1	RAD51 AND MITOTIC FUNCTION OF MUS81 ARE ESSENTIAL FOR RECOVERY FROM
2	LOW-DOSE OF CAMPTOTHECIN IN THE ABSENCE OF THE WRN EXONUCLEASE
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Stabilisation of the stalled replication fork is crucial to prevent excessive fork reversal or degradation, 29 which can undermine genome integrity. The WRN protein is a human RecQ helicase that participates 30 in the processing and recovery of perturbed replication forks. WRN is unique among the other human 31 RecQ family members to possess exonuclease activity. However, the biological role of the WRN 32 exonuclease is poorly defined, and little is known about an involvement in the response to perturbed 33 replication. Recently, the WRN exonuclease has been linked to protection of stalled forks from 34 MRE11-dependent degradation in response to clinically-relevant nanomolar doses of the 35 Topoisomerase I inhibitor camptothecin. Alternative processing of perturbed forks has been 36 37 associated to chemoresistance of BRCA-deficient cancer cells, thus, we used WRN exonucleasedeficiency as a model to investigate the fate of perturbed replication forks undergoing degradation, 38 39 but in a BRCA wild-type condition. We find that, upon nanomolar doses of camptothecin, loss of WRN exonuclease stimulates fork inactivation and accumulation of parental gaps, which engages 40 41 RAD51. Such alternative mechanism affects reinforcement of CHK1 phosphorylation and causes persistence of RAD51 during recovery from treatment. Notably, in WRN exonuclease-deficient cells, 42 persistence of RAD51 correlates with elevated mitotic phosphorylation of MUS81 at Serine 87, which 43 is essential to avoid accumulation of mitotic abnormalities. Altogether, these findings indicate that 44 aberrant fork degradation, in the presence of a wild-type RAD51 axis, stimulates RAD51-mediated 45 post-replicative repair and engagement of the MUS81 complex to limit genome instability and cell 46 death. 47

49 AUTHOR SUMMARY

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51 Correct progression of the molecular machine copying the chromosomes is threatened by multiple 52 causes that induce its delay or arrest. Once the replication machinery is arrested, the cell needs to 53 stabilise it to prevent DNA damage. Many proteins contribute to this task and the Werner's syndrome 54 protein, WRN, is one of them.

Defining what happens to replication machineries when they are blocked is highly relevant. Indeed, 55 destabilised replication machineries may form upon treatment with anticancer drugs and influence 56 57 the efficacy of some of them in specific genetic backgrounds. We used cells that lack one of the two enzymatic functions of WRN, the exonuclease activity, to investigate the fate of destabilised 58 59 replication machineries. Our data show that they are handled by a repair pathway normally involved in fixing DNA breaks but, in this case, recruited to deal with regions of the genome that are left 60 unreplicated after their destabilisation. This alternative mechanism involves a protein, RAD51, which 61 tries to copy DNA from the sister chromosome. In so doing, however, RAD51 produces a lot of DNA 62 63 interlinking that requires upregulation of a complex, called MUS81/EME1, which resolves this interlinking prior cell division and prevents accumulation of mitotic defects and cell death. 64

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The response to perturbed replication is crucial for the maintenance of genome integrity [1–5]. In humans, the proper handling of perturbed replication forks is also linked to cancer avoidance and many proteins involved in this process act also as onco-suppressors [3–6]. The importance of dealing correctly with perturbed replication forks is also demonstrated by the existence of several human genetic diseases caused by mutations in factors involved in sensing, processing and recovering replication forks [7].

The Werner's syndrome protein (WRN) is one of the key factors of the response to perturbed 78 replication and participates directly to the checkpoint in S-phase [8,9]. WRN is mutated in the genetic 79 80 disease Werner's syndrome (WS), which is characterized by cancer predisposition and premature aging [9], and its loss confers sensitivity to several DNA-damaging agents inducing replication stress 81 82 [8,10]. From an enzymatic point of view, WRN is both a DNA helicase and exonuclease; however, while its helicase activity has been linked to processing of reversed or collapsed replication forks 83 84 [2,9], little is known about the biological relevance of the exonuclease activity. We recently reported that the exonuclease activity of WRN is involved in protecting replication forks that are perturbed by 85 treatment with the Topoisomerase I poison Camptothecin (CPT) in the nanomolar range of 86 concentration [11]. Exposure to low doses of CPT, as opposed to high doses, does not induce DSBs 87 in the short-term but stimulates greatly formation of reversed forks [12,13]. Reversed replication forks 88 are versatile yet vulnerable structures and several proteins participate in their stabilisation [14–16]. 89 Two proteins, BRCA2 and RAD51, are the most crucial for the stabilisation of reversed forks 90 [14,16,17]. Thus, cells depleted of each of these two proteins have been used as a prototypical model 91 to assess the consequences of inaccurate handling of reversed forks. However, BRCA2 and RAD51 92 may also participate in DNA repair, which may be used to fix damage generated by fork instability 93 [17-19]. Loss of WRN exonuclease determines a rapid MRE11-dependent degradation of the nascent 94 strand, most likely after fork reversal, and affects correct replication recovery [11]. However, cells 95 expressing the exonuclease-dead WRN retain ability to restart replication and are not overtly sensitive 96 97 to low doses of CPT, suggesting that alternative mechanisms can be activated as a back-up. Since nanomolar doses of CPT are clinically-relevant in cancer therapy, cells expressing a catalytically-98 99 inactive WRN exonuclease can be used as a model to investigate the fate of CPT-perturbed replication 100 forks undergoing pathological degradation but in a BRCA2-RAD51 wild-type background.

Here, we report that, upon prolonged exposure to nanomolar dose of CPT, loss of WRN exonuclease
 channels replication forks through a pathological RAD51-dependent mechanism that makes
 perturbed replication forks resistant to breakage. However, engagement of RAD51 and persisting

104 RAD51 foci make WRN exonuclease-deficient cells reliant on the mitotic function of the MUS81 105 complex, which mitigates mitotic abnormalities deriving from accumulation of RAD51-dependent 106 intermediates. Furthermore, our data suggest that enhanced accumulation of ssDNA and recruitment 107 of RAD51 interfere with correct activation of CHK1, which provides a positive feedback to the 108 formation of nascent ssDNA.

Loss of WRN exonuclease activity leads to a persisting and unusual formation of nascent ssDNA which compromises formation of DSBs in response to a low-dose of camptothecin

113 Treatment with nanomolar concentrations of CPT does not induce DSBs immediately but initially 114 stimulates fork reversal [12]. Under such conditions, loss of WRN exonuclease activity results in the 115 rapid (< 2h) degradation of both nascent strands by the MRE11-EXO1 nucleases [11]. In this regard, 116 WRN exonuclease deficiency is a useful model to determine what happens to nuclease-targeted forks 117 after prolonged treatment with low dose of CPT. This prompted us to analyse the processing of 118 perturbed replication forks beyond 2h of treatment.

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120 We first examined the presence of nascent ssDNA as a sign of fork degradation by the native IdU assay. As expected, WS cells expressing the exo-dead WRN protein (WS^{E84A}) showed less nascent 121 ssDNA then the corrected wild-type counterpart (WS^{WT}) at 1h of treatment (Fig. 1A). Surprisingly, 122 the amount of nascent ssDNA in WRN exonuclease-deficient cells increased greatly over time and 123 largely exceeded the wild-type level at 4h, while it increased only slightly in cells expressing the 124 125 wild-type WRN (Fig. 1A). Since nascent strand degradation is MRE11-dependent in the absence of the WRN exonuclease while it is DNA2-dependent in wild-type cells (refs), we next examined if 126 chemical inhibition of those nucleases might reduce the idiosyncratic accumulation of ssDNA 127 detected at 4h of treatment with nanomolar CPT in WSE84A cells. Mirin treatment, which inhibits 128 MRE11, barely reduced ssDNA detected at 4h of treatment in wild-type cells (Fig. 1B). Surprisingly, 129 130 Mirin did not decrease the formation of ssDNA in WRN exonuclease-deficient cells but rather increased it even further (Fig. 1B). Inhibition of DNA2 by the small-molecule inhibitor C5 [20] 131 increased formation of nascent ssDNA in wild-type but not in WRN exonuclease-deficient cells (Fig. 132 1B). In contrast, concomitant inhibition of DNA2 and MRE11 was ineffective in modulating nascent 133 ssDNA formation in wild-type cells while decreased its level in cells expressing the exo-dead form 134 of WRN (Fig. 1B). Of note, ssDNA derived from end-resection of DSBs induced by a micromolar 135 dose of CPT was efficiently reduced by DNA2 inhibitor C5 (Fig. S1A, B), providing a functional 136 proof of the inactivation of the nuclease activity by the C5 compound in our cell model. These results 137 indicate that different sets of nucleases are involved in the degradation of nascent ssDNA in WRN 138 exonuclease-deficient cells when treatment is prolonged. 139

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141 DNA breakage can occur even in response to low-doses of CPT if treatment is sufficiently prolonged 142 [12,13]. Thus, to further investigate the origin of the late nascent ssDNA in the WRN exonuclease-143 deficient cells and the role of the different nucleases, we analysed the presence of DSBs after treatment with nanomolar CPT by neutral Comet assay. As shown in Figure 1C, treatment with 50nM CPT for 4h is able to induce some DSBs in wild-type cells, although they are very low compared with those generated by the 5µM reference dose. In contrast, no DSBs were detected in WRN exonuclease-deficient cells after treatment with the low-dose of CPT, even if they were readily seen in response to the high-dose of the drug (Fig. 1C). Interestingly, pre-treatment with Mirin, which enhances ssDNA in WS^{E84A} cells (Fig. 1B), resulted in DSBs (Fig. 1C). In contrast, formation of DSBs was

of MRE11 and DNA2 also reduced DSBs in wild-type cells (Fig. 1C). The analyses of DSBs and ssDNA suggest that the increase in nascent ssDNA induced by MRE11 inhibition in WRN exonuclease-deficient cells derives from end-resection of DSBs. In contrast, the reduction of nascent ssDNA induced by concomitant treatment with C5 and Mirin would indicate that DNA2 and MRE11

unaffected in wild-type cells by Mirin while it was reduced by the DNA2 inhibitor (Fig. 1C).

