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#### XLF and APLF bind to Ku80 on two remote sites to ensure DNA repair by 1 non-homologous end-joining 2 3 **AUTHORS** 4 Clement Nemoz<sup>1\*</sup>, Virginie Ropars<sup>1\*</sup>, Philippe Frit<sup>2,3\*</sup>, Amandine Gontier<sup>1</sup>, Pascal Drevet<sup>1</sup>, 5 Jinchao Yu<sup>1</sup>, Raphaël Guerois<sup>1</sup>, Aurelien Pitois<sup>1</sup>, Audrey Comte<sup>1</sup>, Christine Delteil<sup>2,3</sup>, 6 Nadia Barboule<sup>2,3</sup>, Pierre Legrand<sup>4</sup>, Sonia Baconnais<sup>5</sup>, Yandong Yin<sup>6</sup>, Satish Tadi<sup>7</sup>, 7 Emeline Barbet-Massin<sup>8</sup>, Imre Berger<sup>9</sup>, Eric Le Cam<sup>5</sup>, Mauro Modesti<sup>7</sup>, Eli Rothenberg<sup>6</sup>, 8 Patrick Calsou<sup>2,3§</sup>, Jean-Baptiste Charbonnier<sup>1§</sup> 9 10 11 **INSTITUTIONS** <sup>1</sup> Institute for Integrative Biology of the Cell (I2BC), Institute Joliot, CEA, CNRS, Univ. 12 Paris-Sud, Université Paris-Saclay, 91198, Gif-sur-Yvette cedex, France 13 <sup>2</sup> Institut de Pharmacologie et Biologie Structurale, IPBS, Université de Toulouse, CNRS, 14 UPS, Toulouse, France 15 <sup>3</sup> Equipe Labellisée Ligue Contre le Cancer 2013 and 2018 16 <sup>4</sup> Synchrotron Soleil, L'Orme des Merisiers, Saint-Aubin - BP 48, 91192 Gif-sur-Yvette 17 18 Cedex, France. <sup>5</sup> Signalisations, Novaux et Innovations en Cancérologie, UMR 8126, CNRS, Université 19 Paris-Sud, Gustave Roussy, Université Paris-Saclay, 94805 Villejuif, France. 20 <sup>6</sup> New York University School of Medicine, Perlmutter Cancer Center, USA 21 <sup>7</sup> Cancer Research Center of Marseille, CNRS UMR7258, Inserm U1068, Institut Paoli-22 Calmettes, Aix-Marseille Université UM105, Marseille, France. 23 <sup>8</sup> Dynamic Biosensors GmbH, Martinsried, Germany 24 <sup>9</sup> BrisSynBio Centre, The School of Biochemistry, Faculty of Biomedical Sciences, 25 26 University of Bristol, Bristol, United Kingdom. 27 28 \* These authors contributed equally § Corresponding authors : jb.charbonnier@cea.fr and calsou@jpbs.fr 29 30 31 32

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#### 33 ABSTRACT

34 The Ku70-Ku80 (Ku) heterodimer binds rapidly and tightly to ends of DNA double-strand 35 breaks and recruits several factors of the Non-Homologous End Joining (NHEJ) pathway 36 through molecular mechanisms that remain unclear. Here, we describe the crystal structures 37 of the Ku-binding motifs (KBM) of the NHEJ proteins APLF (A-KBM) and XLF (X-KBM) 38 bound to a Ku-DNA complex. The two KBMs motifs bind on remote sites of Ku80  $\alpha/\beta$ 39 domain. The X-KBM occupies an internal pocket formed after an unprecedented large 40 outward rotation of the Ku80  $\alpha/\beta$  domain. We reveal independent recruitment at laser-41 irradiated sites of the APLF-interacting protein XRCC4 and of XLF through the respective 42 binding of A- and X-KBMs to Ku80. Finally, we show that mutations on the X-KBM and A-43 KBM binding sites in Ku80 compromises efficiency and accuracy of end-joining and cellular 44 radiosensitivity. A- and X-KBMs may represent two initial anchorage points necessary to 45 build the NHEJ intricate interactions network. 46 47 Keywords : DSB repair, NHEJ, X-ray crystallography, induced fit, laser micro-irradiation,

48 super resolution microscopy, switchSENSE, microcalorimetry

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# 51 **INTRODUCTION**

52

53 In mammals the majority of DNA double-strand breaks (DSBs) is repaired by the nonhomologous end-joining (NHEJ) pathway <sup>1,2</sup>. The Ku70-Ku80 heterodimer (Ku) rapidly and 54 tightly interacts as a preformed ring with DSBs extremities in a non-sequence specific manner 55 <sup>3</sup>. Ku serves as a hub for the recruitment of several NHEJ factors <sup>4,5</sup>. Among them, Ku recruits 56 the XRCC4-LIG4-XLF ligation complex through interaction with XRCC4-LIG4<sup>6,7</sup> and with 57 XLF, the latter relying on a Ku-binding motif (KBM) localized at the XLF extreme C-58 terminus (thereafter named X-KBM)<sup>8,9</sup> (Figure 1a and Supplementary Figure 1). The ligation 59 complex organizes into filaments both in vitro and in cells <sup>10-14</sup>. 60

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62 Interestingly, Ku interacts also with a number of accessory NHEJ factors. The APTX and PNKP-like factor (APLF) binds poly(ADP)-ribosylated proteins near DSBs sites <sup>15,16</sup>, and has 63 been reported to have nuclease activity <sup>16,17</sup>. APLF tightly interacts with Ku through a KBM 64 (thereafter named A-KBM) that is located in its central region 9,18 (Figure 1a and 65 66 Supplementary Figure 1). This interaction has been mapped to the periphery of the Ku80 von Willebrand A domain (vWA)<sup>5</sup>. Ku-APLF interaction was shown to facilitate recruitment of 67 the APLF-partner XRCC4 at damaged sites <sup>9</sup> and was proposed to stabilize the assembly of 68 NHEJ factors around the DSB<sup>19</sup>. Notably, an A-KBM-like domain is present at the N-69 terminus of a recently identified inhibitor of the NHEJ pathway, CYREN(MRI), that also 70 71 interacts with Ku80<sup>20</sup> (Figure 1a). Ku also associates with the Werner syndrome protein (WRN) that is involved in many aspects of DNA metabolism including NHEJ<sup>21</sup>. Two motifs 72 in the C-terminus of WRN cooperate for interaction with Ku, one being A-KBM like, and the 73 74 other resembling the X-KBM present on XLF (Figure 1a). In addition, we and others showed 75 recently that PAXX (Paralog of XRCC4 and XLF) interacts with the Ku70 subunit through a third type motif that is located in its C-terminus <sup>22,23</sup> (Figure 1a). Despite identification of 76 77 KBMs in several NHEJ factors, their respective contribution to the efficiency of DSB repair is 78 not fully understood. For example, the puzzling observations that KBM deletion in XLF or 79 APLF depletion in human cells lead to null or intermediate repair defect deserve further investigations 9,24-27. 80

81

The interactome of Ku thus defines a large ensemble of motifs and proteins that could potentially compete or act synergistically. However, despite important structural and biophysical studies on NHEJ complexes <sup>28</sup>, the absence of high resolution structures of KuKBMs complexes limits our understanding of the roles and specificity of the different
molecular interactions in the recruitment of NHEJ factors to DSBs. Mapping KBM-binding
sites in structures is also needed to clarify potential competition of all the Ku interacting
factors on limited positions (Supplementary figure 1).

89

90 Here, we establish the structural and functional basis of Ku unique modes of interaction with 91 two factors within the NHEJ repair pathway. We describe the first crystal structures of Ku70-92 Ku80-DNA complex in interaction with the A-KBM (KBM of APLF) and with the X-KBM 93 (KBM of XLF), revealing that the two KBMs occupy remote interaction sites on the Ku80 94 vWA domain. The functional context of our structural data was determined using cell based 95 assays to visualize the recruitment of wild-type and mutant X-KBM motifs or XLF proteins as 96 well as of the APLF-partner XRCC4 to DSBs sites induced by micro-irradiation in wild-type 97 or mutant Ku backgrounds. Our data provide new mechanistic insights on the function of XLF 98 and APLF in the NHEJ process.

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- 100

101 **RESULTS** 

102

# The KBM of APLF tightly interacts with a highly conserved site of Ku80 vWA domain.

105 The APLF factor contains a conserved Ku binding motif (A-KBM, aa 179-192) (Figure 1a). We co-crystallized a 18-mer A-KBM peptide (aa 174-191) with a Ku form deleted for the C-106 terminal regions (Ku<sub>cc</sub>), and an hairpin DNA (hDNA)<sup>3</sup> (Supplementary Figure 2a). In the 107 crystal structure at 3.0 Å resolution (Table 1), the A-KBM peptide is well defined and is 108 positioned at the periphery of the vWA domain of Ku80 (aa T61<sup>80</sup>-C156<sup>80</sup>) (thereafter, 109 superscript <sup>80</sup> stands for Ku80) (Figure 1b, and Supplementary Figure 2b). It is located at 110 111 more than 50 Å from the DNA binding site of Ku80. The A-KBM adopts an extended 112 conformation in a pocket delineated by the helices  $\alpha 4$  and  $\alpha 5$  and the loop located between the 113  $\beta$ -strands B and C of Ku80 (Figure 1c). The hydrophobic part of the A-KBM, located in the 114 C-terminal part of the motif, is composed by the amino acids I<sub>185</sub>LPTWML<sub>191</sub> and is buried in a hydrophobic pocket formed by the Ku80 residues  $L68^{80}$ ,  $I112^{80}$ ,  $M115^{80}$ ,  $I149^{80}$  and  $I150^{80}$ 115 116 (Figure 1d). The N-terminal part of the A-KBM contained a patch of three consecutive basic residues and an acid residue (E181RKR184 in human sequence) (Figure 1e). It forms salt 117

bridges and charged hydrogen bonds with respectively the side chains of  $D106^{80}$ ,  $D109^{80}$ ,

119 Q73<sup>80</sup> and S145<sup>80</sup>, and the main chain of K144<sup>80</sup> and S143<sup>80</sup>.

120

121 Isothermal Titration Calorimetry (ITC) showed that the interaction between A-KBM and Ku 122 had a Kd of  $33 \pm 10$  nM (Table 2, and Supplementary Figure 2e). We measured a nanomolar Kd for the interaction between Ku and a 18bp DNA as already reported <sup>29</sup> and found a similar 123 Kd for the interaction between the A-KBM and Ku alone or Ku bound to a 18bp DNA (Table 124 125 2). The interaction of the A-KBM with Ku<sub>cc</sub> showed Kd and enthalpy values similar to full-126 length Ku (KuFL). Thus, the core heterodimeric region of Ku is likely sufficient for the 127 interaction with the A-KBM. Notably, these affinities are stronger than the ones previously 128 reported by fluorescence polarization with a labelled A-KBM peptide (Kd of 580 nM)<sup>9</sup>. 129 130 Mapping the conservation rate of the residues at the surface of Ku80 shows that this pocket is

the main conserved pocket together with the DNA binding pocket (Figure 1f). The residues L68<sup>80</sup>, Y74<sup>80</sup> and I112<sup>80</sup> make tight interactions with the hydrophobic part of the A-KBM motif. Mutations at these positions were reported to greatly reduce or disrupt the interactions with APLF in yeast two-hybrid experiments or in EMSA <sup>9</sup>. We produced the Ku I112R mutant that, as expected had no residual interaction with the A-KBM motif by ITC (Table 2).

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# 138 X-KBM creates an outward rotation of the vWA domain and a large groove in Ku80.

