1 <b>C</b>	nserved	Methy	yltransferase	Spb1	Targets
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# 2 mRNAs for Regulated Modification

## 3 with 2'-O-Methyl Ribose

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#### 19 SUMMARY

- 20
- 21 Non-coding RNAs contain dozens of chemically distinct modifications, of which only a
- 22 few have been identified in mRNAs. The recent discovery that certain tRNA modifying
- 23 enzymes also target mRNAs suggests the potential for many additional mRNA
- 24 modifications. Here, we show that conserved tRNA 2'-O-methyltransferases Trm3, 7,13
- and 44, and rRNA 2'-O-methyltransferase Spb1, interact with specific mRNA sites in

26	yeast by crosslinking immunoprecipitation and sequencing (CLIP-seq). We developed					
27	sequencing of methylation at two prime hydroxyls (MeTH-seq) for transcriptome-wide					
28	mapping of 2'-O-methyl ribose (Nm) with single-nucleotide resolution, and discover					
29	thousands of potential Nm sites in mRNAs. Genetic analysis identified hundreds of					
30	mRNA targets for the Spb1 methyltransferase, which can target both mRNA and non-					
31	coding RNA for environmentally regulated modification. Our work identifies Nm as a					
32	prevalent mRNA modification that is likely to be conserved and provides methods to					
33	investigate its distribution and regulation.					
34						
35						
36	KEYWORDS					
37	2'-O-methyltransferase, 2'-O-methylribose, CLIP-seq, MeTH-seq, mRNA modifications					
38						
39	HIGHLIGHTS					
40	MeTH-seq identifies 2'-O-methylribose genome-wide at single-nucleotide					
41	resolution					
42	Five conserved methyltransferases interact with yeast mRNA					
43	Spb1 is a major mRNA 2'-O-methyltransferase, and targets most ribosomal					
44	protein mRNAs					
45	• SPB1 expression is required to maintain normal levels of Spb1 target mRNAs					
46						
47	INTRODUCTION					
48						
49	Non-coding RNAs, in particular transfer RNAs (tRNA), are heavily modified with more					
50	than 100 distinct post-transcriptional modifications that affect RNA structure, stability,					

51	and function (Machnicka et al., 2013). The recent development of protocols for
52	transcriptome-wide mapping of certain RNA modifications has revealed a complex and
53	dynamic mRNA 'epitranscriptome' that includes N6-methyladenosine (m <sup>6</sup> A), 5-
54	methylcytidine (m <sup>5</sup> C), inosine (I), pseudouridine ( $\Psi$ ), 5-hydroxymethylcytidine (hm <sup>5</sup> C),
55	and N1-methyladenosine (m <sup>1</sup> A) in eukaryotic cells (Gilbert et al., 2016; Li et al., 2016;
56	Schaefer et al., 2017). Emerging evidence links regulated mRNA modifications, in
57	particular m <sup>6</sup> A, to post-transcriptional gene regulation in diverse biological systems
58	(Roignant and Soller, 2017; Yue et al., 2015). However, the full extent and diversity of
59	mRNA modifications is unknown.
60	
61	The discovery that some tRNA modifying enzymes, specifically pseudouridine synthases
62	(Pus), also target mRNAs (Carlile et al., 2014; Lovejoy et al., 2014; Schwartz et al.,
63	2014), suggests mRNAs might contain many more modifications than are currently
64	known. Notably, the conserved tRNA 2'-O-methyltransferase, Trm44, was identified by
65	proteomic analysis of highly purified UV cross-linked $poly(A)^+$ RNPs in yeast (Beckmann
66	et al., 2015), together with Pus1 and Pus7, which modify dozens of mRNA targets
67	(Carlile et al., 2014; Schwartz et al., 2014). Although no specific mRNA binding sites
68	were identified for Trm44, its association with polyadenylated RNA in vivo suggests it
69	may also target mRNAs in addition to tRNAs.
70	
71	2'-O-methyl ribose can occur on any base (Nm) and is an abundant and highly

conserved modification found at multiple locations in tRNA (Figure S1A), ribosomal RNA
(rRNA), and small nuclear RNA (snRNA). Yeast tRNAs are 2'-O-methylated by four
conserved Trm proteins. Trm3, orthologous to human TARBP1, modifies G18 or G19 in
the D-loop of 12 different tRNAs (Cavaillé et al., 1999). Trm7 (FTSJ1) modifies two
positions in the anticodon loops of tRNA<sup>Phe</sup>, tRNA<sup>Trp</sup>, and tRNA<sup>Leu</sup> (Pintard et al., 2002a).

77 Trm13 (CCDC76) modifies tRNA<sup>Gly</sup>, tRNA<sup>His</sup>, and tRNA<sup>Pro</sup> in the acceptor stem at position 4 or 5 (Wilkinson et al., 2007), and Trm44 (METTL19) modifies tRNA<sup>Ser</sup> at 78 79 position 44 or 45 in the variable loop (Kotelawala et al., 2008). Four additional conserved 80 2'-O-methyltranserases target cytosolic and mitochondrial ribosomes. Three of these are 81 protein-only enzymes: Spb1 (FTSJ3), which modifies Gm2922 of the 25S rRNA 82 (Lapeyre and Purushothaman, 2004), and Mrm1 (MRM1) and Mrm2 (MRM2), which 83 modify Gm2270 and Um2791, respectively, of the 21S rRNA (Pintard et al., 2002b; 84 Sirum-Connolly and Mason, 1993). The fourth, Nop1 (FBL), forms RNP complexes 85 containing additional proteins and C/D-box small nucleolar RNAs (snoRNA) that direct 86 methylation of dozens of sites on the 18S and 25S rRNAs by base pairing with the guide 87 RNA (Kiss-László et al., 1996; Kiss, 2002). Unlike the snoRNA-guided enzymes, the 88 basis for substrate specificity is incompletely understood for the protein-only 2'-O-89 methyltransferases. Thus, it is not possible to predict their RNA targets by bioinformatic 90 analysis.

91

92 2'-O-methylation – and regulation of this modification – has the potential to broadly affect 93 mRNA metabolism through effects on RNA structure, stability, RNA-protein interactions, 94 and translation. 2'-O-methyl ribose increases the thermodynamic stability of RNA:RNA 95 base pairs and stabilizes the C3' endo conformation of ribose found in A-form RNA 96 duplexes (Inoue et al., 1987; Kawai et al., 1992; Majlessi et al., 1998; Tsourkas et al., 97 2002). In addition, many RNA tertiary structures involve the 2' hydroxyl groups of ribose 98 (Butcher and Pyle, 2011), which can be disrupted by 2'-O-methylation (Lebars et al., 99 2008). Methylation of 2' hydroxyls can also inhibit RNA-protein interactions through steric 100 effects (Hou et al., 2001; Lacoux et al., 2012) or by impacting hydrogen bonding, which 101 frequently involves the 2' hydroxyl group (Jones et al., 2001; Treger and Westhof, 2001). 102 2'-O-methylated nucleotides are also resistant to a variety of nucleases (Sproat et al.,

103 1989). Finally, 2'-O-methylation of synthetic mRNAs interferes with translation and

104 induces site-specific ribosome stalling in vitro (Dunlap et al., 1971; Hoernes et al., 2015).

105 Thus, it is of particular interest to know which natural mRNAs contain Nm and what

106 enzymes install this modification in messenger RNAs.

107

108 Recent evidence suggests 2'-O-methyl ribose could be abundant in human mRNA, but 109 the methyltransferases were not identified (Dai et al., 2017). Here we determine mRNA 110 interaction sites genome-wide for five conserved RNA 2'-O-methyltransferases in yeast 111 by UV crosslinking and immunoprecipitation (CLIP-seq). We then develop MeTH-seq, a 112 genome-wide, single-nucleotide-resolution method to determine the locations of 2'-O-113 methyl ribose (Nm). We validate MeTH-seg by detecting known sites in non-coding 114 RNAs and conservatively identify 690 novel methyltransferase-dependent Nm sites in 115 veast mRNAs by comprehensive profiling following deletion or depletion of conserved 116 methyltransferases Trm3, 7, 13, and 44, which canonically target tRNA, and Spb1, which 117 targets rRNA. We identify novel targets for each methyltransferase and find that Spb1 is 118 the predominant mRNA methyltransferase in yeast. We further show that SPB1 is 119 required to maintain normal expression levels of a coherent regulon encoding proteins 120 necessary for ribosome assembly. Finally, we show that mRNA Nm sites are regulated 121 in response to environmental signals. Our results identify Spb1 as the primary 122 methyltransferase installing Nm as a widespread, regulated mRNA modification that is 123 likely to be conserved from yeast to man. 124

125 **RESULTS** 

126

127 Trm44 tRNA Methyltransferase Interacts with Specific mRNAs In Vivo

128 The yeast genome encodes more than 60 proteins characterized as tRNA modifying 129 enzymes (Phizicky and Hopper, 2010), of which several have recently been shown to 130 target mRNAs (Carlile et al., 2014; Lovejoy et al., 2014; Schwartz et al., 2014). We 131 hypothesized that mRNAs may be recognized and modified by additional enzymes 132 among this cohort. As a first step towards identifying likely mRNA modifying factors, we 133 examined published mRNA interactome data obtained by UV crosslinking of living yeast 134 cells followed by stringent purification of poly(A)<sup>+</sup> mRNPs and mass spectrometry 135 (Beckmann et al., 2015). Just four tRNA modifying enzymes – Pus1, Pus7, Dus3 and 136 Trm44 – were found to be mRNA-associated. Two of these, the pseudouridine 137 synthases Pus1 and Pus7, are known to modify dozens of mRNA targets (Carlile et al., 138 2014; Schwartz et al., 2014). In contrast, the known targets of Dus3 and Trm44 are 139 restricted to tRNAs. Given the broad potential for mRNA 2'-O-methylation to affect 140 mRNA metabolism, we selected Trm44 for further study. 141 142 To identify specific mRNAs that interact with Trm44, we performed in vivo UV

143 crosslinking (CL) followed by limited RNase digestion, immunoprecipitation (IP) of HPM-

tagged protein expressed from the endogenous *TRM44* locus (Trm44-HPM; Table S1),

and Illumina sequencing of the associated RNA fragments (Figure 1A; STAR Methods).

As expected, radiolabeling of the crosslinked immunoprecipitated RNA-protein

147 complexes (RNP) showed a prominent band of the predicted size for Trm44-HPM bound

to an intact tRNA, which was not present in control IPs from an untagged strain (Figure

149 S1B). We generated CLIP libraries from these samples using an enhanced protocol

150 (eCLIP-Seq) that increases the number of non-PCR duplicate reads (Van Nostrand et

al., 2016). In parallel, we prepared a size-matched input (SMI) library to control for

- abundant RNA fragments that contribute to non-specific background signal (STAR
- 153 Methods). This comparison is important for identifying true binding sites as normalization

of eCLIP reads to SMI has been shown to eliminate >90% of putative RBP binding sites
as false positives (Conway et al., 2016; Van Nostrand et al., 2016).

