1	Protomer alignment modulates specificity of RNA substrate
2	recognition by Ire1
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## 23 Abstract

- 24 The unfolded protein response (UPR) maintains protein folding homeostasis in the endoplasmic
- 25 reticulum (ER). In metazoan cells, the Ire1 branch of the UPR initiates two functional outputs—
- 26 non-conventional mRNA splicing and selective mRNA decay (RIDD). By contrast, Ire1 orthologs
- 27 from Saccharomyces cerevisiae and Schizosaccharomyces pombe are specialized for only
- 28 splicing or RIDD, respectively. Previously, we showed that the functional specialization lies in
- 29 Ire1's RNase activity, which is either stringently splice-site specific or promiscuous (W. Li et al.,
- 30 2018). Here, we developed an assay that reports on Ire1's RNase promiscuity. We found that
- 31 conversion of two amino acids within the RNase domain of *S. cerevisiae* Ire1 to their *S. pombe*
- 32 counterparts rendered it promiscuous. Using biochemical assays and computational modeling,
- 33 we show that the mutations rewired a pair of salt bridges at Ire1 RNase domain's dimer
- 34 interface, changing its protomer alignment. Thus, Ire1 protomer alignment affects its substrates
- 35 specificity.

#### 36 Introduction

37 In eukaryotes, about one third of all proteins are folded in the endoplasmic reticulum 38 (ER). The protein folding homeostasis of the ER is monitored and tightly regulated by a 39 collective of signaling pathways, known as the unfolded protein response (UPR) (Hetz, Zhang, 40 & Kaufman, 2020; Walter & Ron, 2011). The most evolutionarily conserved branch of the UPR 41 is initiated by Ire1, an ER-transmembrane kinase/endoribonuclease (RNase). In response to 42 accumulated unfolded proteins in the ER, Ire1 forms oligomers (Aragon et al., 2009; Korennykh 43 et al., 2009; H. Li, Korennykh, Behrman, & Walter, 2010) and carries out two functional outputs. 44 First, Ire1 initiates non-conventional splicing of HAC1 (in S. cerevisiae) or XBP1 (in metazoans) 45 mRNA (Cox, Shamu, & Walter, 1993; Mori, Ma, Gething, & Sambrook, 1993; Sidrauski & Walter, 46 1997; Yoshida, Matsui, Yamamoto, Okada, & Mori, 2001). After cleavage by Ire1 and removal of 47 the intron, the severed exons are ligated by tRNA ligase (Jurkin et al., 2014; Kosmaczewski et 48 al., 2014; Lu, Liang, & Wang, 2014; Peschek, Acosta-Alvear, Mendez, & Walter, 2015; Peschek 49 & Walter, 2019; Sidrauski, Cox, & Walter, 1996). The spliced mRNAs are translated into Hac1 50 and Xbp1 proteins, both of which are transcription factors that induce the UPR gene expression 51 program in the nucleus (Calfon et al., 2002; Travers et al., 2000; Van Dalfsen et al., 2018; 52 Yoshida et al., 2001). Second, Ire1 selectively cleaves a set of mRNAs that encode ER-targeted 53 proteins. The cleaved mRNAs are subsequently degraded by the cellular RNA decay machinery 54 (Guydosh, Kimmig, Walter, & Green, 2017). As a result, this process, known as regulated Ire1-55 dependent mRNA decay (RIDD), restores homeostasis of the ER by reducing the protein folding 56 burden (Bae, Moore, Mella, Hayashi, & Hollien, 2019; Hollien et al., 2009; Hollien & Weissman, 57 2006; Kimmig et al., 2012; Moore & Hollien, 2015). 58 While metazoan Ire1 performs both functions, the Ire1 orthologs in Saccharomyces

59 cerevisiae and Schizosaccharomyces pombe are functionally specialized: S. cerevisiae Ire1

60 initiates splicing of HAC1 mRNA as its singular target in the cell (Niwa, Patil, DeRisi, & Walter,

61 2005), and S. pombe Ire1 exclusively performs RIDD (Guydosh et al., 2017; Kimmig et al.,

62 2012). Our previous study reported that the functional specialization of Ire1 is achieved through

63 diverged RNase specificities (W. Li et al., 2018). S. cerevisiae Ire1 has a stringent RNase,

restricting it to HAC1 mRNA. In contrast, S. pombe Ire1 has a promiscuous (i.e. broadly specific)

65 RNase, enabling cleavage of a wide range of mRNA RIDD targets. Which structural

66 determinants on Ire1 influence RNase specificity remained unknown. Here, we addressed this

67 question by mutagenesis-guided biochemical analyses and structural modeling.

- 69 Results
- 70

## 71 RNase activity of *S. pombe* Ire1 is toxic to bacterial cells

72 We recently purified and characterized recombinant S. cerevisiae (Sc) and S. pombe 73 (Sp) Ire1 kinase/RNase (KR) domains (W. Li et al., 2018). During the protein expression 74 process, we noticed that the presence of plasmids bearing the genes encoding Sc and Sp Ire1-75 KR under the control of the T7 promoter differently affected growth of the *E. coli* host cells. 76 Growth curves revealed that *E. coli* cells bearing a plasmid containing the *Sp IRE1-KR* barely 77 grew within the monitored 5 h time window, even in the absence of the isopropyl  $\beta$ -d-1-78 thiogalactopyranoside (IPTG) inducer (Figure 1A, blue filled triangles). By contrast, *E. coli* cells 79 bearing plasmids containing Sc IRE1-KR grew normally with a growth rate comparable to that of 80 a control strain bearing an empty plasmid. Because the T7 promoter is known to exhibit 81 background expression even in the absence of IPTG (Rosano & Ceccarelli, 2014), we reasoned 82 that the observed gene toxicity of Sp IRE1-KR might result from the enzyme's promiscuous 83 RNase activity, which may degrade endogenous *E. coli* RNAs required for viability. By contrast, 84 Sc Ire1-KR might be tolerated, because of its exquisite substrate specificity for Sc HAC1 mRNA 85 splice junctions (Niwa et al., 2005). To test this notion, we used the Ire1 RNase inhibitor 4µ8C 86 (Cross et al., 2012). As expected,  $4\mu$ 8C inhibited cleavage of a 21 base-pair stem-loop 87 substrate derived from the 3' splice junction of XBP1 mRNA for both Sp and Sc Ire1-KR (Figure 88 1B and C, Figure 1-figure supplement 1 A-C), as well as RIDD activity in Sp cells (Figure 1-89 figure supplement 1D). Importantly, when added to the cultures of cells bearing plasmids 90 encoding Sp Ire1-KR,  $4\mu$ 8C restored normal growth (Figure 1A). In further agreement with the 91 notion that Sp RNase activity was the culprit of reduced E. coli growth, an RNase-dead mutant 92 of Sp Ire1-KR harboring the H1018A mutation (Kimmig et al., 2012) was not toxic (Figure 1A). 93 These results suggest that *E. coli* growth was inhibited by the endonuclease activity, rather than 94 by indirect effects, such as protein misfolding or aggregation. Moreover, we established that 95 bacterial growth can be exploited to assess substrate specificity of Ire1's RNase. 96 97 Ire1's RNase domain confers promiscuous RNase activity

We reasoned that this assay might allow us to glean insights into Ire1's substrate specificity. Using structure-guided sequence comparison of Ire1 RNase domains, we picked a total of seventeen residues whose common features include that they are i) part of an oligomerization interface or located within 18 Å from the helix-loop element (HLE), which contains a positively charged loop (N1036 to K1042 on *Sc* Ire1) that engages the RNA 103 substrates (Korennykh et al., 2009; Korennykh et al., 2011; Lee et al., 2008), and ii) divergent 104 amino acids between Sc and Sp but conserved within the Schizosaccharomyces genus. Among 105 the seventeen amino acids, seven are located near the HLE, five are located at the RNase-106 RNase interface within the back-to-back dimer (previously defined as interface IF1<sup>C</sup> (Korennykh 107 et al., 2009)), and five are located at the RNase-RNase interface in the active Ire1 oligomer 108 (previously defined as interface IF2<sup>C</sup> (Korennykh et al., 2009)) (Figure 2A & B, Figure 2 – source 109 data 1).

