Previous studies suggest that snf2 $\Delta$  increases splicing by altering ribosomal protein 122 gene expression and  $ume6\Delta$  allows expression of genes that are usually repressed. We also include set1 $\Delta$  and set2 $\Delta$ , deletion mutations in histone methyltransferase genes. Finally, since 123 we previously showed that the DEAH protein Prp43 contributes to disassembly of suboptimal 124 125 spliceosomes using a prp43 DAMP allele [13], we included a temperature sensitive mutant of 126 prp43, prp43-1, as well as set1 $\Delta$  prp43-1 and set2 $\Delta$  prp43-1. All set1 $\Delta$ , set2 $\Delta$ , and prp43-1 127 strains contribute two samples to our workflow, one grown at 25° and one at 37°. Taken together, these datasets represent 29 samples across a variety of Saccharomyces cerevisiae 128 129 strains and growth temperatures. By leveraging a large number of datasets, we are able to 130 discover more novel splice sites and determine cellular conditions that lead to changes in

131 alternative splice site usage.

	Read	Aligned read
Strain	counts	counts
Wildtype	20395478	14160763
xrn1∆	36132564	25787231
upf1∆	32005528	22034981
swr1∆	38830226	21964031
htz1∆	35969862	18508451
swr1 $\Delta$ xrn1 $\Delta$	86744422	50807249
swr1 $\Delta$ upf1 $\Delta$	11369368	6729660
htz1Δ xrn1Δ	52891892	35039812
htz1 $\Delta$ upf1 $\Delta$	49460562	25424884
Wildtype	35059908	27722481
snf2∆	34016283	25657587
upf1∆	21328241	14465110
upf1 $\Delta$ snf2 $\Delta$	31564582	23487340
xrn1∆	29688304	20282088
xrn1 $\Delta$ snf2 $\Delta$	34289534	27043467
ume6∆	40071056	30161996

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ume6 $\Delta$ snf2 $\Delta$	22534282	17424922
Wildtype 25° C	126355512	97015734
Wildtype 37° C	140962504	82278728
set1∆ 25° C	120413012	75538896
set1∆ 37° C	150376422	84044314
set2∆ 25° C	115930956	61396568
set2∆ 37° C	105592626	63182213
prp43-1 25° C	113138574	78671916
prp43-1 37° C	93850396	54354046
prp43-1 set1∆		
25° C	180103858	107502423
prp43-1 set1∆		
37° C	116668118	70474357
prp43-1 set2∆		
25° C	129498126	93547450
prp43-1 set2∆		
37° C	107964066	68127119

132

133 Using our method, we discover evidence for 944 novel splice events across 408 transcripts with

p < 0.05 (Table 1). Of these novel events, the majority are novel splice products found within

135 known ICGs, using either the annotated 5' splice site with a novel 3' splice site (Table S2,

n=537) or using the annotated 3' splice site with a novel 5' splice site (Table S2, n=129).

137 Additionally, we found several cases of novel splicing that do not use any annotated splice sites

138 within annotated ICGs (Table S2, n=22) and in unannotated ICGs (Table S2, n=198). We also

139 find evidence for splice forms that are antisense to known transcripts (Table S2, n=50).

		All	р	<0.05	р	<0.01	р
	Events	Transcripts	Events	Transcripts	Events	Transcripts	Events
All	6187	1937	944	408	852	386	586
Antisense	2504	1110	50	41	46	37	33
Intronless	2709	1141	198	149	168	132	107
New 3'	561	196	537	194	502	191	368
New 5'	140	95	129	90	113	84	63
New both	273	120	22	18	15	12	7

140

141 Many of the events we discover using our method are low abundance. About half of the total

events are represented by fewer than five read counts across our datasets (figure S2A).

143 However, these low abundance events are high confidence due to our approach's stringent

144 discovery protocol and p-value cutoff. Furthermore, these high confidence, low abundance

novel splice events reveal splicing that is unlikely to be found in the high abundance data.