Interestingly, inhibition of both MRE11 and DNA2, which decreases ssDNA formation in WSE84A

cells (Fig. 1B), reduced significantly the DSBs generated by Mirin (Fig. 1C). Concomitant inhibition

- ssDNA induced by concomitant treatment with C5 and Mirin would indicate that DNA2 and MRE11
 are involved in the formation of the late nascent ssDNA observed in WS^{E84A} cells independently of
- detectable DSBs. Finally, we tested whether MUS81 was required for late DSB formation [21,22].
- As shown in Fig. S1C, DSBs derived from prolonged treatment with a low-dose of CPT in wild-typecells are not affected by depletion of MUS81.
- 161 Collectively, our results indicate that loss of the WRN exonuclease leads to accumulation of nascent 162 ssDNA when treatment with nanomolar CPT is prolonged beyond 2h and that it makes cells refractory 163 to formation of DSBs by CPT. Our data also suggest that the late accumulation of nascent ssDNA in 164 WS^{E84A} cells is related to the activity of multiple nucleases.
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166 Loss of WRN exonuclease stimulates engagement of RAD51 after CPT-induced fork 167 perturbation

In WRN exonuclease-deficient cells, late reappearance and accumulation of nascent ssDNA together with absence of DSBs after CPT might correlate with engagement of an alternative or aberrant fork processing mode over time. Extended regions of ssDNA are a substrate for RAD51 during recombination, thus, we investigated whether loss of WRN exonuclease could affect recruitment of RAD51 after treatment with CPT.

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To evaluate recruitment of RAD51, we first prepared chromatin from cells treated with nanomolar CPT and determined the amount of RAD51 present by Western blotting. In wild-type cells, RAD51 barely increased its chromatin association after CPT treatment (Fig. 2A). Expression of the exo-dead

177 WRN, however, greatly increased the amount of RAD51 in chromatin both in untreated cells and

CPT-treated cells. Treatment with CPT led to a minimal increase over untreated (about 20%; Fig. 178 2A). As a further control, we also measured the level of RPA32, a subunit of the RPA heterotrimer 179 that binds to ssDNA, in chromatin. RPA32 increased after CPT in wild-type cells (Fig. 2A). In 180 contrast, the amount of RPA32 in chromatin was low in the absence of the WRN exonuclease and 181 did not show any increase after treatment (Fig. 2A). RAD51 participates in multiple pathways [23,24] 182 and chromatin recruitment may reflect such pleiotropy of roles. To further verify the increased 183 recruitment of RAD51 in cells expressing the exo-dead WRN, we performed a quantitative 184 immunofluorescence analysis of the RAD51 foci (Fig. 2B). We evaluated the number of foci by 185 analysing the intensity of fluorescence in the RAD51 foci-positive nuclei. Consistent with 186 biochemical data, loss of WRN exonuclease increased recruitment of RAD51 (Fig. 2B). Moreover, 187 CPT treatment led to a higher number of RAD51 foci in WRN exonuclease-deficient cells compared 188 with the wild-type (Fig. 2B) 189

190 These results suggest that, in the absence of the WRN exonuclease, RAD51 takes over the normal 191 mechanism handling CPT-perturbed replication forks, perhaps to provide a backup mechanism for 192 recovery.

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194 To test if RAD51 participated in replication fork restart, we analysed whether, in cells expressing the exo-dead WRN, RAD51 chromatin levels remained elevated also during recovery from CPT. To this 195 end, we treated cells with CPT and allowed them to recover for 2 and 4h prior to preparing chromatin 196 fractions. As shown in Figure 3A, while RAD51 levels, as determined by Western blotting, tended to 197 remain low during recovery in wild-type cells, they remained elevated in WRN exonuclease-deficient 198 cells. Of note, the amount of chromatin-bound RPA32 and RPA70, two subunits of trimeric RPA, 199 remained low in WS^{E84A} cells while it increased greatly during recovery in cells expressing the wild-200 type WRN (Fig. 3A). To confirm this result, we performed quantitative immunofluorescence analysis 201 202 of the formation of RAD51 foci during recovery from CPT. A reduction of RAD51 foci was observed during recovery from CPT in wild-type cells (Fig. 3B). In contrast, the number of RAD51 foci 203 increased in WRN exonuclease-deficient cells (Fig. 3B), confirming that a RAD51-dependent 204 205 pathway is over-activated in response to prolonged treatment with low-dose of CPT in these cells.

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The persistently-high levels of RAD51 observed in the absence of the WRN exonuclease even during recovery prompted use to determine if they were correlated with the increased formation of nascent ssDNA. Having demonstrated that concomitant inhibition of MRE11 and DNA2 restores wild-type levels of nascent ssDNA in WRN exonuclease-deficient cells (Fig. 1C), we analysed RAD51 focusforming activity after pre-treatment of cells with C5 and Mirin. As expected, RAD51 recruitment in foci was elevated in WRN exonuclease-deficient cells after treatment and even more during recovery (Fig. 3C). Interestingly, concomitant pre-treatment with C5 and Mirin significantly reduced the formation of RAD51 foci in WS^{E84A} cells (Fig. 3C), suggesting that it depends on late accumulation of nascent ssDNA observed in the absence of WRN exonuclease.

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The elevated levels of RAD51 during recovery might be related to the presence of under-replicated 217 DNA that requires recombination to be replicated or repaired [25]. Thus, we analysed whether cells 218 recovering from CPT treatment presented under-replicated regions of DNA. Since under-replication 219 220 is expected to leave regions of ssDNA in the parental strand (parental gaps) behind perturbed forks, we performed native IdU assay after a 24h treatment with IdU to label all parental DNA prior to add 221 222 CPT and perform recovery (Fig. 3D). As shown in Figure 3E, little parental ssDNA was detectable in wild-type cells after treatment and its amount decreased substantially during recovery. In WS^{E84A} 223 224 cells, however, parental ssDNA was higher at the end of treatment and remained elevated during recovery (Fig. 3E). Interestingly, the increased amount of parental ssDNA paralleled that of RAD51 225 226 foci, suggesting that RAD51 may be recruited to deal with under-replicated regions.

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228 The presence of under-replicated DNA and the increased recruitment of RAD51 might indicate that perturbed replication forks get inactivated upon prolonged exposure to CPT in the absence of the 229 WRN exonuclease. Thus, we carried out a single-molecule analysis of DNA replication by the DNA 230 fiber assay (Fig. 4A). As observed previously [12,13], treatment with nanomolar CPT did not induce 231 fork arrest but rather a delay in fork progression (Fig. S2). Indeed, the length of the CldU tract is 232 increased during extended periods of treatment. Of note, length of the CldU tract was similar between 233 WS^{WT} and WS^{E84A} cells at 1h and 4h of treatment (Fig. S2). When we analysed the ability to replicate 234 after treatment, we found that most of perturbed replication forks remained active after 1h of CPT in 235 wild-type cells, as indicated by the low level of stalled forks (red-only tracks) (Fig. 4B-D). After 236 prolonged treatment, the number of stalled forks increased by around 2-fold, but perturbed forks 237 remained mostly active (Fig. 4B). Notably, the number of stalled forks was higher in WRN 238 exonuclease-deficient cells and around the 50% of the forks got inactivated at 4h of treatment (Fig. 239 4B-D). Interestingly, in the absence of the WRN exonuclease, treatment with CPT stimulated firing 240 of new origins (green-only tracks), which increased with time of treatment (Fig. 4C, D). In contrast, 241 limited new origin firing was detectable in wild-type cells (Fig. 4C, D). Surprisingly, and in spite of 242 the elevated levels of RAD51 in WS^{E84A} cells, neither the percentage of inactive forks nor that of new 243 origins was affected by inhibition of RAD51 during recovery (Fig. 4B-D). In contrast, we observed 244

that both fork inactivation and firing of new origins were increased by inhibition of RAD51 in wild-type cells (Fig. 4B-D).