156

157 Consistent with the known methylation targets of Trm44, reads mapping to tRNA were 158 enriched >10-fold in the Trm44-HPM eCLIP libraries compared to SMI while reads 159 mapping to rRNA were depleted >50-fold (Table S2). Among tRNA eCLIP reads there was a striking enrichment for 6 tRNAs compared to SMI: 5 tRNA<sup>Ser</sup> that are known to be 160 161 methylated by Trm44 and 1 computationally predicted tRNA of undetermined specificity 162 that is very similar to serine tRNAs (Figure 1B) (Kotelawala et al., 2007; Lowe and Eddy, 163 1997). Next we applied the CLIPper algorithm (Lovci et al., 2013) to identify clusters of 164 CLIP-Seg reads and computed the enrichment of each cluster in IP compared to SMI 165 (STAR Methods). Using stringent enrichment criteria ( $p \le 0.001$  and  $\ge 4$ -fold enriched 166 versus SMI), we identified 33 high-confidence Trm44 binding sites, 60% of which were 167 located in various tRNA<sup>Ser</sup>. 3 stringent clusters were identified in mRNA, with 26 168 additional mRNA clusters showing enrichment at relaxed cutoffs (Figure 1C: Table S3). 169 As a further control, we compared Trm44-HPM eCLIP peaks to Pus1-HPM, which bound 170 a large set of tRNAs as expected and did not significantly enrich the Trm44 mRNA 171 binding sites (Figure S1C; Table S4). These results demonstrate that Trm44 interacts 172 with specific mRNA sites in vivo and suggests that mRNA could be 2'-O-methylated. 173 174 Discovery of mRNA Binding Sites for Trm3, Trm7 and Trm13 by eCLIP-Seq 175 Because mRNP interactome analysis did not identify all tRNA modifying pseudouridine 176 synthases with known mRNA pseudouridylation targets, we reasoned that additional

177 tRNA 2'-O-methyltransferases might associate with specific mRNAs in vivo. We

178 therefore performed eCLIP-Seq on yeast strains expressing HPM-tagged versions of

179 Trm3, Trm7 and Trm13 under their native promoters with SMI controls for each (Figure

180 S1D). As expected, tRNA mapping reads were enriched in these eCLIP libraries (Table 181 S2). CLIPper peak discovery and normalization to SMI identified stringent peaks for 182 each Trm protein ( $p \le 0.001$  and  $\ge 4$ -fold enriched versus SMI; STAR Methods), and 183 most known methylation targets were enriched including tRNA<sup>His</sup>, tRNA<sup>Leu</sup>, tRNA<sup>Ser</sup>, and tRNA<sup>Tyr</sup> for Trm3; tRNA<sup>Leu</sup>, tRNA<sup>Phe</sup>, and tRNA<sup>Trp</sup> for Trm7; and tRNA<sup>Gly</sup>, tRNA<sup>His</sup>, and 184 185 tRNA<sup>Pro</sup> for Trm13 (Figures S1E-S1G; Tables S5-S7). In addition, these eCLIPs each 186 produced numerous stringent peaks that mapped to specific mRNA sites distinct from 187 one another and from Trm44-HPM (Figures 1D-1I; Tables S3 and S5-S7). Thus, each of 188 the tRNA 2'-O-methyltransferases interacts with specific mRNAs in vivo in addition to 189 their canonical tRNA targets.

190

191 We also identified stringent peaks in non-coding RNAs not known to be methylation 192 targets of the Trm enzymes. A substantial fraction of significantly enriched non-coding 193 RNA peaks occurred in other tRNAs (Tables S3 and S5-S7). Moreover, in a few cases, 194 such as tRNA<sup>Ser</sup> crosslinking to Trm7 and tRNA<sup>Val</sup> crosslinking to Trm13, non-target 195 tRNAs showed reproducible enrichment that was comparable to canonical targets 196 (Figure S1F and S1G; Tables S6 and S7). Overall, each of the Trm proteins displayed in 197 vivo tRNA binding specificity that was similar but not identical to their known substrate 198 specificities for tRNA methylation. There is precedent for tRNA modifying enzymes to 199 bind with high affinity to tRNAs they do not modify (Keffer-Wilkes et al., 2016; Müller et 200 al., 2013) and our data highlight particular tRNAs for future investigations of the tRNA 201 features that distinguish modification substrates from non-substrates. A few Trm 202 crosslink peaks were observed in nuclear non-coding RNAs (Tables S5-S7). However, it 203 has been noted that certain non-coding RNAs, particularly snoRNAs, appear as frequent 204 hits in eCLIP-Seq experiments even after SMI normalization, and so such peaks should 205 be interpreted with caution (Van Nostrand et al., 2016). Together, these transcriptomewide crosslinking results show that the known tRNA 2'-O-methyltransferases in yeast

207 interact with diverse RNAs in vivo and have the potential to modify hundreds of

additional substrates including mRNAs.

209

#### 210 Transcriptome-wide Mapping of 2'-O-Methyl Ribose with MeTH-seq

211 To determine whether yeast mRNAs contain 2'-O-methyl ribose (Nm) and define the 212 global landscape of ribose methylation, we developed sequencing of methylation at two 213 prime hydroxyls, MeTH-seq, a high-throughput approach to map Nm sites with single-214 nucleotide resolution. Nm causes reverse transcriptase (RT) to pause one nucleotide 3' 215 to the methylated site, a pause that can be selectively enhanced by limiting the 216 availability of deoxyribonucleotides (Maden et al., 1995) or magnesium (Mg<sup>2+</sup>) in the 217 reaction (Figure S2A). We exploited this effect to map the locations of likely Nm sites using next-generation sequencing to identify Mg<sup>2+</sup>-sensitive RT pause sites (Figure 2A; 218 219 STAR Methods). Different reverse transcriptases and reaction conditions were 220 compared to optimize the sensitivity and specificity of MeTH-seq to identify known 2'-O-221 methylated sites in rRNA (Figure S2A and S2B). Limiting Mg<sup>2+</sup> produced slightly better 222 results than limiting dNTPs (Figure S2A). tRNA sites were not considered in this analysis 223 as tRNAs are poorly captured by most RNA-Seq library protocols including ours. 224

MeTH-seq produced clear peaks of reads one nucleotide 3' of known locations of Nm in rRNA (Figure 2B-2D). Deletion of specific methylation-directing small nucleolar RNAs (snoRNA) eliminated MeTH-seq peaks at their corresponding rRNA target sites (Figure S2C), further confirming the ability of our method to detect Nm sites with singlenucleotide resolution. We used the rRNA data from wild type cells to establish stringent criteria for de novo Nm identification from MeTH-seq data (STAR Methods). 33 out of 55 annotated Nm sites in rRNA were called under these criteria (Table S8; annotations from 232 (Piekna-Przybylska et al., 2007) with the addition of Gm562 in 18S identified by mass 233 spectrometry (Yang et al., 2015). As expected, the MeTH-seq peak caller missed some 234 Nm sites immediately 5' of other Nm sites, such as Gm805/Am807 in 25S, due to 235 'shadowing' from the downstream site (Table S8). MeTH-seg profiling of wild type yeast 236 ribosomes revealed two likely Nm misannotations: instead of 18S Gm1428 and 25S 237 Am1449, MeTH-seq produced clear signals one nucleotide 5' to the annotated sites, at 238 Am1427 and Um1448 respectively (Figure S2D). Inspection of previously published 239 primer extension gels supports these changes to the annotations, while a third potential 240 correction (Cm1639 to Gm1638) could not be confirmed or ruled out (gel images 241 available at http://lowelab.ucsc.edu/snoRNAdb/Sc/Sc-snos-bysite.html). After these 242 corrections, Nm sites in rRNA were called with an observed false positive rate of 0.35% 243 (Table S8; STAR Methods). The false positive peaks at Um795 in the 18S and Am816 in 244 the 25S rRNA can be attributed to 'stuttering' of RT at Am796 and Am817 (Motorin et al., 245 2007). A strong Mg<sup>2+</sup>-sensitive RT pause site was observed at 18S U1191, which 246 contains a complex RNA modification, m1acp3Y. A few false positive peaks were found 247 within broader regions of frequent RT pauses that may be due to stable RNA structure. 248 The remaining false positive peaks were gualitatively and guantitatively indistinguishable 249 from peaks at known Nm (Figure S2E and data not shown; Table S8). Importantly, none 250 of the false positive peaks was affected by deletion of snoRNAs (snR72-78). Thus, in 251 subsequent analysis, we required genetic dependence on a 2'-O-methyltransferase to 252 confidently classify a MeTH-seq peak as an Nm site.

253

Peak heights in rRNA were reproducible between replicates but varied 10-fold between
Nm sites whereas the level of methylation likely differs by no more than 2-fold based on
mass spectrometry (Figure S2F (Yang et al., 2015). Sequence context is thought to
affect the extent of RT pausing at 2'-O-methylated nucleotides (Motorin et al., 2007), and

capture biases during RNA-seq library preparation are well known (Raabe et al., 2014).
In particular, our approach likely underestimates methylation at NmC and NmG sites
(Figure S2G), which may be due to known CircLigase sequence preferences (Lamm et al., 2011). Thus, MeTH-seq peak heights cannot be meaningfully compared between
different Nm sites but may be used for relative quantitation of methylation at a given site
under different cellular conditions or genetic backgrounds.

264

#### 265 **Thousands of Candidate Sites for Regulated 2'-O-Methyl Ribose within mRNAs**

266 Having demonstrated the suitability of MeTH-seq for de novo discovery of Nm sites, we 267 performed MeTH-seq on poly(A)-selected yeast mRNAs during exponential growth in 268 rich medium ( $A_{600 \text{ nm}} = 1.0$ ). Given that the stoichiometry of methylation at known rRNA 269 sites is near 100% (Yang et al., 2015), we reasoned that requiring a peak height  $\geq 4$ 270 would miss partially methylated sites. The minimum peak height was therefore set as 2.0 271 for subsequent analyses of potential mRNA 2'-O-methylation. 6,734 sites in 1,947 272 mRNAs showed a MeTH-seq peak height ≥2.0 in at least 13 out of 16 independent 273 experiments (Figure 3A and 3B; Table S9). However, because we expect more false 274 positives with lower peak heights (Table S10), we subsequently imposed the additional 275 requirement that a peak be genetically dependent on a 2'-O-methyltransferase to be 276 called an Nm site. Relaxing the criteria for Nm identification further, by reducing the 277 requirements for minimum peak height or number of replicates, identified thousands of 278 additional candidate sites (data not shown). Overall, global MeTH-seq profiling suggests 279 Nm may be a prevalent modification in mRNA as well as non-coding RNA, a view 280 supported by genetic evidence as described below.

282	Pseudouridine and m <sup>6</sup> A are dynamically regulated mRNA modifications that respond to
283	nutrient availability and other stresses in yeast (Carlile et al., 2014; Schwartz et al.,
284	2013, 2014). To investigate the possibility of regulated Nm sites, we compared MeTH-
285	seq profiles from exponential and post-diauxic phases of growth, two growth states that
286	differ substantially in gene expression and metabolic activity (Figure S3A and S3B). De
287	novo Nm discovery in post-diauxic cultures ( $A_{600 \text{ nm}} = 12$ ) identified 7,900 candidate sites
288	in mRNAs with a MeTH-seq peak $\geq$ 2.0 in at least 16 out of 20 independent libraries
289	(Table S11), of which 3,981 were also identified in log phase cultures (Figures 3C).
290	Differences in mRNA abundance do not explain condition-dependent detection of most
291	Nm sites (Figure S3C). Together, these MeTH-seq data suggest extensive 2'-O-
292	methylation of mRNA targets that can be regulated in response to environmental
293	changes.
294	
295	Identification of Trm-Dependent 2'-O-Methyl Ribose Sites in mRNAs
296	The widespread distribution of MeTH-seq peaks together with the evidence that Trm
297	methyltransferases interact with hundreds of yeast mRNAs in vivo suggests that mRNAs
298	could be heavily decorated with 2'-O-methyl ribose. However, the unexplained MeTH-
299	seq peaks (Mg <sup>2+</sup> -sensitive RT pause sites) in rRNA indicates the potential for detection
300	of RNA features or modifications other than Nm in mRNAs. We therefore sought genetic
201	ovidence that mPNAs are modified by $2^{\prime}$ O methyltransference at the sites identified by

301 evidence that mRNAs are modified by 2'-O-methyltransferases at the sites identified by

302 MeTH-seq.

303

We profiled deletion strains lacking each of the four tRNA 2'-O-methyltransferases and
identified high-confidence Trm-dependent Nm sites based on reproducible loss of
MeTH-seq signal in independent biological replicates of *TRM* deletion mutants (STAR
Methods). As expected, deletion of *TRM44* (*trm44Δ*) eliminated signal from Um44 in

tRNA<sup>Ser</sup> (Figure 3D). *TRM13*-dependent methylation of the acceptor stems of tRNA<sup>Gly</sup>,
tRNA<sup>His</sup>, and tRNA<sup>Pro</sup> could not be assessed because the target nucleotides were too
close to the 5' ends of the tRNAs. Likewise, known Trm7 tRNA target sites were not
called as MeTH-seq peaks, likely due to the presence of RT-blocking m1A modifications
3' to Nm32 and Nm34. However, these technical limitations should not affect genetic
assignment to a particular 2'-O-methyltransferase of sites already identified by MeTHseq.

315

316 MeTH-seq profiling of *trm*<sup>Δ</sup> strains identified new RNA methylation targets for each of 317 the Trm proteins in log phase and post-diauxic cells. The largest total number of mRNA 318 candidate Nm sites were assigned to Trm44 (Figures 3D and 3E; Table S12), which is 319 consistent with the presumed greater abundance of Trm44 in poly(A)<sup>+</sup> RNPs (Beckmann 320 et al., 2015). The relatively low number of TRM3-dependent sites, particularly in log 321 phase, was unexpected given the widespread mRNA association observed by eCLIP 322 (Figure 1H). This may reflect technical limitations of the MeTH-seq approach or binding 323 interactions that do not result in modification (Discussion). Overall, our data show that, 324 like the Pus proteins, yeast Trm proteins target specific mRNAs for modification. Thus, 325 'moonlighting' activity towards mRNAs may be a common characteristic among tRNA 326 modifying enzymes.