146 Different classes of novel splicing are more prevalent in our low abundance novel splicing data.

147 Specifically, novel splicing within annotated ICGs is more common in our high abundance data,

148 while novel splicing within unannotated ICGs and antisense RNAs are more common in our low

abundance data (figure S2C).

150

#### 151 Validation of novel splicing

152 In order to best validate our method by RT-PCR, we chose candidates with a variety of read counts and splice site scores representing each category of novel splicing we find, including 153 154 novel 5' splice site with annotated 3' splice site, novel 3' splice site with annotated 5' splice site, 155 both novel splice sites within an annotated ICG, novel splicing in an unannotated ICG, and 156 splicing of transcripts that are antisense to annotated genes. In addition, we selected some 157 denes which show only a single novel splice form and others with several. The results of our 158 validation are shown in Figure 2 using oligonucleotides listed in table S3. BIG1, an integral membrane protein gene of the endoplasmic reticulum, shows a single novel splice event 159 160 utilizing a novel 3' splice site with an annotated 5' splice site in our data with 19 total read 161 counts spread roughly evenly across our 29 samples and a splice site score p-value of 0.0003. The novel splice event in *BIG1* is in-frame with the annotated form, and would be predicted to 162 163 generate a protein product that has 6 additional amino acids. SIM1, a gene thought to 164 participate in DNA replication, shows a total of four novel splice forms, each utilizing an 165 annotated 3' splice site and novel 5' splice site with total read counts ranging from 2 to 35 and splice site p-values ranging from 0.018 to 0.0004. The intron in SIM1 is located in the 5' UTR, 166 therefore the novel splice events would not be expected to yield a new protein product, although 167 168 potential regulation in the 5' UTR could be altered. MCR1, a gene involved in ergosterol biosynthesis in mitochondria, shows a single novel splice form utilizing both novel 5' and 3' 169 splice sites within an annotated ICG with 49 total read counts, 9 of which are found across our 170 171 snf2 $\Delta$  samples, and a splice site p-value of 0.004. The annotated and novel introns for MCR1 are both found in the 5' UTR and are therefore unlikely to yield different protein products. 172 173 However, we previously showed that changes in SNF2 expression can affect splicing of others transcripts to alter mitochondrial function [15]. SPF1, which encodes an ion transporter of the 174 175 ER membrane, is an unannotated ICG that shows five novel splice forms in our data with read counts ranging from 1 to 4 across all of our samples and p-value scores ranging from 0.004 to 6 176  $x \, 10^{-6}$ . Of the five novel splice forms found in *SPF1*, only one is in frame, which would produce 177 a protein product that has 134 fewer amino acids. Finally, we find evidence for splicing in the 178 179 antisense direction to LEU4, with a total of six splice forms with counts ranging from 1 to 33 across all samples and splice site p-values ranging from 0.0001 to 2.8 x 10<sup>-6</sup>. Not surprisingly, 180 of the 48 sequence reads that derive from spliced reads antisense to LEU4, 28 come from 181 182 samples which are deleted of SET2, a histone methyltransferase that has previously been 183 implicated in suppressing antisense transcription [16]. Together, we validate each category of novel splicing we observe in our data and validate candidates ranging from a single read count 184 185 to 49 read counts and from splice site p-values ranging from 0.018 to 2.8 x  $10^{-6}$ .

Primer	Sequence
BIG1 F	GTTGCTTATTATGTGTGGAAGCTTTTG
BIG1 R	CACCTGGTCTACGTTACAATACTCC

SPF1 F	CCCAGAGA GCCACAAGTT GATCTTG
SPF1 R	CATTTAGAGGCAATCTTGACCTGTTGC
SIM1 F	GTGCTA CCCAACTACT TACATTCCT
SIM1 R	CTTGGCTAAGGCAGCAGATGAAAC
MCR1 F	CCTTGATTGGTGTCTTGTCGAGAGAG
MCR1 R	ACTCACAGGAGTGTATGGTCTCACC
LEU4	GAGAGTATTATTGCTCTTGCTGAGC
antisense	
LEU4 R	CTTGTTGGGAAGGGTCATCCTTAG