These results suggest that RAD51 is not required to promote replication recovery but rather to promote repair of parental gaps left behind inactive forks after resumption of synthesis when the exonuclease activity of WRN is absent.

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251 Loss of WRN exonuclease reduces activation of CHK1

Our data indicate that loss of the WRN exonuclease results in accumulation of ssDNA and RAD51 252 accompanied by a concomitant decrease of RPA. Since RPA-coated ssDNA is required for 253 checkpoint signalling upon replication fork perturbation, we investigated whether the functionality 254 255 of the WRN exonuclease might also affect activation of the replication checkpoint in response to nanomolar concentration of CPT. As a readout of the activation of the ATR-dependent checkpoint 256 257 response, we analysed phosphorylation of CHK1 at S345 by Western blotting. In wild-type cells, treatment with a nanomolar dose of CPT induced a time-dependent phosphorylation of CHK1 which 258 is also readily observed in cells expressing the helicase-dead form of WRN (WSK577M) (Fig. 5A). In 259 contrast, CPT-induced phosphorylation of CHK1 was reduced in cells expressing the exonuclease-260 261 dead mutant of WRN, and this phenotype was more evident at 4 and 6h of treatment (Fig. 5A). The requirement of the WRN exonuclease for correct CHK1 phosphorylation was specific for the low-262 dose CPT treatment as it was not observed after 5µM of CPT (Fig. S3A). 263

To confirm that loss of WRN exonuclease affected CHK1 phosphorylation by an independent assay, we monitored the status of S345 of CHK1 by immunofluorescence. As shown in Fig. 5B, a reduced phosphorylation of CHK1 at S345 was readily detected also by immunofluorescence in WRN exonuclease-deficient cells.

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Next, we wanted to analyse whether decreased activation of CHK1 correlated with reduced activation 269 of ATR-dependent signalling. To assess activation of ATR, we monitored phosphorylation of the 270 activating site T1989 by immunofluorescence. Despite the defective phosphorylation of CHK1, ATR 271 272 was phosphorylated similarly in wild-type cells and in cells expressing the exonuclease-dead form of WRN (Fig. 5C). Since loss of WRN exonuclease affects recruitment of RPA but did not affect 273 274 activation of ATR, we analysed whether it could influence recruitment of other factors modulating ATR-checkpoint function. As loss of WRN exonuclease leads to accumulation of nascent ssDNA at 275 4h of CPT (Fig. 1A), we analysed the presence of TopBP1 and RAD9, which associates with TopBP1 276 [26,27], specifically at nascent ssDNA by our recently described IdU-PLA assay [11]. In parallel, we 277 278 evaluated the presence of the total amount of ssDNA by IdU assay. Association of TopBP1 or RAD9

with nascent ssDNA was not detected under untreated conditions (data not shown), however,
treatment with 50nM CPT for 4h resulted in recruitment of both factors at nascent ssDNA in wildtype cells (Fig. 5D, E). Of note, loss of WRN exonuclease reduced the recruitment of TopBP1 but
not that of RAD9 at the nascent ssDNA (Fig. 5D, E) although in these conditions the IdU assay
detected 2-times more ssDNA (Fig. 5F).

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Loss of the WRN exonuclease leads to accumulation of nascent ssDNA, which is targeted by RAD51 285 and not by checkpoint factors, possibly resulting in reduced CHK1 phosphorylation. Thus, we 286 investigated whether pre-treatment with the RAD51 inhibitor B02 or treatment before sampling could 287 re-establish a normal CHK1 activation in WRN exonuclease-deficient cells (Fig. S3B). Of note, 288 289 inhibition of RAD51 further decreased the phosphorylation of CHK1 regardless the way it was added (Fig. S3B). This was surprising but prompted us to evaluate whether reduced CHK1 activation could 290 291 be implicated in the enhanced accumulation of nascent ssDNA. To test this potential feedback effect, we expressed the S317/S345D CHK1 phosphomimetic mutant [28] in WS^{E84A} cells (Fig. 6A) before 292 293 evaluating the formation of nascent ssDNA by the IdU assay. As shown in Figure 6B, the phosphomimetic CHK1 was efficiently expressed in the cells and its expression was able to 294 substantially reduce the amount of nascent ssDNA in WRN exonuclease-deficient cells restoring the 295 wild-type levels. 296

Altogether, our results show that loss of WRN exonuclease activity affects proper activation of CHK1 in response to a low-dose of CPT and that reduced phosphorylation of CHK1 probably correlates with reduced recruitment of checkpoint factors at ssDNA. They also suggest that reduced CHK1 phosphorylation contributes to the accumulation of ssDNA in the nascent strand, possibly as part of a positive feedback loop.

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303 WRN exonuclease-deficient cells need the mitotic function of MUS81 to counteract mitotic 304 aberration and mis-segregation

Inaccurate processing of perturbed replication forks, elevated under-replicated DNA and RAD51 305 306 levels observed in the absence of WRN exonuclease could threaten mitosis because of DNA interlinking as shown in BRCA2-deficient cells [29]. Mitotic resolution of DNA interlinked 307 intermediates involves the MUS81/EME1 complex [30,31]. Thus, we evaluated whether WRN 308 exonuclease-deficient 309 cells accumulated active MUS81 in mitosis by performing immunofluorescence with an antibody directed against the pS87-MUS81, which we have 310 demonstrated to be a readout of the active MUS81/SLX4 complex [32]. In wild-type cells, little 311 312 MUS81 phosphorylation at S87 was detectable either under unperturbed cell growth or in response

to low-dose of CPT (Fig. 7A). In contrast, many pS87-MUS81-positive nuclei were detectable in 313 cells expressing the exonuclease-dead WRN already in unperturbed conditions (Fig. 7A). Notably, in 314 WRN exonuclease-deficient cells, pS87-MUS81 levels were further enhanced by CPT and remained 315 elevated also after recovery (Fig. 7A). Interestingly, and consistent with our previous results [32], 316 phosphorylation of MUS81 never occurred in S-phase cells labelled with EdU (Fig. 7A). To 317 determine whether elevated MUS81 activation correlated with under-replication and enhanced 318 RAD51 recruitment, we asked if inhibition of RAD51 by B02 (ref) reverted the pS87-MUS81 levels 319 in WRN exonuclease-deficient cells. As shown in Figure 7B, inhibition of RAD51 greatly decreased 320 activation of the MUS81 complex after recovery from CPT, as indicated by reduction in pS87-321 MUS81-positive nuclei. 322

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Enhanced activation of MUS81 in mitosis in WSE84A cells might be indicative of persistence of 324 unresolved DNA intermediates, which could induce mitotic abnormalities or segregation defects. To 325 assess if loss of WRN exonuclease during the response to low-dose of CPT could result in segregation 326 327 defects, we first analysed the presence of bulky anaphase bridges in DAPI-stained cells (Fig. 7C). Interestingly, anaphase cells were highly enriched in WRN exonuclease-deficient cells (Fig. 7C; see 328 329 numbers above the bars), suggesting that these cells may have a delayed exit from anaphase. Of note, the number of anaphases with bridges was very low in WRN exonuclease-deficient cells as compared 330 to cells expressing WRN wild-type (Fig. 7C). Since delayed exit from anaphase might derive from 331 mitotic defects and could result in post-mitotic abnormalities, we decided to evaluate the presence of 332 aberrant mitoses, multinucleated cells and micronuclei. As shown in Figure 7D, we found that both 333 aberrant mitosis and cells with micronuclei were increased by loss of WRN exonuclease activity. In 334 contrast, no difference in multinucleated cells was found between cells expressing wild-type or exo-335 dead WRN (Fig. 7D). 336

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The presence of under-replicated DNA or the persistence of unresolved DNA intermediates in G2/M 338 triggers the formation of 53BP1 NBs in the subsequent G1 phase [33]. Since WRN exonuclease-339 340 deficient cells showed persistence of under-replicated DNA, we investigated whether they accumulated 53BP1 NBs. To this end, we performed immunofluorescence against 53BP1 and 341 CyclinA in cells after recovery from 4h treatment with CPT and scored the number of 53BP1 NBs-342 positive cells in the CyclinA-negative population (i.e. G1 cells). As shown in Figure 7E, the number 343 of 53BP1 NBs in WS^{E84A} cells was higher than in wild-type cells even in untreated conditions. 344 Treatment with CPT enhanced the number of 53BP1 NBs in wild-type and in WRN exonuclease-345

deficient cells; however, the increase was substantially higher in cells expressing the exo-dead WRN

347 (Fig. 7E and Fig. S5).

Notably, inhibition of RAD51 in WRN exonuclease-deficient cells resulted in persistent accumulation of mitotic cells in pro-metaphase (Fig. S4). This result suggested that loss of RAD51 undermines correct mitotic progression in WRN-exonuclease-deficient cells but prevented assessment of any correlation between enhanced engagement of RAD51 and mitotic defects or formation of 53BP1 NBs.