110 To examine possible effects of these residues on RNase specificity, we cloned and 111 purified an Sc Ire1-KR mutant with all 17 residues replaced by their S. pombe counterparts (Sc 112 Ire1-KR-mut17). We tested its RNase activity using four previously characterized stem-loop 113 RNA substrates derived from the Sc HAC1 mRNA 3' splice site that is exclusively cleaved by Sc 114 Ire1, as well as the Ire1 cleavage sites of the Sp BIP1, PLB1 and SPAC4G9.15 mRNAs that are 115 exclusively cleaved by Sp Ire1 (W. Li et al., 2018). We chose the three Sp RNA substrates 116 because their Ire1 cleavage sites vary in predicted loop sizes (9-, 7- and 3-membered loops, 117 respectively). Cleavage activity towards stem-loop RNAs with variable loop sizes is one of the 118 characteristic features of Ire1 promiscuity (W. Li et al., 2018). In agreement with previous results, 119 wildtype (WT) Sc Ire1-KR cleaved the HAC1 mRNA 3' splice site but none of the Sp stem-loop 120 RNA substrates. Remarkably, Sc Ire1-KR-mut17 efficiently and specifically cleaved all four 121 stem-loop RNA substrates (Figure 2 C-F) with comparable kinetics (Figure 2G & Figure 2-figure 122 supplement 1). These results show that introducing the 17 mutations made Sc Ire1 more 123 "pombe-like" regarding its acceptance of variable-loop RIDD substrates. When tested in the E. 124 coli growth assay, the expression of Ire1-KR-mut17 proved toxic and  $4\mu$ 8C alleviated the toxicity 125 (Figure 2H), confirming the notion that toxicity results from the enzyme's broadened substrate range.

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#### 128 Residues at Ire1's dimer interface confer RNase promiscuity

129 The above results demonstrate that bacterial growth can be a useful readout for Ire1's 130 RNase promiscuity. Hence, we used this assay to identify the residue(s) that were causal in 131 conferring the broadened substrate specificity of Sc Ire1-KR-mut17. To this end, we created 132 single revertants, each with one of the seventeen mutations was converted back to the original 133 amino acid in Sc (Figure 3-source data 1). We expected that when we revert a mutation that 134 contributes to the enzyme's broadened substrate specificity, the revertant would be stringent 135 and less toxic to *E. coli* cells. We found that 3 revertants (K992, H1044, and Y1059) showed 136 markedly reduced toxicity, whereas the other 14 revertants remained toxic (Figure 3A). Next, we 137 cloned and purified a mutant protein, *Sc* Ire1-KR(K992D,H1044D,Y1059R), in which we

- 138 combined the three identified mutations. *Sc* Ire1-KR(K992D,H1044D,Y1059R) cleaved both *Sp*
- 139 stem-loop and *Sc HAC1* 3' splice-site substrates efficiently (Figure 3 B-E) and with similar rates
- 140 (Figure 3F & Figure 3-figure supplement 1). These results narrowed the list of candidate amino
- 141 acid changes that confer RNase promiscuity down to three.

142 Based on the locations of these three residues, we divided them into two groups. Sc Ire1 143 K992 and Y1059 (corresponding to D950 and R1016 of Sp Ire1) are located at Ire1's back-to-144 back dimer interface, while Sc Ire1 H1044 (corresponding to Sp Ire1 D1001) is located two 145 amino acids C-terminal of the HLE. We cloned and purified two Ire1 mutants, Sc Ire1-146 KR(K992D,Y1059R) and Sc Ire1-KR(H1044D). Recombinantly expressed and purified Sc Ire1-147 KR(K992D, Y1059R) cleaved both Sc and Sp stem-loop RNA substrates with efficiencies 148 comparable to those of Sc Ire1-KR(K992D,H1044D,Y1059R) (Figure 3B-F). By contrast, Sc 149 Ire1-KR(H1044D) cleaved the cognate Sc RNA substrate but none of the Sp RNA substrates 150 (Figure 3 B-F), suggesting that the two cerevisiae-to-pombe mutations at Ire1's RNase-RNase 151 dimer interface confer Ire1 RNase promiscuity. By contrast, H1044D appears a false positive, 152 likely isolated because the bacterial assay cannot distinguish stringent RNase from inactive 153 RNase as neither is toxic to the bacterial cells.

154

## 155 S. cerevisiae Ire1-KR(K992D,Y1059R) recognizes RNA substrates with reduced

#### 156 stringency

157 Sc Ire1 displays a strong preference for RNA substrates that contain a consensus 158 sequence within a stem-loop structure (Gonzalez, Sidrauski, Dorfler, & Walter, 1999; Hooks & 159 Griffiths-Jones, 2011; W. Li et al., 2018; Oikawa, Tokuda, Hosoda, & Iwawaki, 2010). We next 160 characterized the RNA motif recognized by Sc Ire1-KR(K992D,Y1059R) and compared it to 161 those recognized by WT Sc and Sp Ire1-KR. To this end, we examined Ire1 cleavage 162 efficiencies on a series of HAC1- and BIP1-derived mutant stem-loop RNAs, in which each loop 163 residue was individually changed into the three other possible ribonucleotides (Figure 4A & B). 164 Using the HAC1-derived mutant substrates, we showed that WT Sc Ire1-KR showed specificity 165 for the sequence motif CNGI(C/A)NGN, in close agreement with previous findings (Gonzalez et 166 al., 1999). By comparison, Sc Ire1-KR(K992D,Y1059R) recognized a less-stringent sequence 167 motif, CNGINNGN, in particular tolerating base substitutions in the +1 position (Figure 4A). 168 To assess the effects of RNA loop size variation on Ire1 cleavage efficiency, we 169 engineered mutations at positions -4 to break the base-pairing at the tip of the stem and enlarge

170 the loop from 7 to 9 nucleotides. The 9-membered stem loops were not cleaved by WT Sc Ire1-

KR, in line with previous study (Gonzalez et al., 1999). By contrast, the same RNAs were
cleaved by *Sc* Ire1-KR(K992D,Y1059R) (Figure 4A). The results suggest that the two interface
mutations in *Sc* Ire1-KR(K992D,Y1059R) render the enzyme more tolerant to both RNA
sequence and loop size variations.

175 The Ire1 cleavage site in Sp BIP1 mRNA contains a 9-nucleotide loop with a UGIC 176 cleavage site shifted by one nucleotide (Figure 4B). We confirmed that a stem-loop RNA 177 substrate containing this site was efficiently cleaved by Sp Ire1-KR but not by Sc Ire1-KR, in 178 agreement with our previous reports (Guydosh et al., 2017; Kimmig et al., 2012; W. Li et al., 179 2018). By contrast, Sc Ire1-KR(K992D,Y1059R) cleaved the motif efficiently with the additional 180 tolerance of any nucleotide at position -2, thus reducing the required sequence motif to only G 181 and C nucleotides flanking the cleavage site (Figure 4B). GIC is likewise present at the PLB1 182 and SPAC4G9.15 mRNA cleavage sites embedded in 7- and 3-membered loops, respectively, 183 which also proved to be substrates of Sc Ire1-KR(K992D,Y1059R) (Figure 3B-E). In further 184 support of the notion that Sp Ire1-KR and Sc Ire1-KR(K992D,Y1059R) are tolerant to loop size 185 variation, two of our stem-loop RNA substrates harboring mutations,  $U_{-5} \rightarrow A$  and  $U_{+4} \rightarrow A$ , 186 respectively, which are predicted to contract the 9-membered loop to a 7-membered one, were 187 efficiently cleaved by both enzymes (Figure 4B). Together, these data affirm the notion that Sp 188 Ire1-KR and Sc Ire1-KR(K992D,Y1059R) are promiscuous enzymes that recognize short RNA 189 sequence motifs and can accept a range of loop sizes.

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## 191 Salt-bridge rewiring at Ire1's dimer interface changes Ire1's protomer alignment

192 To understand how the interface mutations confer promiscuous RNase activity, we 193 explored structural differences between Sp Ire1, Sc Ire1 and Sc Ire1-KR(K992D,Y1059R) using 194 molecular modeling. Active Sc Ire1 oligomers are composed of multiple Ire1 back-to-back 195 dimers that stack in a helical arrangement (Korennykh et al., 2009). K992 and Y1059 are 196 located at the RNase-RNase interface of the back-to-back assembly of Ire1 protomers in PDB 197 3FBV (protomer A and B in Figure 5A). To build a structural model of Sc Ire1-198 KR(K992D,Y1059R), we introduced K992D and Y1059R onto the Sc Ire1 dimer structure and 199 performed energy minimization to optimize distances and resolve steric clashes. We followed 200 this calculation with molecular dynamics (MD) simulations, comparing Sc Ire1-201 KR(K992D,Y1059R) and WT Sc Ire1-KR. Analysis was performed from 10 ns to 20 ns, and the 202 simulation structures reached equilibrium within 10 ns (Figure 5-figure supplement 1). The 203 convergent structure model of Sc Ire1-KR(K992D,Y1059R) predicts a structural rearrangement

at Ire1's dimer interface. Specifically, whereas in the WT Sc Ire1-KR dimer residues E988 and

205 K992 of both protomers form a symmetric pair of salt bridges across the dimer interface (Figure 206 5B), these salt bridges are absent in the dimer of Sc Ire1-KR(K992D,Y1059R) due to the charge 207 reversal introduced by the K992D mutation. More interestingly, the model predicts the formation 208 of a new pair of salt bridges in Sc Ire1-KR(K992D,Y1059R), connecting the two newly 209 introduced amino acids D992 and R1059 across the protomer/protomer interface (Figure 5C). 210 The MD simulation predicts these new bonds as stable features (Figure 5-figure supplement 2). 211 Thus, molecular modeling suggests a structural rearrangement, resulting from the two interface 212 mutations in Sc Ire1, which, we propose, allows Sc Ire1 to assume the promiscuous "pombe-like" 213 state (Figure 5D).