186

#### 187 Comparison with other sequence-based approaches for alternative splicing

188 Even though our method filters (1) putative splice forms with strong similarity to the

189 Saccharomyces cerevisiae genome, (2) known transcripts, and (3) those found outside of

190 genes, the splice sites identified by our method include many reported in recent studies. Of the

- 191 522 novel splice sites described in Kawashima et al. [6], 420 are discoverable by our method,
- 192 236 are found in our raw data and 189 pass our p < 0.05 filter. Of those described in Schreiber
- et al. [7], 248 out of 314 of the described splice events are discoverable by our method, 214 are
- in our data and 185 pass our p < 0.05 filter. Qin et al. [10] describe a method of identification of</li>
   novel splice forms by lariat sequencing, a process which reveals 5' splice sites and branch
- points but not 3' splice sites. Of the 45 novel 5' splice sites found in their work, 11 are present
- in our raw data and 9 pass our p < 0.05 filter with at least one corresponding 3' splice site.
- 198 Gould et al. [9] combined lariat sequencing with RNA-seq to identify both 5' and 3' splice sites
- along with novel branch point sequences. Of the 213 novel splice sites they report, 194 are
- discoverable by our method, 133 are found in our raw data, and 114 pass our p < 0.05 filter.
- 201 The overlap between our work and these previous studies illustrates the power of our approach.
- None of the splice sites described by Qin et al. are found in any of the three other studies, while
- 203 each of Kawashima et al., Gould et al., and Schreiber et al. have greater overlap with our work

than with one another (figure 3). Taken together, of the 944 splice junctions predicted with p < 1000

205 0.05 by our method, 268 have been previously described and 676 are novel.

206

# 207 Saccharomyces cerevisiae contains antisense transcripts that undergo splicing

208 In addition to discovering novel splice sites within genes already known to contain introns and 209 unannotated ICGs, our method allows us to discover splicing that is antisense to annotated 210 transcripts, primarily in the degradation mutant strains  $xrn1\Delta$  and  $upf1\Delta$  and the histone 211 methyltransferase mutant set2 $\Delta$ . Previous studies report that Set2 suppresses antisense 212 transcription [16]. We find evidence for 50 antisense novel splice events spread across 41 213 transcripts that pass our statistical criteria. Of these 41 transcripts, 37 show a single novel 214 splice event, two transcripts have two unique novel splice events, one transcript has 3 separate events, and a single transcript, which is antisense to the LEU4 gene, has 6 unique novel splice 215 216 events. While most of the antisense splice events in our data have low read counts, the LEU4 217 isoforms together account for 51 reads across 11 samples, including wildtype, set2 $\Delta$ , and set2 $\Delta$ 218 prp43-1 at both 25° and 37°, set1 $\Delta$  and set1 $\Delta$  prp43-1 at 37° only, and xrn1 $\Delta$ , swr1 $\Delta$  upf1 $\Delta$ , and 219  $upf1\Delta snf2\Delta$ . Together, LEU4 antisense splicing represents over 20% of our total antisense 220 read counts across all samples. These six isoforms arise from 3 different 5' splice sites and 4 3'

splice sites. 5 out of 6 of these isoforms generate similar mature mRNAs with an intron in the

size range from 115 nucleotides to 129 nucleotides and are therefore indistinguishable by RT-

PCR (figure 2E). The remaining form is generated from a unique 5' splice and 3' splice and

- causes an intron of 464 nucleotides, and is low abundance in our RNA sequence data, with only
- a single read count in a single sample,  $set2\Delta$  at 37°.

226 In unannotated ICGs, we find 149 transcripts that show novel splicing in the sense direction and 227 37 that undergo novel splicing in the antisense direction. Interestingly, the number of transcripts 228 that show novel splicing in both the sense and antisense direction is just one, DJP1. If a set of 229 149 genes and a set of 37 genes are each randomly chosen from all Saccharomyces cerevisiae 230 transcripts, the expected value of the overlap is a single transcript, suggesting that within unannotated ICGs, presence of novel splicing in the sense direction does not significantly 231 232 impact the chances of novel spicing of an antisense transcript or vice versa. Within annotated ICGs, 220 transcripts undergo novel splicing in the sense direction and 4 have novel splicing in 233 234 the antisense direction, indicating no correlation between the splicing of annotated intron-235 containing genes and their corresponding antisense transcripts.