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As in WRN exonuclease-deficient cells engagement of RAD51 is functionally related to elevated 354 levels of S87-MUS81 phosphorylation (Fig. 7B), we next analysed whether inactivation of the mitotic 355 356 function of MUS81 by overexpression of the unphosphorylable S87A-MUS81 mutant (ref) in these cells could aggravate the mitotic defects. Interestingly, over-expression of the S87A-MUS81 mutant 357 in wild-type cells did not affect the percentage of anaphase bridges and micronuclei, while it increased 358 the number of multinucleated cells (Fig. 8A). In sharp contrast, expression of the S87A-MUS81 359 360 mutant substantially aggravated the mitotic defects in WRN exonuclease-deficient cells (Fig. 7A). Indeed, expression of the S87A-MUS81 protein increased the number of anaphase bridges and 361 362 micronucleated cells. Similarly, expression of the S87A-MUS81 mutant enhanced the presence of 53BP1 NBs in WS^{E84A} cells (Fig. 8B). 363

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To further assess the biological significance of the MUS81 hyperactivation observed in mitosis in the absence of the WRN exonuclease, we evaluated the sensitivity to nanomolar doses of CPT by clonogenic survival. As shown in Figure 8C, WRN exonuclease-deficient cells were slightly more sensitive to CPT then wild-type cells. Overexpression of the wild-type MUS81 resulted in a mild increase in sensitivity in wild-type cells but not in cells expressing the exo-dead WRN (Figure 7C). In contrast, overexpression of the S87A-MUS81 resulted in a substantial increase in the sensitivity of WRN exonuclease-deficient cells to CPT (Fig. 8C).

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Altogether, our data indicate that the enhanced engagement of RAD51 observed in the absence of the WRN exonuclease requires the increased activation of the MUS81 complex in mitosis. Therefore, expression of a MUS81 mutant that disables mitotic activation of the MUS81/EME1 complex increases mitotic abnormalities and sensitivity to CPT of WRN exonuclease-deficient cells. Thus, in the absence of the WRN exonuclease, hyperactivation of the MUS81 complex functions as a fail-safe system that maintains mitotic abnormalities at low levels, allowing survival.

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In recent years, an increasing interest arose around alternative mechanisms of fork processing and 382 fork degradation since they correlate with response to chemotherapeutics in cells that are deficient 383 for the primary pathway(s) as described in the absence of BRCA1/2 [15,17]. Most of these studies 384 focused on the early events occurring in the absence of BRCA1 or BRCA2, but few of them 385 investigated mechanisms involved during recovery from replication stress [29,34]. Furthermore, loss 386 of BRCA1 or BRCA2 affects recombination as well as fork protection and this prevents the 387 investigation of the role of recombination for the recovery of replication forks undergoing 388 degradation. Recently, we reported that the WRN exonuclease activity protects against fork 389 390 degradation when cells are treated with clinically-relevant doses of CPT [11]. Here, we used WRN exonuclease-deficient cells as a model to assess what happens at destabilised perturbed forks when 391 392 treatment with nanomolar doses of CPT is prolonged. We find that, in the absence of the WRN exonuclease, nascent strands undergo continuous degradation that produces accumulation of ssDNA. 393 394 This late accumulation of ssDNA follows its disappearance at early time points because of the activities of MRE11 and EXO1 [11]. Interestingly, the late wave of ssDNA at perturbed forks is only 395 396 minimally affected by inactivation of each single exonuclease acting at perturbed forks, and is only reduced when MRE11 and DNA2 are both inhibited. This suggests that multiple nucleases take over 397 with time at forks destabilised by the absence of WRN exonuclease while most of the degradation 398 observed in cells deficient of BRCA1/2, or other factors assisting RAD51, seems to involve only 399 MRE11-EXO1 [35-38]. Treatment with nanomolar doses of CPT does not induce DSBs unless 400 treatment is prolonged [12,13]. Interestingly, loss of the WRN exonuclease makes cells resistant to 401 402 the induction of DSBs after prolonged treatment with nanomolar CPT. Induction of DSBs in response to nanomolar doses of CPT has been correlated with activation of RECQ1 possibly to promote restart 403 of those forks that failed to be processed otherwise [13]. Loss of the ability to induce DSBs at forks 404 would be consistent with engagement of a distinct fork recovery mechanism in cells expressing the 405 exo-dead WRN protein. Indeed, WRN exonuclease-deficient cells do not show RECQ1 PARylation 406 407 [11], which is required to avoid unscheduled RECQ1 activation [13], which, together with the absence of DSBs support a pathway switch at CPT-perturbed forks. 408

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Consistent with the pathway switch, inhibition of MRE11 is sufficient to restore DSBs after prolonged
treatment with a low-dose of CPT in WRN exonuclease-deficient cells, suggesting that formation of
DSBs does not necessarily occur downstream of pathological fork processing. Interestingly, RAD51

is strongly accumulated in the absence of the WRN exonuclease and persists during recovery from

treatment. An elevated engagement of RAD51 in the absence of the WRN exonuclease has been 414 reported in Drosophila [39], suggesting that the role of WRN exonuclease at perturbed forks is 415 conserved. Similarly, unscheduled exonuclease-mediated processing of perturbed forks in yeast has 416 been recently shown to engage a RAD51-mediated pathway [40]. Furthermore, parental ssDNA, a 417 readout of template gaps, also accumulates in the absence of WRN exonuclease. RAD51 binds to 418 ssDNA and initiates recombination [19,25]. The concomitant accumulation of ssDNA and elevated 419 recruitment of RAD51 in the absence of the WRN exonuclease would be consistent with the 420 engagement of gaps left behind inactivated forks in a template-switch mode of replication recovery, 421 as shown after DNA damage in Xenopus egg extracts [23]. Consistent with this possibility, WRN 422 exonuclease-deficient cells show enhanced inactivation of CPT-perturbed forks and new origin firing. 423 424 Moreover, although RAD51 has been implicated in fork restart [24,41], in WRN exonucleasedeficient cells, RAD51 is not involved in fork reactivation after CPT treatment. Indeed, its inhibition 425 does not increase fork inactivation. This result is in agreement with the participation of RAD51 in 426 "gap repair" and supports the notion that prolonged treatment with nanomolar CPT channels 427 428 perturbed forks into alternative fork processing pathways if the function of WRN exonuclease is 429 absent. Consistent with this, in wild-type cells, inhibition of RAD51 reduces fork reactivation. This 430 is not unexpected since RAD51 plays crucial roles in both fork remodelling and stability [14,16]. In addition, as RAD51 and WRN have been proposed to cooperate during recovery from fork arrest 431 [42], it is reasonable to speculate that loss of WRN function also compromises the normal activity of 432

- 433 RAD51 at fork.
- 434

Loss of WRN exonuclease results in a mild defect in the activation of CHK1. Activation of the ATR-435 dependent checkpoint requires formation of ssDNA [43–45]. In the absence of the WRN exonuclease 436 ssDNA accumulates but is hijacked by RAD51 and it is not completely free for the binding of 437 checkpoint factors. Indeed, in WRN exonuclease-deficient cells, TopBP1 and its binding factor 438 RAD9 are not more highly associated with ssDNA as compared with the wild-type. Notably, 439 phosphorylation of ATR, a readout of its activation, is indistinguishable from the wild-type. It 440 441 suggests that ATR also gets activated independently from ssDNA. Alternatively, a hyperactivation of ATR is also prevented by sequestering of ssDNA by RAD51. Indeed, overexpression of RAD51 442 has been shown to affect checkpoint activation [46]. Notably, bypassing of the CHK1 activation 443 defect by expression of a phosphomimetic CHK1 mutant in WRN exonuclease-deficient cells restores 444 normal levels of ssDNA. This suggests that accumulation of ssDNA is also unleashed by reduced 445 CHK1 activation through a positive feedback loop. 446

The observed elevated recruitment of RAD51, which is used during recovery in the absence of the 448 WRN exonuclease to deal with under-replicated DNA, also leads to elevated phosphorylation of 449 MUS81 at S87. Phosphorylation of MUS81 at S87 occurs in G2/M and is related to resolution of 450 recombination intermediates [32]. Consistently, inhibition of RAD51 reduces S87 phosphorylation 451 in WRN exonuclease-deficient cells. Thus, engagement of RAD51-dependent fork recovery, possibly 452 by template switch since DSBs do not form, results in an increased number of interlinked 453 intermediates calling for resolution by the MUS81 complex. Our data indicate that activation of 454 MUS81 complex in G2/M is essential to overcome segregation defects arising from excessive 455 RAD51-dependent recombination and support proliferation upon treatment with CPT. Indeed, 456 expression of the unphosphorylable S87A-MUS81 mutant increases abnormal mitosis and sensitizes 457 458 WRN exonuclease-deficient.