214 The predicted new salt bridges in Sc Ire1-KR(K992D,Y1059R) are mediated by 215 guanidinium-carboxylate bidentate interactions, which are among the strongest non-covalent 216 interactions in proteins and are considerably stronger than the ammonium-carboxylate 217 interaction seen in the salt bridges in WT Sc Ire1-KR (Masunov & Lazaridis, 2003). Given that 218 the interaction at the dimer interface is predicted to be stronger in mutant Sc Ire1-219 KR(K992D, Y1059R) than in WT Sc Ire1-KR, the mutant enzyme should be more prone to form 220 dimers/oligomers than WT Sc Ire1-KR. Indeed, we confirmed this notion using protein 221 crosslinking *in vitro*, followed by SDS-PAGE, showing that WT Sc Ire1-KR was mostly 222 monomeric, while Sc Ire1-KR(K992D,Y1059R) formed mostly dimers and tetramers (Figure 5E, 223 compare lanes 2 and 3 with lanes 6 and 7). We further verified this result using sedimentation 224 equilibrium analytical ultracentrifugation (Figure 5-figure supplement 3). For Sc Ire1-225 KR(K992D,Y1059R), we calculated a dissociation constant ( $K_D$ ) of 0.98  $\mu$ M, which is about 60-226 fold smaller than that of WT Sc Ire1-KR ( $K_D = 57 \mu$ M). Based on these results, we consider it 227 likely that the estimated gain in free energy of the predicted new salt bridges results in higher 228 affinity within the back-to-back dimer, thus increasing the propensity of Ire1 to oligomerize.

229 Next, we experimentally tested the predicted salt bridges using mutagenesis. To this end, 230 we first engineered Sc Ire1-KR(K992D,Y1059A), introducing an alanine at position 1059. This 231 Ire1 mutant can neither form S. cerevisiae-like nor S. pombe-like salt bridges (Figure 5D). Thus, 232 as expected, Sc Ire1-KR(K992D,Y1059A) did not form dimers (Figure 5E, lanes 8 and 9) and 233 displayed ~100-fold reduced cleavage efficiency on HAC1-derived and an additional 100-fold 234 (i.e., overall ~10,000-fold) reduced cleavage rate on BIP1-derived RNA substrates (Figure 5-235 figure supplement 4). Thus, surprisingly, breaking the pombe-like salt bridge arrangement 236 restored Sc Ire1's ability to discriminate between substrate RNAs by ~100-fold (Figure 5F). 237 Breaking the predicted salt bridges on Sc Ire1-KR(K992D,Y1059R) by mutating 238 aspartate 992 to alanine also abolished Ire1's RNase activity (Sc Ire1-KR(K992A, Y1059R);

Figure 5-figure supplement 4), in this case reducing activity towards both *HAC1-* and *BIP1*derived substrate RNAs beyond our detection limit (>10,000-fold). Thus, we were not able to assess substrate specificity for this mutant.

242 Finally, we generated a charge-reversal mutant of Sc Ire1-KR(K992D,Y1059R) by 243 changing aspartate 992 to arginine and arginine 1059 to aspartate (Sc Ire1-KR(K992R,Y1059D). 244 We expected these two mutations to restore the salt bridges predicted for Sc Ire1-245 KR(K992D,Y1059R) but with reversed polarity. We found that, while the overall cleavage rate 246 remained 100-fold suppressed for both RNA substrates, Sc Ire1-KR(K992R,Y1059D) regained 247 activity towards the BIP1-derived substrate (Figure 5F, Figure 5-figure supplement 4) and 248 formed dimers (Figure 5E, lanes 10 and 11). Together, these experiments validate the predicted 249 salt bridges on Sc Ire1-KR(K992D,Y1059R) and further underscore the notion that salt bridge 250 rewiring to a pombe-like arrangement confers promiscuity to Ire1's RNase activity.

251 The importance of both types of salt bridges is further highlighted by a sequence 252 comparison of 230 Ire1 orthologs from yeast to human (see Figure 5G for a partial list of the Ire1 253 orthologs; a complete list is included in Figure 5-figure supplement 5 and its figure source file). 254 We compared residues at three positions corresponding to the Sc Ire1 E988, K992 and Y1059. 255 We found that 175 out of 230 of the Ire1 orthologs, including human Ire1 $\alpha$  and Ire1 $\beta$ , have the S. 256 pombe-like pattern, characterized by significant amino acid variation at position 988, a 257 negatively charged amino acid (aspartate or glutamate) at position 992, and a positively 258 charged amino acid (lysine or arginine) at position 1059. The apparent co-evolution of position 259 992 and 1059 further supports the existence of an inter-molecular salt bridge. Thirty-three out of 260 230 of the Ire1 orthologs have the S. cerevisiae-like pattern—with a negatively charged amino 261 acid (aspartate or glutamate) at position 988, a positively charged amino acid (lysine or arginine) 262 at position 992 and, in most cases, a tyrosine at position 1059. Co-evolution of position 988 and 263 992 supports their interaction at the dimer interface.

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### 265 Interface mutations change the protomer alignment in Ire1 dimer

To gain an appreciation of how the interface mutations affect Ire1's active RNase site in the dimer, we compared the structure of WT *Sc* Ire1-KR dimer with the predicted structure of *Sc* Ire1-KR(K992D,Y1059R) dimer after aligning the two KR dimers by one protomer (Figure 6A and B for front and bottom-up view, respectively). Interestingly, we observed a rocking motion between the RNase domains of the protomers (Figure 6B). Specifically, the salt bridge between D992 and R1059' in *Sc* Ire1-KR(K992D,Y1059R) reduced the distance between the two juxtaposed  $\alpha$ -helices from which their sidechains protrude ( $\alpha$ 1-helix: aa 983-998;  $\alpha$ 4-helix: aa 273 1048-1064, as named in (Lee et al., 2008)) (Figure 6C), while, concomitantly, the loss of the salt 274 bridges between E988 and K992 present in WT Sc Ire1-KR allows an increase in the distance 275 between  $\alpha$ 1-helices from which both of these amino acid side chains protrude (Figure 6D). We 276 used two metrics to quantify this change. First, we measured the distance between the centers 277 of mass of the two  $\alpha$ 1-helices throughout the 20 ns simulation trajectories, which was increased 278 by about 3 Å in Sc Ire1-KR(K992D,Y1059R) compared to WT Sc Ire1-KR. This measurement 279 reflects the changes in protomer alignment caused by the salt bridge rewiring. Second, we 280 measured the distance between the  $\alpha$  carbons of R1039 involved in RNA substrate binding and 281 H1061' involved in phosphodiester bond hydrolysis (Korennykh et al., 2011), which was 282 decreased by about 5 Å in Sc Ire1-KR(K992D,Y1059R) compared to WT Sc Ire1-KR (Figure 6E, 283 F). Therefore, the mutations that increase Ire1's RNase promiscuity are predicted to change 284 both the RNase-RNase interface and the relative alignment of important elements in the 285 catalytic site of the Ire1 dimer.

286 The kinase/RNase domain of Ire1 is homologous to the kinase homology (KH) and 287 kinase extension nuclease (KEN) domains of the Ribonuclease L (RNase L), which mediates 288 the antiviral and apoptotic effects of interferons in mammalian cells (Chakrabarti, Jha, & 289 Silverman, 2011). Upon activation, RNase L forms homodimers (Han et al., 2014; Huang et al., 290 2014) and cleaves mRNAs with a sequence motif of UNIN, where N can be any ribonucleotide 291 (Han et al., 2014). Since RNase L has promiscuous RNase activity, we wondered if its protomer 292 alignment would resemble that of the Sc Ire1-KR(K992D,Y1059R). To test this notion, we 293 compared the crystal structure of RNase L (PDB: 4OAV) with the structures of Sc Ire1-KR and 294 Sc Ire1-KR(K992D,Y1059R). At the RNase-RNase dimer interface of RNase L, N601' and E671 295 form a pair of intermolecular hydrogen bonds between RNase L's  $\alpha$ 1- and  $\alpha$ 4-helix (Figure 6-296 figure supplement 1A) (Han et al., 2014) (in RNase L the  $\alpha$ 1-helix spans as W589-V599 and the 297  $\alpha$ 4-helix aa V659-H672, as defined in (Huang et al., 2014)). Thus, RNase L and Sc Ire1-298 KR(K992D,Y1059R) share a conserved arrangement by which their  $\alpha$ 1- and  $\alpha$ 4-helix are 299 connected. By contrast, Sc Ire1-KR is different in that two  $\alpha$ 1-helices, one contributed by each 300 protomer, are connected instead. In further support of this notion, we analyzed RNase L using 301 the metrics defined in Figure 6E and found that the protomer alignment of RNase L closely 302 resembles that of Sc Ire1-KR(K992D,Y1059R) (Figure 6E, Figure 6-figure supplement 1A & B). 303 Thus, RNase L and Sc Ire1-KR(K992D,Y1059R), both of which have promiscuous RNase 304 activity, share a similar protomer alignment.