236

### 237 **Prp43 is required for efficient splicing of annotated and novel introns**

238 Several of the strains in our analysis include *prp43-1*, a temperature sensitive mutation that is

viable at 25° C but not at 37° C. Prp43 is an RNA helicase that has a vital role in spliceosome

disassembly and is required for efficient mRNA splicing in *Saccharomyces cerevisiae*. *PRP43* 

has also been implicated in ribosome biogenesis [17], and we previously showed that

decreasing levels of *PRP43* using a DAMP allele can suppress splice defects [13]. To

characterize the consequences of the *prp43-1* mutation in splicing, we compared the *prp43-1* strain to a wildtype strain, a set1 $\Delta$  prp43-1 strain to a set1 $\Delta$  strain, and a set2 $\Delta$  prp43-1 strain to

244 strain to a set  $2\Delta$  strain, a set  $\Delta$  pipes i strain to a set  $\Delta$  strain, and a set  $\Delta$  pipes i strain to a set  $2\Delta$  strain at both 25° and 37° C. As expected from its role in splicing, prp43-1 shows a

decrease in the splicing of annotated introns in *Saccharomyces cerevisiae* (figure 4A).

247 Interestingly, despite earlier findings that reducing levels of wildtype Prp43 can suppress splice

defects and promote splicing of weak introns [13], we find that strains with *prp43-1* show less

novel splicing than their counterparts with wildtype *PRP43* (figure 4B).

250

### 251 Elevated temperature favors novel introns that are longer than their annotated introns

252 Across our datasets, we detect fewer novel splice events in high temperature samples than 253 would be expected by sequence depth alone. This is unsurprising given that many of our lower 254 temperature samples are mutants that lead to accumulation of normally degraded products, 255 such as  $xrn1\Delta$  and  $upf1\Delta$ . While the total number of novel splice events is underrepresented at 256 high temperature, the novel splice products generated differ in intron length. When cells are 257 grown at 37° C, novel splicing within annotated ICGs tends to favor intron sizes that are larger 258 than novel splice forms from cells that are grown at 25° or 30° C (figure 5A). This can be explained by effects observed in our most common class of novel splice events, those that use 259 an annotated 5' splice site with a novel 3' splice site. At higher temperature, these novel 3' 260 splice sites tend to be further downstream than the annotated splice sites. We find that most of 261 262 the temperature-enriched splice events are more commonly found in our samples with wildtype 263 prp43, since prp43-1 decreases both annotated and novel splicing (figure 5B). These results

are consistent with work that finds that intron structure can function to control alternative splicingin yeast [18]. Nonetheless, our data reveal a number of previously unreported events.

266 As an example, TMH18, a mitochondrial membrane protein gene, has an annotated intron that 267 is 96 nucleotides long and two common novel splice isoforms discovered by our method, both of which generate longer introns. The isoform that is highly enriched in high temperature samples 268 contains a 161 nucleotide intron, while the isoform that is not favored at high temperature 269 270 contains a 128 nucleotide intron. We speculate that the increase in temperature destabilizes 271 the secondary structure of some pre-mRNA molecules to allow access to normally inaccessible splice sites. This may lead to an increase in use of distant novel splice sites. To confirm this, 272 273 we analyzed the predicted pre-mRNA secondary structure using MFOLD [19]. The predicted secondary structure with the most favorable free energy shows that the annotated 5' splice site, 274 275 the annotated 3' splice site, and the novel 3' splice site that is not favored at high temperature 276 are all readily accessible, while the novel 3' splice site that is favored only at high temperature is 277 not (figure 5C).

278 Interestingly, these distant splice sites may be under evolutionary pressure to be unable to code 279 for protein, as 74/84 (88%) of these temperature-enriched splice sites contain premature 280 termination codons. This is similar to the fraction of PTC-generating events found in the totality 281 of our novel splicing events in annotated intron-containing genes, with 572/636 (90%) containing 282 premature termination codons. This agrees with results described by Kawashima et al. [6] that 283 find that stress conditions, including heat shock, cause an increase in non-productive novel 284 splice usage. Together these data suggest that alternative splicing may be a mechanism for 285 reconfiguring the transcriptome in response to stress.