139

140 We then determined the crystal structures of the Ku70-Ku80-hDNA complex bound with peptides derived from the XLF X-KBM ((L281<sup>X</sup> to S299<sup>X</sup>) and (S287<sup>X</sup>-S299<sup>X</sup>) peptides) 141 142 (Supplementary Figure 2c). The crystal structures at 2.8 and 2.9 Å resolution (Table 1) show 143 an unprecedented large outward rotation of the Ku80 vWA (Figure 2a, b). This movement 144 forms a large groove between the Ku80 vWA and the rest of the Ku heterodimer. We 145 therefore termed this conformation the open state of Ku, in contrast to the closed state 146 observed in the three other crystal structures reported (Ku alone (1JEQ), Ku-hDNA (1JEY) 147 and Ku-hDNA-A-KBM (this study)). The conformational change of Ku80 vWA does not 148 affect Ku interaction with the duplex DNA (Supplementary Figure 2d)

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150 The X-KBM is located on the Ku80 vWA face of the newly created groove in a pocket 151 delineated by 4 strands ( $\beta_A$ ,  $\beta_D$ ,  $\beta_E$  and  $\beta_{E'}$ ) and 3 helices ( $\alpha_2$ ,  $\alpha_{7'}$ ,  $\alpha_{7''}$ ). The motif is positioned

closer to the DNA (at 12Å) than the A-KBM, though not in direct contact. We can model 152 153 eight residues of the X-KBM (292KKPRGLFS299) in the crystal obtained with the 19mer or 154 14mer X-KBM. These residues are the last eight residues of the XLF sequence. The residues 155  $_{296}$ GLFS<sub>299</sub> of the X-KBM occupy a hydrophobic pocket delineated by Ku80 residues L12<sup>80</sup>, V37<sup>80</sup>, F41<sup>80</sup>, F135<sup>80</sup>, F164<sup>80</sup>, Y225<sup>80</sup>, and L234<sup>80</sup> (Figure 2c). In the closed state, these Ku 156 157 residues are buried and mediate intramolecular contacts (Figure 2d). The X-KBM may thus 158 stabilize a transient open conformation of Ku80 in equilibrium with the closed state under 159 basal conditions. In addition, we observed that the Ku80 vWA opening comes along with important secondary structure changes in the linker region (R232<sup>80</sup>-E241<sup>80</sup>) that separates the 160 161 vWA and the rest of the Ku heterodimer (Figure 2c, d).

162

163 ITC measurements showed a moderate affinity of X-KBM for Ku, with a Kd of  $4.4 \pm 0.2 \mu$ M 164 (about 200-fold weaker than the affinity of the A-KBM) (Table 2, and Supplementary Figure 2f). We observed similar affinities and thermodynamic parameters for the X-KBM with a Ku-165 166 DNA complex or with Kucc (Table 2). Notably, this micromolar interaction was not detected in a previous study using fluorescence polarization with labelled peptides <sup>9</sup>. We also measured 167 168 similar affinities for the interactions between XLF homodimer and Ku alone or Ku bound to a 169 18bp DNA (Table 2). These data show that in absence of DNA or with a short DNA protected 170 by Ku ring, XLF and its X-KBM interact similarly with Ku.

171

EMSA analyses confirmed an interaction between XLF and Ku complexed with a 50bp DNA in the  $\mu$ M range (Supplementary Figure 3a-b). Competitions experiments showed that the X-KBM peptide (pXLF) competes in the  $\mu$ M range with the XLF protein and that the A-KBM peptide (pAPLF) does not displace XLF, supporting remote sites of interactions (Figure 2e and Supplementary Figure 3c, d). Also, the C-terminus of PAXX (pPAXX) does not compete with XLF binding, in agreement with previous studies that report an interaction between PAXX C-terminus and Ku70 subunit <sup>22,23</sup> (Supplementary Figure 3e).

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# 182 The outward rotation of the vWA domain of Ku80 is mediated by Glu133.

183

To evaluate if the outward rotation of Ku80 observed in the crystal structure with the X-KBM peptide was present in solution, we performed SAXS analyses. The SAXS data with the A- 186 KBM peptide were comparable with those of the Ku-hDNA sample without any peptide 187 (respective Rg of 53.1Å and 53.7) (Figure 2f). We measured an increase of the Rg to 59.0Å in 188 presence of the X-KBM, corresponding to the opening of Ku80 observed in the crystal 189 (Figure 2f). Comparison of the Dmax values provided further corroboration for the opening of

- 190 the Ku molecule with an increase of 30 Å.
- 191

The acid residue E133<sup>80</sup> buried in Ku80 is well positioned to act as a spring facilitating the 192 Ku80 opening (Figure 2c, d). The glutamate E133<sup>80</sup> is buried in the closed conformation of 193 the Ku80 vWA and its pKa value is estimated by the PDB2POR-2.0 server <sup>30</sup> at a value of 9.1, 194 195 far from the normal pKa of 4.5 for a glutamate in solvent. The outward rotation of Ku80 vWA 196 should be energetically facilitated by the solvation of this Glu following the displacement of Ku80 residues V236<sup>80</sup>, F237<sup>80</sup> and I240<sup>80</sup> away from E133<sup>80</sup> carboxylate function (Figure 2d). 197 Multiple sequence alignments show that E133<sup>80</sup> position and the residues surrounding are 198 199 well conserved in mammalian and saurian and that Ku70 has no equivalent buried acidic 200 residue at this position (Supplementary Figure 4a-b).

201

#### 202 Molecular bases of the specificities of A-KBM and X-BKM motifs binding to Ku80.

203

204 The A-KBM and X-KBM present sequence similarities with a basic patch in their N-terminus followed by a hydrophobic patch <sup>9</sup> (Supplementary Figure 1). Comparison of the crystal 205 206 structures of Ku bound to these two motifs suggests that the high affinity of the A-KBM relies on the tryptophan W189<sup>A</sup> in place of the Leu297<sup>X</sup> in X-KBM. Thus, we used ITC to measure 207 the interaction of the X-KBM motifs with the mutation L297W (LW) or with a non-208 209 conservative L297E mutation (LE). The (LW) peptide has a Kd of 0.12  $\pm$  0.03  $\mu$ M, an 210 interaction 40-fold tighter than wild-type X-KBM (Table 2 and Supplementary figure 2g). 211 The LE mutant presents no detectable interaction with Ku (Table 2). Competition experiments 212 with Ku saturated with the A-KBM showed that the X-KBM (L297W) no longer interacts 213 with Ku, suggesting that the sole L297W mutation is able to redirect the X-KBM towards the 214 A-KBM binding site on Ku80 (Table 2).

215

Then live cell imaging was used to monitor the recruitment of CFP-fused A-KBM and X-KBM fragments to DSBs sites induced with laser micro-irradiation. Under conditions of similar damage yield (Supplementary Figure 5e), the A-KBM motif transfected in U2OS cells was strongly recruited but the W189G mutation impaired both its nuclear localization and

recruitment to laser sites (Figure 3a-b), as reported <sup>9,18</sup>. X-KBM live recruitment was impaired 220 by L297E mutation but not L297W mutation (Figure 3c), corroborating ITC data. We then 221 used U2OS cells expressing an inducible shRNA against Ku80<sup>31</sup> (Supplementary Figure 5a) 222 223 that were complemented with wild-type or I112R mutant Ku80 (Supplementary Figure 5c-d). 224 The I112R mutation impaired A-KBM recruitment, as expected, but not that of X-KBM 225 (Figure 3d, e). Notably, I112R Ku80 mutant specifically lowered the recruitment of L297W 226 X-KBM (compare Figures 3c and 3f). Conversely, APLF knock-down boosted the 227 recruitment of the LW mutant peptide above that of wild-type X-KBM (Figure 3g and 228 Supplementary Figure 5b for control of shAPLF efficiency). Together, these data in cells 229 support that the LW mutation redirects the X-KBM fragment to the APLF-binding site in 230 Ku80 and point out the W189 residue as a key determinant for APLF specific interaction with 231 Ku80.

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# 234 X-KBM mutations impair XLF recruitment and XRCC4-XLF filament stability.

235

236 We then investigated the properties of the interaction between full-length XLF protein and Ku. We first used the SwitchSENSE approach <sup>32</sup> in which oligonucleotide nanolevers labelled 237 with a fluorescence probe are bound to a gold surface (Supplementary Figure 3f). Ku bound 238 onto 48bp DNA nanolevers with a Kd in the nM range as already reported <sup>29</sup> and a long 239 240 dissociation time (Supplementary Figure 3g). Wild-type XLF onto the Ku-DNA complex showed a rapid k<sub>on</sub>  $(4.7 \pm 1.7 \ 10^5 \ M^{-1} s^{-1})$  followed by a rapid dissociation  $(k_{off} = 0.09 \pm 0.004)$ 241 s<sup>-1</sup>) and a corresponding Kd of  $0.19 \pm 0.07 \mu M$  (Figure 4a). This affinity is about 10 fold 242 stronger than the one measured by ITC with a smaller DNA and may reflect additional 243 interactions of XLF with DNA emerging from Ku ring as observed with PAXX <sup>22</sup>. LW and 244 245 LE mutants showed a 2.3- and 5.1-fold weaker affinity than WT protein, respectively (Kd of 246  $0.45 \pm 0.26 \ \mu\text{M}$  for LW and Kd of  $0.98 \pm 0.15 \ \mu\text{M}$  for LE) (Supplementary Figure 3h). As 247 compared with ITC and recruitment data with X-KBM LW mutant peptide, this suggests that 248 the LW mutation cannot redirect the XLF protein to the APLF binding site in Ku80.

249

250 We then measured the recruitment of the CFP-tagged full-length XLF protein (CFP-XLF)

251 expressed in human XLF-defective BuS cells (SV40T-transformed, telomerase immortalized

252 fibroblasts derived from the XLF-deficient P2 patient - homozygous C622T nonsense mutant

253 (R178X))<sup>33</sup> (Figure 4b). Wild-type XLF showed a rapid mobilization to irradiated nuclear

sites, as reported <sup>34</sup>. Both LE and LW mutations impaired to various extents XLF protein 254 255 recruitment (Figure 4b), again indicating that LW mutation cannot redirect the XLF protein to 256 the APLF binding site in Ku80. In addition, we observed an important reduction of the 257 recruitment of G296W, S299E or F298G XLF mutants in the extreme C-terminus 258 (Supplementary Figure 5f-h). We also questioned the contribution of APLF-XRCC4 complex to XLF recruitment by expressing L115D XLF mutant unable to interact with XRCC4 <sup>35,36</sup>. 259 We measured an efficient recruitment of L115D XLF to laser-induced DSBs that was 260 261 insensitive to APLF knock-down but impaired with the XLF L115D/L297E XLF double 262 mutant (Supplementary Figure 5i). Together, these data support a major role for Ku80 263 interaction with L297 and extreme C-terminal residues for XLF recruitment at DSBs in cells.

264

Multi-color super-resolution localization microscopy (STORM) allows characterizing 265 formation of XRCC4-XLF filaments close to Ku foci and DSBs ends<sup>14</sup>. On DSBs induction 266 267 with the radiomimetic drug neocarzinostatin, extended XLF filaments close to Ku80 foci were 268 observed in XLF complemented BuS cells, whereas cells harboring (LE) and (LW) mutants 269 showed slightly smaller and more punctuated XLF structures (Figure 4c and Figure 4d-e for 270 quantification). In contrast to these data, evaluation of the effect of mutations in the X-KBM 271 on cell radiosensitivity showed that (LW) or (LE) XLF mutants were associated with 272 respectively no or minor radiosensitization, compared to high radiosensitivity of BuS cells 273 and full restoration of radioresistance on expression of wild-type XLF (Figure 4f).

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# 276 APLF and XLF binding to Ku80 promote DSB repair and cell survival to IR.

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278 Finally, we questioned the discrepancy between subnormal cell survival and defective XLF 279 recruitment and filaments formation associated with mutations in X-KBM. The outward rotation in Ku80 upon X-KBM binding more likely relies on E133<sup>80</sup>, the equivalent position 280 281 of which in Ku70 is a methionine (M167) (Supplementary Figure 4b). Therefore, we designed 282 E133M mutant in Ku80 and Q162E mutation that may alter the charge environment of E133 283 (Supplementary Figure 5c). E133M or Q162E mutation in Ku80 negatively impacted X-KBM 284 recruitment but not that of control A-KBM (Supplementary Figures 6a and 6b). This result 285 further supports the independent binding sites on Ku80 of the A-KBM and X-KBM motifs 286 and validates E133 and Q162 positions in Ku80 as key residues for X-KBM binding.

287

288 Then, mutations in the Ku80 binding sites for X- and A-KBM were combined in the 289 I112R/E133M Ku80 double mutant (Supplementary Figure 5d). This combination of 290 mutations clearly impaired the recruitment of both CFP-(X- and A-KBM) peptides (Figures 291 5a and 5b) while the recruitment of the mutant CFP-Ku fusions was preserved 292 (Supplementary Figure 6c-d). We also measured the recruitment of the APLF partner XRCC4 293 and of XLF, expressed as CFP-tagged full-length proteins (Figure 5c-d). Strikingly, only the 294 double E133M/I112R mutation strongly but not completely impaired the recruitment of both 295 proteins (Figure 5c-d). This result supports the independent binding of APLF-XRCC4 and 296 XLF proteins on Ku80.