306 **Discussion** 

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308 From both an evolutionary and mechanistic angle, it has long been a puzzle how two 309 modalities of Ire1 function arose and are structurally implemented. At one extreme lies Ire1 from 310 S. cerevisiae, which is highly specific, precisely cleaving but a single mRNA in the cell (HAC1 311 mRNA) at its two splice junctions to excise the intron and initiate mRNA splicing. At the other 312 extreme lies Ire1 from S. pombe, which is highly promiscuous cleaving numerous mRNAs at 313 recognition sites that share but a three-nucleotide consensus in a variably sized loop. Cleavage 314 in this case initiates mRNA breakdown by RIDD. We previously showed by domain swapping 315 experiments that Ire1's cytosolic kinase/RNase domains determine whether Ire1 works in the 316 specific S. cerevisiae-like or relatively non-specific S. pombe-like modality. Using a growth 317 assay based on heterologous expression of Ire1 kinase/RNase domains in bacteria that reports 318 on Ire1's RNase promiscuity, we found that the S. cerevisiae Ire1's RNase specificity becomes 319 promiscuous when only two amino acids, K992 and Y1059, are replaced by aspartate and 320 arginine respectively, which are the corresponding amino acids in S. pombe Ire1. While these 321 replacements rendered Sc Ire1 more promiscuous, they did not entirely switch Sc Ire1's 322 substrate RNA profile to that of Sp Ire1: Sc Ire1-KR(K992D,Y1059R) retained activity towards 323 HAC1 mRNA-derived stem-loops, which is inert to cleavage by Sp Ire1 (W. Li et al., 2018). The 324 K992D and Y1059R mutations therefore rendered Sc Ire1 even more promiscuous than Sp Ire1.

325 MD simulations and biochemical assays revealed that the identified residues cause 326 rewiring of two symmetry-related inter-molecular salt-bridges at Ire1's RNase-RNase interface 327 within the back-to-back dimer. Sequence comparison of Ire1 orthologs showed that this rewiring 328 is conserved and, where known, correlates with reported functional outputs (Cheon et al., 2011; 329 Hollien et al., 2009; Hollien & Weissman, 2006; Miyazaki & Kohno, 2014; Miyazaki, Nakayama, 330 Nagayoshi, Kakeya, & Kohno, 2013). By this criterion, the vast majority of species is predicted 331 to have RIDD-enabled Ire1s as indicated by their S. pombe-like salt bridge pattern. Even though 332 human Ire1 $\alpha$  and Ire1 $\beta$  have different RNase specificity (Han et al., 2014; Imagawa, Hosoda, 333 Sasaka, Tsuru, & Kohno, 2008), both have a S. pombe-like interface pattern. This characteristic 334 is consistent with both human Ire1 isoforms being able to perform RIDD (Hollien et al., 2009; 335 Iwawaki et al., 2001). Of note, the ability of Sc Ire1-KR(K992D,Y1059R) to conduct both HAC1-336 specific and promiscuous cleavage resembles that of human lre1 $\alpha$  in its fully phosphorylated. 337 oligomeric state, while the more restricted activity of Sc Ire1 resembles that of dimeric human 338 Ire1 $\alpha$  (Le Thomas et al., 2021).

339 Surprisingly, our work identified the RNase-RNase interface, rather than regions 340 involved in substrate binding or catalysis, as a determinant for Ire1's RNase specificity. We 341 show evidence that conserved salt bridges determine the relative protomer alignment. In the 342 composite RNase active site of the back-to-back dimer, the relative distance of residues that 343 contribute to cleavage from both protomers is changed (Korennykh et al., 2011). The small 344 molecule, guercetin, which stabilizes S. cerevisiae Ire1's dimers/oligomers and increase its 345 RNase activity (Wiseman et al., 2010), binds to the same site where the two mutations identified 346 in this study are located. However, guercetin binding does not change Ire1 protomer alignment 347 and hence is not expected to alter Ire1's RNase specificity (Wiseman et al., 2010). Nevertheless, 348 modulation of the RNase selectivity by targeting the guercetin pocket is conceivable. In human 349 Ire1, crystal structures showed that the two RNase domains in the dimer are further apart in the 350 inactive state than in the active state (Joshi et al., 2015), and our data similarly indicate that 351 breaking the salt bridges in Sc Ire1-KR(K992D,Y1059R) by either changing the aspartate or the 352 arginine to alanine leads to profound reduction of activity. Related work demonstrates that the 353 more promiscuous RIDD modality of human Ire1 $\alpha$  requires phosphorylation-driven 354 oligomerization, which can be prevented by an oligomer-disrupting mutation at the RNase-355 RNase interface within the back-to-back dimers (Le Thomas et al., 2021). Thus together, the 356 data presented here demonstrate that Ire1 RNase domain's dimer interface is a dynamic site 357 through which both activity and substrate specificity can be regulated.

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- 367

## 368 **Competing Financial Interests**

369 The authors declare no competing financial interests.

## 370 Materials and Methods

371

## 372 Recombinant protein expression and purification

373 All the plasmids used in this study are listed in Table 1. The cytoplasmic portion of Sc or 374 Sp Ire1 containing its kinase and RNase domains (Ire1-KR) was expressed and purified from 375 BL21-CodonPlus (DE3)-RIPL Escherichia coli. We used an expression vector which fuses a 376 PreScission Protease cleavage site between the Ire1-KR and glutathione S-transferase (GST) 377 domains of the recombinant polypeptide and was regulated by a T7 promoter. The expression 378 cassette was transformed into E. coli cells. The WT Sc Ire1-KR was expressed as described 379 previously (Korennykh et al., 2009). For *E. coli* cells transformed with plasmids containing the 380 Sp Ire1-KR or Sc Ire1-KR mutant, all colonies on the transformation plate were collected 16 h 381 after transformation and mixed with 50 mL of LB medium. After 3 h incubation at 37°C, the 382 sample was diluted to 12 L of LB medium and further incubated at 37°C until optical density 383 reached 1. The incubation temperature was reduced to 25°C and protein expression was 384 induced by adding IPTG to a final concentration of 0.5 mM. After 4 h of growth at 25°C, the cells 385 were pelleted by centrifugation.

386 Cells were resuspended in GST binding buffer (50 mM Tris-HCl pH 7.5, 500 mM NaCl, 2 387 mM Mg(OAc)<sub>2</sub>, 2 mM DTT, 10% glycerol) and homogenized using high-pressure homogenizer 388 (EmulsiFlex). The cell lysate was applied to a GST-affinity column and eluted with GST elution 389 buffer (50 mM Tris-HCl pH 7.5, 200 mM NaCl, 2 mM Mg(OAc)<sub>2</sub>, 2 mM DTT, 10% glycerol, 10 390 mM reduced glutathione). The column elution was treated with GST-tagged HRV 3C protease 391 (PreScission Protease, GE Health) to cleave off the GST tag. At the same time, the sample was 392 dialyzed to remove glutathione in the elution buffer. After 12 h dialysis, the sample was further 393 purified through negative chromatography by passing through a GST-affinity column (GSTrap 394 FF Columns, GE Healthcare Life Sciences) to remove free GST, residual GST-fused Ire1 KR, 395 and GST-tagged protease, and a Q FF anion exchange column (GE Healthcare Life Sciences) 396 to remove contaminating nucleic acids. The flow-through containing untagged Ire1 KR was 397 further purified by applying it to a Superdex 200 16/60 gel filtration column (GE healthcare) and 398 then concentrated to 20-40 µM in storage buffer (50 mM Tris-HCl pH 7.5, 500 mM NaCl, 2 mM 399  $Mg(OAc)_2$ , 2 mM TCEP, 10% glycerol) and flash-frozen in liquid nitrogen. The final purity, as 400 well as purity at intermediate steps, was assessed by SDS-PAGE using Coomassie blue 401 staining.