286

#### 287 Xrn1 deletion increases splice forms that do not use annotated splice sites

288 Novel splicing is found most commonly in strains in which xrn1 has been deleted. These strains account for approximately 9% of our total sequence depth but 19% of our total novel splice 289 290 counts. Interestingly, xrn1 deletion mutants are particularly enriched in novel splice forms that do not use any annotated splice sites. Roughly 18% of novel splice form counts that use either 291 292 an annotated 3' or 5' splice site derive from an xrn1 deletion mutant, while 31% of those that 293 utilize two novel splices are found in an xrn1 deletion strain. Two examples of unannotated 294 ICGs impacted strongly by xrn1 deletion are AGC1 and MRM2, both involved in mitochondrial 295 function. Interestingly, these genes' expression increases in  $xrn1\Delta$ , and they each have their 296 highest RPKM values in the five samples in which xrn1 is deleted [12, 13]. The 50 unannotated ICGs that are most enriched in our xrn1 $\Delta$  strains only have a single GO term in common, 297 298 "intracellular membrane-bounded organelle," further suggesting an impact on transcripts 299 important for mitochondrial function. Interestingly, a 2012 study found that Xrn1 is critical for the 300 translation of genes necessary for mitochondrial function in Saccharomyces cerevisiae [20]. 301 Together, this suggests a role for Xrn1 in the regulation of alternative splice products of 302 mitochondrial transcripts.

303

# 304 Ume6Δ-derived increase in IC-RPG expression leads to increase in IC-RPG alternative

305 splicing

306 Our  $ume6\Delta$  strain is highly enriched in novel splice events which utilize an annotated 3' splice 307 site with a novel 5' splice site. Of the newly discovered splice sites which utilize a novel 5'

splice site and a canonical 3' splice site. 59% are in intron-containing ribosomal protein genes

(IC-RPGs). However, of the events that are found disproportionately more frequently in *ume6* $\Delta$ ,

310 70% are in IC-RPGs. Our RNA sequence data suggests that IC-RPG expression increases

slightly in *ume6Δ*. Novel splicing in IC-RPGs increases by 70%, and 5' novel IC-RPG splicing

increases by 91% in  $ume6\Delta$ -containing samples relative to samples with wildtype ume6.

Previous studies have shown that Ume6 is degraded under meiotic conditions [21]. This raises

the interesting possibility that genetic manipulations that remove Ume6 may lead to changes in

the unannotated splicing landscape that are related to what occurs under meiotic conditions.

- 316 Ongoing experiments will test this model.
- 317

# 318 **Discussion**

319

In this study we show that there are many rarely utilized splice sites in Saccharomyces 320 321 cerevisiae. Our methodology is capable of discovering many novel splice forms by utilizing a 322 large number of RNA-seq datasets. We are able to do this in a robust, high-confidence manner by excluding reads that can be explained by sequencing error or genomic DNA contamination 323 324 and filtering based on existing splice site census sequence data. Overall, we find a strong 325 preference for novel splicing using a known 5' splice site and a novel 3' splice site within 326 annotated ICGs, which represent over half of our statistically significant novel splice products. 327 However, due to our high total sequencing depth and variety of strains and experimental 328 conditions we are still able to find a relatively large number of novel splice forms that utilize 329 known 3' splice sites with novel 5' splice sites, those that utilize novel 3' and 5' splice sites 330 within annotated ICGs, those that utilize two novel splice sites within unannotated ICGs, and 331 splicing of transcripts that are antisense to annotated genes. The large number of novel splice events that we discover allow us to correlate changes in splice site preferences with different 332 333 mutant strains and experimental conditions. While it has been tempting to call splicing events that have not been previously annotated as "errors," we believe that these data actually reveal 334 335 the remarkable substrate flexibility of an evolutionarily conserved enzyme. Moreover, it stands to reason that if splicing is to contribute to adaptation and, ultimately, evolution of multicellular 336 organisms, then an array of sequences, not simply those that match a strong consensus, need 337 338 to be recognized and spliced. The results described here provide a window into this sequence 339 landscape. In the adjoining manuscript [22], the authors find 229 "protointrons" - rapidly evolving, 340

inefficiently spliced introns. Of these, we define 60 as novel introns with an additional 9 found in
 our RNA sequence data but filtered out due to low splice site scores. The limited overlap

343 between the methods highlights that neither of our methods has reached saturation.