327

#### 328 Conserved rRNA 2'-O-Methyltransferase Spb1 Targets mRNAs

The majority of candidate Nm peaks were unaffected by deletion of TRMs, suggesting additional 2'-O-methyltransferases may have a role in modifying mRNA. We considered each of the four yeast proteins with validated rRNA 2'-O-methylation target sites: Nop1, Spb1, Mrm1 and Mrm2. Nop1 selects its 18S and 25S rRNA targets through base pairing with a C/D snoRNA guide, which allows computational prediction of potential 334 target sites. All mRNA candidate Nm sites from OD1 and OD12 were examined for 335 possible base pairing with one of 48 snoRNAs followed by filtering for correct 336 positioning of the Nm site at +5 with respect to the snoRNA D box (STAR Methods) 337 (Kiss-László et al., 1996; Nicoloso et al., 1996). Sites positioned at +6 were also 338 included as they are consistent with known snoRNA-directed methylation (e.g. 25S 339 Cm650 targeted by snR18 and Gm1450 targeted by snR24 (Piekna-Przybylska et al., 340 2007)). Altogether, 10 novel Nm sites were identified as plausible targets of 7 snoRNAs 341 (Figures 4A and 4B; Table S13), suggesting some limited interactions of yeast C/D 342 snoRNAs outside their canonical rRNA targets.

343

344 We selected Spb1 for further characterization of the potential for mRNA methylation by 345 rRNA modifying enzymes. Spb1 localizes to the nucleoplasm in addition to the nucleolus 346 and thus has the potential to interact with nuclear mRNA (Kressler et al., 1999). In 347 contrast, Mrm1 and Mrm2 localize to mitochondria where they methylate 21S 348 mitochondrial rRNA (Breker et al., 2013; Pintard et al., 2002b; Sirum-Connolly and 349 Mason, 1993). In addition to its canonical methylation target, 25S rRNA, we found by 350 eCLIP that Spb1 crosslinked to numerous mRNAs (Figure S4A; Table S14). To identify 351 SPB1-dependent sites of 2'-O-methylation, the essential SPB1 gene was placed under 352 control of a repressible GAL1 promoter (pGAL-SPB1) and grown in glucose to inhibit 353 SPB1 expression prior to global MeTH-seg analysis in log phase (OD1) and post-diauxic 354 cells (OD12) (Figure 4C: STAR Methods). As expected, prolonged cell culture in the 355 absence of Spb1 reduced MeTH-seq signal at the known 25S target site, Gm2922, with 356 a peak height of 0.9-1.3 in Spb1-depleted cells compared to 2.6±0.7 across all SPB1<sup>+</sup> 357 libraries (Bonnerot et al., 2003; Lapeyre and Purushothaman, 2004). In addition, signal 358 at 393 (OD1) and 129 (OD12) Nm sites was reproducibly diminished following depletion 359 of Spb1 (Figure 4D; Tables S15 and S16). 87.5% of SPB1-dependent Nm sites mapped

360 to mRNAs of which 88.9% were Um (Figures 4E and 4F), which is consistent with the 361 known ability of Spb1 to methylate uridine (Bonnerot et al., 2003; Lapeyre and 362 Purushothaman, 2004). mRNA signal was strongly reduced following Spb1 depletion 363 with 181/393 OD1 and 114/129 OD12 sites decreased ≥4-fold (Tables S15 and S16). In 364 contrast, rRNA sites affected by Spb1 depletion showed modest though reproducible 365 reductions only in log phase, e.g. 25S Um1888 was reduced by ~20% (Figure S4B-S4D); 366 Table S14). Given that Um1888 and similarly affected rRNA sites are known targets of 367 C/D snoRNAs, these reductions in methylation are likely to be indirect effects of Spb1 368 depletion although we cannot exclude redundant targeting by Spb1 and snoRNAs 369 (Bonnerot et al., 2003; Lapeyre and Purushothaman, 2004). Of the five tested 2'-O-370 methyltransferase, our data identify Spb1 as the predominant enzyme acting on mRNAs 371 (Figure 4G).

372

## 373 Distribution of 2'-O-Methyl Ribose within mRNA Coding Sequences

374 SPB1-dependent MeTH-seq peaks were found in all regions of mRNAs with significant 375 enrichment of sites in CDS (Bonferroni corrected Fisher's exact test p < 8.1e-9) and 376 depletion of sites from 3' UTRs (p < 1.4e-6) (Figures 5A and 5B). Within coding regions 377 Nm was detected in 41 out of 61 sense codons and 1 out of 3 stop codons with 378 enrichment of specific A, I, L, V and M codons (Figure 5C, 5D and S5A; STAR Methods). 379 18/33 methylated M codons were AUG initiation codons, accounting for the higher 380 density of Nm at the start of CDS regions (Figure 5B). Nm sites were more frequently 381 observed in the 3<sup>rd</sup> position of sense codons, a distribution of codon positions that 382 differed significantly from a uniform background (Chi Square p<2.2e-16) (Figure 5E). 383 Based on reported position-specific translational effects of Nm on translation in E. coli 384 lysates (Hoernes et al., 2015), we analyzed ribosome footprint profiling data from wild 385 type yeast or  $dom34\Delta$  mutants defective in No-Go Decay (Guydosh and Green, 2014)

but found no evidence of translation pausing at Nm sites either within ORF bodies or at
start codons (Figure S5B-S5D; STAR Methods).

388

389 To explore potential mechanisms underlying mRNA recognition by Spb1, we examined 390 its targets for enriched RNA motifs using MEME (Bailey et al., 2015). A motif rich in UG 391 dinucleotides was notably enriched within a 50-nucleotide window surrounding SPB1-392 dependent Nm sites in mRNA (E value = 1.7e-40; Figure 5F: STAR Methods). This motif 393 was present in 71/409 mRNA target sites while other significantly enriched motifs 394 occurred much less frequently (Figure S5E). The UGNUGN motif was located at variable 395 distances from the modified site (Figure S5F), which was almost invariantly U (Figure 396 4F). Non-coding RNA targets of FtsJ methyltransferases, including Spb1, are modified 397 on unpaired nucleotides within stem-loop structures (Guy and Phizicky, 2014), but the 398 importance of this structure for Spb1 activity is unknown (Bonnerot et al., 2003; Lapeyre 399 and Purushothaman, 2004). Using RNAsubopt to identify the probable structures of 400 Spb1 targets (STAR Methods), we found that the modified sites were significantly likelier 401 to be in the loop of a hairpin than a background set of sites. Similarly, positions >= 4 nt 402 upstream or downstream of the site were significantly likelier to be involved in base 403 pairing (Figures 5G and S5G). The basis for site-specific methylation by Spb1 remains to 404 be determined and may involve distinct co-factors for different sites as shown for the 405 related methyltransferase Trm7 (Guy and Phizicky, 2014). 406

100

407 Spb1 Methylates and Maintains Normal Levels of mRNAs Required for Ribosome
408 Assembly

To gain insight into the biological roles of mRNA methylation by Spb1, gene ontology

410 (GO) enrichment analysis was performed. GO terms related to ribosomes, ribosome

411 biogenesis, and translation were almost exclusively over-represented (Figure 6A; Table

412 S17). These related GO term enrichments were driven by the presence of SPB1-

- 413 dependent Nm sites in 97/139 (69%) mRNAs encoding cytoplasmic ribosomal proteins
- 414 (RP mRNA). "Translational elongation" was also significantly enriched due to
- 415 methylation of mRNAs encoding elongation factors EF-1 alpha (*TEF1/2*), EF-1B (*TEF4*),
- 416 EF-2 (*EFT1*), and eIF-5A (*HYP2*) in addition to proteins of the ribosomal stalk (*RPP1A/B*
- 417 and *RPP2A/B*) that promote recruitment of elongation factors to ribosomes (Gonzalo and
- 418 Reboud, 2003). Thus, Spb1 targets a coherent regulon of factors required for ribosome
- 419 assembly and function that includes both its canonical pre-rRNA substrate and
- 420 numerous RP mRNAs.
- 421

422 What might be the function of mRNA modification by Spb1? To determine whether loss 423 of Spb1 affects the abundance of its target RNAs we performed RNA-seq on Spb1-424 depleted and control cells (STAR Methods). In wild type cells, Spb1 target mRNAs were 425 highly expressed with a median TPM (transcripts per million) of 608.5 compared to 15.5 426 TPM for all genes with adequate read coverage for MeTH-seg analysis (KS test p < p427 2.2e-16; Figure S6A). Depletion of Spb1 lead to significantly decreased abundance of 428 most targets, with somewhat larger changes for mRNAs containing multiple Nm sites 429 (Figure 6B and 6C). RP target mRNAs showed the largest reductions (Figure 6D). 430 Although some of this difference may stem from global effects of Spb1 depletion on RP 431 expression (Figure S6B), target RP mRNAs were significantly reduced compared to non-432 target RP mRNAs (KS test, p < 5.9e-15; Figure S6C). This reduction in mRNA target 433 levels following methyltransferase depletion was specific to Spb1: Trm44 targets 434 increased slightly in *trm44* $\Delta$  (KS test, *p* < 5.7e-7; Figure S6D and targets of the other 435 methyltransferases were not significantly affected by the corresponding knockout (data 436 not shown). These data show that SPB1 is required to maintain the levels of its target 437 mRNAs and suggest Spb1-dependent mRNA methylation may promote mRNA stability.

- 438 Because a majority of cytoplasmic ribosomal protein mRNAs are targets of SPB1-
- 439 dependent methylation, the role of *SPB1* in ribosome biogenesis is likely to be broader
- 440 than previously appreciated (Discussion).
- 441

#### 442 **DISCUSSION**

- 443 Here we identify Spb1 as a conserved methyltransferase that modifies hundreds of yeast
- 444 mRNAs with 2'-O-methyl ribose (Nm). We further show that SPB1 expression is required
- to maintain normal levels of Spb1 target RNAs including most mRNAs encoding
- 446 ribosomal proteins. Our results expand the known epitranscriptome and establish
- 447 methods to discover Nm sites transcriptome-wide and illuminate the mechanisms
- 448 underlying regulation of this novel mRNA modification that is likely to be conserved
- throughout evolution.
- 450

#### 451 **Detection of 2'-O-methyl Ribose – Methods and Limitations**

452 Three next-generation sequencing methods for Nm profiling have recently been 453 described. Our MeTH-seq library preparation is very similar to the 20Me-seq method 454 used to identify new Nm sites in mammalian ribosomes (Incarnato et al., 2017), but limits 455 Mg<sup>2+</sup> rather than dNTPs during reverse transcription, which we found to modestly 456 improve selectivity for Nm. The RiboMethSeq approach exploits the resistance of 2'-O-457 methylated sites to hydrolysis (Krogh et al., 2016; Marchand et al., 2016). An advantage 458 of this method is the possibility to determine the absolute stoichiometry of modification at 459 specific positions by quantifying the depletion of 3' and 5' read ends. However, very high 460 read depth is required to observe depletion of reads at an Nm site making it costly for 461 transcriptome-wide profiling. The Nm-seq method exploits the differential reactivity of 462 Nm to periodate oxidation for selective capture of RNA fragments with Nm 3' ends (Dai 463 et al., 2017). Pre-enrichment of Nm-containing RNA fragments (e.g. with an antibody)

464 prior to MeTH-seq, 2OMe-seq or RiboMethSeq would allow comprehensive Nm 465 discovery, with single-nucleotide resolution, at much lower sequencing depth. However, 466 enrichment of modified RNA fragments prior to sequencing, which is standard in m6A 467 and m1A profiling studies, may result in capture of RNAs that are modified at very low 468 stoichiometry and have uncertain physiological relevance. In contrast, as implemented 469 here, our approach requires RT to pause on a substantial fraction of total RNA 470 molecules to detect a peak for a given Nm site.