402

## 403 In vitro RNA cleavage assay

404 Short RNA oligos derived from the Ire1 cleavage sites on Sp BIP1 mRNA, SPAC4G9.15 405 mRNA, *PLB1* mRNA, and the Sc HAC1 mRNA 3' splice site were purchased from Dharmacon, 406 Inc. The sequence of stem-loop RNA substrates ordered were the following: Sp BIP1 cleavage 407 site: 5'CGCGAGAUAACUGGUGCUUUGUUAUCUCGCG, Sp SPAC4G9.15 cleavage site: 408 5'CCACCACCGAGUAUGCUACUCGGUGGUGG, Sp PLB1 cleavage site: 409 5'ACGGCCUUUGUUGCAAAAGGGUCGU (25bp), and Sc HAC1 3' splice site: 410 5'GCGCGGACUGUCCGAAGCGCAGUCCGCGC. RNA oligos were gel extracted, acetone 411 precipitated, and resuspended in RNase-free water. The oligos were 5'-end radiolabeled with gamma-[<sup>32</sup>P]-ATP (Perkin Elmer) using T4 polynuclotide kinase (NEB) and cleaned using 412 413 ssDNA/RNA Clean and Concentrator kit (Zymo Research D7010).

414 To fold the RNA, we heated the RNA oligos to 90°C for 3 min and slowly cooled them 415 down at a rate of 1°C per minute until the temperature reached 10°C. In the Ire1 cleavage 416 assays, the reaction samples contained 12.5 µM of Ire1-KR. The cleavage reaction was 417 performed as described previously (W. Li et al., 2018) by incubating at 30°C in reaction buffer 418 (50 mM Tris/HCl pH 7.5, 200 mM NaCl, 2 mM Mq(OAc)<sub>2</sub>, 2 mM TCEP, 10% glycerol). For 419 reactions in Figure 1B & C, 200 µM of 4µ8C (Sigma-Aldrich) was added. At each time point, an 420 aliquot was transferred to 1.2x STOP buffer (10 M urea, 0.1% SDS, 1 mM EDTA, trace amounts 421 bromophenol blue). RNAs were separated using denaturing 15% Novex<sup>™</sup> TBE-Urea Gels 422 (ThermoFisher) and transferred to Amersham Hybond-N+ membranes. Radioactive RNA 423 membranes were imaged with a Phosphorimager (Typhoon FLA 9500, GE Health). The band 424 intensities were quantified using ImageJ. The cleaved portion was calculated as the cleaved 425 band intensity divided by the sum of the cleaved band and uncleaved band intensities. The  $k_{obs}$ 426 values were obtained by fitting the data to first-order 'one-phase' decay equations using Prism 427 (GraphPad).

## 428 In vitro RNA cleavage assay of *HAC1-* and *BIP1-*derived RNA mutants

In vitro transcription of the mutant RNA stem-loops derived from the HAC1 3' splice site
 and *BIP1* cleavage site were carried out as follows. Singe-stranded DNA oligonucleotides were
 used as templates to which the 18mer 5'TAATACGACTCACTATAG "T7 promoter
 oligonucleotide" was annealed to create a double-stranded T7 RNA polymerase promoter. The
 templates contain the indicated single point mutations from Figure 4 on the following parent

- 434 oligonucleotides: HAC1-27 (encoding wild-type HAC1 3' stem-loop RNA with T7 promoter):
- 435 5'GCGCGGACTGCGTTCGGACAGTCCGCCTATAGTGAGTCGTATTA, and BIP1-32 (encoding

436 wild-type *BIP1* stem-loop RNA with T7 promoter):

437 5'CGCGAGATAACAAAGCACCAGTTATCTCGCGCTATAGTGAGTCGTATTA.

438 A solution containing 5 nM T7 promoter oligonucleotide and 0.75 nM template oligonucleotide 439 was heated to 100°C for 3 min and immediately placed on ice. 20 µL transcription reactions 440 containing 5 µL of the template solution, 1 mM each of ATP, CTP, GTP, and UTP, 1x reaction 441 buffer, and 2 µL T7 RNA Polymerase mix (HiScribe T7 High Yield RNA Synthesis Kit, NEB) 442 were incubated at 37°C for 3 h. RNA oligos were gel extracted in 300 µL RNase-free water. 443 These RNA substrates are not radio labeled. The RNAs are folded, cleaved by Ire1-KR and 444 separated by TBE-Urea gels in the same way as the radio labeled RNAs. The TBE-Urea gels 445 were stained with SYBR Gold (ThermoFisher) and imaged on the Typhoon with excitation at 446 488 nm. The emission was collected using a band pass filter at 550 nm. Image analysis is the 447 same as radio labeled RNAs.

448

## 449 Bacterial growth assays

450 Expression vectors containing the Sc Ire1-KR, Sp Ire1-KR, or a mutant form of these 451 proteins regulated by a T7 promoter were transformed into BL21-CodonPlus (DE3)-RIPL E. coli 452 cells. Freshly transformed E. coli cells were cultured overnight (~20 h) and then diluted to an 453 OD<sub>600</sub> of 0.02. The cultures were incubated at 37°C and their OD<sub>600</sub> was obtained every 15 min 454 by the Tecan Spark Multimode Microplate Reader (Life Sciences) (60 cycle kinetic loop, 455 continuous shaking, double orbital 2.5 mm, 108 rpm) for 5 h. For cultures containing the Ire1 456 RNase inhibitor 4µ8C, 1 µM of 4µ8C (Sigma-Aldrich) was added into both the overnight culture 457 and the diluted culture. It is important to use freshly transformed (transformed within 72 h) E. 458 coli cells as the toxicity of Sp Ire1-KR and mutants of Sc Ire1-KR accumulates over time.

459

## 460 Crosslinking gel

461 Each Ire1 construct was buffer-exchanged three times with Zeba spin desalting columns 462 (ThermoFisher Scientific 89882) into a buffer containing 50 mM HEPES, 200 mM NaCl, 2 mM 463 Mg(OAc)<sub>2</sub>, 2 mM TCEP, and 10% Glycerol. 8 µL crosslinking reactions containing 12.5 µM Ire1, 464 2 mM ADP, and 1 mM BS3 crosslinker (ThermoFisher Scientific 21580) were carried out on ice 465 for two hours and guenched by adding concentrated Tris-HCI to a final concentration of 60 mM. 466 The entire reaction was separated on an SDS-PAGE gel, stained with SYPRO Ruby 467 (ThermoFisher Scientific S21900) or coomassie blue overnight and scanned with the Typhoon 468 FLA 9500 or Gel Documentation system (Bio-Rad) respectively.

## 470 Analytical Ultracentrifugation

471 Experiments were performed in a Beckman Coulter Optima XL-A analytical 472 ultracentrifuge equipped with UV-visible absorbance detection system using a 4-hole An-60 Ti 473 analytical rotor. Multi-speed sedimentation equilibrium experiments were carried out at 20°C 474 and 7,000, 10,000, and 14,000 rpm until equilibrium was reached for 110 µL samples of 475 concentrations of 10  $\mu$ M, 5  $\mu$ M, and 2.5  $\mu$ M protein. Samples were dialyzed overnight into 476 analysis buffer (50 mM Tris/HCl, pH 7.5, 200 mM NaCl, 2 mM Mg(OAc)<sub>2</sub>, 2 mM TCEP) to 477 remove glycerol. Analysis buffer without protein was used as reference. Measurements were 478 made at 280 nm in the absorbance optics mode. Raw data were trimmed using WinReEdit (Jeff 479 W. Lary, University of Connecticut) and globally fitted to a self-association equilibrium model 480 using WinNonlin (Johnson, Correia, Yphantis, & Halvorson, 1981) using all concentrations and 481 speeds for each protein sample.

482

## 483 Molecular dynamics simulation

484 All simulations were performed using the Amber suite (Case et al., 2014). Initial structure 485 and topology files were prepared in LEaP (Sastry, Adzhigirey, Day, Annabhimoju, & Sherman, 486 2013) using the Amber ff12SB force field, the general Amber force field (gaff) and the phosaa10 487 parameters for phosphorylated amino acids. The structural model included an inhibitor bound to 488 the kinase domain that was kept in place for the simulations. Inhibitor parametrization was 489 performed using AnterChamber (Wang, Wang, Kollman, & Case, 2006; Wang, Wolf, Caldwell, 490 Kollman, & Case, 2004). We solvated the system with TIP3P water molecules in a periodic 491 cubic box such that the closest distance between the periodic boundary and the closest atom in 492 the protein was 10 Å. We added counterions to neutralize the box.

We minimized the energy of the system, first using harmonic restraints on the protein
backbone (10 kcal mol<sup>-1</sup> Å<sup>-2</sup>) then in an unrestrained minimization, for 500 steps of steepest
descent and 500 steps of conjugate gradient, each at constant volume with a non-bonded cutoff
distance of 9 Å.