297

298 Ku80 mutants expressed in U2OS cells had no detectable impact on XLF filaments 299 (Supplementary Figure 6e-f) but lowered end-joining efficiency on a linear plasmid 300 transfected in U2OS cells (Figure 5e). To assess repair accuracy, we used an assay in which 301 GFP expression from a cassette integrated in cells relied on the loss of a DNA fragment 302 between two cut sites (modified from <sup>37</sup>) (Figure 5f), reported to be favoured upon NHEJ inhibition <sup>38,39</sup>. Strikingly, we observed an increase in fragment loss with the three Ku80 303 304 mutants (Figure 5f), indicating that loose APLF-XRCC4 or XLF interactions with Ku80 305 promote genomic instability. Finally, the combination of E113M and I112R Ku80 mutations 306 had a strong impact on cell radiosensitivity with an additive effect of both mutations (Figure 307 5g). These results indicate that APLF-XRCC4 and XLF interactions with Ku80 cooperate to 308 promote cell survival to IR.

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# 311 **DISCUSSION**

312

Here, we show that each of the A- and X-KBM binds Ku independently of other APLF or XLF protein domains. This is in contrast with the absence of binding reported for the isolated X-KBM in recent studies with fluorescent polarization <sup>9,40</sup> that may be due to lower sensitivity, steric hindrance by the fluorescence probe or differences in the Ku constructs used. Therefore, the present results rule out the conclusion that Ku-XLF interaction necessarily needs XLF binding to DNA (even though DNA could stabilise the interaction, as deduced from our switchSENSE data) or to DNA-PKcs <sup>40</sup>.

320

321 Despite the A-KBM and X-KBM motifs have important sequence similarities (Supplementary 322 figure 1), they target different regions of Ku80 with different modes of actions, a more rigid 323 one for the A-KBM and an induced fit one for the X-KBM. In contrast to the L297W X-KBM 324 peptide, the (LW) full-length XLF mutant protein cannot be redirected to the APLF-binding 325 site on Ku80. This may be explained by steric constraints either intrinsic to the protein 326 structure or due to interactions with other partners in the NHEJ complex. It was reported that 327 a X-KBM in WRN protein functions cooperatively with an A-KBM located upstream (1403-1412) in binding Ku complexes <sup>40</sup> (Figure 1a). Our structural data allow proposing the first 328 model of Ku interacting with the tandem sequence of the WRN A-KBM and X-KBM, 329 330 including the central linker (Supplementary Figure 4c). This model now awaits validation by 331 the crystal structure of the tandem motif of WRN on Ku.

332

333 XRCC4 and XLF organize into filaments both *in vitro* and in cells <sup>10-14</sup> and can also assemble 334 as sliding sleeves-like structures on broken DNA *in vitro* even without Ku <sup>41</sup>. In this study, 335 cells containing single or double mutations on the A-KBM and/or X-KBM binding sites of 336 Ku80 show intact filaments while mutations on the X-KBM of XLF induce a slight reduction 337 in the filaments size. These data suggest that interactions additional to Ku80-XLF may 338 participate in filament formation, like the XLF C-terminal DNA-binding domain that spans 339 the X-KBM <sup>42</sup>.

340

How is NHEJ ligation complex assembled at broken DNA? APLF FHA domain interacts with 341 XRCC4<sup>17,26</sup> and XLF establishes head to head contact with XRCC4<sup>10-13</sup>. Here, we show that 342 343 the two remote APLF- and XLF-KBM binding sites in Ku80 promote the independent 344 recruitment of XRCC4 and XLF at broken DNA and that disruption of XRCC4-XLF 345 interaction (through L115D mutation) does not compromise XLF recruitment. Destabilisation 346 of APLF or XLF interactions with Ku80 impairs repair efficiency and cell survival to DSBs 347 and also favours genome instability associated with distal end-joining. These effects are most 348 likely explained by loose assembly of the NHEJ apparatus at break ends. Therefore, we 349 propose a model in which APLF and XLF KBMs represent two initial anchorage points for 350 the rapid and independent recruitment of APLF-XRCC4 and XLF on Ku (Figure 5h). After 351 initial recruitment, interactions additional to Ku80-XLF contacts may stabilize XLF at DBS 352 sites (Figure 5h): our recruitment data show a stable interaction of XLF at DSB sites in cells 353 that differs from the rapid dissociation from Ku-DNA observed *in vitro* by switchSENSE; 354 also, although XRCC4 is dispensable for XLF initial recruitment, it has been shown to

stabilize XLF at damaged sites <sup>34</sup>. Thus, additional contacts stabilizing XLF may include 355 XLF-DNA <sup>34,42,43</sup> and/or XLF-XRCC4 interactions, including filaments formation <sup>10-14</sup> and/or 356 interaction with the LIG4 BRCT1 domain <sup>44</sup>. In addition, after XRCC4 recruitment through 357 358 APLF binding to Ku, XRCC4 may also stabilize in turn APLF at damaged sites since FHA 359 mutants of APLF that do not interact with XRCC4 show a reduced retention after laser micro-360 irradiation <sup>18</sup>. Moreover, links with DNA-PK may further properly stabilize and/or position the NHEJ ligation complex at DSBs (Figure 5h): Ku directly interacts with the XRCC4-LIG4 361 complex <sup>7</sup> through either XRCC4 <sup>45</sup> or LIG4 <sup>6</sup> and DNA-PKcs also directly contacts XRCC4 362 <sup>46-48</sup>. Intimate links between the DNA-PK and ligation complexes are illustrated by the 363 364 requirement of an intact XLF-XRCC4-LIG4 complex to ensure optimal DNA ends synapsis 49,50 365

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367 The intricate network linking end-recognition and ligation NHEJ complexes may allow compensation of partially defective individual components. Indeed, we found that separate or 368 369 even combined Ku80 mutations do not completely abolish XRCC4 and XLF recruitment and 370 do not radiosensitize cells as much as XLF complete defect that abolishes all XLF functions in NHEJ <sup>51</sup>. In that view, only a mild, if any, repair defect in human cells has been associated 371 with X-KBM deletion <sup>27</sup> or APLF depletion <sup>9,24-26</sup>: in case of individual absence of XLF C-372 373 terminal tail or of APLF, the other intact partner would still bind to Ku80 and be able, 374 although with a slower kinetics, to recruit the other components to achieve ligation.

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376 Finally, our present study adds a new aspect to the DNA-PK-ligation complex interaction 377 network that is the swing of the Ku80 vWA domain upon XLF binding (Figure 5h). From our structural and mutagenesis approaches, we propose that the outward swing of the vWA 378 domain of Ku80 is mainly dependent on the conserved acidic residue E133<sup>80</sup>. Although this 379 380 swing does not impact the A-KBM binding site nor the DNA binding domain, it exposes a 381 large groove between the vWA and the ring domain of Ku80 that may non-exclusively 382 reinforce XRCC4-LIG4 and/or XLF interaction with Ku or attract yet unknown components. Interestingly, XLF deficiency impacts on the activity of the Ku partner terminal 383 deoxynucleotidyl transferase during V(D)J recombination <sup>52</sup>. Mutagenesis studies on the 384 385 conserved positions that delineated the unmasked surface of this groove will help to define 386 precisely the role of this swing in the NHEJ reaction.

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In conclusion, the present data further substantiate the emerging model that several NHEJ factors bearing a limited repertoire of KBMs recognize a limited number of KBM-binding sites on Ku<sup>16</sup>. The complex regulation of Ku sites occupancy by NHEJ factors during the repair process deserves further investigations.

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ACCESSION CODES. Crystal structures are deposited at the pdb with the following codes
Ku-DNA-pAPLF (6ERF), Ku-DNA-pXLF (6ERH) and Ku-DNA-pXLFshort (6ERG)

- 395
- 396 ACKNOWLEDGMENTS
- 397

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414

## 415 AUTHORS CONTRIBUTIONS

P.C. and J-B.C conceived this study. C.N, V.R, A.G, A.P, A.C, S.B expressed and purified
recombinant proteins with help from P.D, E.L.C and I.B. C.N and V.R produced crystals and
collected crystallographic data with help from P.L. C.N, V.R, P.L and J-B.C carried out the
crystallographic analysis and interpreted the results. C.N, A.G, S.B, E.L.C and J-B.C
designed, performed, and analyzed microcalorimetry and biophysical experiments. E.B-M,
A.C and J-B.C designed, performed, and analyzed switchSENSE experiments. S.T and M.M

- 422 designed, performed, and analyzed electromobility shift assays. P.F and P.C designed and
- 423 constructed vectors and cell lines for life cell imaging and radiosensitivity. P.F, C.D, N.B and
- 424 P.C designed, performed and analyzed western blot analyses of Ku variants, life cell imaging,
- 425 DNA repair and radiosensitivity experiments. Y.Y and E.R designed, performed and analyzed
- 426 the super resolution microscopy. J.Y and R.G performed bioinformatic analyses. All the
- 427 authors discussed the data. C.N, P.C and J-B.C wrote the manuscript with input from V.R,
- 428 P.F, P.D, R.G, M.M, E.B-M, Y.Y and E.R.
- 429

# 430 COMPETING FINANCIAL INTERESTS

- 431 The authors declare no competing financial interests.
- 432
- 433

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# 569 FIGURE LEGENDS

570

# 571 Figure 1. Crystal structure of the APLF KBM (A-KBM) bound to the Ku80 vWA 572 domain.

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V(D)J recombination. Blood 128, 650-9 (2016).

573 (a) Positions of the A-KBM (magenta) and X-KBM (blue) motifs in APLF, XLF, WRN and 574 CYREN. The C-terminal domain of PAXX contains a P-KBM that interacts with Ku70 575 subunit. NTD: N-terminal domain. (b) Overall view of the quaternary complex 576 Ku70/Ku80/hDNA/(APLF peptide). The A-KBM (magenta) binds at the periphery of the 577 Ku80 (light green) vWA domain. The Ku70 subunit and hDNA are represented respectively 578 in orange and red. The hairpin part of the DNA has been removed for clarity. (c) The N-579 terminal part of the A-KBM motif has an extended conformation whereas the C-terminal 580 residues form a turn. (d-e) Zoom of the interactions made by (d) the hydrophobic patch and 581 (e) the basic patch of the A-KBM. (f) The A-KBM binding site is delineated by conserved 582 residues of Ku80 vWA domain. The binding site is represented in surface mode with amino 583 acids colored according to their conservation rate: red (highly conserved) to white (not 584 conserved)). The conservation rate was measured using sequences of metazoan Ku80. The 585 orientation is the same as in (c).

586

# 587 Figure 2. Crystal structure of the XPLF KBM (X-KBM) bound to the Ku80.

588 (a) Crystal structure of the quaternary complex Ku70-Ku80-DNA-(X-KBM peptide). The X-589 KBM (blue) binds in an internal site of the Ku80 subunit created upon an outward rotation of 590 the vWA domain. The Ku80 vWA opening creates a large groove between the Ku80 vWA 591 and the rest of the heterodimer. (b) The crystal structure of Ku70/Ku80/DNA in presence of 592 the A-KBM is shown with the same orientation. (c-d) Comparison of the X-KBM binding site 593 in presence of X-KBM (c) or A-KBM (d) peptides. The X-KBM interacts with Ku80 residues 594 involved in Ku intramolecular contacts in the closed state of Ku observed with the A-KBM or 595 with no peptide. The last GLFS residues of the X-KBM interact with the bottom of the groove 596 formed in the open state. The glutamic acid presents an atypical hydrophobic environment 597 and could be at the origin of the vWA instability. The X-KBM residues occupy the position of

the helix 236-241 of Ku80 in the closed conformation and some X-KBM side chains (R295<sup>X</sup>), 598 L297<sup>X</sup> and F298<sup>X</sup>) mimic the intramolecular interactions made by Ku80 residues with the 599 600 vWA domain. (e) Gel shift assay with XLF and Ku in presence of a 50bp DNA with a FAM 601 in 5' and competition with pXLF containing the X-KBM motif. The arrow indicates the XLF-602 Ku-DNA complex. Uncropped gel image is shown in Supplementary Data Set 1. (f) The pair distributions P(r) obtained in solution by SAXS analysis indicates an opening of the 603 604 Ku70/Ku80/DNA complex with higher Dmax and Rg in presence of the X-KBM (blue line) 605 compared to the Ku/DNA complex without peptide (grey line) and to the A-KBM complex 606 (magenta line). Values deduced from SAXS analysis are reported beside the curves.

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# Figure 3. Life cell imaging of A-KBM and X-KBM recruitment after nuclear micro irradiation.