471

472 MeTH-seq may not capture all 2'-O-methylated sites. It is notable that MeTH-seq 473 identified relatively few TRM3-dependent Nm sites despite the widespread mRNA 474 association of Trm3 observed by eCLIP. Remarkably, >96% of high-confidence binding 475 sites identified in the Trm3 eCLIP libraries mapped to mRNAs. Altogether, we identified 476 1611 stringent peaks in 975 mRNAs in addition to 29 stringent peaks that mapped to 19 477 tRNAs. Such pervasive mRNA binding by Trm3 was unexpected, and the basis for it is 478 not immediately apparent. Trm3 does not have known RNA-binding domains outside the 479 conserved methyltransferase catalytic domain. Furthermore, Trm3 is not more highly 480 expressed than the other tRNA 2'-O-methyltransferases (Ghaemmaghami et al., 2003; 481 Kulak et al., 2014), and all four HPM-tagged Trm proteins were immunoprecipitated with 482 similar efficiency. Given that Trm3 methylates its canonical tRNA target sites in the 483 context GmG, this discrepancy between detection of binding and modification may 484 reflect false negatives due to sequence capture bias in the MeTH-seq protocol. 485 Specifically, truncated cDNAs ending in C are inefficiently circularized by CircLigase 486 (Lamm et al., 2011). In addition, the true methylation targets may interact with Trm3 very 487 transiently and therefore be poorly captured by CLIP. Alternatively, Trm3 crosslinking 488 may enrich for deadenylated mRNAs that were not examined by our profiling of Nm sites 489 in poly(A)<sup>+</sup> RNA. There is also precedent for tRNA modifying enzymes to bind with high

490 affinity to non-substrate RNAs, perhaps to affect RNA folding (Keffer-Wilkes et al., 2016;
491 Müller et al., 2013).

492

493 The majority of highly reproducible MeTH-seq peaks (those present in at least 13 494 independent samples) identified here were not affected by depletion of known 2'-O-495 methyltransferases. As the TRM methyltransferases and Spb1 do not modify one 496 another's non-coding RNA targets in vivo, we consider widespread redundant targeting 497 of mRNA sites by these enzymes to be unlikely. The mitochondrial enzymes Mrm1 and 498 Mrm2, whose modification targets were not determined in our study, conceivably 499 methylate cytoplasmic mRNAs either en route to the mitochondria or from a cytoplasmic 500 pool of protein (Breker et al., 2013). Consistent with this possibility, it is interesting that 501 the ortholog of Mrm1 crosslinks to  $poly(A)^+$  RNA in human cells (Baltz et al., 2012; 502 Castello et al., 2012). In addition, the S. cerevisiae genome encodes multiple predicted 503 RNA methyltransferases of unknown activities (Wlodarski et al., 2011), which potentially 504 include additional mRNA 2'-O-methyltransferases. 505 506 However, although other known or putative RNA methyltransferases may install Nm in 507 mRNA, it is likely that other RNA features, such as structure, can give rise to 508 reproducible Mg<sup>2+</sup>-sensitive RT pause sites in the absence of 2'-O-methylation. In any 509 sequencing library preparation there is the potential for robust artefacts, such as 510 mispriming during reverse transcription (Gillen et al., 2016), which may explain the 511 enrichment of a primer sequence motif among candidate Nm sites proposed from 512 Nmseq data (Dai et al., 2017). Therefore, we do not interpret MeTH-seq peaks as Nm 513 sites in the absence of genetic validation. The remaining potential mRNA 2'-O-514 methyltransferases need to be tested before the complete Nm landscape can be known

515 in yeast.

516

#### 517 **Conservation of mRNA 2'-O-methylation**

518 While our study was being finalized, He and colleagues reported the presence of Nm in 519 mRNA from human cells based on mass spectrometry and sequencing-based 520 approaches, although they did not identify the methyltransferases responsible (Dai et al., 521 2017). Each of the mRNA modifying yeast methyltransferases characterized here is 522 conserved to man. Notably, FTSJ3, which is the human ortholog of Spb1, here identified 523 as a predominant mRNA methyltransferase in yeast, has been found in the poly(A)<sup>+</sup> 524 mRNP interactomes of multiple human cell lines (Baltz et al., 2012; Beckmann et al., 525 2015; Castello et al., 2012). Proteomic characterization of FTSJ3 co-immunoprecipitates 526 identified 10 hnRNP proteins providing further evidence that FTSJ3, like Spb1, may 527 interact with nuclear mRNA (Simabuco et al., 2012). Our analysis also suggests some 528 mRNAs are targeted by the snoRNA-directed methyltransferase Nop1, whose human 529 counterpart fibrillarin was likewise found to crosslink to poly(A)<sup>+</sup> human RNA. Although 530 the reported locations of Nm in human mRNA have been called into question based on 531 the apparent mispriming artefact described above, mass spec analysis establishes 2'-O-532 methylated nucleotides as abundant in purified bulk human mRNA (Dai et al., 2017). 533 Furthermore, Um was by far the most abundant with a molar ratio of 0.15% for Um/U, 534 which is consistent with the predominance of Um among genetically validated Spb1-535 dependent Nm sites. Thus, we conclude that 2'-O-methyl ribose is conserved as an 536 mRNA modification in diverse eukaryotes and suggest that the mRNA methylation 537 activity of Spb1/FTSJ3 in particular is likely to be conserved.

538

## 539 **Functional consequences of Nm in mRNA**

540 We found that depletion of Spb1 led to substantial reductions in the levels of its mRNA 541 targets, which suggests endogenous Nm could promote mRNA stability. 2'-O-methylated 542 nucleotides are widely used in synthetic small interfering RNA (siRNA) due to their 543 resistance to degradation by various nucleases (Czauderna et al., 2003; Sproat et al., 544 1989). However, it is unclear how the presence of sparse Nm residues within coding 545 sequences would stabilize an mRNA against exonucleolytic decay from the 5' or 3' end 546 of the transcript. If Nm increases steady-state mRNA levels by conferring resistance to 547 endonucleolytic cleavage at the modified site, this uncharacterized decay pathway must 548 play a sizable role in yeast mRNA homeostasis. Alternatively, Nm may recruit a protein 549 factor that promotes mRNA stabilization. Such a mechanism would be opposite to 550 mRNA destabilization by m6A, which occurs through direct binding of YTHD family 551 proteins to modified mRNA and subsequent recruitment of decay factors (Ke et al., 552 2017; Wang et al., 2014). Any Nm 'reader' likely recognizes the modified nucleotide in a 553 broader sequence or structural context; recognition of additional RNA features could 554 explain why loss of Nm affects the abundance of Spb1 targets specifically whereas loss 555 of Nm did not decrease the levels of mRNAs targeted by other methyltransferases.

556

557 Consistent with this possibility, the presence of Nm has been found to affect interactions 558 with several RNA binding proteins in both natural and artificial contexts (Devarkar et al., 559 2016; Lacoux et al., 2012; Lavoie and Abou Elela, 2008; Simon et al., 2011; Tian et al., 560 2011). With the exception of PAZ domains binding the 2'-O-methylated 3' ends of small 561 RNAs, most characterized RNA binding protein interactions with 2'-O-methylated RNAs 562 show reduced protein binding to modified RNA. In contrast, m<sup>6</sup>A is known to promote 563 binding to multiple 'reader' proteins to mediate a variety of downstream effects on mRNA 564 metabolism (Roignant and Soller, 2017; Yue et al., 2015). Our findings motivate the 565 search for Nm readers and their functional interactions with mRNA decay pathways.

567 We anticipated that mRNA 2'-O-methylation might impede translation elongation based 568 on results with synthetic Nm containing mRNAs (Dunlap et al., 1971; Hoernes et al., 569 2015). We therefore analyzed ribosome footprint profiling data to determine ribosomal A 570 site occupancy at modified codons. No pausing was observed either in wild type cells or 571 in *dom34* mutants that are defective in ribosome stalling-dependent No-Go Decay 572 (Guydosh and Green, 2014) nor did deletion of DOM34 increase steady-state levels of 573 translating Nm-containing mRNAs. Given that Spb1 methylation targets are among the 574 most highly expressed and efficiently translated mRNAs in growing yeast, it would be 575 surprising if their expression were inhibited at the level of translation elongation. 576 Furthermore, depletion of Spb1 decreased target mRNA levels, which is the opposite of 577 what would be expected for a modification that slowed elongation (Hanson and Coller, 578 2017). The lack of detectable ribosome pausing at Nm codons in vivo may reflect more 579 robust translation elongation in cells compared to lysates. However, CAmA, which 580 caused a 10-fold reduction in synthesis of full-length protein in E. coli lysates (Hoernes 581 et al., 2015), was found infrequently among Nm-containing codons in yeast and the 582 ribosome profiling sequencing depth was not sufficient to assess pausing at this small 583 number of codons in isolation. Thus, we cannot exclude the possibility that specific Nm 584 modified codons could inhibit elongation in cells.

585

#### 586 A Broader Role for Spb1 in Ribosome Biogenesis?

587 A wealth of genetic and biochemical evidence implicates SPB1 in ribosome biogenesis,

588 but the essential function of the Spb1 protein in this process remained unclear. SPB1

589 was originally characterized as one of seven Suppressors of Poly(A) Binding Protein that

affect the levels of 60S ribosomal subunits in yeast (Sachs et al., 1987). Depletion of

591 Spb1, which co-purifies with 66S pre-ribosomal particles, leads to accumulation of rRNA

592 processing intermediates and depletion of mature 25S rRNA (Harnpicharnchai et al.,

593 2001; Kressler et al., 1999). The subsequent discovery that Spb1 methylates 25S rRNA 594 revealed a molecular function for its conserved methyltransferase domain (Bonnerot et 595 al., 2003; Lapeyre and Purushothaman, 2004). Mutating the predicted catalytic aspartate 596 (spb1-D52A) abolished the 25S Gm2922 modification and produced a viable but 597 severely growth impaired strain with greatly reduced 60S levels (Lapeyre and 598 Purushothaman, 2004). However, it is remarkable for a single Nm site to be critical for 599 ribosome biogenesis. Indeed, most yeast C/D snoRNAs can be deleted with little effect 600 on growth. Thus, the apparent importance of Spb1-dependent RNA methylation for 601 ribosome production and cell growth was puzzling. 602 603 Our work identified hundreds of new RNA targets for 2'-O-methylation by Spb1 that 604 include the majority of mRNAs encoding cytoplasmic ribosomal proteins. Depletion of 605 Spb1 led to significantly reduced levels of its mRNA targets consistent with a positive 606 role for SPB1 – and presumptively RP mRNA 2'-O-methylation – in ribosomal protein 607 production. Spb1 modification targets included RP mRNAs encoding proteins of both 608 ribosomal subunits, which could contribute to the reduction in 40S observed following 609 longer depletion of Spb1 (Kressler et al., 1999). Of note, although the human ortholog 610 FTSJ3 is also required for ribosome biogenesis, it contributes primarily to 40S 611 production (Morello et al., 2011). It will be important to learn whether FTSJ3 similarly 612 modifies RP mRNAs and determine how mRNA 2'-O-methylation affects RNA levels in 613 human cells.