We performed a three-step equilibration: 1. heating the system to 300 K at constant volume with harmonic restraints on the protein backbone (10 kcal mol<sup>-1</sup> Å<sup>-2</sup>) at constant volume using the SHAKE algorithm (Ciccotti & Ryckaert, 1986) constraining bonds involving hydrogens and the Andersen thermostat for 20 ps; 2. constant pressure of 1 bar with lower restraints on the protein backbone (1 kcal mol<sup>-1</sup> Å<sup>-2</sup>) for 20 ps with consistent parameters; 3. the restraints on the protein backbone were released and the system was equilibrated at constant pressure for 1 ns.

- 503 We seeded the production runs with random new velocities at constant pressure of 1 bar 504 and a non-bonded cutoff distance of 9 Å and ran the molecular dynamics simulations for 20 ns 505 with a 2 fs time step. Coordinates and energy were saved every picosecond (500 steps). We 506 assessed the convergence of the simulations by examining backbone root-mean-square 507 deviation (RMSD) plots with particular focus on the RNase domain.
- 508 Visual inspection of molecular models was performed using ChimeraX (Goddard et al., 509 2018), a virtual reality implementation of the traditional molecular visualization program.
- 510 Molecular graphics and analyses performed with UCSF Chimera (Pettersen et al., 2004),
- 511 developed by the Resource for Biocomputing, Visualization, and Informatics at the University of
- 512 California, San Francisco, with support from NIH P41-GM103311.
- 513

## 514 Ire1 deep alignment

515 Ire1 orthologs were identified by searching the Pfam Ribonuc\_2-5A hidden Markov 516 model (HMM) with HMMSEARCH from HMMER 3.1b2 (Eddy, 2011) against all proteins from 517 RefSeq fungal genomes downloaded from GenBank 1/17/2017 plus six plant genomes (C. 518 elegans, D. melanogaster, D. rerio, M. musculus, M. mulatta, and H. sapiens) and six animal 519 genomes (A. thaliana, C. reinhardtii, P. patens, O. sativa, S. lycopersicum, and P. glauca) 520 downloaded from GenBank 7/19/2019. This gave predominantly hits to Ire1 orthologs. Outlier 521 hits to RNaseL and difficult to place microsporidian RNases were removed. A guide multiple 522 alignment was generated by PROBCONS 1.12 alignment (Do, Mahabhashyam, Brudno, & 523 Batzoglou, 2005) of the Ire1 hits from plants and animals plus 14 fungal species (H. capsulatum, 524 A. nidulans, N. crassa, F. graminearum, Y. lipolytica, S. cerevisiae, C. albicans, K. lactis, S. 525 pombe, P. carnii, C. neoformans, U. maydis, P. graminis, and M. verticillata). HMMBUILD was 526 used to generate an HMM from the conserved regions of the PROBCONS alignment, spanning 527 the kinase and RNase domains. HMMALIGN was then used to realign the full set of Ire1 hits to 528 the new HMM. A maximum likelihood phylogeny was estimated from the aligned positions with 529 FASTTREE 2.1.8 (Price, Dehal, & Arkin, 2010). The resulting tree was rendered with an in-530 house PYTHON script.

- 531
- 532

#### 533 Figure 1. The promiscuous RNase activity of *S. pombe* Ire1 causes toxicity to 534 bacterial cells. (A) Growth curves of bacterial cells expressing various Ire1 kinase-RNase (KR) 535 domains. Optical densities at 600 nm (OD<sub>600</sub>) were measured every 15 min for 5 h. Bacterial 536 cells expressing S. cerevisiae (Sc) or S. pombe (Sp) Ire1-KR were incubated at 37°C. In the 537 indicated samples, 1 µM of the Ire1's RNase inhibitor 4µ8C was added to the culture. The Sp 538 Ire1-KR(H1018A) has a catalytically inactive RNase. (B, C) In vitro RNA cleavage assays with 539 or without 200 µM of 4µ8C. 5' radio-labeled stem-loop RNA substrates, which are derived from 540 the XBP1 mRNA 3' splice site, were incubated with 12.5 µM of Sp (B) or Sc (C) Ire1-KR at 30°C 541 for the indicated time.

#### 542 Figure 2. S. cerevisiae Ire1-KR-mut17 has a promiscuous RNase activity. (A) 543 Sequence alignment of the RNase domains of Ire1 orthologs from Saccharomyces cerevisiae, 544 Schizosaccharomyces pombe, Schizosaccharomyces octosporus, Schizosaccharomyces 545 cryophilus. A total of 17 residues (green) were selected as the candidate residues that may 546 regulate Ire1's RNase promiscuity. These candidate residues are located at back-to-back dimer 547 interface (marked with \*), oligomer interface (marked with +) or near the helix loop element 548 (marked with ^). Sequence of the helix loop element (HLE) is underlined. (B) The location of the 549 17 candidate residues (green) on the back-to-back dimer structure of the Ire1 cytosolic domain 550 (PDB: 3FBV) with kinase domain in yellow and RNase domain in purple. The dimer interface, 551 oligomer interface and HLE regions are indicated in dashed boxes. (C-F) In vitro RNA cleavage 552 assays with 12.5 µM of wildtype (WT) Sc Ire1-KR or Sc Ire1-KR-mut17. The stem-loop RNA 553 substrates are derived from the Sc HAC1 mRNA 3' splice site (C), Sp BIP1 (D), PLB1 (E) and 554 SPAC4G9.15 (F) mRNA cleavage sites. Experimental conditions are the same as in Figure 555 1B&C. Predicted RNA secondary structures are illustrated. Ire1 cleavage sites are marked with red dashed lines. (G) Comparison of the kobs of WT Sc Ire1-KR, Sc Ire1-KR-mut17 and Sp Ire1-556 557 KR. The kobs of WT Sc Ire1-KR and Sc Ire1-KR-mut17 were calculated from experiments in (C-558 F). The k<sub>obs</sub> of Sp Ire1-KR was measured under the same condition from our previous study (W. 559 Li et al., 2018). "BD" indicates cleavage activity below detection limit. Experiments were 560 performed in duplicates. (H) Bacterial growth assay for WT Sc Ire1-KR, Sc Ire1-KR-mut17 and 561 Sp Ire1-KR. Experimental conditions are the same as in Figure 1A. In the indicated samples, 1 562 $\mu$ M of 4 $\mu$ 8C was added. 563

## 564 Figure 3. Two residues at Ire1's RNase-RNase dimer interface regulate Ire1's

565 **RNase promiscuity.** (A) Bacterial growth assay for *Sc* Ire1-KR revertants. Conditions are the

566 same as in Figure 1A. OD<sub>600</sub> at 5-hour time-point was measured. Experiments were performed

- 567 in duplicates. Dashed line marks the threshold used to separate toxic and non-toxic Ire1
- 568 constructs. (B-E) In vitro cleavage assays of Sc Ire1-KR(K992D,H1044D,Y1059R), Sc Ire1-
- 569 KR(K992D,Y1059R) and Sc Ire1-KR(H1044D) on Sc HAC1 mRNA 3' splice site (B), Sp BIP1
- 570 (C), *PLB1* (D) and *SPAC4G9.15* (E) mRNA cleavage sites. Experimental conditions are the
- same as in Figure 1C. (F) Comparison of the  $k_{obs}$  that are calculated from results in (B-E). "BD"
- 572 indicates cleavage activity below detection limit. Experiments were performed in duplicates.

## 574 Figure 4. S. cerevisiae Ire1-KR(K992D,Y1059R) has a promiscuous RNase activity.

575 (A-B) A series of twenty-four (A) and twenty-seven (B) stem-loop RNA substrates, which are

- 576 derived from the *S. cerevisiae HAC1* mRNA 3' splice site (A) or the *S. pombe BIP1* mRNA
- 577 cleavage site (B), are in vitro transcribed. Each of the substrate carries a single-point mutation,
- 578 which is located on the loop or at the end position of the stem. The sequence of the various
- 579 point mutations is indicated next to each residue (above the line). Listed below these sequences
- are the cleavage efficiencies, at which each mutant RNA substrate was cleaved by *Sc* Ire1-KR
- 581 (first row below the line in panel A), *Sp* Ire1-KR (first row below the line in panel B) or *Sc* Ire1-
- 582 KR(K992D,Y1059R) (second rows below the lines in panels A and B). k<sub>obs</sub> of mutant stem-loop
- 583 RNAs is normalized to  $k_{obs}$  of WT HAC1 (A) or BIP1 (B) stem-loop RNAs that are cleaved by the
- 584 corresponding Ire1. (-) is no cleavage; (-/+) is 1-10%; (+) is 10-60%; (++) is 60-120%; (+++) is >
- 585 120%. Ire1 cleavage sites are marked with red dashed lines. Yellow squares (in A), blue
- 586 squares (in B) and grey circles (in A and B) mark the positions, at which specific residues are
- 587 required to achieve efficient cleavages by Sc Ire1-KR, Sp Ire1-KR and Sc Ire1-
- 588 KR(K992D,Y1059R), respectively. Experimental conditions are the same as in Figure 1C.