611 (a) Wild-type (WT) and mutant CFP-(A-KBM) behaviour at 0 s and 50 s after laser nuclear 612 micro-irradiation. The white rectangle and arrows mark irradiated areas. Magnification: X40. 613 (b) Dynamics of wild-type and mutant CFP-(A-KBM) at laser-induced damage sites in U2OS 614 cells. Mean values of relative fluorescence with s.e.m. were calculated from data obtained in 615 several individual cells: n=23 and 19 cells for WT and mutant A-KBM, respectively). p 616 values at last time point were calculated using unpaired two-tailed t-test: WT vs W189G 617 p < 0.0001. (c) Dynamics of wild-type and mutant CFP-(X-KBM) at laser-damaged sites as in 618 b). n=27, 21, and 24 cells for WT, L297E and L297W X-KBM, respectively. p values at last 619 time point : WT vs L297W p=0.8574; WT vs L297E p=0.0021. (d-e) Dynamics of CFP-(A-620 KBM) (d) and (X-KBM) (e) at laser damaged sites in cells expressing wild-type or I122R mutant Ku80 as in b). n=20, 13 cells for A-KBM in WT or I122R Ku80, and n=48 and 39 621 622 cells for X-KBM in WT or I122R Ku80, respectively. p values at last time point : (d) WT vs 623 I112R p=0.0002; (e) WT vs I112R p=0.5692 (f-g) Dynamics of wild-type and mutant CFP-624 (X-KBM) at laser-damaged sites in cells expressing I112R mutant Ku80 (f) or treated with a 625 shAPLF (g) as in b). n=26, 28, and 21 cells for WT, L297E and L297W X-KBM in (f), and 626 n=15 cells for each of WT, L297E and L297W X-KBM in (g). p values at last time point : (f) 627 WT vs L297W p=0.023; WT vs L297E p=<0.0001; (g) WT vs L297W p=0.0144; WT vs 628 L297E *p*=0.2654. 629

630

#### 631 Figure 4. Biophysical and cellular analyses of XLF mutants in X-KBM.

632 (a) SwitchSENSE kinetic analysis of the WT XLF interaction with Ku-DNA complexes. 633 Solid grey lines represent raw data (from 1 to 8  $\mu$ M; light grey to dark grey; averages of 634 triplicates). Global fitting was performed, following a single-exponential function (solid orange lines) yielding kinetic rate constants;  $k_{ON}=4.7 \pm 1.7 \ 10^5 \ M^{-1}s^{-1}$  and  $k_{OFF}=9.1 \pm 0.4 \ 10^{-2}$ 635  $s^{-1}$  for XLF(wt). (b) Dynamics of wild-type and mutant CFP-XLF at laser-damaged sites in 636 BuS cells as in Figure 3b. n=20 cells for WT, L297E and L297W XLF. p values at last time 637 638 point : WT vs L297W p=0.0093; WT vs L297E p<0.0001. (c) Representative super-resolution 639 images of WT, L297E mutant, and L297W mutant BuS nucleus, with XLF and Ku displayed 640 in green and magenta, respectively (scale = 2500 nm). Right: zoomed-in areas (scale = 250 nm). (d) Representative pair correlation function calculated from the  $8x8 \ \mu m^2$  center square of 641 642 one XLF nucleus image of WT (green), L297E (red), and L297W (blue) mutants. WT XLF 643 shows bigger correlation radius (arrow). (e) Statistics of XLF foci size. Each plot represents 644 the average XLF foci size (indicated as radius translated from the correlation radius) in one 645 nucleus. Box's height displays the s.d. with the mean value labelled in the middle. n=116, 95, 646 104 nuclei for WT, L297E, and L297W. The two-sample unpaired *t*-test between WT and L297E is  $p=10^{-13}$  while that between WT and L297W is p=0.03. (f) Cell survival of BuS cells 647 648 complemented with vector (EV) or WT or mutated XLF. y axis is log scale. Error bars 649 represent s.d., n=5 to 6 independent experiments. p values were calculated using unpaired 650 two-tailed *t*-test: WT vs EV p=1.788e-06; WT vs LW p=0.068; WT vs LE p=0.021. 651 Significant *p*-values are indicated as follows: p < 0.05, p < 0.01, p < 0.01, p < 0.001.

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#### 653 Figure 5. Effects of Ku80 mutations in APLF and XLF binding sites.

654 (a-b) Dynamics of CFP-(A-KBM) (a) and (X-KBM) (b) at laser damaged sites as in Figure 655 3b, in U2OS cells expressing wild-type or I112R/E133M mutant Ku80. n=20 and 9 cells for 656 WT and mutant Ku80 in (a) and n=48 and 11 cells for WT and mutant Ku80 in (b). p values 657 at last time point: (a) WT vs I112R/E133M p=0.001; (b) WT vs I112R/E133M p=0.0111. (c-658 d) Dynamics of CFP-XRCC4 (c) and XLF (d) at laser-damaged sites in cells expressing wild-659 type, I112R, E133M or I112R/E133M mutant Ku80. n=38, 27, 28, and 24 cells for WT, 660 E133M, I112R and I112R/E133M Ku80 conditions in (c) and n=24, 26, 20 and 23 cells for 661 I112R, WT, I112R/E133M and E133M Ku80 conditions in (d). p values at last time point: (c) 662 WT vs E133M p=0.532; WT vs I112R p=0.0133; WT vs I112R/E133M p=0.0048; (d) WT vs 663 1112R p=0.246; WT vs E133M p=0.0048; WT vs I112R/E133M p=0.0248. (e) End-joining 664 activity in U2OS cells expressing mutated or WT Ku80. Error bars represent s.d., n=4

- 665 independent experiments. p values were calculated using unpaired two-tailed t-test: WT vs 666 E133M p=0.0004; WT vs I112R p=0.0052; WT vs I112R/E133M p=0.0002. (f) Distal end-667 joining in U2OS cells containing mutated or WT Ku80. Error bars represent s.d., n=7 668 independent experiments. p values were calculated using unpaired two-tailed t-test: WT vs 669 E133M p=7.49 e-05; WT vs I112R p=2.21 e-06; WT vs I112R/E133M p=4.05 e-06. (g) 670 Survival of U2OS cells expressing WT or mutated Ku80. y axis is log scale. Error bars represent s.d., n=7 to 10 independent experiments. p values were calculated using unpaired 671 672 two-tailed *t*-test: WT vs I112R p=1.47 e-06; WT vs E133M p=6.32 e-05; WT vs 673 I112R/E133M p=2.52 e-13; I112R vs E133M p=0.011. Significant p-values are indicated as 674 follows: \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001. (h) Model for APLF and XLF KBMs function
- 675 during NHEJ.

# 676 Table 1 Data collection and refinement statistics

	Ku-pAPLF (PDB 6ERF)	Ku-pXLF (PDB 6ERH)	Ku-pXLFs (PDB 6ERG)
Data collection		· · ·	
Space group	P <sub>1</sub>	P2 <sub>1</sub>	P2 <sub>1</sub>
Cell dimensions :			
<i>a</i> , <i>b</i> , <i>c</i> (Å)	98.9, 140.8, 150.3	111.8, 118.9, 128.2	111.7, 114.3, 127.2
α, β, γ (°)	68.6, 80.8, 81.2	90.0, 93.1, 90.0	90.0, 93.1, 90.0
Resolution (Å)	49.5-3.0 (3.28-3.0)	49.01-2.8 (3.16-2.8)	50-2.9 (3.09-2.9)
Anisotropy resolution limits $(Å)^{\$}$	2.9, 3.9, 3.3	2.7, 4.4, 3.4	2.8, 3.8, 3.0
Resolution limit overall (Å)§	3.15	3.11	3.0
R <sub>merge</sub>	0.056 (0.79)	0.164 (1.35)	0.171 (2.48)
R <sub>meas</sub>	0.072 (0.93)	0.182 (1.47)	0.182 (2.56)
$R_{\rm pim}$	0.051 (0.66)	0.094 (0.76)	0.068 (0.94)
Ι/σ (Ι)	11.8 (1.3)	9.1 (1.5)	9.7 (1.0)
$CC_{1/2}$	0.99 (0.60)	0.997 (0.60)	0.996 (0.51)
Completeness (spherical, %) <sup>§</sup>	66.5 (14.3)	51.3 (8.5)	72.9 (21.0)
Completeness (ellipsoidal, %) <sup>§</sup>	92.1 (70.1)	92.4 (69.6)	94.6 (68.5)
Redundancy	3.4 (3.5)	7.1 (6.9)	14.0 (14.2)
Refinement			
Resolution (Å)	49.47-3.01	49.38-2.8	48.76-2.9
No. reflections	90993	41644	51644
$R_{ m work}$ / $R_{ m free}$	0.209/0.227	0.225/0.252	0.218/0.244
No. Atoms	35410	18810	18989
Protein	33225	16649	16821
DNA	2178	2118	2133
$SO_4^{2-}$	0	10	15
Water	7	33	20
B factors ( $Å^2$ )			
Protein	112	76	94
DNA	201	109	126
R.m.s. deviations			
Bond lengths (Å)	0.008	0.007	0.008
Bond angles (°)	0.96	0.95	0.95

677 \*Values in parentheses are for highest-resolution shell. <sup>§</sup> Values from STARANISO, Global Phasing Ltd.

679 Table 2 : Interactions measured by microcalorimetry between Ku70/Ku80, A-KBM, X-KBM and XLF

680

678

	Protein in the measurement cell	Ligand in the syringe	K <sub>d</sub> (μM)	ΔH (kcal.M <sup>-1</sup> )	Remarks
1	Ku <sub>FL</sub>	pAPLF	$0.033 \pm 0.01$	$-13.3 \pm 0.1$	A-KBM (APLF 174-191)
2	Ku <sub>FL</sub>	DNA 18bp	$0.0041 \pm 0.0007$	$+5.3\pm0.2$	DNA <sub>18bp</sub> versus Ku
3	Ku <sub>FL</sub> +DNA-18bp	pAPLF	$0.023\pm0.002$	$-16.0 \pm 0.4$	in presence of DNA <sub>18bp</sub>
4	Ku <sub>CC</sub>	pAPLF	$0.020\pm0.002$	$-18.4 \pm 0.7$	Ku without Cter domains
5	Ku <sub>FL</sub> I112R	pAPLF	NI <sup>(a)</sup>	NI	Ku80 mutant on APLF site
6	Ku <sub>FL</sub>	pXLF	$4.4\pm0.2$	$-2.8 \pm 0.2$	X-KBM (XLF 281-299)
7	Ku <sub>FL</sub> +DNA-18bp	pXLF	$2.4 \pm 0.1$	$-8.1 \pm 0.6$	in presence of DNA <sub>18bp</sub>
8	Ku <sub>CC</sub>	pXLF	$2.2\pm0.9$	$-3.1 \pm 1.2$	Ku without Cter domains
9	Ku <sub>FL</sub>	XLF	$1.0 \pm 0.1$	-9.0 <sup>(b)</sup>	XLF versus Ku
10	Ku <sub>FL</sub> +DNA-18bp	XLF	$2.35\pm0.1$	$-8.1 \pm 0.6$	XLF versus Ku/DNA <sub>18bp</sub>
11	Ku <sub>FL</sub>	pXLF(LW)	$0.12\pm0.03$	$-12.1 \pm 2.0$	X-KBM (L297W)
12	Ku <sub>FL</sub>	pXLF(LE)	NI	NI	X-KBM (L297E)
13	Ku <sub>FL</sub> +pAPLF	pXLF(LW)	NI	NI	L297W in presence of A-KBM

681

682 (a) NI means no interaction; (b) The enthalpy value was deduced from the first injection point in absence of lower plateau;

683 The thermograms and isotherms of titration of the experiments corresponding to lines 1, 6 and 11 are presented in Supplementary Figure 2e-g.

Interactions in lines 1, 4, 6, 8, 9, 11 were measured in triplicate, and the mean value with standard deviation is reported. Interactions in lines 2, 3,

685 4, 6, 10 were measured in duplicate, and the mean value with variation between min and max values is reported.

686

#### 687 ONLINE METHODS

688

#### 689 **DNA preparation**

The 500 bp linear dsDNA molecules were amplified from the 3516-4016 region of pBR322
plasmid with biotinylated primers (5'-bGGATCTCAACAGCGGTAA-3' and 5'
bCTTTATCCGCCTCCATCC-3'). DNA fragments were purified on a MiniQ anion exchange
column with a chromatography SMART system (GE Healthcare), ethanol precipitated and

- resuspended in a 10 mM Tris-HCl, pH 7.5, 1 mM EDTA buffer.
- 695

#### 696 Oligonucleotides for ITC and EMSA

- 697 CN1 34bp : CGCGCCCAGCTTTCCCAGCTAATAAACTAAAAAC
- 698 CN2 21bp : GTTTTTAGTTTATTGGGCGCG
- 699 CN3 18bp up : GTTATCCGAGCGTGAGAC
- 700 CN4 18bp down : GTCTCACGCTCGGATAAC
- 701 NLB48 : TAG TCG TAA GCT GAT ATG GCT GAT TAG TCG GAA GCA TCG AAC
- 702 GCT GAT
- 703 MM 50bp up : FAM-TAAATGCCAATGCTGCTGATACGTACTCGGACTGATTCGGAACTGTAACG
- 704 MM 50bp do : CGTTACAGTTCCGAATCAGTCCGAGTACGTATCAGCAGCATTGGCATTTA
- 705

## 706 **Peptides and Proteins:**

The synthetic peptides containing the KBM and XLM motifs were purchased from Genecust at 95% purity, and the concentrations of the stock peptide solutions were determined by amino acid composition. The oligonucleotides used for ITC and crystallization were synthesized by Sigma-Aldrich and Eurogentec.