614

615

#### 616 **ACKNOWLEDGEMENTS**

We thank E. Van Nostrand and G. Yeo for eCLIP advice; B. Waldman for help with
MeTH-seq feasibility studies; C. Burge, E. Phizicky and members of the Gilbert Lab for

- discussion. The sequencing was performed at the MIT BioMicro Center under the
- direction of S. Levine. This work was supported by grants from The American Cancer
- 621 Society Robbie Sue Mudd Kidney Cancer Research Scholar Grant (RSG-13-396-01-
- 622 RMC) and the National Institutes of Health (GM094303, GM081399) to W.V.G. T.M.C.
- and K.M.B. were supported by fellowships from The American Cancer Society. C.S. was
- 624 supported by an NSF GRF.
- 625

## 626 AUTHOR CONTRIBUTIONS

- 627 K.M.B. designed research and performed and analyzed MeTH-seq experiments. T.M.C.
- 628 developed the MeTH-seq data analysis pipeline. C.S. designed and performed all other
- 629 computational analyses. W.V.G. conceived the project, supervised research, designed
- 630 research, and performed eCLIP. K.M.B., C.S. and W.V.G. interpreted the results. W.V.G.
- 631 wrote the paper with input from all authors.
- 632

## 633 **DECLARATION OF INTERESTS**

- 634 The authors declare no competing interests.
- 635

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1005	
1006	Material and Methods
1007	
1008	CONTACT FOR REAGENT AND RESOURCE SHARING
1009	
1010	Further information and requests may be directed to and will be fulfilled by the Lead
1011	Contact, Wendy Gilbert (wendy.gilbert@yale.edu).
1012	
1013	EXPERIMENTAL MODEL AND SUBJECT DETAILS
1014	
1015	Yeast strains and growth
1016	All yeast strains are Saccharomyces cerevisiae BY4741 or BY4742 derivatives
1017	(BY4742:wild type (YWG11), BY4741:wild type (YWG506), TRM44-HPM (YWG1354),
1018	PUS1-HPM (YWG1357, YWG1358), TRM3-HPM (YWG1348, YWG1349), TRM7-HPM
1019	(YWG1350, YWG1351), TRM13-HPM (YWG1352, YWG1353), SPB1-FLAG (YWG1359,
1020	YWG1523), snr72-78∆ (YWG318, YWG372) <i>, trm3</i> ∆ (YWG1320, YWG1321) <i>, trm</i> 7∆
1021	(YWG1322, YWG1323), <i>trm13</i> ∆ (YWG1326, YWG1327), <i>trm44</i> ∆ (YWG1324,
1022	YWG1325), <i>trm</i> 732∆ (YWG1332, YWG1333), <i>rtt10</i> ∆ (YWG1330, YWG1331), <i>pGAL1-</i>
1023	HA-SPB1 (YWG1522, YWG1523). See Table S1 for complete genotypes. snoRNA
1024	deletions, <i>trm13</i> $\Delta$ and <i>pGAL-HA-SPB1</i> strains were made using PCR-based deletion
1025	cassettes (Longtine et al., 1998). Other deletion strains were obtained from the Yeast
1026	Deletion Collection (Winzeler et al., 1999). His-PrecissionProtease-MYC (HPM) tagged
1027	strains were made using PCR-based cassettes (Graumann et al., 2004). All strains, with
1028	the exception of pGAL-HA-SPB1, were grown at 30°C in YPAD (1% yeast extract, 2%
1029	peptone, 0.01% adenine hemisulphate, 2% glucose). Cultures for MeTH-seq were

1030 harvested by centrifugation in log phase ( $A_{600nm} \approx 1.0$ ) or at high density ( $A_{600nm} \approx 12.0$ ).

1031 High density *trm7* $\Delta$  cells were grown to A<sub>600nm</sub> $\approx$ 8.0.

1032

- 1033 The *pGAL-HA-SPB1* depletion strain was maintained at 30°C in YPARG (YPA
- 1034 with 20% raffinose, 30% galactose). Cells were transferred to YPAD to inhibit expression
- 1035 of SPB1. Log phase cells: Cells were grown to an A<sub>600nm</sub>≈0.3, centrifuged and
- 1036 resuspended in YPAD media for an additional 4-6 hours until A<sub>600nm</sub>≈1.0. High density
- 1037 cells: Cells were grown to an  $A_{600nm} \approx 3.0$ , centrifuged and resuspended in YPAD media
- 1038 for an additional 8-12 hours until  $A_{600nm} \approx 12.0$ .
- 1039
- 1040 For ePAR-CLIP assays yeast were grown at 30°C in minimal media with reduced

1041 uracil (SC<sub>Ura120µM</sub>) to an A<sub>600nm</sub>=0.4-0.5, supplemented with 4-thiouracil to a final

1042 concentration of 500µM and cultured for 3 more hours before harvest as described

1043 (Beckmann, 2017).

1044

1045

1046 **METHOD DETAILS** 

1047

#### 1048 Yeast PAR-CLIP

1049 Yeast strains expressing tagged methyltransferases from their endogenous loci were

1050 cultured in the presence of 500µM 4-thiouracil. Cells were harvested by centrifugation,

- 1051 resuspended in ice cold water, and UV irradiated on ice in a Stratalinker (365 nm,
- 1052 energy =  $7.2 \text{J/cm}^2$ ). Crosslinked cells were washed in ice water, flash frozen in liquid N<sub>2</sub>,
- and stored at -80°C until lysis. Cells were resuspened in 1.5 mL iCLIP lysis buffer (50
- 1054 mM Tris-HCl pH7.4, 100 mM NaCl, 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate,

- 1055 1X protease inhibitor cocktail (Millipore), 400U/mL Murine RNase inhibitor (NEB)) per g
- 1056 of cell pellet and lysed by vortexing with glass beads (Manufacturer). Lysates were
- 1057 clarified by centrifugation, flash frozen in liquid  $N_2$  and stored at -80°C.
- 1058

#### 1059 eCLIP library preparation

1060 eCLIP libraries were prepared as described (Van Nostrand et al., 2016). Briefly, lysates 1061 were diluted in iCLIP lysis buffer to 26 ODU/2 mL and treated with Turbo DNase 1062 (Lifetech) and RNase I (Lifetech) for 15min at 22°C before placing on ice. Treated 1063 lysates were centrifuged (15,000 x g 15min) and added to Protein G magnetic beads 1064 (Dynabeads) pre-bound with 10 µg of antibody (anti-Myc or anti-FLAG, Sigma) and 1065 rotated for 2.5hr at 4°C. 4% of the binding reaction was saved as input before extensive 1066 washing of bead-bound protein-RNA complexes. Bound RNA 3' ends were 1067 dephosphorylated with FastAP (Lifetech) and PNK (NEB) before on-bead ligation of a 3' 1068 RNA linker with T4 RnI (NEB) at 16°C overnight. After washing, 10% of bound RNA was labeled by PNK with <sup>32</sup>P ATP for diagnostic gels. The remainder was eluted from the 1069 1070 beads by incubation in 1X NuPAGE loading buffer with 0.1M DTT at 70°C for 10min. 1071 CLIP eluates and paired inputs were electrophoresed on 4-12% Bis-Tris NuPAGE and 1072 transferred to nitrocellulose membranes. Tagged protein MW was determined by 1073 western blot of 2% input before CLIP and size-matched input (SMI) membranes were 1074 cut, taking a 75kDa region just above the target protein size. RNA was released with 1075 proteinase K, extracted with phenol:chloroform:isoamy alcohol, and purified on RNA 1076 Clean & Concentrator-5 columns (Zymo). SMI RNA was dephosphorylated, a 3' linker 1077 ligated, and purified on silane beads (Dynabeads). CLIP and SMI RNA was reverse 1078 transcribed with SuperScript IV, hydrolyzed with NaOH, neutralized with HCI, and cDNA 1079 was purified on silane beads. A 5' linker was ligated with T4 RnI (NEB) overnight at 1080 16°C. Linker-ligated cDNA was purified on silane beads before diagnostic PCRs to

- 1081 determine the minimum number of cycles. Final PCR products were gel-purified,
- 1082 precipitated and sequenced on an Illumina HiSeq 2000.
- 1083

#### 1084 **RNA isolation**

- 1085 Total yeast RNA was isolated by hot acid phenol extraction followed by isopropanol
- 1086 precipitation as described (Carlile et al., 2015). PolyA+ mRNA was isolated from 10-
- 1087 15mg of yeast total RNA by two rounds of selection on oligo-dT cellulose beads
- 1088 according to the manufacturer's instructions (NEB).
- 1089

#### 1090 **MeTH-seq library preparation**

- 1091 RNA fragmentation was performed in 10mM ZnCl<sub>2</sub> at 95°C for 55s. RNA fragments were
- 1092 precipitated, dephosphorylated with CIP and PNK, and separated by denaturing (8%
- 1093 urea-TBE) polyacrylamide gel electrophoresis (PAGE). Gel-purified RNA fragments (60-
- 1094 70nt, 70-80nt, 80-90nt) were eluted overnight with rocking in RNA elution buffer (300mM
- 1095 NAOAc pH 5.5, 1mM EDTA, 100 U/ml RNasin (Promega) and ligated to a pre-
- adenylated adaptor (/5Phos/TGGAATTCTCGGGTGCCAAGG/3ddC/) (IDT) using T4
- 1097 RNA ligase (NEB) at 22°C for 2.5h, followed by precipitation. Reverse transcription was
- 1098 completed using AMV-RT (Promega) and the reverse transcription primer
- 1099 (/5Phos/GATCGTCGGACTGTAGAACTCTGAACCTGTCGGTGGTCGCCGTATCATT/iS
- 1100 p18/CACTCA/iSp18/GCCTTGGCACCCGAGAATTCCA) (IDT) using the following
- 1101 conditions. RNA and RT primer were denatured and annealed in reverse transcription
- 1102 buffer (50 mM Tris-Cl pH 8.6, 60mM NaCl, 10mM DTT). After annealing, the reverse
- 1103 transcription reaction was split in equal halves for the restrictive and permissive
- $1104 \qquad \text{conditions. The restrictive reaction received a final concentration of 2.5 mM dNTPs, 6 mM \\$
- 1105 MgCl<sub>2</sub> and 7.1% DMSO, whereas the permissive reaction received a final concentration
- 1106 of 4mM dNTPs, 20mM MgCl<sub>2</sub> and 7.1% DMSO. Reactions were carried out at 42°C for

- 1107 1h, prior to degradation of RNA with the addition of NaOH at 85°C. Truncated cDNAs
- 1108 were size-selected and purified on an 8% urea-TBE PAGE gel, followed by elution from
- gel slices in DNA elution buffer (300mM NaCl, 10mM Tris-HCl, pH 8.0) overnight at room
- 1110 temperature. cDNAs were circularized with circLigase (Epicentre) and amplified by PCR
- 1111 (Phusion; NEB) with the forward primer (AATGATAC GGCGACCACCGA) and a
- 1112 barcoded reverse primer (CAAGCAGAAGAC
- 1113 GGCATACGAGATXXXXXGTGACTGGAGTTCCTTGGCACCCGAGAATT CCA) (IDT).
- 1114 PCR products were gel-purified, precipitated and sequenced on an Illumina HiSeq 2000.
- 1115

# 1116 Ribo-seq

- 1117 Ribosome footprint profiling was performed essentially as described in Thompson et al.
- 1118 (2016), with the following minor modifications. Yeast cells were grown to an  $A_{600nm} \approx 1.0$
- 1119 or A<sub>600nm</sub>≈12.0, harvested by vacuum filtration (http://bartellab.wi.mit.edu/protocols.html.),
- 1120 lysed using a Cryomill and cycloheximide was added to the lysis buffer (0.1 mg/mL) prior
- 1121 to centrifugation. rRNA was subtracted from mRNA libraries as described (Brar et al.,
- 1122 2012).
- 1123

# 1124 QUANTIFICATION AND STATISTICAL ANALYSES

- 1125
- 1126 Most statistical tests were carried out in R, except for the codon enrichment analysis,
- 1127 where the tests were carried out using the scipy package in python. Plots were
- generated using matplotlib in python, or the ggplot2 package from the tidyverse in R.
- 1129

# 1130 eCLIP-seq data analysis

- 1131 Our analysis pipeline is based on the one described by the Yeo lab (Van Nostrand et al,
- 1132 2016). For full code, see <GitHub repo>. Reads were first demultiplexed by removing the

1133 first 10-nt of the forward read and adding them to the sequence name for later use: 1134 these 10-nt correspond to the random barcode added as part of the 5' linker. The 1135 adapter sequence was trimmed from raw reads using cutadapt (version 1.7.1), requiring 1136 a minimum trimmed read length of 18-nt. Reads were mapped using STAR (version 1137 2.5.1b) to a repeat-masked yeast genome. This genome was generated by masking 1138 repetitive regions and non-coding RNA loci (particularly tRNA genes and rRNA repeats) 1139 that had at least one identical sequence elsewhere in the genome. To ensure unique 1140 copies of these genes, we included a pseudo-chromosome that concatenated single 1141 copies of the masked non-coding RNA genes. We used the 10-nt barcodes to collapse 1142 PCR duplicates, merging any reads that mapped to the same position and shared the 1143 same barcode. Peaks were then called using the CLIPper algorithm. The pulldown and 1144 size-matched input libraries were processed separately in this way, and finally peaks 1145 from pulldown libraries were normalized to peaks in the input library using a 1146 normalization pipeline (Van Nostrand et al 2016).

1147

# 1148 **RNA-seq data analysis**

RNA-seq data was analyzed with in-house Bash and Python scripts. For a given MeTHSeq experiment, we used the reads from the permissive library to quantify gene
expression. Reads were trimmed using cutadapt, requiring at least 5 nucleotides of
overlap between the read and the adapter, and requiring trimmed reads to be at least
18-nt long. Trimmed reads were mapped to yeast mature mRNA sequences (obtained
from SGD) and quantified using kallisto (version 0.43.1), using options --single -I 40 -s
Differential expression was calculated using DESeq2 (version 1.14.1).