590 Figure 5. Structural re-arrangement at Ire1 dimer interface regulates the RNase 591 promiscuity. (A) Back-to-back dimer structure of WT Sc Ire1 cytosolic domain (PDB: 3FBV) 592 with kinase domain in yellow and RNase domain in purple. K992 and Y1059 are colored in 593 green while E988 is colored in blue. Side chain labels on protomer B are marked with '. (B) 594 Close-up view focusing on the interface region of WT Sc Ire1 dimer. Dashed lines indicate salt 595 bridges. (C) Close-up view focusing on the interface region of the predicted dimer structure of 596 Sc Ire1-KR(K992D,Y1059R), which was generated by a 20-ns molecular dynamics (MD) 597 simulation from an initial structural model that was built from the WT Sc Ire1 dimer (PDB: 598 3FBV). The final frame of the simulation was illustrated. D992 and R1059 are colored in green 599 while E988 is colored in blue. Dashed lines indicate salt bridges. (D) Illustration of the residues 600 at Sc Ire1 position 988, 992 and 1059 (or Sp Ire1 position 946, 950 and 1016). Dashed lines 601 indicate salt bridges. (E) Crosslinking gel for various Ire1-KR constructs. Indicated Ire1-KR (12.5 602 µM) constructs were incubated with or without 1 mM of crosslinker bissulfosuccinimidyl suberate 603 for 2 h on ice before being separated on an SDS-PAGE gel and stained by coomassie blue. (F) 604 Ire1's ability to distinguish Sc HAC1- and Sp BIP1-derived RNA substrates is measured by the 605 ratio of their corresponding k<sub>obs</sub>. (G) Evolutionary comparison of Ire1 orthologs from various 606 species. The analysis focuses on three residues, which correspond to position 988, 992 and 607 1059 on Sc Ire1. The S. cerevisiae-like pattern (yellow) has an E/D at 988, K/R at 992 and Y at 608 1059. The S. pombe-like pattern (blue) has a varying amino acid (aa) at 988, D/E at 992 and 609 K/R at 1059.

611 Figure 6. Interface mutations change the protomer alignment in Ire1 dimer. (A) 612 Structure alignment of WT Sc Ire1-KR and Sc Ire1-KR(K992D,Y1059R). 20-ns MD simulations 613 were performed on both WT Sc Ire1-KR and Sc Ire1-KR(K992D,Y1059R). The last simulation 614 frame was used for structure alignment. The protomer A of the two dimers were aligned with 615 minimal root mean square deviation. Sc Ire1-KR(K992D,Y1059R) is in grey. WT Sc Ire1-KR has 616 its kinase domain in yellow and RNase domain in purple. The  $\alpha$ 1-helix at position 983-998,  $\alpha$ 4-617 helix at position 1048-1064, H1061, and R1039 are in red, pink, yellow and cyan, respectively. 618 (B, C) Bottom-up view of the aligned RNase domains (B), and the  $\alpha$ 1- and  $\alpha$ 4-helix (C). Side 619 chains of D992 and R1059 are shown. Dashed lines are salt bridges. (D) Side view of  $\alpha$ 1-620 helices. Color coding are the same as in (A). Side chain labels on protomer B are marked with '. 621 (E) Measuring Ire1 protomer alignment in the MD simulation. Y-axis is the distance between the 622 centers of mass of the two  $\alpha$ 1-helices. X-axis is the distance between the  $\alpha$ -carbons of R1039 623 on protomer A and H1061' on protomer B. Each dot represents a time point in the MD 624 simulation. Measurements of WT Sc Ire1-KR are in yellow while measurements of Sc Ire1-625 KR(K992D,Y1059R) are in grey. The green dot is the measurement from the crystal structure of 626 human RNase L (PDB: 4OAV). The R1039-H1061' distance on Sc Ire1 corresponds to the 627 R651-H672' distance on RNase L. The  $\alpha$ 1-helix on RNase L is from W589 to V599. (F) Cartoon 628 illustration showing the protomer alignment of WT Sc Ire1-KR and Sc Ire1-KR(K992D,Y1059R). 629 Their RNase domains were zoomed in with double-arrow lines showing the distances being 630 measured in (E). 631

## 632 Figure 1-figure supplement 1. $4\mu$ 8C inhibits the RNase activity of *S. pombe* and *S.*

633 *cerevisiae* Ire1. (A-B) Results from the in vitro cleavage assays in Figure 1B & C were

634 quantified using ImageJ. Cleaved portion was calculated as: cleaved RNA / (cleaved RNA +

635 uncleaved RNA). For cleaved portion  $\ge$  0.1 at time point 30 min, the data were fitted to a first-

order exponential function. For cleaved portion < 0.1 at time point 30 min, the data were fitted to

637 a linear function. Experiments were performed in duplicates. (C) Bar graph showing the  $k_{obs}$  of

638 the indicated Ire1 with or without  $4\mu$ 8C. (D)  $4\mu$ 8C inhibits S. *pombe* Ire1's RNase activity in vivo.

- 639 200 ul of WT S. *pombe* cell culture at OD<sub>600</sub> 0.5 was evenly spread onto YE5S plate, which does
- or doesn't contain 0.05  $\mu$ g/ml of tunicamycin. Circular Whatman paper (diameter ~1 cm) was

641 placed onto the YE5S plates. 5 ul of DMSO or  $4\mu$ 8C (at concentration of 1 mM) was spotted

onto the Whatman papers. Plates were incubated at 30°C for 3 days.

- Figure 2-figure supplement 1. Quantification of in vitro cleavage assays. Results
   from the in vitro cleavage assays in Figure 2C-F were quantified and analyzed using the same
- 646 methods as in Figure 1-figure supplement 1A.

- 648 Figure 2-source data 1. A list of the 17 candidate residues on *S. cerevisiae* and *S.*
- 649 *pombe* Ire1. The oligomer interface is referred as the interface IF2<sup>C</sup> and the back-to-back dimer
- 650 interface is referred as the interface IF1<sup>c</sup> in (Korennykh et al., 2009).

- 652 **Figure 3-figure supplement 1. Quantification of in vitro cleavage assays.** Results
- 653 from the in vitro cleavage assays in Figure 3B-E were quantified and analyzed using the same
- 654 methods as in Figure 1-figure supplement 1A.

- 656 Figure 3-source data 1. The detailed sequence information of *S. cerevisiae* Ire1-KR
- 657 **constructs used in this study.** "Y" indicates the mutation is present in the corresponding Ire1
- 658 construct. "N" indicates the mutation is absent in the corresponding Ire1 construct.

- 660 Figure 5-figure supplement 1. The root-mean-square deviation (RMSD) of atomic
- 661 positions of Ire1 RNase domain during the simulation. Result of WT Sc Ire1-KR is in yellow
- and Sc Ire1-KR(K992D,Y1059R) is in grey.

**Figure 5-figure supplement 2. Time fraction of the MD simulation during which the indicated salt bridges are present.** A 20-ns MD simulation was performed on the dimer structure of WT *Sc* Ire1 KR or *Sc* Ire1-KR(K992D,Y1059R). At a given time point during the simulation, if the distance between the two indicated residues is smaller than 2.5 Å, the two residues are considered to be connected by a salt bridge. The bar graph shows the accumulative time fraction during which each indicated salt bridge is present. Side chain on protomer B are marked with '.

## **Figure 5-figure supplement 3. Sedimentation equilibrium analytical**

673 ultracentrifugation (SE-AUC) analysis of Sc Ire1-KR and Sc Ire1-KR(K992D,Y1059R). Each 674 protein was examined under nine conditions: at three protein concentrations (2.5, 5 and 10  $\mu$ M) 675 and three centrifugal speeds (7,000, 10,000, and 14,000 rpm). All nine sets of data for each 676 protein were fitted to an equilibrium model. (A) WT Sc Ire1-KR was mostly monomeric at the 677 concentrations tested and had a monomer-dimer equilibrium constant of 57 µM. (B) Sc Ire1-678 KR(K992D,Y1059R) showed a higher propensity to form dimers/tetramers than WT Sc Ire1 KR, 679 with a monomer-dimer equilibrium constant of 0.98 µM. (C) Plot of the natural logarithm of the 680 absorbance at 280 nm versus the square of the spin radius for the 10  $\mu$ M Sc Ire1-KR and Sc 681 Ire1-KR(K992D,Y1059R) samples at 7,000 rpm.