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Loss of the WRN exonuclease although resulting in fork degradation does not induce MUS81 460 activation in S-phase, which is observed in the absence of BRCA2 [37]. However, the persistence of 461 462 under-replicated DNA and requirement of MUS81 complex activity in G2/M shown by WRN exonuclease-deficient cells are also characteristic of BRCA2-deficient cells [34]. Thus, it is tempting 463 to speculate that elevated fork degradation correlates with inability to replicate all the genome. 464 Notably, BRCA2-deficient cells show much more severe mitotic defects [29,34]. In WRN 465 exonuclease-deficient cells, mitotic abnormalities are increased by disabling MUS81 function in 466 mitosis but are likely increased also by impairing RAD51 function during recovery, since RAD51 467 inhibition results in increased accumulation of parental ssDNA and induces a significant mitotic 468 block. As BRCA2 deficiency also interferes with the post-replicative function of RAD51 [23] it is 469 tempting to speculate that the elevated mitotic defects might be the end-result of combined fork 470 deprotection and recombination defects. Indeed, it is recently demonstrated that mitotic abnormalities 471 in BRCA2-deficient cells are primarily linked to loss of the recombination function of RAD51 using 472 separation-of-function mutants [29]. 473

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Collectively, our data show that WRN exonuclease-deficient cells can be a useful model to investigate the fate of deprotected or destabilised replication forks under a clinically-relevant, specific type of replication stress; and, together with published data, they can be summarised in the model shown in Figure 9. In response to nanomolar CPT, perturbed replication forks rapidly undergo fork reversal [12]. The WRN exonuclease is required somehow at this stage to prevent MRE11-dependent degradation. In wild-type cells, with time, reversed forks degenerate into DSBs, possibly because of unscheduled RECQ1-mediated fork restoration [13]. In the absence of WRN exonuclease, perturbed

replication forks undergo a further cycle of degradation of nascent strand by MRE11 and/or DNA2, 482 which leads to ssDNA accumulation and engagement of RAD51. The accumulation of ssDNA and 483 possibly engagement of RAD51 make perturbed replication forks resistant to DSBs and interfere with 484 checkpoint signalling, resulting in a mild defect in CHK1 activation. In the absence of WRN 485 exonuclease, more perturbed forks become inactivated over-time and RAD51 is required also during 486 recovery from CPT to support repair at template gaps left behind the inactive forks. Engagement of 487 RAD51 during recovery results in elevated activation of the MUS81 complex in G2/M to deal with 488 intermediates, and to limit mitotic defects and cell death. As CPT is a chemotherapeutic, our data also 489 490 indicate that tumors with impaired function of the WRN exonuclease can be sensitized to treatment by genetic or chemical interference with the MUS81 complex in mitosis, which is less relevant for 491 492 survival in cells expressing the WRN wild-type.

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496 MATERIALS AND METHODS

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498 Cell lines and culture conditions

The SV40-transformed WRN-deficient fibroblast cell line (AG11395) was obtained from Coriell Cell Repositories (Camden, NJ, USA). To produce stable cell lines, AG11395 (WS) fibroblasts were transduced with retroviruses expressing the full-length cDNA encoding wild-type WRN (WS^{WT}), exonuclease-dead (WS^{E84A}), or helicase-dead (WS^{K577M})[47]. All the cell lines were maintained in Dulbecco's modified Eagle's medium (DMEM; Life Technologies) supplemented with 10% FBS (Boehringer Mannheim) and incubated at 37 °C in a humidified 5% CO₂ atmosphere.

505

506 Chemicals

507 Camptothecin (ENZO Lifesciences) was dissolved in DMSO and a stock solution (10 mM) was 508 prepared and stored at -20°C. Mirin (Calbiochem), an inhibitor of MRE11 exonuclease activity, was 509 used at 50 μ M; the B02 compound (Selleck), an inhibitor of RAD51 activity, was used at 27 μ M. C5 510 (ref), DNA2 inhibitor C5 was dissolved in DMSO and used at final concentration of 300 μ M [20]. 511 IdU and CldU (Sigma-Aldrich) were dissolved in sterile DMEM at 2.5mM and 200mM respectively 512 and stored at -20°C.

513

514 Plasmids transfection

Plasmid expressing the phospho - mimic (Flag-CHK1^{317/345D}) mutant form of CHK1, a kind gift from
Professor K.K. Khanna (Queensland Institute of Medical Research, Australia) was generated as
described [28]. To express the plasmids, cells were transfected using the NeonTM Transfection System
Kit (Invitrogen), according to the manufacturer's instructions.

519

520 Immunofluorescence assays

Cells were grown on 35-mm coverslips and harvested at the indicated times after treatments. For 521 RAD51 IF, after further washing with PBS, cells were pre-extracted with 0,5% TritonX-100 and fixed 522 523 with 3% PFA / 2% sucrose at RT for 10min. After blocking in 3% BSA for 15 min, staining was performed with rabbit monoclonal anti-RAD51 (Bioss, 1:100) diluted in a 1% BSA / 0,1% saponin 524 in PBS solution, for 1h at 37° in a humidified chamber. For 53BP1, pS87MUS81 and α-tubulin 525 staining, cells were fixed with 4% PFA at RT for 10 min. Cells were subsequently permeabilized with 526 0,4% Triton-X100. Staining with primary antibodies diluted in a 1% BSA / 0,1% saponin in PBS 527 solution was carried out for 1h at RT. After extensive washing with PBS, specie-specific fluorophore-528 529 conjugated antibody (Invitrogen) was applied for 1h at RT followed by counterstaining with 0.5 mg/ml DAPI. Secondary antibody was used at 1:200 dilution. Images were acquired as greyscale files
using Metaview software (MDS Analytical Technologies) and processed using Adobe Photoshop

- 532 CS3 (Adobe). For each time point, at least 200 nuclei were examined, and foci were scored at $40\times$.
- 533 Only nuclei with > 5 foci were considered as positive and were quantified using ImageJ.
- 534

535 EdU incorporation assay

To label replicated DNA, cells were incubated with 10 μ M EdU for 30 minutes. Samples were fixed with 4% PFA at RT for 10 min and cells were subsequently permeabilized with 0,5% Triton-X100. EdU incorporation was detected using the Click-It Edu Alexa Fluor 488 Imaging Kit (Invitrogen) according to the manufacturer's instructions.

540

541 Antibodies

The primary antibodies used were: anti-pS10H3 (1:1000, Santa Cruz Biotechnologies), anti-Cyclin
A (IF: 1:100, Santa Cruz Biotechnologies), anti-53BP1 (1:300, Millipore), anti-BrdU (1:80, Abcam;
CldU detection), anti-BrdU (1:50, Becton Dickinson; anti-IdU detection), anti-pS87MUS81 (ref
1:200), anti-RAD51 (1:1000, Bioss Antibodies), anti-αTubulin (1:50, Sigma-Aldrich) and anti-Lamin
B1 (1:10000, Abcam).

547

548 Chromatin fractionation and Western blot analysis

Chromatin fractionation experiments were performed as previously described [48]. Western blotting 549 was performed using standard methods. Blots were incubated with primary antibodies against: rabbit 550 anti-pCHK1(S345) (Cell Signalling Technology), mouse anti-CHK1 (Santa Cruz Biotechnology), 551 rabbit anti-RAD51 (Bioss Antibodies), mouse anti-RPA32 (Calbiochem), rabbit anti-RPA70 552 (GeneTex), mouse anti-GAPDH (Santa Cruz Biotechnology) and rabbit anti-Lamin B1 (Abcam). 553 After incubations with horseradish peroxidase-linked secondary antibodies (1:20000, Jackson 554 Immunosciences), the blots were developed using the chemiluminescence detection kit 555 WesternBright ECL HRP substrate (Advansta) according to the manufacturer's instructions. 556 557 Quantification was performed on scanned images of blots using the Image Lab software, and the values shown on the graphs represent normalization of the protein content evaluated through 558 LaminB1 or GAPDH immunoblotting. 559

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564 Clonogenic survival

Cells were plated onto 35mm dishes, after 24h they were treated with different doses of CPT. After
18, cells were washed, trypsinized and seeded in 60mm dishes. After 14-21 days, plates were stained
with crystal violet and colonies counted.

568

569 **DNA fibres analysis**

570 DNA fibres were prepared, spread out and immunodecorated as previously described [11]. Images 571 were acquired randomly from fields with untangled fibres using Eclipse 80i Nikon Fluorescence 572 Microscope, equipped with a VideoConfocal (ViCo) system. The length of labeled tracks were 573 measured using the Image-Pro-Plus 6.0 software. A minimum of 100 individual fibres were analysed 574 for each experiment and the mean of at least three independent experiments presented.

575

576 Detection of nascent single-stranded DNA

To detect nascent single-stranded DNA (ssDNA), cells were plated onto 22x22 coverslips in 35mm 577 dishes. After 24h, the cells were labelled for 15 min before the treatment with 250µM IdU (Sigma-578 Aldrich), cells were then treated with CPT 5µM for different time points. Next, cells were washed 579 580 with PBS, permeabilized with 0.5% Triton X-100 for 10 min at 4°C and fixed wit 2% sucrose, 3% PFA. For ssDNA detection, cells were incubated with primary mouse anti-BrdU antibody (Becton 581 Dickinson) for 1h at 37°C in 1%BSA/PBS, followed by Alexa Fluor488-conjugated goat-anti-Mouse 582 (Invitrogen), and counterstained with 0.5µg/ml DAPI. Slides were analysed with Eclipse 80i Nikon 583 Fluorescence Microscope, equipped with a VideoConfocal (ViCo) system. For each time point, at 584 least 100 nuclei were scored at 60×. Parallel samples either incubated with the appropriate normal 585 serum or only with the secondary antibody confirmed that the observed fluorescence pattern was not 586 attributable to artefacts. Fluorescence intensity for each sample was then analysed using ImageJ 587 588 software.