# 1157 Identification of Nm sites

1158 Sites of 2'-O-methylation were identified as previously described for pseudouridine 1159 (Stanley et al., 2016) with modifications. MeTH-seq signal was calculated as follows. For 1160 each position *i* in a 51 nt window centered at a given genomic position, the fraction of 1161 reads in the window whose 5' ends map to i was calculated. MeTH-seq signal is the 1162 difference in fractional reads between the restrictive and permissive libraries, multiplied 1163 by the window size. For an identified Nm, the reported MeTH-seq signal corresponds to 1164 the RT stop position 1 nt 3' of the site. All genomic positions with coverage of  $\geq 0.25$ 1165 reads per nt in the above described window within annotated features (downloaded from 1166 SGD on 9/2/2011) were considered, including 5' and 3'UTRs identified in (Pelechano et 1167 al., 2014). For Nm calling we required a reproducible peak height  $\geq$  2.0 with 1168 reproducibility in at least 13 of 16 log phase libraries and at least 16 of 20 high density 1169 libraries. A window size of 51 nt was examined in all libraries and only windows that 1170 surpassed the read cutoff of 0.25 reads per nt were considered. 1171

1172 **Methyltransferase assignment** 

1173 MeTH-seg peak heights were compared between methyltransferase deletion strains and 1174 all libraries from a given growth condition from cells wild type for a given 1175 methyltransferase. An Nm site was called genetically dependent on a specific Trm 1176 enzyme if the peak heights in both  $trm\Delta$  replicates were 1.5 standard deviations below 1177 the mean across all libraries. This threshold was set based on the observed reductions in peak heights at positive control tRNA<sup>ser</sup> Um44 sites in *trm44*<sup>Δ</sup> libraries. Spb1 target 1178 1179 sites were called at a more stringent threshold requiring peak heights at least 2.0 1180 standard deviations below the mean in both Spb1-depleted libraries and coverage of  $\geq$ 1181 0.25 reads per nt in the 51nt window surrounding the Nm site in both Spb1-depleted 1182 replicates.

# 1184 MeTH-seq signal plots and ROC curves

1185	MeTH-seq signal plots show the average of the MeTH-seq signal over the 51 nt window
1186	(described above) for all Nm included in the analysis. To generate receiver operating
1187	characteristic (ROC) curves for a given permissive/restrictive library pair, MeTH-seq
1188	signal was calculated for each position within the rRNA. A range of 5000 equally spaced
1189	cutoff scores were chosen spanning the range of observed MeTH-seq signal values. At
1190	each cutoff score, the true positive and false positive rates were calculated, and plotted.
1191	
1192	SnoRNA target analysis
1192 1193	SnoRNA target analysis In this analysis, we did not attempt to predict potential target sites across the
1193	In this analysis, we did not attempt to predict potential target sites across the
1193 1194	In this analysis, we did not attempt to predict potential target sites across the transcriptome; instead, we aimed to identify which of the putative sites Nm sites were
1193 1194 1195	In this analysis, we did not attempt to predict potential target sites across the transcriptome; instead, we aimed to identify which of the putative sites Nm sites were consistent with modification by snoRNAs. To do this, we generated a fasta file of the 30-

1199 queries and the options -task blastn-short -strand minus. We filtered the list of hits based on

1200 the mechanism of C/D box targeting: we required that the aligment include the putative

1201 Nm position, and that this position were 5 or 6 nt downstream of the position opposite a

1202 D-box.

1203

# 1204 **Codon enrichment analysis**

1205 We calculate background frequencies for each codon in each of the N wild-type libraries

used for peak calling at OD1. To calculate the frequency of a codon *i* in a given library,

1207 we weigh the expression (RPKM) of each gene by the number of times it contains codon

*i*, then sum across these and normalize by the total.

1209  $codon.expression_i = \Sigma_i expression_i \times count_{i,i}$ 

1210 
$$codon.frequency_i = codon.expression_i / \Sigma_i codon.expression_i$$

1211

1212

- 1213 We compare the frequency of each codon among Spb1 target sites to its background
- 1214 distribution using the SciPy implementation of the Student's T test.
- 1215 We calculate the reported fold-change as the ratio of the codon frequency in Spb1 sites
- 1216 and the average frequency in the background distribution.
- 1217

# 1218 Motif detection

- 1219 MEME (version 4.12.0) was used to find enriched motifs among Spb1-dependent mRNA
- 1220 sites. Detection was carried out on the 50-nt sequences surrounding each of the Spb1-
- 1221 dependent sites, calculating motif enrichment relative to a 2-order background model
- 1222 that summarized trinucleotide frequencies in the set of target sequences. The motif
- 1223 search required that any identified motifs be at least 3-nt and at most 15-nt long. The
- 1224 background model was generated using the fasta-get-markov script that is included with
- 1225 the MEME package, using the following options: -m 2 -rna -norc.
- 1226

## 1227 **Pairing probability distribution**

1228 Pairing probabilities of 20-nt regions surrounding Spb1-dependent 2'OMe sites were

1229 compared to a background set consisting of randomly selected sites from genes that

1230 contained at least one Spb1-dependent site. For each sequence, RNAfold (ViennaRNA

- 1231 package version 2.3.3) was used to obtain a pairing probability matrix. We then summed
- 1232 across the columns of the matrix to calculate the probability that each position is
- 1233 involved in base-pairing. To obtain a background distribution of average pairing
- 1234 probabilities, the following was done for each of 100 iterations: randomly chose 391 sites

- 1235 from the background sets (such that each background set is as large as the target set),
- 1236 then folded and obtained pairing probabilities as described above, and finally calculated
- 1237 average pairing probability per position in this set and stored the result.
- 1238
- 1239 Structure context of Spb1 targets
- 1240 For each 20-nt sequence surrounding either a target site or a background site, we used
- 1241 RNAsubopt (ViennaRNA package version 2.3.3) to obtain an ensemble of 100
- 1242 suboptimal structures. These structures were labeled as follows: S for base-paired
- 1243 positions, B for unpaired positions within stems, L for positions in the loop of a hairpin
- 1244 loop, and F for unpaired positions at the ends of the sequence. For each position in the
- 1245 sequence, we calculated the frequency with which it assumes an S, F, L or B and used
- these as approximations of the probability that the position assumes each of these
- 1247 conformations.
- 1248

# 1249 GO Analyses

- 1250 We used the statistical overrepresentation test in PANTHER (v13.0) to identify GO terms
- 1251 enriched in Spb1 target genes. As a background list of genes, we used all genes
- 1252 containing at least one site that met the MeTH-Seq read coverage threshold, regardless
- 1253 of peak height.
- 1254

# 1255 DATA AND SOFTWARE AVAILABILITY

- 1256 eCLIP-Seq and MeTH-Seq data may be downloaded from the Gene Expression
- 1257 Omnibus database, under accession numbers GSE109927. Analysis pipelines are
- 1258 available in <GitHub repository>.
- 1259

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## 1260 **FIGURE LEGENDS**

- 1261 Figure 1
- 1262 tRNA 2'-O-methyltransferases Associate with mRNAs In Vivo
- 1263 (A) Schematic of eCLIP procedure on an HPM-tagged methyltransferase (MTase)
- 1264 (B) Overlap between Trm44 non-coding RNA eCLIP targets and methylation substrates.
- 1265 The fraction of the library mapped to each feature is shown for two eCLIP replicates and
- 1266 SMI control. Genes are ordered by their abundance in SMI. tRNA<sup>Ser</sup> that are known to be
- 1267 modified by Trm44 are highlighted in blue. tRNA<sup>XXX</sup> (\*) is a computationally predicted
- 1268 tRNA<sup>Ser</sup> of unknown modification status (Lowe and Eddy, 1997).
- 1269 (C) Distribution of significantly enriched eCLIP peaks for Trm44. Top: ≥ 4-fold versus
- 1270 SMI. Bottom:  $p \le 0.001$  and  $\ge 4$ -fold-enriched versus SMI. Numbers for the bottom pie
- 1271 chart indicated in parentheses.
- 1272 (D-G) Examples of mRNA eCLIP peaks for Trm44 (D), Trm3 (E), Trm7 (F), and Trm13
- 1273 (G).
- (H) Number of significant eCLIP peaks in tRNA, mRNA, rRNA, and snoRNA for Trm3, 7,
- 1275 13 and 44;  $p \le 0.001$  and  $\ge 4$ -fold enriched versus SMI.
- (I) Venn diagram illustrates lack of overlap between mRNAs CLIPed to different Trmproteins.
- 1278

1279 Figure 2

- 1280 Illumina Sequencing-Based Detection of 2'-O-Methyl Ribose with Single-Nucleotide
- 1281 Resolution
- 1282
- 1283 (A) Schematic of MeTH-seq procedure. nt, nucleotides.
- (B) MeTH-seq reads mapping to a 235 nt region of RDN25-1 (chrXII: 452,800-453,035)
- 1285 containing 8 known Nm sites (black bars). Peaks of Nm-dependent reads are indicated
- 1286 with dashed red lines.
- 1287 (C) Meta plot of MeTH-seq signal for 55 known Nm sites in rRNA.
- 1288 (D) Receiver operating characteristic curve of MeTH-seq signal for all known Nm in
- 1289 rRNA.
- 1290
- 1291 Figure 3
- 1292 2'-O-Methylation of mRNAs
- 1293
- 1294 (A) Distribution of reproducible MeTH-seq peaks between RNA types in exponentially
- 1295 growing yeast. Shown are sites with a peak height  $\geq$ 2.0 in at least 13 out of 16
- 1296 independent experiments.
- 1297 (B) Example MeTH-seq peaks in mRNA. Top: genome browser views of reads from
- restrictive (rest) and permissive (perm) RT conditions. Bottom: MeTH-seq signal plots.

- 1299 (C) Overlap between MeTH-seq peaks identified in exponential (OD1) and post-diauxic
- 1300 (OD12) growth. OD12 sites had peak heights ≥2.0 in at least 16 out of 20 independent
- 1301 experiments.
- 1302 (D) MeTH-seq signal plots of TRM-dependent Nm sites.
- 1303 (E) Summary of Nm sites identified by MeTH-seq as TRM-dependent.
- 1304
- 1305 Figure 4
- 1306 mRNA Methylation by Spb1 Is Widespread
- 1307
- 1308 (A) Predicted base pairing between C/D snoRNAs and Nm sites in mRNAs is
- 1309 comparable to canonical snoRNA target sites in rRNA.
- 1310 (B) MeTH-seq reads at putative snoRNA-dependent Nm sites.
- 1311 (C) Depletion of Spb1 by promoter shutoff. Top: The essential SPB1 gene was placed
- 1312 under control of a repressible GAL1promoter (GALpSPB1) and grown in glucose to
- 1313 inhibit Spb1 expression. Bottom: Tagged Spb1 protein was barely detectable by Western
- 1314 blot after 2 hours. Asc1 loading control was detected as described (Coyle et al., 2009).
- 1315 (D) MeTH-seq signal plots of *SPB1*-dependent Nm sites.
- 1316 (E) Distribution of *SPB1*-dependent Nm sites between RNA types.
- 1317 (F) Distribution of *SPB1*-dependent Nm sites between nucleotides.

1318 (G) Methyltransferase dependence of Nm sites in mRNA.

1319

1320 Figure 5

1321 Features of mRNA Target Sites of Spb1

1322

1323 (A) SPB1-dependent Nm sites are enriched in CDS (\*\*\*p < 8.1e-9) and depleted from 3'

1324 UTRs (\**p* < 1.4e-6). Bonferroni corrected *p*-values from Fisher's exact test.

1325 (B) Distribution of SPB1-dependent Nm sites with respect to transcription start sites,

1326 translation start and stop sites. Numbers indicate distance in nucleotides from landmark.

1327 (C-D) Amino acid and codon frequencies among SPB1-dependent Nm sites compared

to the background of all codons with adequate read coverage for MeTH-seq analysis

1329 (STAR Methods). (C) Amino acids A, I, L, M and V are significantly overrepresented. (D)

1330 Codon table colors reflect the fold increase or decrease in Nm site frequency compared

to background. Codons that did not contain any identified sites are indicated in grey. \* *p* 

1332 < 0.01, \*\*\* *p* < 0.0001, Student's T test.

1333 (E) Bias in codon positions methylated by Spb1 (Chi Square p<2.2e-16).

1334 (F) Web logo of top motif enriched in Spb1 target sites. E value = 1.7e-40 (STAR
1335 Methods).