- 711 The full length Ku70(1-609)/Ku80(1-732) heterodimer and a truncated version of the
- heterodimer deleted Ku70(1-544)/Ku80(1-551) were cloned in the Multibac vectors with a
- 713 10-His tag and a TEV site on the Ku80 N-terminus <sup>53</sup>. Each plasmid was integrated in a
- 714 Yellow Green Protein (YFP) containing bacmid by transformation in EMBACY E.coli stain
- 715 (kind gift from Imre Berger, Bristol University). The resulting recombinant bacmids were
- visual relation relat
- 717 of viruses were titrated by the dilution limit method using YFP as marker for infected cells
- 718 and Mac Grady table. Production was initiated in Sf21 cells culture by infection with
- 719 baculovirus at MOI of  $5 \times 10^{-3}$ . Insect cells were collected 5-6 days after the infection (3-4 days
  - 23

720 after the proliferation arrest). Cells were sonicated and the supernatant was incubated with 721 Benzonase (300 units for 30 min at 4°C). The Ku heterodimer was purified on a NiNTA-722 Agarose affinity column (Protino, Macherey Nagel) with a 1M NaCl wash step to remove 723 DNA excess. The eluted Ku was then bound onto an anion exchange column (Resource Q, 724 GE Healthcare) equilibrate with buffer Q (20 mM Tris pH 8.0, 50 mM NaCl, 50 mM KCl, 10 725 mM  $\beta$ -mercaptoethanol). Final yield of the Ku heterodimer was typically 35 mg of purified 726 heterodimer by liter of culture. The full length 10-His tagged XLF protein was produced in 727 insect cells with a similar protocols with a yield of 50mg of purified protein by L of culture. A 728 truncated version of XLF(1-224) was also produced in E. coli .

729

# 730 Crystallization of the Ku70/Ku80/hDNA complexes with A-KBM and X-KBM peptides

The DNA used in this study is the hairpin DNA previously used by Walker et al <sup>3</sup>. It was 731 732 obtained using HPLC-purified oligonucleotide of 34 and 21 nucleotides (see oligonucleotides 733 above). The oligonucleotides were annealed and added in 1.1-fold molar excess to Ku 734 heterodimer. The peptides containing the A-KBM motif (18mer, 174-192), the long X-KBM 735 motif (19mer, 281-299) or the short X-KBM motif (13mer 287-299) were added respectively 736 with a 1.1, 2 and 2 fold excess. Crystallization screenings on the Ku<sub>CC</sub>-hDNA-peptides were 737 performed on the HTX platform (EMBL, Grenoble) with an automatic visualization at 4°C. 738 The crystals of Ku<sub>cc</sub>-hDNA-pAPLF were reproduced and optimized in the laboratory at 20 °C 739 using the sitting drop method by mixing 1.5µL of the 20 mg/mL Ku-DNA-peptides 740 complexes with 1.5  $\mu$ L of the solution containing 13% polyethylene glycol (PEG) 3350, 150 741 mM NaNO<sub>3</sub>, and 100 mM Bis-Tris-Propan (pH 6.5). The crystals (100x150x1000µm) grew in 742 5-6 hours and were frozen in a solution of the mother liquor with 20% glycerol. The  $Ku_{cc}$ -743 hDNA-pXLF and Kucc-DNA-pXLFs complexes were crystallized at 20 °C by mixing 1.5µL 744 of the 7 mg/mL complex solution with 1.5  $\mu$ L of a solution containing 18% polyethylene 745 glycol (PEG) 3350, 150 mM Na<sub>2</sub>SO<sub>4</sub>, and 100 mM Bis-Tris-Propane (pH 8,5). The crystals 746 (100x200x50µm) grew in 5-6 days and were frozen with 20% glycerol.

747

#### 748 **Determination of the crystal structures**

Diffraction data were collected at the Proxima 1 and Proxima 2 beamlines at the *synchrotron* SOLEIL. The datasets were indexed and integrated using the XDS package <sup>54</sup>, the XDSME package (XDS Made Easier, https://github.com/legrandp/xdsme) and the CCP4 suite <sup>55</sup>. The crystals present a highly anisotropic diffraction (between 2.85Å and 4.25Å resolution according to the axes). The anisotropy of Kucc-hDNA-pAPLF crystals was treated with the

STARANISO program (http://staraniso.globalphasing.org/). The software performs an 754 755 anisotropic cut-off of merged intensity data, a Bayesian estimation of the structure 756 amplitudes, and applies an anisotropic correction to the data. The structure of the Ku70/80-757 hDNA-pAPLF was determined by molecular replacement with the program MOLREP using 758 the structure of Ku70/80 (pdb 1JEY) without the DNA coordinates <sup>3</sup>. Four molecules of Ku 759 were consecutively positioned. Electron density for the hDNA was clearly visible in the position previously reported by Walker. Refinement was performed using BUSTER<sup>56</sup> and 760 PHENIX <sup>57</sup>. The models were built with Coot <sup>58</sup>. After DNA building, an electron density was 761 visible on the Ku80 vWA near the Ku80 amino acids identified by Grundy et al by 762 763 mutagenesis. The final statistics are presented below. In the final model, the following regions 764 of Ku70/Ku80 are not visible: Ku70 1-33, 535-544 and Ku80 1-5, 170-181, 190-191, 543-765 551.. The quality of the model was assessed using Molprobity <sup>59</sup>.

766 The crystal structures of the Ku70/80-hDNA complexed with the long and short pXLF were solved by molecular replacement. Firstly, the coordinates of Ku70/80 and the hairpin DNA 767 768 present in pdb 1JEY were used as model. We positioned two molecules in the asymmetric 769 unit with clear electron density except on the vWA region of Ku80 (region aa 6 to 242). We 770 then performed the molecular replacement with the same coordinates deleted of the Ku80 771 vWA region. The electron density for the secondary structure elements of the Ku80 vWA was 772 clearly visible though at a different position than in 1JEY. A second molecular replacement 773 step was performed to position the vWA domain of the two Ku80 molecules in the 774 asymmetric unit. The structure of the Ku70/80-DNA-short pXLF was solved in a similar 775 manner. An electron density was visible in an internal position of Ku80 vWA that could be 776 attributed to pXLF. The final statistics are presented in Table1. In the final model with the 777 long pXLF, the following regions of Ku70/Ku80 are not visible (Ku70 1-34, 535-554 and 778 Ku80 543-544). In the final model with the short pXLF, the following regions of Ku70/Ku80 779 are not visible (Ku70 1-34; 535-554 and Ku80 171-194, 300-301, 542-544).

An additional electron density was observed in the long and short pXLF complex structures located close to the extreme N-terminus of Ku80 and to the hairpin DNA. This electron density was successfully modeled with the first missing residues of Ku80, the TEV site sequence preceding (ENLYFQG) and seven histidines from the 10-His tag. To evaluate the influence of the tag present on the N-terminus of Ku80 on the Ku-XLF interaction, we prepared a digested form of Ku with the TEV protease (Ku<sup>TEV</sup>) and measured by ITC its interaction with the X-KBM. The ITC shows that thermodynamic parameters of the Ku<sup>TEV</sup> are similar to the Ku and that the tag does not significatively influence the binding of the XLFmotif.

- 789
- 790
- 791

792 **Small-angle Xray scattering:** Several data of Ku<sub>cc</sub>-hDNA complex with a protein-DNA ratio 793 (1:1.2) were collected alone or in presence of pXLF or pAPLF peptides. Data were collected 794 on the SWING beamline (SOLEIL synchrotron) at a 1.8m sample-detector distance. The 795 complexes were prepared at 1.0, 3.0 and 5mg/mL and spun for 10 minutes at 13000 rpm prior 796 to SAXS analysis to eliminate aggregates. Volumes of 40µL of each sample and buffer (20 797 mM Tris pH 8.0, 150mM NaCl, 5mM β-mercaptoethanol) were injected into the SAXS 798 capillary cell and collected continuously, with a frame duration of 0.5 s and a dead time 799 between frames of 0.5 s. Data reduction to absolute units, frame averaging and subtraction were done using FOXTROT <sup>60</sup>, a dedicated home-made application. All subsequent data 800 801 processing and analysis steps were carried out with PRIMUS and other programs of the ATSAS suite <sup>61</sup>. The program GNOM <sup>62</sup> was used to compute the pair-distance distribution 802 803 functions, P(r).

804

# 805 Isothermal titration Calorimetry (ITC)

806 Interactions between Ku70/80 wild-type and Ku70/80cc and the different peptides containing 807 the KBM and XLM motifs were determined by isothermal titration calorimetry (ITC) using a 808 VP-ITC calorimeter (Malvern). Prior to measurements, all solutions were degassed under 809 vacuum. The reaction cell of the ITC (volume 1.8 mL) was loaded with Ku heterodimers 810 alone or complexed with DNA or peptides for competition experiments. Proteins were 811 extensively dialyzed against buffer I (20 mM Tris, pH 8.0, 150 mM NaCl, and 5 mM β-812 mercaptoethanol). Peptides and DNA were prepared at high concentrations. The syringe (290 813  $\mu$ L) was filled with the different peptides at concentration between 20  $\mu$ M to 200  $\mu$ M. The Ku 814 heterodimer present in the cell was titrated by automatic injections of 6-10µL of the different peptides. Enthalpy  $\Delta H$  (in kcal.mol<sup>-1</sup>), stoichiometry of the reaction N, and association 815 constant Ka (in M<sup>-1</sup>) were obtained by nonlinear least-squares fitting of the experimental data 816 817 using the single set of independent binding sites model of the Origin software provided with 818 the instrument. The free energy of binding ( $\Delta G$ ) and the entropy ( $\Delta S$ ) were determined using 819 the classical thermodynamic formula,  $\Delta G = -RT \ln(Ka)$  and  $\Delta G = \Delta H - T\Delta S$ . All binding

820 experiments were performed in duplicate or triplicate at 25°C. Control experiments were

821 performed with peptides injected into the buffer to evaluate the heat of the dilution.

822

# 823 EMSA:

824 Binding reactions (10 µL) were performed by incubating the annealed oligonucleotides

825 (oligonucleotides used in this study are listed in Table Sxx and indicated in the figure 826 legends) at a final concentration of 25 nM, with the indicated final concentrations of proteins 827 in 75 mM KCl, 10 mM Tris (pH 7.5), 0.5 mM EDTA, 0.5 mM DTT, 0.5 mg/mL acetylated-828 BSA, and 5% glycerol. Reactions were incubated at room temperature for 1 hr and 829 fractionated by 6% PAGE (29%/1% [w/v] Acrylamide:Bis-acrylamide) in 0.53 standard Tris-830 borate-EDTA (TBE) buffer at 80 V for 45 min to 1 hr. After electrophoresis, DNA was 831 visualized using a ChemiDoc MP imaging system (Bio-Rad), either by direct detection of the fluorescently labeled DNA (FAM) or after staining with 0.2 mg/mL EtBr. Data were 832 833 processed and quantified with the Image Lab software version 5.2.1 (Bio-Rad).

834

#### 835 switchSENSE measurements :

836 All switchSENSE measurements were carried out on a DRX 2400 instrument, using a multi-837 purpose 48bp chip (both Dynamic Biosensors GmbH; Planegg, Germany). The sample and 838 running buffer was Tris 10mM pH 7.4, 140mM NaCl, 0.05% Tween20, 50µM EDTA, 50µM 839 EGTA. In all kinetics experiments, complementary DNA to the strand immobilized on the 840 chip was first hybridized on the measurement electrodes. On top of the 48bp complementary 841 to the chip, this DNA carried a 32bp-overhang hybridized with its 32bp complementary 842 strand. Second, the Ku70/80 protein was associated at 100nM for 3min (reaching saturation) 843 to the immobilized dsDNA. Finally, the association and dissociation of the XLF protein was 844 measured using triplicates of each concentration, at a flow rate of 2mL/min. For each mutant, 845 the surface was not regenerated in between concentrations - Ku does not dissociate in the 846 timescale of the XLF measurement, and XLF fully dissociates for each concentration. The 847 electrodes were only regenerated between the measurements of different mutants of XLF. 848 Kinetics values were determined using Origin software.