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590 Statistical analysis

All the data are presented as means of at least two independent experiments. Statistical comparisons of WS^{WT} or WRN-mutant cells to their relevant control were analysed by ANOVA or Mann-Whitney test. P < 0.5 was considered as significant.

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603

604 AUTHOR CONTRIBUTIONS

F.A.A. performed the analysis of CHK1 phosphorylation, fork recruitment by PLA and chromatin fractionation, and performed experiments to determine DNA damage. A.P. performed the analysis of MUS81 phosphorylation and experiments to evaluate mitotic abnormalities. E.M. analysed the persistence of RAD51 and parental ssDNA. F.A.A., A.P., E.M. analysed data, contributed to designing the experiments and writing the manuscript. A.F. and P.P. designed experiments, analysed data and wrote the paper. L.Z., J.L.C. and B.H.S. provided the DNA2 inhibitor C5, advised the relevant experiments, and revised the manuscript. All authors approved the paper.

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613 CONFLICT OF INTEREST

The authors declare that they do not have any conflict of interest.

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617 REFERENCES

- Técher H, Koundrioukoff S, Nicolas A, Debatisse M. The impact of replication stress on
 replication dynamics and DNA damage in vertebrate cells. Nat Rev Genet. Nature Publishing
 Group; 2017; doi:10.1038/nrg.2017.46
- Franchitto A, Pichierri P. Replication fork recovery and regulation of common fragile sites
 stability. Cell Mol Life Sci. 2014;71: 4507–4517. doi:10.1007/s00018-014-1718-9
- Zeman MK, Cimprich KA. Causes and consequences of replication stress. Nat Cell Biol.
 2014;16: 2–9. doi:10.1038/ncb2897
- 4. Macheret M, Halazonetis TD. DNA Replication Stress as a Hallmark of Cancer. Annu Rev
 Pathol Mech Dis. 2015;10. doi:10.1146/annurev-pathol-012414-040424
- Magdalou I, Lopez BS, Pasero P, Lambert S a E. The causes of replication stress and their
 consequences on genome stability and cell fate. Semin Cell Dev Biol. Elsevier Ltd; 2014;30:
 154–164. doi:10.1016/j.semcdb.2014.04.035
- 6. Hills SA, Diffley JFX. DNA Replication and Oncogene-Induced Replicative Stress. Curr
 Biol. Elsevier Ltd; 2014;24: R435–R444. doi:10.1016/j.cub.2014.04.012
- 633 7. Ciccia A, Elledge SJ. The DNA Damage Response: Making It Safe to Play with Knives. Mol
 634 Cell. Elsevier Inc.; 2010;40: 179–204. doi:10.1016/j.molcel.2010.09.019
- 8. Pichierri P, Ammazzalorso F, Bignami M, Franchitto A. The Werner Syndrome protein:
- Linking the replication checkpoint response to genome stability. Aging (Albany NY).
 2011;3: 311–318. doi:100293 [pii]
- 638 9. Rossi ML, Ghosh AK, Bohr V a. Roles of Werner syndrome protein in protection of genome
 639 integrity. DNA Repair (Amst). Elsevier B.V.; 2010;9: 331–44.
- 640 doi:10.1016/j.dnarep.2009.12.011
- Franchitto A, Pichierri P. Understanding the molecular basis of common fragile sites
 instability: Role of the proteins involved in the recovery of stalled replication forks. Cell
 Cycle. 2011;10: 4039–4046. doi:10.4161/cc.10.23.18409
- Iannascoli C, Palermo V, Murfuni I, Franchitto A, Pichierri P. The WRN exonuclease
 domain protects nascent strands from pathological MRE11/EXO1-dependent degradation.
 Nucleic Acids Res. 2015;43: 9788–803. doi:10.1093/nar/gkv836
- 12. Ray Chaudhuri A, Hashimoto Y, Herrador R, Neelsen KJ, Fachinetti D, Bermejo R, et al.
- 648
 Topoisomerase I poisoning results in PARP-mediated replication fork reversal. Nature
- 649 Structural & Molecular Biology. Nature Publishing Group; 2012. pp. 417–423.
- 650 doi:10.1038/nsmb.2258

Berti M, Ray Chaudhuri A, Thangavel S, Gomathinayagam S, Kenig S, Vujanovic M, et al.
Human RECQ1 promotes restart of replication forks reversed by DNA topoisomerase I
inhibition. Nat Struct Mol Biol. Nature Publishing Group; 2013;20: 347–54.

654 doi:10.1038/nsmb.2501

- Bhat KP, Cortez D. RPA and RAD51: fork reversal, fork protection, and genome stability.
 Nat Struct Mol Biol. 2018;25: 446–453. doi:10.1038/s41594-018-0075-z
- Quinet A, Lemaçon D, Vindigni A. Replication Fork Reversal: Players and Guardians. Mol
 Cell. 2017;68: 830–833. doi:10.1016/j.molcel.2017.11.022
- Kolinjivadi AM, Sannino V, de Antoni A, Técher H, Baldi G, Costanzo V. Moonlighting at
 replication forks a new life for homologous recombination proteins BRCA1, BRCA2 and
 RAD51. FEBS Lett. 2017; 1–18. doi:10.1002/1873-3468.12556
- Feng W, Jasin M. Homologous Recombination and Replication Fork Protection: BRCA2 and
 More! Cold Spring Harb Symp Quant Biol. 2017;82: 329–338.
- doi:10.1101/sqb.2017.82.035006
- 18. Costanzo V. Brca2, Rad51 and Mre11: Performing balancing acts on replication forks. DNA
 Repair (Amst). Elsevier B.V.; 2011;10: 1060–1065. doi:10.1016/j.dnarep.2011.07.009
- Pellegrini L, Venkitaraman A. Emerging functions of BRCA2 in DNA recombination.
 Trends Biochem Sci. 2004;29: 310–316. doi:10.1016/j.tibs.2004.04.009
- Liu W, Zhou M, Li Z, Li H, Polaczek P, Dai H, et al. A Selective Small Molecule DNA2
 Inhibitor for Sensitization of Human Cancer Cells to Chemotherapy. EBioMedicine. The
 Authors; 2016;6: 73–86. doi:10.1016/j.ebiom.2016.02.043
- Murfuni I, Nicolai S, Baldari S, Crescenzi M, Bignami M, Franchitto a, et al. The WRN and
 MUS81 proteins limit cell death and genome instability following oncogene activation.

```
674 Oncogene. Nature Publishing Group; 2012;32: 610–20. doi:10.1038/onc.2012.80
```

- 475 22. Hanada K, Budzowska M, Davies SL, van Drunen E, Onizawa H, Beverloo HB, et al. The
 476 structure-specific endonuclease Mus81 contributes to replication restart by generating
- double-strand DNA breaks. Nat Struct Mol Biol. 2007;14: 1096–1104.
- doi:10.1038/nsmb1313
- Hashimoto Y, Chaudhuri AR, Lopes M, Costanzo V. Rad51 protects nascent DNA from
 Mre11-dependent degradation and promotes continuous DNA synthesis. Nat Struct Mol Biol.
 2010;17: 1305–1311. doi:10.1038/nsmb.1927
- Petermann E, Orta ML, Issaeva N, Schultz N, Helleday T. Hydroxyurea-Stalled Replication
 Forks Become Progressively Inactivated and Require Two Different RAD51-Mediated
 Pathways for Restart and Repair. Mol Cell. 2010;37: 492–502.

685 doi:10.1016/j.molcel.2010.01.021

- 686 25. Carr AM, Lambert S. Replication stress-induced genome instability: The dark side of
- replication maintenance by homologous recombination. J Mol Biol. Elsevier Ltd; 2013;425:
 4733–4744. doi:10.1016/j.jmb.2013.04.023

689 26. Lee J, Kumagai A, Dunphy WG. The Rad9-Hus1-Rad1 checkpoint clamp regulates

interaction of TopBP1 with ATR. J Biol Chem. 2007;282: 28036–28044.