1336 (G) Structural context of Spb1 sites. Each panel plots the probability that a position in

1337 Spb1 targets (orange) or the background set (grey) assumes a loop, a stem, or an

- 1338 otherwise unpaired position. Top and bottom edges of the ribbons indicate 75<sup>th</sup> and 25<sup>th</sup>
- 1339 percentiles respectively. *p*-values for each position (KS-test) are indicated in the
- 1340 structure diagrams below—left: loops, right: stems.

1341

- 1342 Figure 6
- 1343 Spb1 Methylates and Maintains Normal Levels of mRNAs Necessary for Ribosome
- 1344 Biogenesis

1345

- 1346 (A) Log10 *p*-values for GO term enrichments among Spb1 target genes. Displayed
- 1347 categories were enriched ≥4-fold compared to the background of all genes with
- adequate read coverage for MeTH-seq analysis in Spb1-depleted cells. GO enrichment
- 1349 *p*-values were obtained using PANTHER v13.0.
- (B) Volcano plot of RNA-seq data shows reduced levels of Spb1 target mRNAs in Spb1-depleted cells.
- 1352 (C) mRNAs with multiple SPB1-dependent Nm sites show greater reductions in Spb1-
- 1353 depleted cells. KS test \* p<0.0166, \*\*\* p<2.2e-16.
- 1354 (D) Spb1 target mRNAs encoding ribosomal proteins showed the largest reductions. KS
- test p < 4.31e-08 For RNA-seq analysis, n=2 for Spb1-depleted cells and n=6 for wild type control cells.
- 1357

1358 **TABLES** 

- 1359 Table S1: Yeast strains used in this study.
- 1360 Table S2: Sequencing read summary for eCLIP libraries.
- 1361 Table S3: Trm44 eCLIP clusters: positions, enrichments and *p*-values.
- 1362 Table S4: Pus1 eCLIP clusters: positions, enrichments and *p*-values.
- 1363 Table S5: Trm3 eCLIP clusters: positions, enrichments and *p*-values.
- 1364 Table S6: Trm7 eCLIP clusters: positions, enrichments and *p*-values.
- 1365 Table S7: Trm13 eCLIP clusters: positions, enrichments and *p*-values.
- 1366 Table S8: MeTH-seq peaks in rRNA, peak height  $\geq$  4.0.
- 1367 Table S9: All MeTH-seq peaks  $\geq$  2.0 in log phase (OD1).
- 1368 Table S10: MeTH-seq peaks in rRNA,  $4.0 \ge$  peak height  $\ge 2.0$ .
- 1369 Table S11: All MeTH-seq peaks  $\geq$  2.0 in post-diauxic phase (OD12).
- 1370 Table S12: TRM-dependent Nm sites (OD1 and OD12).
- 1371 Table S13: Summary of predicted snoRNA-dependent Nm sites in mRNA.
- 1372 Table S14: Spb1 eCLIP clusters: positions, enrichments and *p*-values.
- 1373 Table S15: SPB1-dependent Nm sites in log phase (OD1).
- 1374 Table S16: SPB1-dependent Nm sites in post-diauxic phase (OD12).
- 1375 Table S17: GO term summary: Spb1 target mRNAs.
- 1376

# 1377 SUPPLEMENTAL FIGURE LEGENDS

- 1378
- 1379 Figure S1
- 1380 eCLIP Analysis of tRNA 2'-O-Methyltransferases, Related to Figure 1
- 1381

1382 (A) Summary of canonical non-coding RNA targets of yeast 2'-O-methyltransferases.

- (B) 32P labeled RNA immunoprecipitated with anti-Myc from cross-linked Trm44-HPMand untagged control.
- 1385 (C) Read coverage in Pus1 and Trm44 eCLIP libraries for significantly enriched Trm44
- 1386 mRNA eCLIP peaks ( $p \le 0.001$  and  $\ge 4$ -fold enriched versus SMI).
- 1387 (D) Anti-Myc immunoprecipitates from cross-linked Trm3-HPM, Trm7-HPM, and Trm13-
- 1388 HPM. Left: 32P labeled co-immunoprecipitated RNA. Right: Western blots. \* background
- 1389 band.
- 1390 (E-G) Overlap between non-coding RNA eCLIP targets and canonical methylation
- 1391 substrates of Trm3 (E), Trm7 (F) and Trm13 (G). The fraction of the library mapped to
- each feature is shown for two eCLIP replicates and SMI control. Genes are ordered by
- their abundance in SMI. tRNAs that are known to be modified are highlighted in blue.

1394

1395 Figure S2

1396 Controls for Transcriptome-wide Mapping of 2'-O-Methyl Ribose with MeTH-seq, Related1397 to Figure 2

- 1399 (A) Left: Meta plot of MeTH-seq signal at rRNA Nm sites detected by primer extension
- 1400 with limiting dNTPs (purple line) or limiting Mg<sup>2+</sup> (green line). AMV was the reverse
- 1401 transcriptase used. Right: Receiver operating characteristic curves for MeTH-seq signal
- 1402 at all known Nm in rRNA obtained using different reaction conditions.

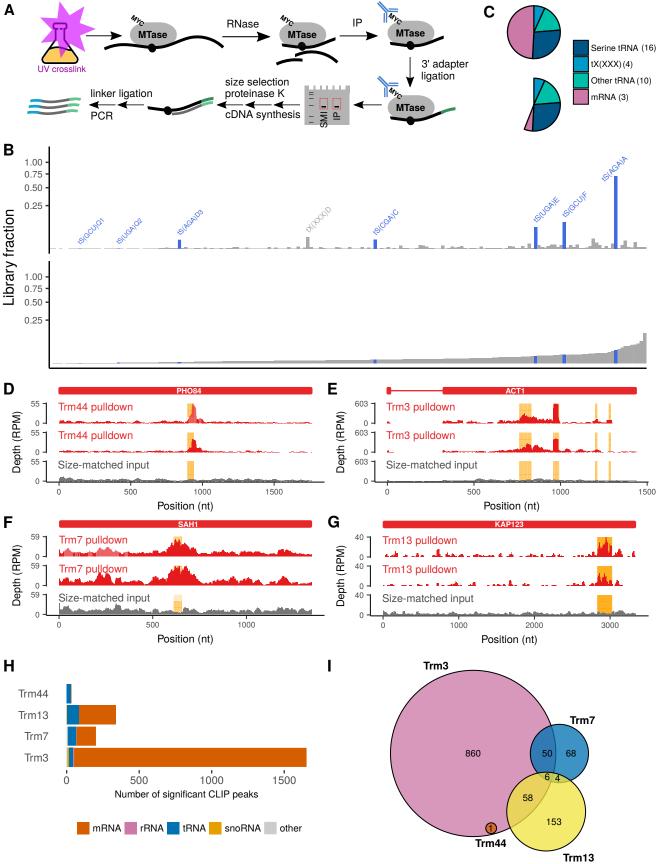
- 1403 (B) Left: Meta plot of MeTH-seq signal at rRNA Nm sites detected by primer extension
- 1404 using AMV (purple line) or SSIII (green line) reverse transcriptases. In both reactions,
- 1405 magnesium was limiting. Right: Receiver operating characteristic curves for MeTH-seq
- signal at all known Nm in rRNA obtained using different reverse transcriptases.
- 1407 (C) Deletion of C/D snoRNAs eliminated MeTH-seq peaks at their corresponding rRNA
- target sites.
- 1409 (D) Apparent Nm site misannotations identified by MeTH-seq.
- 1410 (E) Examples of false positive MeTH-seq peaks in rRNA.
- 1411 (F) Reproducibility of MeTH-seq peak heights at known Nm in rRNA.
- 1412 (G) Sequence bias in Nm site signal from MeTH-seq.
- 1413
- 1414 Figure S3
- 1415 Conditions of Regulated mRNA 2'-O-Methylation, Related to Figure 3
- 1416
- 1417 (A) Yeast growth curve. Cells were harvested for MeTH-seq analysis in mid exponential
- growth (OD1) and when cultures reached OD12, after the diauxic shift and before
- 1419 stationary phase.
- 1420 (B) Ribosome footprint profiling reveals widespread differences in gene expression
- 1421 between log phase (OD1) and post-diauxic growth (OD12). Ribosome profiling results
- 1422 were reproducible,  $R^2=0.-0.$ . Rpkm, reads per kilobase per million mapped reads.

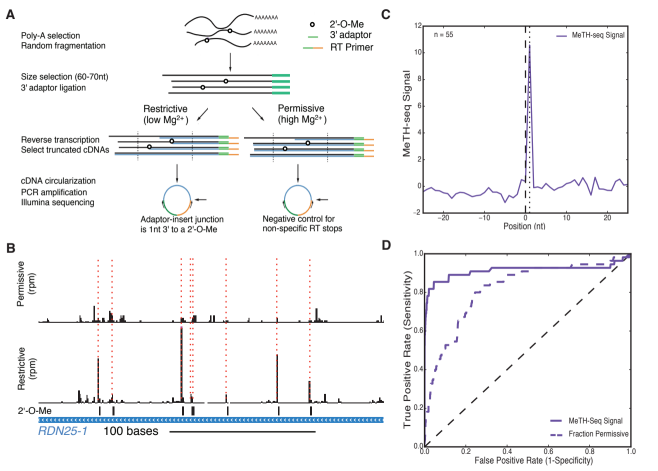
- 1423 (C) RNA-seq data showing that expression levels minimally affect identification of
- 1424 regulated Nm sites in mRNAs. Average log-transformed expression levels (TPM) from
- 1425 two growth conditions for all mRNAs in which MeTH-seq identified Nm sites in both log
- 1426 phase and post-diauxic growth (red), log phase only (blue), and post-diauxic only
- 1427 (green). n=6 replicates for each condition.
- 1428
- 1429 Figure S4
- 1430 eCLIP Analysis of Spb1 and Effects of Spb1 Depletion on rRNA 2'-O-Methylation at
- 1431 snoRNA-Dependent Sites, Related to Figure 4
- 1432
- 1433 (A) Distribution of significantly enriched eCLIP peaks for Spb1,  $p \le 0.001$  and  $\ge 4$ -fold
- 1434 enriched versus SMI.
- 1435 (B-D) MeTH-seq signal plots of rRNA Nm sites with reproducible slight reductions of
- signal in SPB1-depleted cells. These sites are methylated by Nop1 guided by snR48 (B),
- 1437 snR62 (C), and snR48 (D).
- 1438
- 1439 Figure S5
- 1440 Codon and Motif Features of mRNA Target Sites of Spb1, Related to Figure 5

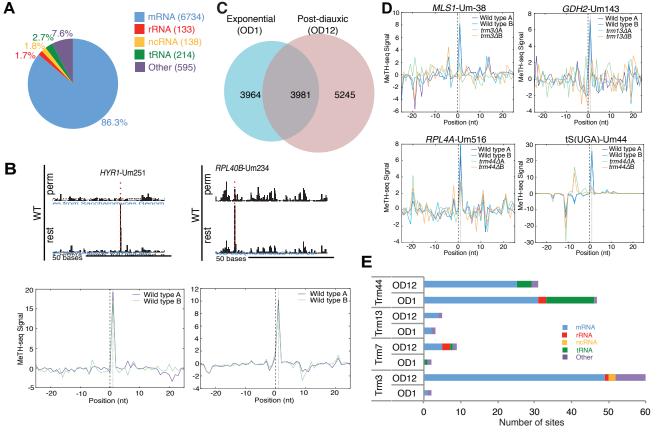
- 1442 (A) Frequency of codons among *SPB1*-dependent Nm sites compared to the
- background of all codons with adequate read coverage for MeTH-seq analysis (STARMethods).
- 1445 (B-D) Ribosome footprint distributions. Ribosomal A sites aligned to Nm sites from wild
- 1446 type cells (B), *dom34* mutants defective in No-Go Decay (C), and His codons in cells
- 1447 treated with 3-AT (D). Ribosome profiling data from (Guydosh and Green, 2014).
- 1448 (E) Web logos of significantly enriched motifs within a 50-nucleotide window surrounding
- novel *SPB1*-dependent Nm sites. The first motif (top) was found in 71/409 mRNA sites.
- 1450 The second was found exclusively in pre-tRNAs and the third was found in 15 mRNA
- 1451 sites.
- 1452 (F) Relative distances between *SPB1*-dependent Nm sites and the center of UGNUGN
- 1453 motifs (as called by MEME).
- (G) Average pairing probability in the 20-nt surrounding Spb1-dependent Nm sites(orange) or a background set of sites (grey).
- 1456
- 1457 Figure S6
- 1458 Global Effects of Spb1 Depletion on mRNA Abundance, Related to Figure 6