- 683 Figure 5-figure supplement 4. RNA cleavage efficiencies of Sc Ire1-KR mutants
- 684 bearing mutations at back-to-back dimer interface. k<sub>obs</sub> of the indicated Sc Ire1-KR
- 685 constructs cleaving stem-loop RNA substrates derived from *Sc HAC1* mRNA 3' splice site or *Sp*
- 686 BIP1 mRNA cleavage site. "BD" indicates cleavage activity below detection limit.
- 687

688 Figure 5-figure supplement 5. Evolutionary comparison of Ire1 orthologs. We 689 generated a deep alignment of 230 Ire1 orthologs from fungi (211), representative plants (9) and 690 animals (10). We compared residues at three positions, which correspond to the S. cerevisiae 691 Ire1 E988, K992 and Y1059. Among the 230 orthologs being analyzed, 175 of them have the S. 692 pombe-like pattern (blue) – with a varying amino acid at position 988, a negatively charged 693 amino acid (aspartate or glutamate) at position 992 and a positively charged amino acid (lysine 694 or arginine) at position 1059. 33 of the Ire1 orthologs have the S. cerevisiae-like pattern-with a 695 negatively charged amino acid (aspartate or glutamate) at position 988, a positively charged 696 amino acid (lysine or arginine) at position 992 and mostly tyrosine at position 1059. Tyrosine at 697 position 1059 may orchestrate the positioning of lysine/arginine side chain at position 992 (Lee 698 et al., 2008). The S. cerevisiae-like pattern is restricted to the Saccharomycotina. In 6 Ire1 699 orthologs, the S. cerevisiae- or S. pombe-like inter-molecular salt bridge are replaced with a 700 putative hydrogen bond (marked with x). Ire1 orthologs from higher plants exhibit a pattern 701 distinct from S. cerevisiae and S. pombe. One interesting case (Table insert A) is with the Ire1 702 from Yarrowia lipolytica, whose location on the evolutionary tree is at the boundary between S. 703 cerevisiae-like and S. pombe-like species. Yarrowia lipolytica Ire1 is in a transitioning state from 704 the S. cerevisiae-like into the S. pombe-like Ire1-its amino acids at position 988 (aspartate) 705 and 1059 (tyrosine) remain S. cerevisiae-like whereas its amino acid at position 992 (glutamate) 706 changed from S. cerevisiae-like (positively charged) into S. pombe-like (negatively charged). 707 The case of Yarrowia lipolytica Ire1 suggests that salt-bridge rewiring occurs during the 708 evolutionary process. Another interesting case (Table insert B) is with the Ire1 from Capronia 709 epimyces. While its neighboring species have S. pombe-like pattern, Capronia epimyces Ire1 710 undergoes a unique evolution with arginine, aspartate and methionine at positions 988, 992 and 711 1059, making it a charge reversal (at positions 988 and 992) of S. cerevisiae-like Ire1. The 712 detailed sequences are listed in Figure 5-source data 1.

Figure 5-source data 1. In this table, 230 Ire1 orthologs were compared. Their residues
at three positions, which correspond to the *S. cerevisiae* Ire1 E988, K992 and Y1059, are listed.

Figure 6-figure supplement 1. Protomer alignment of RNase L. The dimer structure of WT *Sc* Ire1-KR (as used in Figure 6A) and human RNase L (PDB: 4OAV) were compared. The protomer A of the *Sc* Ire1-KR and human RNase L dimers were aligned with minimal root mean square deviation. The RNase L is in light brown. The color coding of *Sc* Ire1-KR is the same as in Figure 6. (A) Bottom-up view of the  $\alpha$ 1- and  $\alpha$ 4-helix; (B) Side view of  $\alpha$ 1-helices. **Table 1. Plasmids used in this study.** In all of the plasmids, a GST and an HRV 3C

722 protease site are N-terminally fused to Ire1-KR.

plasmid		
number	description	source
nPW1477	Sc Ire1-KB on nGEX6P-2	Korennykh et al.,
pPW3205	So Ire1-KB on pGEX6P-2	Lietal 2018
pPW3244	Sc Ire1-KB-mut17 on pGEX6P-2	This study
pPW3262	Sc Ire1-KR(K992D,Y1059R) on pGEX6P-2	This study
pPW3263	Sc Ire1-KR(K992D,H1044D,Y1059R) on pGEX6P-2	This study
pPW3256	revertant 1 (K992) on pGEX6P-2	This study
pPW3245	revertant 2 (N1001) on pGEX6P-2	This study
pPW3246	revertant 3 (M1010) on pGEX6P-2	This study
pPW3247	revertant 4 (T1032) on pGEX6P-2	This study
pPW3248	revertant 5 (F1033) on pGEX6P-2	This study
pPW3257	revertant 6 (E1038) on pGEX6P-2	This study
pPW3258	revertant 7 (R1039) on pGEX6P-2	This study
pPW3259	revertant 8 (H1044) on pGEX6P-2	This study
pPW3260	revertant 9 (S1045) on pGEX6P-2	This study
pPW3249	revertant 10 (M1049) on pGEX6P-2	This study
pPW3250	revertant 11 (Y1059) on pGEX6P-2	This study
pPW3261	revertant 12 (F1062) on pGEX6P-2	This study
pPW3251	revertant 13 (M1063) on pGEX6P-2	This study
pPW3252	revertant 14 (I1069) on pGEX6P-2	This study
pPW3253	revertant 15 (A1070) on pGEX6P-2	This study
pPW3254	revertant 16 (E1071) on pGEX6P-2	This study
pPW3255	revertant 17 (L1109) on pGEX6P-2	This study
pPW3441	Sc Ire1-KR(K992D,Y1059A) on pGEX6P-2	This study
pPW3442	Sc Ire1-KR(K992A,Y1059R) on pGEX6P-2	This study
pPW3443	Sc Ire1-KR(K992R,Y1059D) on pGEX6P-2	This study
pPW3275	Sc Ire1-KR(H1018A) on pGEX6P-2	This study

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B RNA: XBP1 3' splice site (21bp) IRE1: Sp Ire1-KR Sp Ire1-KR 4µ8C: + time 0 1 3 10 30 0 1 3 10 30 (min) С RNA: XBP1 3' splice site (21bp) IRE1: Sc Ire1-KR Sc Ire1-KR 4µ8C: + \_ time 0 3 10 30 0 1 3 10 30 1 (min)





А

HAC1 mRNA 3'splice site



residue mutation Sc Ire1-KR cleavage efficiecy Sc Ire1-KR(K992D,Y1059R) cleavage efficiecy Required for Sc Ire1-KR cleavage Required for Sc Ire1-KR(K992D,Y1059R) cleavage  $\bigcirc$ Required for both +++  $k_{obs}$  ratio  $\geq 120\%$ ++ k<sub>obs</sub> ratio 60-120% k<sub>obs</sub> ratio 10-60%  $^{+}$ k<sub>obs</sub> ratio 1-10% -/+ no cleavage

В

## BIP1 mRNA cleavage site



residue mutation *Sp* lre1-KR cleavage efficiecy

Sc Ire1-KR(K992D,Y1059R) cleavage efficiecy

Required for Sp Ire1-KR cleavage

) Required for Sc Ire1-KR(K992D,Y1059R) cleavage

Required for both

# Figure 5 Li & Crotty et al.

D



	Sc Ire1-KR		Sp Ire1-KR		Sc Ire1-KR (K992D,Y1059R)		Sc lre1-KR (K992D,Y1059A)		Sc lre1-KR (K992A,Y1059R)		Sc lre1-KR (K992R,Y1059D)	
protomer position	A	В	А	В	А	В	А	В	А	В	А	В
Sc 988 (Sp 946)	E۰	, É	D	D	E	E	E	E	E	Е	E	E
Sc 992 (Sp 950)	κ.⁄	``К	D.	<i>, ,</i> D	D,	, D	D	D	A	А	R .	, R
Sc 1059 (Sp 1016)	Y	Y	R ∕ ́	`` R	R ∕ ′	`` <sub>`</sub> R	А	А	R	R	D, '	``D

--- salt bridge





aa position (Sc) Species	988	992	1059	
Kluyveromyces lactis	E	К	Y	Sc-like
Saccharomyces cerevisiae	E	К	Y	Sp-like
Debaryomyces hansenii	E	К	Y	
Candida albicans	E	К	Y	
Aspergillus fumigatus	S	D	R	
Schizosaccharomyces pombe	D	D	R	
Cryptococcus neoformans	G	D	К	
Drosophila melanogaster	S	D	К	
Caenorhabditis elegans	А	D	К	
Homo sapiens Ire1a	Q	D	К	
<i>Homo sapiens</i> Ire1β	Q	D	К	





















B side view of interface α1-helices