Furthermore, the 10 strains in our analysis with the greatest normalized overlap with Talkish et

al. are the 10 strains that include either  $xrn1\Delta$  or  $upf1\Delta$ , suggesting that overlap between the

346 methods is more driven by stabilization of protointron-containing splice products than biological

347 similarity between "hungry spliceosome" conditions and the strains used here. Talkish et al.

confirm by RT-PCR several splice events that would be removed from our analysis due to poor

- 349 splice signals, indicating that our method's stringency likely eliminates many true splice
- 350 products. The approach used here scores putative novel splice sites by learning from
- annotated splice sites, so as additional protointrons with unusual splice sites are discovered and
- validated, our method's ability to discover these forms will increase.
- 353 The analysis of several mutant strains allows insights into splice site selection in
- 354 *Saccharomyces cerevisiae*. We find general trends in our data as well as specific effects for
- 355 particular strains or conditions. For example, elevated temperature leads to an increase in
- novel splice form intron length,  $xrn1\Delta$  leads to a large increase in novel splice forms that do not
- use an annotated splice site, and  $ume6\Delta$  leads to an increase in novel splicing of intron-
- 358 containing ribosomal protein genes.
- 359 Despite the fact that analyzing many mutant strains can increase our understanding of splice
- site selection, every new category and class of splice form that we find is observed in our
- 361 wildtype data. While  $xrn1\Delta$  dramatically increases the number of splice products that arise from
- use of two novel splice sites, we see many examples of splice products that use two novel
- splice sites in our wildtype samples. This holds true for elevated temperature,  $ume 6\Delta$ , and
- $set2\Delta$ , as well as splicing of transcripts that are antisense to known transcripts, splicing that
- uses a single annotated splice site, and splicing in unannotated ICGs. Together, this indicates
   that while mutant strains are useful for correlating genetic changes with splicing changes, even
- wildtype Saccharomyces cerevisiae in normal conditions are capable of producing a large
   variety of splice products. Taken together, these findings illustrate the diversity and depth of
   splicing in Saccharomyces cerevisiae, and also show the presence of latent introns that are
   found across the genome.
- 371

### 372 Methods

373

#### 374 Public datasets

- 375
- 376 Saccharomyces cerevisiae wildtype,  $xrn1\Delta$ ,  $upf1\Delta$ ,  $swr1\Delta$ ,  $htz1\Delta$ ,  $swr1\Delta$   $xrn1\Delta$ ,  $swr1\Delta$   $upf1\Delta$ ,
- htz1Δ xrn1Δ, and htz1Δ upf1Δ strain RNA-seq data were download from GEO (accession
- number GSE97416). Additional wildtype,  $upf1\Delta$ , and  $xrn1\Delta$  RNA-seq data as well as  $snf2\Delta$ ,
- 1379  $ume6\Delta$ ,  $upf1\Delta$   $snf2\Delta$ ,  $xrn1\Delta$   $snf2\Delta$ , and  $ume6\Delta$   $snf2\Delta$  strain RNA-seq data were also
- downloaded from GEO (accession number GSE94404).
- 381

#### 382 Yeast culture and Sequencing

383

Wildtype, set1 $\Delta$ , set2 $\Delta$ , prp43-1, set1 $\Delta$  prp43-1, and set2 $\Delta$  prp43-1 strains were grown at 25°C to OD 0.3. Then, cultures were equally split, half remaining at 25°C and half shifting to 37°C for four hours. 10 mL of cells were pelleted from each sample and RNA extraction was performed. Then, 20 µg total RNA per sample was treated with DNAse I (Roche) and depleted of rRNA with the Ribo-Zero Gold rRNA Removal Kit (Illumina). RNA-seq libraries were prepared using the Illumina TruSeq v2 RNA Kit. 50 base pair paired-end reads were generated on an Illumina
 HiSeq 4000.