849

#### 850 Cell lines and cell culture

U2OS human osteosarcoma cells and immortalized BuS cells (derived from an XLFdeficient patient, gift from Jean-Pierre de Villartay, Institut Imagine, Paris, France) were grown in DMEM and RPMI, respectively. Media were supplemented with 10% fetal calf

854	serum (Eurobio), 125 U/ml penicillin and 125 $\mu g/ml$ streptomycin. Cells were maintained at
855	37°C in a 5% CO2 humidified incubator. Cell lines were tested negative for mycoplasma by
856	PCR. All culture media and antibiotics were from Invitrogen. When necessary (conditional
857	expression of shRNA against Ku80), doxycyclin (Sigma-Aldrich) was added to the medium at
858	a 4 µg/ml final concentration.
859	
860	Expression vectors
861	See Supplementary Data Set 1
862	
863	Cell transfection and transduction
864	Production of lentiviral particles in HEK-293T cells and transduction of U2OS and BuS
865	cells were performed as previously described <sup>31</sup> . Transduced cells were used as populations
866	without clonal selection, except when indicated otherwise.
867	
868	Plasmid recircularization assay
869	U2OS cell populations expressing wild-type or mutated Ku80 protein were seeded to near
870	confluence in 6-well plates and incubated overnight at 37°C. Cells were then transfected with
871	1 $\mu$ g BamHI-linearized pEGFP-N1 plasmid (Clontech) and 1 $\mu$ g pmCherry-C1-3NLS circular
872	plasmid as a transfection control (gift from Dyche Mullins (Addgene #58476), <sup>63</sup> ). Cells were
873	split 24 h later, incubated at $37^{\circ}C$ for two additional days and analyzed by flow cytometry on
874	a Fortessa X-20 (BD Biosciences). For each cell population, the integrated GFP fluorescence
875	signal was normalized to mCherry signal. End-joining activity was set to 100% for Ku80-WT
876	expressing cells. Results were plotted as the mean values of four independent experiments
877	+s d

878

# 879 Distal End-Joining assay

880 U2OS-EJ5 cells (a kind gift from Jeremy Stark, City of Hope, Duarte, USA) containing 881 one integrated copy of a GFP reporter cassette which allows to measure rejoining of two 882 tandem I-SceI cut sites separated by a ~1.8-kb insert, were modified as follows. The cells 883 were first transduced with lentiviruses produced from pLV-tTR-KRAB and pLV3-Tet-RFP-884 ISceI-GRLBD to allow inducible expression and nuclear translocation of the I-SceI yeast 885 meganuclease. The resulting cells were then transduced with lentiviruses prepared from 886 pLVTHM2-shKu80. A positive clone enabling doxycyclin-dependent conditional knockdown 887 of Ku80 expression and subsequent cell death was isolated. These cells were further

888 transduced with lentiviruses produced from pLV3-HA-Ku80-shR-(WT or mutants) plasmids 889 to replace, in the presence of doxycyclin, endogenous Ku80 expression by expression of the 890 various Ku80 constructs described in the study. To perform Distal End-Joining assay, the 891 different U2OS-EJ5 modified cell populations were seeded onto 6-well plates and incubated 892 at 37°C for 24 h. Dexamethasone (Sigma-Aldrich) was then added to a final concentration of 893 250 nM. Cells were washed 24 h later, further incubated at 37°C for two to three days and 894 analyzed by flow cytometry on a Fortessa X-20 analyzer (BD Biosciences). The fraction of 895 GFP-positive cells was measured and normalized to 100% for Ku80-WT expressing cells. 896 Results were plotted as the mean values of seven independent experiments  $\pm$ s.d.

897

#### 898 Ionizing irradiation and cell survival analysis

899 Three to six thousand cells per well were seeded in 6-well plates. Plated cells were 900 exposed 24 h later to various doses of X-ray using a Faxitron RX-650 device (130 kV, 5 mA, dose rate 0.5 Gy.min<sup>-1</sup>). Six to seven days later, cells were washed with PBS, stained 10 min 901 902 with crystal violet (0.1% aqueous solution). Stained cells were extensively washed with water 903 and plates were air dried. Staining was dissolved with 10% acetic acid solution and absorption 904 was measured at 570 nm (Ultrospec-3000 spectrophotometer, Pharmacia Biotech). Results 905 were plotted as mean values of 5-10 independent experiments  $\pm$ s.d. using Microsoft Excel 906 software.

907

## 908 Live-cell microscopy and micro-irradiation

909 U2OS or BuS cells were seeded in 35-mm glass-bottom culture dishes (MatTek) two 910 days prior laser irradiation. Experiments were carried out with a Zeiss LSM-710 confocal 911 laser scanning microscope equipped with a coherent chameleon Vision-II tunable laser (690-912 1080 nm), a 40X/1.3 oil immersion objective and a heated environmental chamber set at  $37^{\circ}C$ 913 in 5% CO2 atmosphere. ECFP was excited using biphotonic laser at 800 nm (1.5% of 914 maximum power). Confocal image series were recorded with a frame size of  $512 \times 512$  pixels. 915 Nuclei micro-irradiation was carried out at 800 nm at 20% of maximum power (mean max power was 3070 mW) in rectangle of 15  $\mu$ m<sup>2</sup> area during 50  $\mu$ s pixel dwell time. Before and 916 917 after micro-irradiation, confocal image series of one mid z-section were recorded at 1.94 s 918 time interval (typically 9 pre-irradiation and 40-45 post-irradiation frames). For evaluation of 919 the recruitment kinetics, fluorescence intensities of the irradiated region were corrected for 920 total nuclear loss of fluorescence over the time course and normalized to the pre-irradiation 921 value. Data from micro-irradiation of individual cells obtained in several independent 922 experiments performed on different days were averaged, analyzed and displayed using

923 PRISM software. *p* values at last time point were calculated using unpaired Student's t-test.

924

# 925 Protein extraction and immunoblotting

926 Sub-confluent cells from 60-mm culture dishes were harvested and washed with PBS. 927 Pellets were resuspended in 100 µl of lysis buffer (50 mM HEPES.KOH pH 7.5, 450 mM 928 NaCl, 1% Tritin-X100, 1 mM EDTA, 1 mM DTT, protease-phosphatase Halt Inhibitor 929 cocktail (Pierce, Thermo Scientific)) and processed by four freeze/thaw cycles. After 930 centrifugation at 14000 g for 10 min, protein concentration was measured in the supernatant 931 with the Bradford assay (Bio-Rad). Proteins from 50 µg of cell extracts were separated in 4-932 15% Mini-Protean TGX precast polyacrylamide gels (Bio-Rad) and transferred to PVDF 933 membrane (Millipore). Immunoblotting analysis was performed with the following 934 antibodies: anti-Ku70 monoclonal antibody (clone N3H10 from NeoMarkers), anti-Ku80 935 monoclonal antibody (clone 111 from Thermo Fisher Scientific), anti-DNA-PKcs monoclonal 936 antibody (clone 18.2 from Abcam), anti-XLF polyclonal antibody (Bethyl Laboratories), anti-APLF polyclonal antibody (SK3595, kind gift from K.W. Caldecott, University of Sussex, 937 Brighton. UK, see <sup>25</sup>), anti-β-Actin monoclonal antibody (clone AC-15 from Ambion). 938

939

#### 940 Cell culture and preparation for super-resolution imaging

BuS cells were grown in RPMI medium with 10% FBS and 100 U/mL PenicillinStreptomysin. For super-resolution imaging, cells were cultured on glass coverslips for 24
hours, followed by serum starvation for 48 hours. The cells were then released into full
medium for 4 hours so that most of the cells were in G1 phase.

The synchronized cells were then washed twice with PBS, and permeabilized with 0.5% Triton X-100 in CSK buffer (10 mM Hepes, 200 mM Sucrose, 100 mM NaCl, and 3 mM MgCl<sub>2</sub>, pH=7.4) for 10 minutes <sup>64</sup>. Cells were then fixed with paraformaldehyde (4%) for 20 mins, and blocked in block solution (2% glycine, 2% BSA, 0.2% gelatin, and 50 mM NH<sub>4</sub>Cl in PBS) overnight at 4°C.

Fixed cells were then immunostained with validated monoclonal antibodies: XLF was stained
with XLF-antibody (3D6, NBP2-03275, NOVUS) for 1 hour at room temperature, followed
by Alexa Fluor 647 conjugated goat-anti-mouse 2nd antibody staining (ab 150115, abcam) for
30 minutes at room temperature. Cells were then stained with Alexa Fluor conjugated antiKu80 antibody (EPR3467, ab202659, abcam) for 1 hour at room temperature.

955 Cells were then mounted onto microscope glass slide, and imaged with freshly mixed imaging
956 buffer (1 mg/mL glucose oxidase, 0.02 mg/mL catalase, 10% glucose, and 100 mM
957 cycteanube (MEA)).

958

## 959 Microscope and Single-Molecule Localization imaging

960 Super-resolution imaging was performed on a custom-built optical imaging platform based on 961 a Leica DMI 300 inverse microscope, equipped with a 488 nm (OBIS, Coherent) and a 639 962 nm laser line (MRL-FN-639-800, CNI). Laser lines were reflected into an HCX PL APO 63X 963 NA=1.47 OIL CORR TIRF Objective (Zeiss) by a penta-edged dichroic beam splitter 964 (FF408/504/581/667/762-Di01-22x29), and the emitted fluorescence was further extended by 965 a 2X lens tube (Diagnostic Instruments), filtered by single-band filters (Semrock FF01-531/40 966 and FF01-676/37 for Alexa Fluor 488 and Alexa Fluor 647, respectively), and collected onto 967 a sCMOS camera (Prim95B, Photometrics). A 405 nm Laser line (MDL-III-405-150, CNI) 968 was also equipped to reactivate Alexa Fluor 647 fluorophores.

For super-resolution imaging, the 488 and 639 laser lines were adjusted to  $\sim 1.0$  and 1.5 kW/cm2, and a Highly Inclined and Laminated Optical sheet (HILO) illumination mode for sample excitation. Alexa Fluor 488 and Alexa Fluor 647 were sequentially exited and their emitted fluorescence was also sequentially collected by switching the single-band filters in a filter wheel. The emitted photons were collected onto a sCMOS camera (Photometrics Prime 95B), and a minimum of 2000 frames at 33 Hz were recorded for each image stack.

975

#### 976 **Reconstruction from Single-Molecule Localization image to super-resolution image**

Each image of the image stack was first of all box-filtered with the box size of 4 times of the FWHM of a 2D gaussian point spread function (PSF). Considering the patterned noise for sCMOS camera construction, each pixel was weighted by the inverse of its noise variance during filtering. The low-pass filtered image was then extracted as the background from the raw image. The local maximums of the image were then recognized and segmented into single PSFs for 2D gaussian fit.

2D gaussian fit of each individual PSF was performed by GPU using the Maximum 284 Likelihood Estimation (MLE) algorithm. Briefly, the likelihood function of each pixel was 285 constructed by convolving the Poisson distribution of the shot noise governed by the photons 286 emitted from fluorophores nearby, and the gaussian distribution of the readout noise of the 287 camera itself <sup>65</sup>. The offsets, variance, and analog-to-digital conversion factor of each pixel of 288 the camera was calibrated beforehand. The position, amplitude, sigma, and background of each individual 2D gaussian PSF were addressed by maximizing the likelihood function of
each pixel given its readout digital value. The fitting accuracy was estimated by Cramér-Rao
lower bound (CRLB) and the accuracy of Alexa Fluor 488 and Alexa Fluor 647 in this work
are ~ 17 and 13 nm, respectively.

993

## 994 Alignment of images of different colors

995 Mapping of the two colors was performed using a polynomial mapping algorithm. Briefly, a spatially separated and broad-spectrum fluorescent bead (TetraSpec, ThermoFisher) slide was 996 997 imaged on both color channels. Mass centers of each same bead but in different channels were 998 recorded pairwise. The coefficient of each term in a 2nd order polynomial function was 999 optimized by training the polynomial regression using all the recorded bead's centers. The 1000 optimized polynomial function was then used for two color channels alignment. We note that 1001 choosing the proper order of polynomial function for optimization depends on the number of 1002 training beads, and higher order but not enough trainees would lead to overfit.

1003

# 1004 **Pair-Correlation analysis**

1005 Coordinates localized within  $\sim 2.5$  times of the averaged localization uncertainty, and from 1006 consecutive frames were considered as artificial blinking and grouped as one coordinate. This 1007 coordinate was calculated by taking the 1/var weighted average of all the coordinates within 1008 the group, where var is the localization uncertainty of each coordinate. The grouped 1009 coordinates were then rendered onto a pixelized image of 5 nm/pixel, and the pair-correlation 1010 was performed on this image by series 2D Fourier and inverse Fourier transfers <sup>66</sup>.