691 doi:10.1074/jbc.M704635200

- Delacroix S, Wagner JM, Kobayashi M, Yamamoto KI, Karnitz LM. The Rad9-Hus1-Rad1
 (9-1-1) clamp activates checkpoint signaling via TopBP1. Genes Dev. 2007;21: 1472–1477.
 doi:10.1101/gad.1547007
- Gatei M, Sloper K, Sörensen C, Syljuäsen R, Falck J, Hobson K, et al. Ataxia-telangiectasiamutated (ATM) and NBS1-dependent phosphorylation of Chk1 on Ser-317 in response to
 ionizing radiation. J Biol Chem. 2003;278: 14806–14811. doi:10.1074/jbc.M210862200
- Feng W, Jasin M. BRCA2 suppresses replication stress-induced mitotic and G1
 abnormalities through homologous recombination. Nat Commun. 2017;8: 525.
 doi:10.1038/s41467-017-00634-0
- Ying S, Minocherhomji S, Chan KL, Palmai-Pallag T, Chu WK, Wass T, et al. MUS81
 promotes common fragile site expression. Nat Cell Biol. Nature Publishing Group; 2013;15:
 1001–7. doi:10.1038/ncb2773
- Naim V, Wilhelm T, Debatisse M, Rosselli F. ERCC1 and MUS81-EME1 promote sister
 chromatid separation by processing late replication intermediates at common fragile sites
 during mitosis. Nat Cell Biol. Nature Publishing Group; 2013;15: 1008–15.
 doi:10.1038/ncb2793
- 708 32. Palma A, Pugliese GM, Murfuni I, Marabitti V, Malacaria E, Rinalducci S, et al.
 709 Phosphorylation by CK2 regulates MUS81/EME1 in mitosis and after replication stress.
- 710Nucleic Acids Res. Oxford University Press; 2018;46: 5109–5124. doi:10.1093/nar/gky280
- 33. Lukas C, Savic V, Bekker-Jensen S, Doil C, Neumann B, Pedersen RS, et al. 53BP1 nuclear
 bodies form around DNA lesions generated by mitotic transmission of chromosomes under
 replication stress. Nat Cell Biol. Nature Publishing Group; 2011;13: 243–253.
- 714 doi:10.1038/ncb2201
- 34. Lai X, Broderick R, Bergoglio V, Zimmer J, Badie S, Niedzwiedz W, et al. MUS81 nuclease
 activity is essential for replication stress tolerance and chromosome segregation in BRCA2deficient cells. Nat Commun. 2017;8: 15983. doi:10.1038/ncomms15983
- 718 35. Schlacher K, Wu H, Jasin M. A Distinct Replication Fork Protection Pathway Connects

- Fanconi Anemia Tumor Suppressors to RAD51-BRCA1/2. Cancer Cell. Elsevier Inc.;
 2012;22: 106–116. doi:10.1016/j.ccr.2012.05.015
- 36. Schlacher K, Christ N, Siaud N, Egashira A, Wu H, Jasin M. Double-Strand Break RepairIndependent Role for BRCA2 in Blocking Stalled Replication Fork Degradation by MRE11.
 Cell. Elsevier Inc.; 2011;145: 529–542. doi:10.1016/j.cell.2011.03.041
- 37. Lemaçon D, Jackson J, Quinet A, Brickner JR, Li S, Yazinski S, et al. MRE11 and EXO1
 nucleases degrade reversed forks and elicit MUS81-dependent fork rescue in BRCA2deficient cells. Nat Commun. 2017;8: 860. doi:10.1038/s41467-017-01180-5
- 38. Kolinjivadi AM, Sannino V, De Antoni A, Zadorozhny K, Kilkenny M, Técher H, et al.
 Smarcal1-Mediated Fork Reversal Triggers Mre11-Dependent Degradation of Nascent DNA
 in the Absence of Brca2 and Stable Rad51 Nucleofilaments. Mol Cell. 2017;67: 867–881.e7.
 doi:10.1016/j.molcel.2017.07.001
- 39. Bolterstein E, Rivero R, Marquez M, McVey M. The Drosophila Werner exonuclease
 participates in an exonuclease-independent response to replication stress. Genetics.
 2014;197: 643–52. doi:10.1534/genetics.114.164228
- García-Rodríguez N, Morawska M, Wong RP, Daigaku Y, Ulrich HD. Spatial separation
 between replisome- and template-induced replication stress signaling. EMBO J. 2018;
 e98369. doi:10.15252/embj.201798369
- Hashimoto Y, Puddu F, Costanzo V. RAD51- and MRE11-dependent reassembly of
 uncoupled CMG helicase complex at collapsed replication forks. Nat Struct Mol Biol. Nature
 Publishing Group; 2011;19: 17–24. doi:10.1038/nsmb.2177
- 42. Sidorova JM, Kehrli K, Mao F, Monnat R. Distinct functions of human RECQ helicases
 WRN and BLM in replication fork recovery and progression after hydroxyurea-induced
 stalling. DNA Repair (Amst). 2013;12: 128–139. doi:10.1016/j.dnarep.2012.11.005
- 43. Shechter D, Costanzo V, Gautier J. Regulation of DNA replication by ATR: signaling in
 response to DNA intermediates. DNA Repair (Amst). 2004;3: 901–8.
- 745 doi:10.1016/j.dnarep.2004.03.020
- Friedel AM, Pike BL, Gasser SM. ATR/Mec1: coordinating fork stability and repair. Curr
 Opin Cell Biol. 2009;21: 237–244. doi:10.1016/j.ceb.2009.01.017
- Flynn RL, Zou L. ATR: A master conductor of cellular responses to DNA replication stress.
 Trends Biochem Sci. 2011;36: 133–140. doi:10.1016/j.tibs.2010.09.005
- 46. Parplys AC, Seelbach JI, Becker S, Behr M, Wrona A, Jend C, et al. High levels of RAD51
- 751 perturb DNA replication elongation and cause unscheduled origin firing due to impaired
- 752 CHK1 activation. Cell Cycle. 2015;14: 3190–202. doi:10.1080/15384101.2015.1055996

- 47. Pirzio LM, Pichierri P, Bignami M, Franchitto A. Werner syndrome helicase activity is
 essential in maintaining fragile site stability. J Cell Biol. 2008;180: 305–314.
 doi:10.1083/jcb.200705126
- 48. Murfuni I, Basile G, Subramanyam S, Malacaria E, Bignami M, Spies M, et al. Survival of
- the Replication Checkpoint Deficient Cells Requires MUS81-RAD52 Function. Maizels N,
- editor. PLoS Genet. 2013;9: e1003910. doi:10.1371/journal.pgen.1003910

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Figure 1. Loss of WRN exonuclease activity leads to formation of nascent ssDNA which 763 compromises formation of DSBs in response to a low-dose of camptothecin (A) Evaluation of 764 ssDNA by anti-IdU immunofluorescence under non-denaturing condition. Nascent DNA was pre-765 labelled for 15 min with IdU before treatment and labelling remained during treatment with CPT. Dot 766 plots show the mean intensity of ssDNA staining for single nuclei from cells expressing the wild-type 767 (WS^{WT}) or the exo-dead form of WRN (WS^{E84A}). Cells were either left untreated or challenged with 768 50 nM CPT for increasing periods, as indicated. The intensity of the anti-IdU immunofluorescence 769 was measured in at least 200 nuclei from three independent experiments. Values are represented as 770 771 means ±SE. Representative images of ssDNA labelling are shown. (B) Evaluation of nascent ssDNA in cells treated with nuclease inhibitors. Cells were treated with Mirin, C5 or both 30 min before IdU 772 773 labelling and 45 min before CPT treatment for 4 h, and then subjected to the ssDNA assay. The graph shows the mean intensity of IdU fluorescence measured from two independent experiments (n=200), 774 775 data are presented as mean ±SE. Statistical analysis in A and B was performed by the Mann–Whitney test (**P < 0.1; ***P < 0.01; ****P < 0.001) (C) Analysis of DSB accumulation by the neutral 776 777 Comet assay. Cells were treated or not with CPT 50 nM for the indicated time, or with 5µM CPT (high-dose) for 1 h, and then subjected to the neutral Comet assay. Where indicated, cells were pre-778 treated with Mirin, C5 or both. In the graph, data are presented as mean tail moment \pm SE from two 779 independent experiments (ns = not significant; **P < 0.1; ***P < 0.01; ****P < 0.001; ANOVA 780 test). Representative images from the neutral Comet assay are shown. 781

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Figure 2. Loss of WRN exonuclease stimulates engagement of RAD51 after CPT. (A) WB 783 analysis of chromatin association of RAD51 and RPA32 in wild-type (WS^{WT}) and in cells expressing 784 the exo-dead mutant form of WRN (WS^{E84A}). Cells were treated or not with CPT for 4 h. LaminB1 785 was used as loading control. The blot is representative of three replicates. The graphs show the 786 quantification of the amount of RAD51 or RPA32 normalised against LaminB1 (mean±SE). (B) 787 Quantitative immunofluorescence analysis of RAD51 foci in WSWT and WSE84A cells. Cells were 788 treated with CPT 50nM for 4h, triton-extracted and subjected to RAD51 immunostaining. Graph 789 shows the intensity of RAD51 immunostaining for each cell with scorable foci (n>3). Values are 790 presented as means \pm SE (*** P < 0.01; **** P < 0.001; Mann–Whitney test). Representative images 791 are shown. 792