391

#### 392 Sequence alignment

393

RNA sequence data were combined and aligned to the Saccharomyces cerevisiae SacCer3 394 genome reference and Ares Lab Yeast Intron Database Version 3 [23] in a single step using 395 STAR [24] allowing up to six mismatches and no unannotated gaps. Sequence reads that fail to 396 397 align in this step are then aligned to the SacCer3 genome again, this time allowing no mismatches, a single gap, at most one alignment locus, and at least 15 nucleotide overhang on 398 399 each end of the alignment. Successful alignments in this step define putative novel introns. In 400 a final alignment step, all reads are aligned to the SacCer3 genome, the Ares Intron Database, 401 and newly defined putative novel introns in one step, allowing at most 1 alignment locus with up 402 to 2 mismatches. Counts for novel splice forms are based on those derived from this alignment

403 step.

404

#### 405 Splice site scoring

406

5' splice site, 3' splice site, and branch point scores were generated based on how closely each splicing signal matches the consensus sequence for that signal. For each position within a splice site, the total number of adenine, cytosine, guanine, and thymine bases present at that position in annotated splice signals was determined based on the Ares Lab Intron Database, and then the proportion of each nucleotide at that position was found by dividing by the total number of annotated splice products. The score for each splice site was calculated as follows:

$$Score = \sum_{i=1}^{k} N_i / X$$

Where i is the position within the splice signal, k is the number of positions in the splice signal, 413 414  $N_i$  is the count of that specific nucleotide at that position, and X is the total counts for the splice 415 signal. To score each putative novel splice form, the 5' and 3' splice sites are determined from 416 RNA-seg data, while branch point sequences were chosen by selecting the highest scoring 417 possible branch point sequence within a maximum distance of 200 nucleotides from the 3' 418 splice site. The score for each putative novel splice site is simply the product of its 5' splice site score, its 3' splice site score, and its branch point score. Putative novel splice sites were 419 420 considered antisense if the score in the antisense direction is higher than the score in the sense 421 direction.

422

#### 423 Statistical analysis of splice sites

424

P-values for putative novel splice sites were generated by comparing the potential splice site to

all possible combinations of 5' splice sites, 3' splice sites, and branch points. Based on a six 426 427 nucleotide 5' splice site, a three nucleotide 3' splice site, and a seven nucleotide branch point, 428 there are 4,294,967,296 possible combinations of splice signals that can be described. To 429 convert our putative novel splice form scores to p-values we compared the score to the scores 430 of all possible combinations of splice signals. The raw p-value is the fraction of these combinations that score equal to or better than the putative novel form, which also represents 431 the chance of seeing a splice score as good or better than this by chance. We then correct the 432 433 raw p-values for multiple hypothesis testing using the Bonferroni correction by multiplying each 434 raw p-value by the number of tests conducted, which is the total number of putative novel splice 435 sites times two, since each splice site score is calculated in both the sense and antisense

- 436 direction. All adjusted p-values greater than one are then set to one.
- 437

425

#### 438 Splicing Efficiency Calculation

439

To calculate the splicing efficiency of annotated splice sites, reads were aligned with STAR [24]

allowing only a single alignment locus, only annotated splice sites, and at most two mismatches.