1011 The correlation was then fitted into two correlation terms: the correlation among coordinates 1012 within localization uncertainty, and that among the coordinates that form a cluster/molecular 1013 assemble/filament. Considering the size of XLF 'clusters' is not randomly distributed, we 1014 fitted the second term into a normal distribution and interpret the fitted sigma as the apparent 1015 radius of the averaged XLF 'cluster' radius (Figure XC) across the image.

1016

#### 1017 Data availability

- 1018 Crystal structures are deposited at the pdb with the following codes Ku-DNA-pAPLF (6ERF),
- 1019 Ku-DNA-pXLF (6ERH) and Ku-DNA-pXLFshort (6ERG)
- 1020 Source data for figures 3, 4 and 5 are available with the paper online
- 1021 Other data that support the findings of this study are available from the corresponding author
- 1022 upon reasonable request.

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M<sup>115</sup>

Y74

68

D116

1149

150

b

A-KBM

Ku80

W<sup>189A</sup>



Ku70













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X-ray (Gy)



# Sup Figure 1



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(a) Scheme of the interactions between Ku70-Ku80 and the NHEJ factors containing an A-KBM (APLF, CYREN), an X-KBM (XLF) and both KBMs (WRN). The interaction of PAXX with Ku70 through its C-terminus is also represented. (b-d) Logo motif of the A-KBM, X-KBM and PAXX motifs obtained from multiple sequences alignment of these proteins as indicated (Crooks, G.E. *et al.*, WebLogo: a sequence logo generator. Genome Res 14, 1188-90 (2004)).



(a) SDS gel showing purified Ku and XLF proteins as indicated. FL: full-length; cc: C-terminal truncation. (b) Electron density of peptide pAPLF. (c) Electron density of peptide pXLF. (c) DNA interactions with Ku in presence of X-KBM of XLF. Ku70-Ku80-hDNA-X-KBM (colored) compared to Ku70/Ku80/hDNA (PDB 1JEY, grey). Front view of Ku70-Ku80-hDNA-X-KBM showing the major deviation of hDNA molecules because of the conformational change of Ku80. (d-e) ITC analyses: representative thermograms and isotherms of titration corresponding to selected measurements from Table 2, as indicated.



(a-e) EMSA analyses: (a, b) Gel shift assays with XLF and Ku at 20nM (a) or 200nM concentrations (b) in presence of a 50bp DNA with a FAM in 5'. (c) pXLF interaction with DNA as a control of the competition experiment in Figure 2e. (d-e) The pAPLF and PAXX Cter do not compete with the Ku-XLF complex. (f-h) switchSENSE analyses: (f) Scheme of the switchSENSE measurement flow: 1) Ku is bound to an 80bp nanolever with a fluorescent probe at position 48; 2) A washing step removes non-specifically bound Ku molecules; 3) XLF is then injected for real-time associations and dissociations at different concentrations followed by normalized changes in the fluorescence. (g) Binding kinetics of the Ku protein on the 80mer double-stranded DNA prior to the interaction with XLF, shown as changes in the dynamic response upwards (between 0 and 4µs). The dynamic response reflects the speed of the switching DNA, which decreases upon binding of the Ku analyte. The dissociation is represented only for one minute, to show that no dissociation of Ku from the DNA occurs while the XLF kinetics is measured. (h) Kinetic analyses of (LW) and (LE)XLF interactions. Solid grey lines represent raw data (from 1 to 8 µM; light grey to dark grey; averages of triplicates). Global fitting was performed, following a single-exponential function (solid orange lines) yielding kinetic rate constants;  $k_{ON} = 4.9 \pm 0.5 10^{-4} \text{ s}^{-1}$  and  $k_{OFF} = 4.8 \pm 0.5 10^{-2} \text{ s}^{-1}$  for XLF(LE) and  $k_{ON} = 1.9 \pm 1.1 10^5 \text{ M}^{-1} \text{s}^{-1}$  and  $k_{OFF} = 8.4 \pm 0.6 10^{-2} \text{ s}^{-1}$  for XLF(LW).



(a) Variations among the sequence motifs observed for the C-terminal tail of XLF and for the seven positions of the Ku80 XBM pocket in various clades of the eukaryotic phylogenetic tree. 10 clades are represented summarizing the properties of 60 Mammalia, 36 Sauria, 31 Neopterygii, 62 Ecdysozoa, 9 Lophotrochozoa, 5 Cnidaria, 150 Fungi, 5 Ciliophora and 4 Dictyostellida sequences of XLF and Ku80. For each clade, web logos of the last 25 C-terminal amino-acids of XLF sequences are represented on top and the web logo of the X-KBM site positions is squared in green. A red star indicates the clades in which the position of Ku80 E133 was conserved as an acidic residue whereas a grey star points out that the acidic character of the residue was not maintained and was generally switched to a hydrophobic residue as observed in Ku70. (b) Superimposition of human Ku70 and Ku80 structures (PDB: 1JEQ) colored in orange and green, respectively, and focused on the region surrounding Ku80 E133 position in the X-KBM site. The red star points out the location of Ku80E133. Residues labelled and shown as sticks are the spatial neighbours of Ku80 E133. The side chain of Ku80 E133 is buried in the hydrophobic core of Ku80 and is not involved in any hydrogen bond or salt-bridge interaction resulting in a predicted pKa above 9.1 in the absence of XLF. (c) Molecular modelling of the interaction between Ku80 and the C-terminus of WRN containing an A-KBM in tandem with a X-KBM. The position of WRN motifs were deduced from the crystal structures presented here with APLF and XLF KBMs. The orientation of the KBMs and the size of the linker between WRN KBMs are compatible with a simultaneous binding of both WRN motifs to Ku80.



(a-d) Western blot of U2OS cell extracts. (a) Whole cell extracts of U2OS shKu80 (U2OS-Ku80<sup>KD</sup>) cells treated with doxycyclin for the indicated time were denatured and separated on 10% SDS-PAGE gel followed by electrotransfer on membrane. The membranes were blotted with the antibodies as indicated. (b) Whole cell extracts of U2OS shAPLF cells expressing WT or mutant CFP-X-KBM as indicated were processed as in (a). (c) Whole cell extracts of U2OS-Ku80<sup>KD</sup> cells treated with doxycyclin for 7 days and expressing WT or mutant HA-Ku80 as indicated were processed as in (a). (d) Whole cell extracts of U2OS-Ku80<sup>kD</sup> cells treated with doxycyclin for 7 days and expressing WT or mutant HA-Ku80 as indicated were processed as in (a). Uncropped blot images are shown in Supplementary Data Set 1. (e) Wild-type (WT) or mutant CFP-(A-KBM) and mCherry-Ku70 simultaneous behaviour after 800 nm pulsed-laser nuclear micro-irradiation assessed in U2OS cells by live cell-imaging at 0 s and 50 s post-irradiation. The white rectangle and arrows mark irradiated areas. (f-h) Dynamics of wild-type (WT) and mutant CFP-tagged full-length XLF at laser-damaged sites in BuS cells. Images were obtained at 1.94 s intervals and fluorescence intensities at the damage sites and in undamaged area were quantified. Mean values of the relative fluorescence with SEM were calculated from 20 independent measurements for each of WT and G296W XLF in (f), from 45, 40 and 20 independent measurements for each of WT, S299A and S299E XLF in (g) and from 45, 36, 20 and 20 independent measurements for each of WT, F298G, F298G/S299A, and F298G/S299E XLF in (h), respectively. p values at last time point: (f) WT vs G296W p<0.0001; (g) WT vs S>A p=0.2785; WT vs S>E p<0.0001. (h) WT vs F>G p<0.0001; WT vs FS>GA p<0.0001; WT vs FS>GE p<0.0001. (i) Dynamics of L115D and L115D/L233E CFP-tagged full-length mutant XLF at laser-damaged sites in BuS cells as in (f). Mean values of the relative fluorescence with SEM were calculated from 11 independent measurements for each of L115D±shAPLF and L115D/L233E XLF conditions. p values at last time point: L115D vs L115D±shAPLF p=0.6113: L115D vs L115D/L233E p=0.0002.



(a-b) Dynamics of CFP-(A-KBM) (a) and (X-KBM) (b) at laser damaged sites in U2OS cells expressing wild-type (WT), E133M or Q162E mutant Ku80 as in Figure 3 b). Mean values of the relative fluorescence with SEM were calculated from 20, 23 and 22 independent measurements for A-KBM with WT, E133M or Q162E mutant Ku80 in (a) and from 48, 29 and 29 independent measurements for X-KBM with WT, E133M or Q162E mutant Ku80 in (b), respectively. *p* values at last time point: (a) WT vs E133M p=0.831; WT vs Q162E p=0.59519; (b) WT vs E133M p=0.0003; WT vs Q162E p=0.0111. (c-d) Dynamics of wild-type (WT) and

mutant CFP-Ku80 at laser damaged sites in U2OS cells. Mean values of the relative fluorescence with SEM were calculated from 25, 24, 20 and 15 independent measurements for WT, I112R, E133M and Q162E mutant Ku80 in (c) and from 25 and 26 independent measurements for WT or I112R/E133M mutant Ku80 in (d), respectively. *p* values at last time point: (c) WT vs Q162E =0.9252; WT vs I112R *p*=0.2734; WT vs E133M *p*=0.1101. (d) WT vs I112R-E1333M *p*=0.5362. (**e**-**f**) Analysis of XLF foci in U2OS cells by super-resolution. (e) Statistics of XLF foci size: each plot represents the average XLF foci size (indicated as radius translated from the correlation radius) in one nucleus. Box's height displays the standard deviation with the mean value labelled in the middle. 87, 110, 64, and 79 nuclei were taken in account for WT, E133M, I112R, and E133M-I112R double-mutant, respectively. The *p*-values were obtained by the t-test; (f) Statistics of the Cross-Pair-Correlation between Ku and XLF: Ku and XLF were stained with antibodies labelled by different fluorophores (Alexa488 conjugated rabbit anti-Ku80, abcam198586, Alexa647 conjugated goat anti-mouse secondary + Mouse anti-XLF, NBP2-03275), and dual-colour super-resolution imaging was performed to examine the cross-correlation between Ku and XLF foci within each nucleus. Each plot represents the cross-correlation amplitude calculated across one nucleus. Box's height displays the standard deviation with the mean value labelled in the middle. 83, 107, 57, and 72 nuclei were taken in account for WT, E133M, I112R, and E133M-I112R mutants respectively. The *p*-value were obtained by the t-test.

1

# **Expression vectors**

All lentiviral vectors derived from pLVTHM (Addgene plasmid #12247) and pLV-tTR-KRAB-Red (Addgene plasmid #12250) plasmids. Both were gifts from Didier Trono<sup>1</sup>.

The pLVTHM2 vector was obtained by digesting pLVTHM with PmeI/SpeI and inserting the PBXS linker (i.e. preannealed PBXS-F/PBXS-R pair of oligonucleotides) in order to remove the GFP coding sequence. To generate lentiviral vectors for conditional expression of shRNA, pLVTHM2 was digested with MluI/ClaI and the following pairs of preannealed oligonucleotides were inserted by ligation: shKu80-F/shKu80-R (target sequence designed by Denis Biard, CEA-DSV, France, personal communication) or shAPLF-F/shAPLF-R (target sequence from <sup>2</sup>) to knockdown the expression of Ku80 or APLF, respectively.

The pLV-tTR-KRAB vector was obtained from pLV-tTR-KRAB-Red by replacing the DsRed coding sequence by the XBES linker at XmaI/SpeI restriction sites.

The pLV-Red vector derived from pLV-tTR-KRAB-Red by replacing the tTR-KRAB coding sequence by the PEKBBMX linker at PmeI/XmaI restriction sites.

The pLV3 vector derived from pLV-tTR-KRAB-Red through the following modifications: first, the NsiI/Kpn2I fragment was replaced by the nPNk linker. The tTR-KRAB coding sequence was then removed by PmeI/XmaI digestion and replaced by the PEKBBMX linker. Finally, the IRES-DsRed fragment was removed by PmeI/SpeI digestion and replaced by the PKXMBBES linker.

To express ECFP-tagged A-KBM (APLF P177 to E193) or X-KBM (XLF S287 to S299) motifs, the A-KBM linker or X-KBM linker, respectively, was inserted into the Acc65I/BamHI restriction sites of the pECFP-C1 plasmid (Clontech) and the AgeI/BamHI fragment (ECFP-A-KBM or ECFP-X-KBM) was subcloned into the Kpn2I/BamHI restriction sites of pLV3. The ECFP-tagged A-KBM-W189G, X-KBM-L297W and X-KBM-L297E expressing vectors were obtained by amplifying by PCR the corresponding cDNAs using the pECFP-A-KBM or pECFP-X-KBM plasmid as a template, as well as CMV-F as forward primer and Bam-A-KBM-W189G-R, XLF-L297W-Bam-R or XLF-L297E-Bam-R as reverse primer, respectively. The PCR fragments were digested with AgeI/BamHI and inserted into Kpn2I/BamHI restriction sites of pLV3.