Figure 3. RAD51 recruitment persisted during recovery from CPT in the absence of the WRN 794 exonuclease. (A) WB analysis of chromatin association of RAD51, RPA70 and RPA32 in wild-type 795 (WS^{WT}) and in cells expressing the exo-dead mutant form of WRN (WS^{E84A}). Cells were treated or 796 not with 50nM CPT for 4h followed by recovery as indicated. LaminB1 was used as loading control. 797 The blot is representative of three replicates. The graphs show the quantification of the amount of 798 RAD51 or RPA32 normalised against LaminB1 (mean±SE). (B) Quantitative immunofluorescence 799 analysis of RAD51 foci in WSWT and WSE84A cells. Cells were treated with CPT 50nM for 4h and 800 recovered or not as indicated. Graph shows the intensity of RAD51 immunostaining for each cell 801 with scorable foci (n>3). Values are presented as means \pm SE (*** P < 0.01; **** P < 0.001; Mann– 802 Whitney test). Representative images are shown. (C) Quantitative immunofluorescence analysis of 803 RAD51 foci in WS^{WT} and WS^{E84A} cells pre-treated with the nuclease inhibitors. Cells were pre-treated 804 with the indicated inhibitors prior to be challenged with CPT 50nM for 4h and recovered or not as 805 806 indicated. Graph shows the intensity of RAD51 immunostaining for each cell with scorable foci (n>3). Values are presented as means \pm SE (** P < 0.1; **** P < 0.001; Mann–Whitney test). (D-E) 807 808 analysis of parental ssDNA. Parental DNA was labelled with IdU as indicated in the experimental scheme (D). (E) The graph shows the amount of parental ssDNA calculated as mean intensity of IdU 809 810 fluorescence measured from two independent experiments (n=200), data are presented as mean \pm SE. Statistical analysis was performed by the Mann–Whitney test (**P < 0.1; ****P < 0.001). 811 Representative images are shown. 812

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Figure 4. RAD51 inhibition does not impair replication fork recovery following treatment with 814 low dose of CPT. (A) Experimental scheme of dual-labelling replication assay with DNA fibres. Red 815 tract: CldU; Green tract: IdU. (B) The graph shows the average number of stalled forks (red only 816 tracts opposed to the active forks marked as red+green tracks) after recovery from 50nM CPT 817 treatment. Where indicated, RAD51 inhibitor (B02; RAD51i) was added to cultures together with 818 CPT and during the IdU pulse. Data are presented as mean ± SE. One-hundred IdU-positive tracts 819 were analysed in each experimental point (n=2). (C) The graph shows the average number of new 820 821 origins (green only tracts) after recovery from 50nM CPT treatment. Where indicated, RAD51 inhibitor (B02; RAD51i) was added to the cultures together with CPT and during the IdU pulse. 822 Values are presented as mean \pm SE. In B and C statistical analysis was performed by Anova test. (D) 823 Representative DNA fibres fields are shown in the images. 824

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Figure 5. Loss of WRN exonuclease activity affects phosphorylation of CHK1. (A) WB analysis
 of CHK1 phosphorylation at S345 in wild-type (WS^{WT}) and in cells expressing the exo-dead mutant

form of WRN (WS^{E84A}) or the helicase-dead form (WS^{K577M}). Cells were treated or not with 50nM 828 CPT as indicated. Total CHK1 and GAPDH were used as loading controls. The blot is representative 829 of three replicates. Below is reported quantification of p345CHK1 phosphorylation normalised 830 against total CHK1. (B) Immunofluorescence analysis of pS345CHK1 in WS^{WT} and WS^{E84A} cells 831 treated with CPT 50nM for 4h. Numbers in insets represent the mean percentage of pS345CHK1-832 positive nuclei (n=2; errors are not shown but are < 15% of the mean). (C) Immunofluorescence 833 analysis of pT1989ATR in WSWT and WSE84A treated with CPT 50nM for 4h. Numbers in insets 834 represent the mean percentage of pS345CHK1-positive nuclei (n=2; errors are not shown but are < 835 15% of the mean). (D-E) Analysis of TopBP1 or RAD9 recruitment at nascent ssDNA by PLA. 836 Nascent strand was labelled with IdU for 15min before cells were treated with 50nM CPT for 4h. 837 838 PLA was performed under native conditions using anti-IdU to detect nascent ssDNA and anti-TopBP1 or RAD9 to detect the protein. Negative controls are from samples processed with anti-IdU 839 840 only. The graphs show the number of PLA spots in each nucleus (n=300 from 3 independent replicates). Statistical analysis was performed by the Mann–Whitney test (ns = not significant; ****P 841 842 < 0.001). Representative images are shown. (F) Duplicated samples from D-E were analysed for the presence of nascent ssDNA by native anti-IdU detection only. The graph shows the mean intensity 843 844 of IdU fluorescence measured from two independent experiments (n=200), data are presented as mean \pm SE. Statistical analysis was performed by the Mann–Whitney test (***P < 0.01). 845

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Figure 6. Expression of a phosphomimic CHK1 mutant restores wild-type levels of nascent ssDNA in WRN exonuclease-deficient cells. (A) WB analysis of FLAG-CHK1^{317/345D} expression in WS^{E84A} cells. (B) Evaluation of nascent ssDNA formation. Cells treated with 50nM CPT for 4h were analysed for the presence of nascent ssDNA by native anti-IdU detection. The graph shows the mean intensity of IdU fluorescence measured from two independent experiments (n=200), data are presented as mean ±SE. Statistical analysis was performed by the Mann–Whitney test (*P < 0.5; ***P < 0.01; ****P < 0.001). Representative images are shown.

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Figure 7. WRN exonuclease-deficient cells show enhanced MUS81 phosphorylation on S87 and mitotic defects. (A) Anti-pS87MUS81 immunofluorescence staining (red) was performed in wildtype and WRN exonuclease-dead expressing cells. The S-phase cells (green) were revealed with short EdU pulse followed by Click-IT reaction. Nuclei were depicted with DAPI staining (blue). The mean frequency (±SE; n=3) of pS87-MUS81-positive nuclei are indicated in the representative images. (B) WS cells expressing the WS^{E84A} mutant were treated with CPT for 4h and then released in fresh medium for 18 hours. The RAD51 inhibitor B02 was added with CPT and during the recovery. The

frequency (± SE; n=2) of pS87-MUS81-positive nuclei are indicated as percentage in the 862 representative images. (C) The graph shows the mean percentage \pm SE of bulky anaphase bridges 863 analysed in untreated and CPT-treated cells expressing WRN wild-type and the exonuclease-deficient 864 mutant. The number of anaphases counted for each experimental point are indicated above as n. 865 Randomly-selected representative anaphases with bridges are shown. (D) Representative images of 866 mitotic aberrations analysed in α-Tubulin (green) and DAPI-stained cells are shown above the graph 867 indicating the frequency of each event after treatment with 50nM of CPT for 4h followed by a 18h 868 recovery. Data are presented as mean±SE. Statistical analysis was performed by the ANOVA test (*P 869 870 < 0.5). (E) Analysis of 53BP1 NBs. Cells were either untreated or treated with 50nM CPT as indicated. Samples were subjected to immunofluorescence using anti-53BP1 and anti-Cyclin A to 871 evaluate 53BP1 fluorescence only in G1 cells (Cyclin A-negative). Graphs show the frequency of 872 each class of nuclei in two independent replicates. Representative images from CPT-treated samples 873 874 are shown.

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Figure 8. MUS81^{S87A} mutant overexpression aggravates the mitotic phenotypes of WRN 876 exonuclease-deficient cells. (A) Bulky bridges, multinucleated and micro-nucleated cells were 877 878 analysed with or without MUS81^{S87A} mutant overexpression, in WRN wild-type and exonucleasedeficient cells. The graph represents the frequency of the aberration analysed in two independent 879 experiments ± SE. (B) Analysis of 53BP1 NBs formation in Cyclin A-negative cells. Representative 880 images of fluorescence cells stained with anti-53BP1 (green) and Cyclin A (red). Nuclear DNA was 881 counterstained with DAPI (blue). For each point at least 300 nuclei were counted and cells with > 5882 53BP1 NBs were considered as positive. The graph shows the quantification of 53BP1-positive G1 883 cells. (C) Clonogenic assay in cells treated with low-doses of CPT. Cells were exposed to different 884 doses of CPT for 18h, re-plated at low density and survival evaluated as percentages of colonises 885 normalised against the untreated. Statistical analyses in A-C were performed by ANOVA test (* P <886 0.5; ** P < 0.1). 887

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Figure 9. Proposed model of the effect of prolonged treatment with nanomolar CPT doses in
absence of WRN exonuclease (see text for details).

892 SUPPORTING INFORMATION LEGENDS

893

894 Supplementary Figures and Legends. The file contains five supplementary figures and their
895 legends.





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Figure 1



В



А





●ws^{w™}





В



С



Figure 3



Е

D





Figure 3 cont'd







D



CPT 1h

А







В

Figure 5

IdU-TOPBP1

CPT



Е

WSE84A















CPT



DAPI/EdU/pS87MUS81

В



DAPI/pS87MUS81

С



4 4 7

Bulky bridges



DAPI/53BP1/CyclinA



Aberrant mitosis



Untreated

E

40

Micronuclei







MUS81^{S87A} + untr CPT DAPI/53BP1/CyclinA

G1 cells with >5 53BP1 NBs



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