Aligned reads within ICGs were categorized as exonic, spliced, or unspliced. We generated

normalized spliced and unspliced read counts by dividing the raw counts in each category by

the number of possible alignments that can fall into that category. This equates to read length

445 minus one for spliced reads and the intron length plus the read length minus one for unspliced

- reads. Splicing efficiency is then computed as normalized spliced counts divided by the sum of
- 447 normalized spliced and normalized unspliced counts.
- 448

### 449 **RNA Folding**

450

To find the optimum secondary structure for TMH18, we used the MFOLD web server with the

- 452 pre-mRNA sequence accessed from the Saccharomyces Genome Database
- 453 (https://www.yeastgenome.org/) and default MFOLD parameters [19;
- http://unafold.rna.albany.edu/?q=mfold]. The optimum secondary structure was visualized using
   MFOLD.
- 456

### 457 **RNA isolation and RT-PCR**

458

459 RNA was isolated from log phase yeast by hot phenol:chloroform:isoamyl alcohol (PCA)

460 extraction with SDS. The RNA was then precipitated with ethanol. 20 μg of total RNA was

461 DNase-treated with 30 U DNase 1 (Roche) for 1 hour at 25°C. 1 μg of DNase-treated RNA was

462 used for cDNA synthesis. cDNA synthesis was performed using the Maxima First Strand cDNA

463 Synthesis Kit (ThermoFisher). 1 μL of cDNA was used for the Taq PCR reaction using gene-

464 specific primers to analyze splicing.

465

# 466 Data Visualization

467

- Venn diagrams to view the overlap between this and previous works were generated using
- Venny 2.1.0 [27]. Box plots and bar graphs were generated using the MATLAB functions
- boxplot and bar, respectively.

471

# 472 Accession numbers

473

Data generated in this study is available under GEO accession number GSE120497.

475

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479

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Figure 1. Workflow for discovery of novel splice forms. Multiple RNA sequence datasets 548 are consolidated then aligned to the Saccharomyces cerevisiae genome and annotated 549 550 transcripts liberally, allowing up to six mismatches but no gaps. Reads that fail to align are 551 again aligned, but now allowing a single gap corresponding to a putative intron and no 552 mismatches. This step defines all putative novel introns. Once again all reads are aligned to 553 the Saccharomyces cerevisiae genome, known transcripts, and putative novel introns allowing 554 up to two mismatches, which determines each putative novel intron's read counts. Consensus 555 sequences are determined from known Saccharomyces cerevisiae 5' splice site, 3' splice site, 556 and branch point sequences, and each putative novel intron is scored based on sequence 557 similarity to the consensus sequences.

558

**Figure 2. RT-PCR of representative novel splice isoforms.** (A) *BIG1*; novel 3' splice site within annotated ICG. (B) *SIM1*; 5' UTR intron, novel 5' splice site within annotated ICG. (C) *MCR1*; 5' UTR intron, novel 5' and 3' splice sites within an annotated ICG. (D) *SPF1*; intron within an unannotated ICG. (E) *LEU4* antisense; antisense transcript of the *LEU4* gene containing an intron.

564

**Figure 3. Agreement with previous studies.** Venn diagram showing the overlap between the novel splice forms discovered here and those described previously.

567

Figure 4. *prp43-1* leads to a decrease in splicing. (A) Strains with the mutated *prp43-1* gene
 show a decrease in splicing efficiency of annotated introns relative to strains with wildtype
 *PRP43.* (B) Evidence of novel splicing is lower in strains with *prp43-1* versus wildtype *PRP43.*

571

Figure 5. Elevated temperature leads to longer introns. (A) Boxplot of temperatureenriched splice forms versus temperature-independent splice forms shows higher temperature
leads to an increase in intron size. Temperature-enriched splice forms have much higher
counts at higher temperature, but are still present at lower temperature. (B) Shifting *prp43-1* to

- 576 higher temperature causes a dramatic decrease in these temperature-enriched splice forms.
- 577 (C) Predicted structure of TMH18 with annotated and novel splice sites labeled.
- 578

579 **Figure S1. Putative novel splice site scores.** Each putative novel splice site was scored by 580 similarity to known *Saccharomyces cerevisiae* splice signals. The splice site score was plotted 581 against putative intron length for novel splice products within annotated ICGs (blue) and

- 582 unannotated ICGs (red).
- 583

**Figure S2. Summary of novel splice products.** (A) Over half of the data have fewer than 6 counts across all 29 of our datasets. (C) Novel splicing outside of annotated intron-containing genes and splicing of RNAs that are antisense to known transcripts is observed.