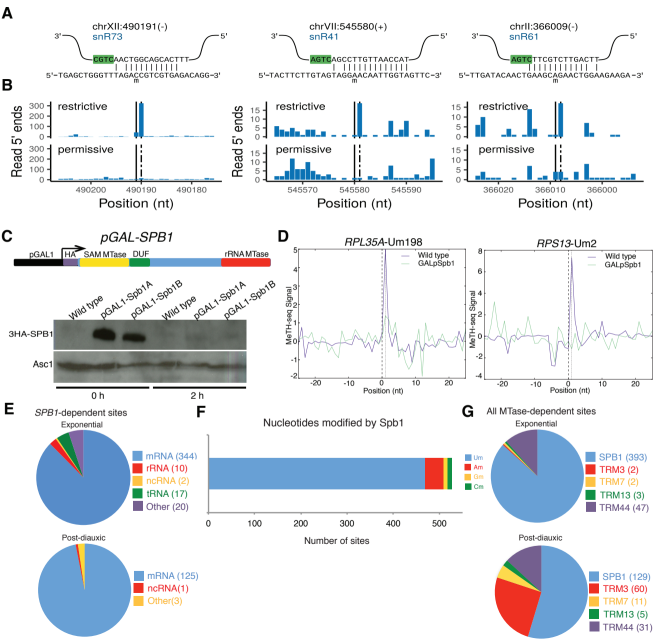
- 1460 (A) Cumulative distributions of mRNA levels. Spb1 target genes were highly expressed
- 1461 compared to the background of all genes with adequate read coverage for MeTH-seq

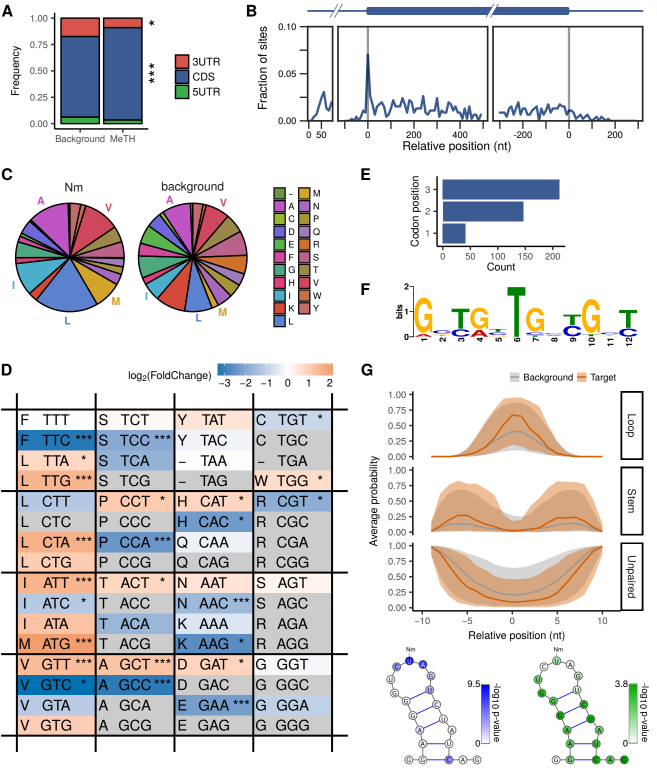
- analysis in wild type controls (n=6) and showed reduced levels in Spb1-depleted cells
- 1463 (n=2). TPM, transcripts per million. KS test *p*-values are indicated.
- 1464 (B) Volcano plot of RNA-seq data shows reduced levels of ribosomal protein mRNAs in
- 1465 Spb1-depleted cells.
- 1466 (C) Levels of ribosomal protein mRNAs that are Spb1 targets were significantly reduced
- 1467 compared to non-targets. KS test, p < 5.9e-15.

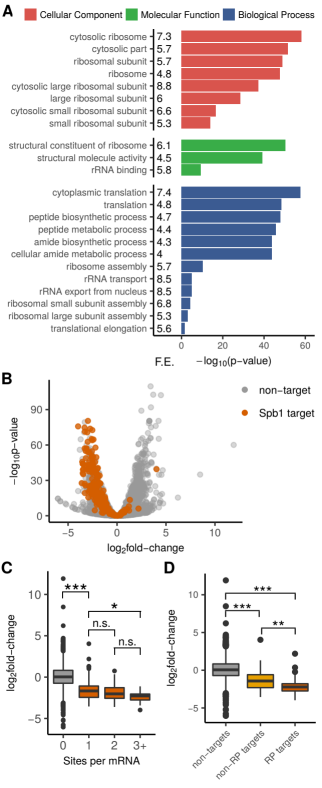


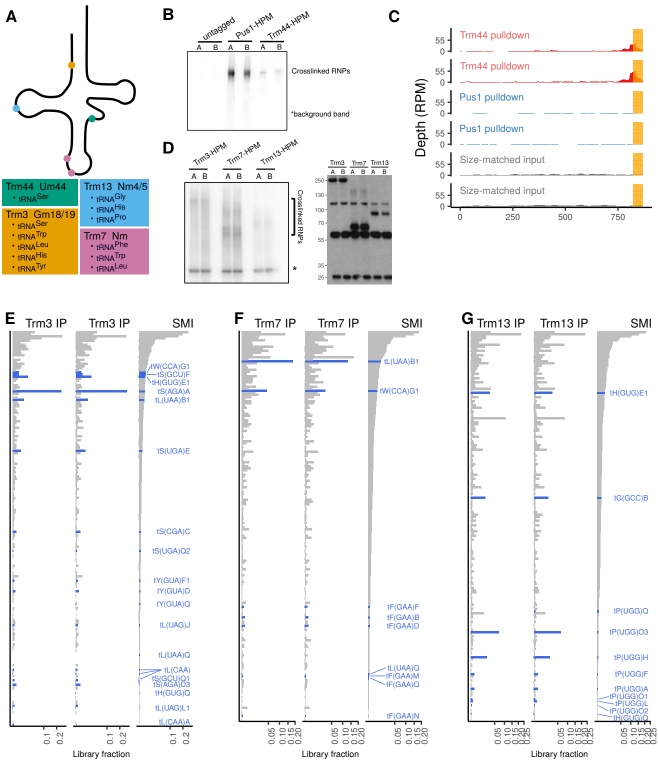


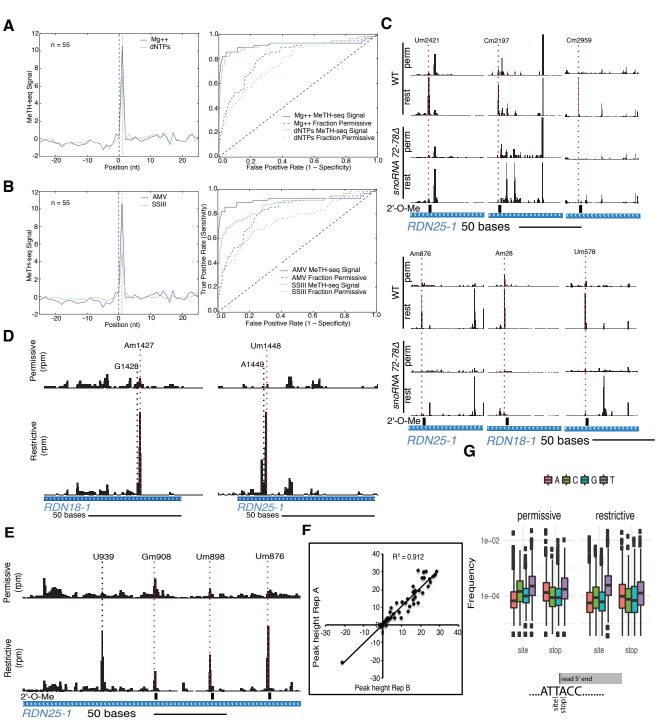


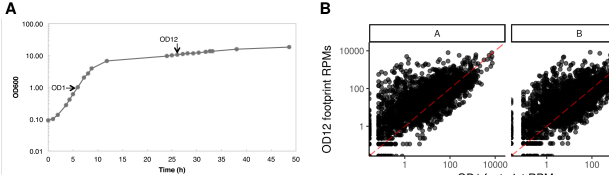








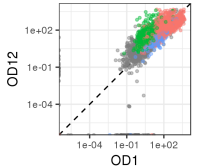


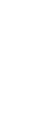


OD1 footprint RPMs

10000

С



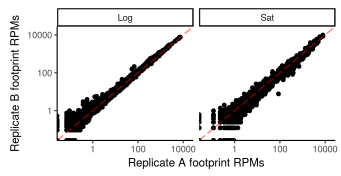


both

sat

log

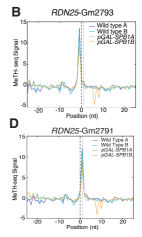
• NA

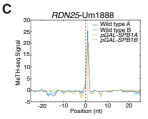


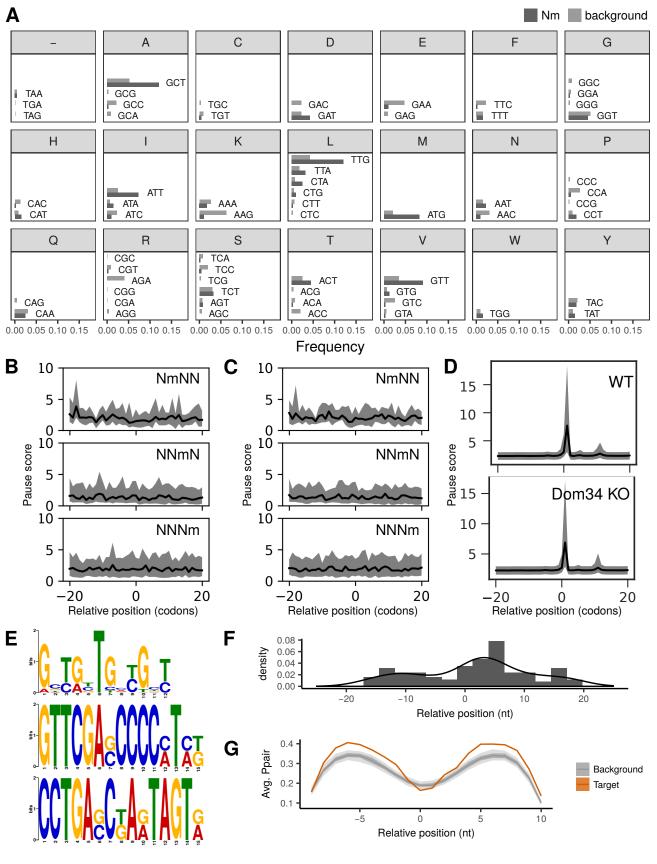
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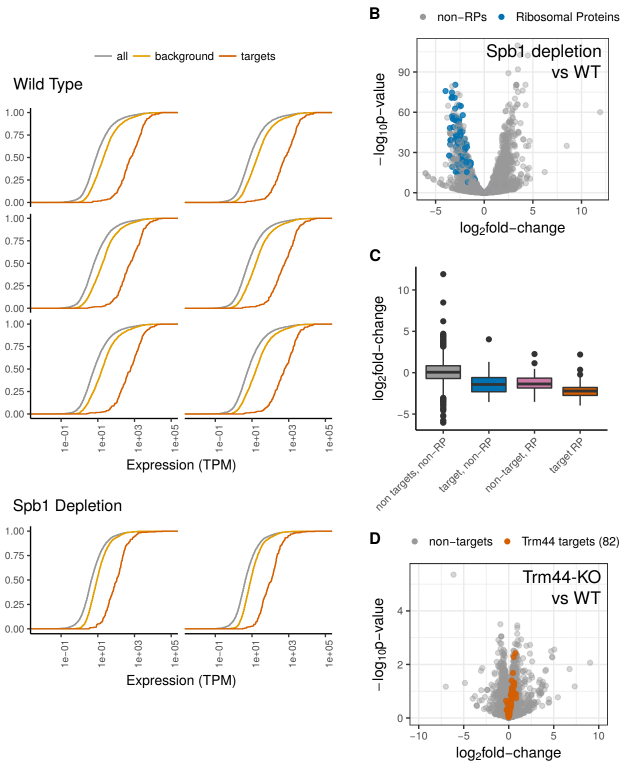


#### Spb1 CLIP targets rRNA (83) other (52) mRNA (108) snoRNA (116)









Cumulative frequency

Cumulative frequency