To express full-length ECFP-tagged XLF protein, human XLF cDNA (a gift from Jean-Pierre de Villartay, Institut Imagine, Paris, France) was amplified by PCR using the XLF-Hind-F and XLF-Bam-R primers. The resulting PCR fragment was digested with HindIII/BamHI and inserted into pECFP-C1. The ECFP-XLF coding fragment was then excised with AgeI/BamHI and inserted into pLV3 at Kpn2I/BamHI restriction sites. ECFP-XLF-L297W, -L297E, -F298G, -S299A, -S299E, -F298G-S299A, and -F298G-S299E mutants were expressed by PCR amplification of the corresponding XLF coding sequences using the pECFP-XLF vector as a template, Kpn2-MCS-F as forward primer, and XLF-L297W-Bam-R, XLF-L297E-Bam-R, XLF-F298G-Bam-R, XLF-S299A-Bam-R, XLF-S299E-Bam-R, XLF-FS298GA-Bam-R, or XLF-FS298GE-Bam-R, as reverse primer, respectively. The resulting fragments were digested with Kpn2I/BamHI and inserted into pLV3-ECFP-XLF to replace the XLF wild-type coding sequence. The XLF-L115D coding sequence was obtained by overlapping PCR mutagenesis on pLV3-ECFP-XLF template using ECFP-Cter-F and XLF-Bam-R oligos as outer primers, and XLF-L115D-F and XLF-L115D-R as mutated inner primers. The PCR product (XLF-L115D) was then digested with Kpn2I/BamHI and inserted into pLV3-ECFP-XLF to replace the XLF-WT coding sequence. The XLF-L115D-L297E double mutant coding sequence was obtained as above, except that the XLF-Bam-R outer primer was replaced by XLF-L297E-Bam-R.

Lentiviral vectors expressing untagged full-length XLF proteins (WT, L297W or L297E) were obtained by subcloning Kpn2I/BamHI fragments from the respective pLV3-ECFP-XLF into the pLV-Red vector.

The ECFP-XRCC4 expressing vector was obtained by excision of a Kpn2I/BamHI fragment containing the XRCC4 coding sequence from the pEGFP-C1-FLAG-XRCC4 plasmid (gift from Steve Jackson (Addgene #46959), <sup>3</sup>). The resulting fragment was inserted into pLV3-ECFP-XLF to replace the XLF coding sequence.

To obtain shRNA-resistant human Ku80 expression vector, HA-Ku80 coding sequence was amplified by overlapping PCR from pICE-Puro-HA-Ku80 (a kind gift from Sébastien Britton, IPBS, Toulouse, France) with Kpn2-HA-F and pICE-Xba-R as outer primers, and Ku80-shRes-F and Ku80-shRes-R as inner primers that introduce silent mutations in the shRNA target sequence. The resulting fragment was then inserted into pLV3 after digestion with Kpn2I/MluI. The Ku80-L112R expressing pLV3 vector was constructed by overlapping PCR from pLV3-HA-Ku80-shR using pLV-F and pLV-R oligonucleotides as outer primers and Ku80-L112R-F and Ku80-L112R-R as mutated inner primers. The PCR product was then digested with Kpn2I/MluI and inserted into pLV3. Other Ku80 single mutant constructs

(E133M and Q162E) were obtained similarly by using the corresponding pairs of inner primers (Ku80-E133M-F/Ku80-E133M-R and Ku80-Q162E-F/Ku80-Q162E-R, respectively). The pLV3-HA-Ku80-shR-L112R-E133M double mutant expressing vector was obtained as above for the E133M single mutant construct, except that pLV3-HA-Ku80-shR-I112R was used as a template for PCR reactions.

Expression vectors for ECFP-tagged WT or mutants Ku80 were obtained by amplifying the ECFP coding sequence by PCR from the pECFP-C1 plasmid with the Pme-Koz-ECFP-F and pme-CFP-80-R primers. The PCR fragment was then inserted at the PmeI restriction site in the various pLV3-HA-Ku80-shR plasmids by use of the Hot-Fusion strategy <sup>4</sup>.

To generate pLV3-mCherry-FLAG-Ku70 vector allowing expression of an mCherrytagged Ku70 protein, the FLAG-Ku70 coding sequence was amplified by PCR using Kpn2-FLAG-F and Mlu-Ku70-R primers, digested with Kpn2I/MluI and cloned into pLV3. The mCherry coding sequence was then inserted at the PmeI restriction site by Hot-Fusion cloning <sup>5</sup> following PCR amplification using Pme-Koz-ECFP-F and Kpn2-pme-mCh-R primers and the pmCherry-C1-3NLS plasmid as a template (gift from Dyche Mullins (Addgene #58476), <sup>6</sup>).

The pLV3-Tet-RFP-ISceI-GRLBD lentiviral vector for conditional expression of I-SceI was prepared as follows : pLV3 was first modified by inserting at the PacI restriction site the Tet-Pac-F/Tet-Pac-R pre-annealed linker which contains two tetracyclin operator DNA elements. The resulting pLV3-Tet plasmid was then digested by Kpn2I/BamHI to receive the AgeI/BamHI fragment from the pISceI-GR-RFP plasmid (gift from Tom Misteli (Addgene #17654), <sup>7</sup>) which contains the coding sequence of DsRed-ISceI-GRLBD.

All oligonucleotides were purchased from Eurofins Genomics (Ebersberg, Germany). Restriction and modifying enzymes (Phusion and T4 DNA Ligase) were from ThermoFisher Scientific (Illkirch, France). All constructs were checked by sequencing (Eurofins Genomics).

	GTACC CCA ATC CTT GCC GAG AGG AAA AGA ATC CTT CCA ACT TGG ATG
A-KBM-F	TTA GCA GAA TAG
	GATCCTA TTC TGC TAA CAT CCA AGT TGG AAG GAT TCT TTT CCT CTC
A-KBM-R	GGC AAG GAT TGG G
nPNk-F	CTCCATCGATCGCCATGGTGA
nPNk-R	CCGGTCACCATGGCGATCGATGGAGTGCA

Oligonucleotides used as linkers (alphabetical order; sequences 5' to 3')

PBXS-F	AAAC CGTACG GATATC T CCCGGG TC A
PBXS-R	CTAGT GA CCCGGG A GATATC CGTACG GTTT
	AAACTACGGGATC GAATTC CTCGCT TCCGGA CTTCGT GGATCC ACTCTC
PEKBBMX-F	CGTACG ACTGCT ACGCGT ACTTCAC
	CCGGGTGAAGT ACGCGT AGCAGT CGTACG GAGAGT GGATCC ACGAAG
PEKBBMX-R	TCCGGA AGCGAG GAATTC GATCCCGTAGTTT
	AAACTACGG GATC TCCGGA CACCTT CCCGGG TCACTC ACGCGT CTCATT
PKXMBBES-F	GGATCC CGTACG GAATTC A
	CTAGT GAATTC CGTACG GGATCC AATGAG ACGCGT GAGTGA CCCGGG
PKXMBBES-R	AAGGTG TCCGGA GATC CCGTAGTTT
	CGCGTCCCC GAA GAA ATC TGC AAA GAT A TTCAAGAGA T ATC TTT GCA
shAPLF-F	GAT TTC TTC TTTTTGGAAAT
	CGATTTCCAAAAA GAA GAA ATC TGC AAA GAT A TCTCTTGAA T ATC TTT
shAPLF-R	GCA GAT TTC TTC GGGGA
	CGCGTCCCC G AAC AAG GAT GAG ATT GCT TTCAAGAGA AGC AAT CTC
shKu80-F	ATC CTT GTT C TTTTTGGAAAT
	CGATTTCCAAAAA G AAC AAG GAT GAG ATT GCT TCTCTTGAA AGC AAT
shKu80-R	CTC ATC CTT GTT C GGGGA
TetO-Pac-F	TCCCTATCAGTGATAGAGATCTCCCTATCAGTGATAGAGAAT
TetO-Pac-R	TCTCTATCACTGATAGGGAGATCTCTATCACTGATAGGGAAT
XBES-F	CCGGG GGATCC CTCGAG GAATTC A
XBES-R	CTAGT GAATTC CTCGAG GGATCC C
X-KBM-F	GTACC TCA AAG GTC AAG AGG AAG AAG CCA AGG GGT CTC TTC AGT TAG
X-KBM-R	GATCCTA ACT GAA GAG ACC CCT TGG CTT CTT CCT CTT GAC CTT TGA G

# **Oligonucleotides used as PCR primers** (alphabetical order; sequences 5' to 3')

Bam-A-KBM-	CGTACGGGATC CTA TTC TGC TAA CAT CCC AGT TGG AAG GAT TCT TTT
W189G-R	CCT C
CMV-F	GTAGGCGTGTACGGTGGGAGG
ECFP-Cter-F	C ATG GTC CTG CTG GAG TTC GTG
Kpn2-HA-F	CTCTGC TCCGGA GCCACC ATG TAC CCC TAC GAT GTG C
Kpn2-FLAG-F	CTCTCGTCCGGAGCCGCACC ATG GAC TAC AAG GAT G
Kpn2-MCS-AKF-F	CTCGCTTCCGGACTCAGATCTCGAGCTC
Kpn2-pme-mCh-R	GGTGCGGCTCCGGAGATCCCGTAGTTTGGACTTGTACAGCTCGTCCATGCCG
Ku80-E133M-F	GAG GCA TAT TAT GAT ATT CAC TGA CCT CAG CAG CCG ATT C
Ku80-E133M-R	GGT CAG TGA ATA TCA TAA TAT GCC TCT TCT CAA ACT TCT TTC CTA TTG
Ku80-I112R-F	C TTC CTG GAT GCA CTA AGA GTG AGC ATG GAT GTG ATT CAA C
Ku80-I112R-R	G AAT CAC ATC CAT GCT CAC TCT TAG TGC ATC CAG GAA GTC
Ku80-Q162E-F	CAT CTC CCT GGA ATT CTT CTT GCC TTT CTC ACT TGG C

Ku80-Q162E-R	GGC AAG AAG AAT TCC AGG GAG ATG TCA CAT TTC TTC AAG C
	GCT GAA AAT AAA GAC GAA ATC GCC TTA GTC CTG TTT GGT ACA GAT
Ku80-shRes-F	GGC
Ku80-shRes-R	GAC TAA GGC GAT TTC GTC TTT ATT TTC AGC AAA CAC CTG TCG CTG TAC
Mlu-Ku70-R	CTCTGCACGCG TCA GTC CTG GAA GTG CTT GGT GAG GGC
pICE-Xba-R	CAGCGGGTTTA TCTAGA CTGCAG ACGCGT GC
pLV-F	CCGATCACGAGACTAGCCTCGAGG
pLV-R	CCAGTCAATCTTTCACAAATTTTGTAATCCAGAGG
pme-CFP-80-R	CATGGTGGCTCCGGAGATCCCGTAGTTTGACTTGTACAGCTCGTCCATGCCG
	CGATCACGAGACTAGCCTCGAGGTTTAAACGCCGCCACCATGGTGAGCAAGG
Pme-Koz-CFP-F	GC
XLF-Bam-R	CTCTC GGATC CTA ACT GAA GAG ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-F298G-Bam-	
R	CTCTC GGATC CTA ACT GCC GAG ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-FS298GA-	
Bam-R	CTCTCGGATC CTA AGC GCC GAG ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-FS298GE-	
Bam-R	CTCTCGGATC CTA CTC GCC GAG ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-Hind-F	CTCTCAAGCTTCCGCCACC ATG GAA GAA CTG GAG CAA GGC CTG
XLF-L115D-F	G CGA AGT GAG CTC TCT GGC GAC CCC TTC TAT TGG AAT TTC C
XLF-L115D-R	G GAA ATT CCA ATA GAA GGG GTC GCC AGA GAG CTC ACT TCG C
XLF-L297E-Bam-R	CTCTC GGATC CTA ACT GAA CTC ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-L297W-Bam-	
R	CTCTC GGATC CTA ACT GAA CCA ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-S299A-Bam-R	CTCTCGGATC CTA AGC GAA GAG ACC CCT TGG CTT CTT CCT CTT GAC C
XLF-S299E-Bam-R	CTCTCGGATC CTA CTC GAA GAG ACC CCT TGG CTT CTT CCT CTT GAC C

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# Uncropped image related to Figure 2e



# Uncropped scans related to Suppl. Fig. 5a-d

