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## CRISPR screens identify genomic ribonucleotides as a source of PARP-trapping lesions

- 2 3
- 4 Michal Zimmermann<sup>1</sup>\*, Olga Murina<sup>2</sup>\*, Martin A. M. Reijns<sup>2</sup>, Angelo Agathanggelou<sup>3</sup>, Rachel
- 5 Challis<sup>2</sup>, Žygimantė Tarnauskaitė<sup>2</sup>, Morwenna Muir<sup>4</sup>, Adeline Fluteau<sup>2</sup>, Michael Aregger<sup>5</sup>,
- 6 Andrea McEwan<sup>1</sup>, Wei Yuan<sup>6</sup>, Matthew Clarke<sup>6</sup>, Maryou Lambros<sup>6</sup>, Shankara Paneesha<sup>7</sup>, Paul
- 7 Moss<sup>8</sup>, Megha Chandrashekhar<sup>5,9</sup>, Stephane Angers<sup>10</sup>, Jason Moffat<sup>5,9,11</sup>, Valerie G. Brunton<sup>4</sup>,
- 8 Traver Hart<sup>12</sup>, Johann de Bono<sup>6</sup>, Tatjana Stankovic<sup>3</sup>, Andrew P. Jackson<sup>2</sup>¶ and Daniel
- 9 Durocher<sup>1,9</sup>¶
- 10
- <sup>1</sup>The Lunenfeld-Tanenbaum Research Institute, Mount Sinai Hospital, 600 University Avenue, Toronto, ON, M5G
   1X5, Canada
- <sup>13</sup><sup>2</sup>MRC Human Genetics Unit, Institute of Genetics and Molecular Medicine, University of Edinburgh, Western
- 14 General Hospital, Crewe Road South, Edinburgh EH4 2XU, UK
- <sup>15</sup> <sup>3</sup>Institute for Cancer and Genomic Sciences, University of Birmingham, Edgbaston, Birmingham, B15 2TT, UK.
- <sup>4</sup>Cancer Research UK Edinburgh Centre, University of Edinburgh, Edinburgh, UK.
- <sup>5</sup>Donnelly Centre, University of Toronto, Toronto, ON, M5S 3E1, Canada.
- <sup>6</sup>The Institute of Cancer Research, London, UK and the Royal Marsden NHS Foundation Trust, London, UK.
- <sup>7</sup>Heartlands Hospital, Bordesley Green East, Bordesley Green, Birmingham, West Midlands, B9 5SS, UK.
- <sup>9</sup>Department of Molecular Genetics, University of Toronto, Toronto, ON, M5S 3E1, Canada.
- <sup>10</sup>Department of Pharmaceutical Sciences, Leslie Dan Faculty of Pharmacy & Department of Biochemistry, Faculty
   of Medicine, University of Toronto, Toronto, ON, M5S 3M2, Canada.
- <sup>24</sup><sup>11</sup>Canadian Institute for Advanced Research, Toronto, ON, M5G 1M1, Canada
- <sup>12</sup>Department of Bioinformatics and Computational Biology, The University of Texas MD Anderson Cancer Center,
- <sup>12</sup>Department of Bio
   Houston, TX, USA
- 27
- 28 \*Equal contribution
- 29
- 30 <sup>¶</sup>Address correspondence to:

Daniel Durocher, Ph.D.

The Lunenfeld-Tanenbaum Research Institute Mount Sinai Hospital, Room 1073 600 University Avenue Toronto, ON M5G 1X5 CANADA Tel: 416-586-4800 ext. 2544 e-mail: durocher@lunenfeld.ca Andrew P. Jackson, MRCP Ph.D. MRC Human Genetics Unit Institute of Genetics and Molecular Medicine University of Edinburgh Western General Hospital Crewe Road South Edinburgh EH4 2XU UK e-mail: andrew.jackson@igmm.ed.ac.uk

#### 32 Summary

33 The observation that BRCA1- and BRCA2-deficient cells are sensitive to poly(ADP-ribose) 34 polymerase (PARP) inhibitors spurred their development into cancer therapies that target homologous recombination (HR) deficiency<sup>1</sup>. The cytotoxicity of PARP inhibitors depends 35 on PARP trapping, the formation of non-covalent protein-DNA adducts composed of 36 inhibited PARP1 bound to DNA lesions of unclear origins<sup>1-4</sup>. To address the nature of such 37 38 lesions and the cellular consequences of PARP trapping, we undertook three CRISPR 39 screens to identify genes and pathways that mediate cellular resistance to olaparib, a clinically approved PARP inhibitor<sup>1</sup>. Here were present a high-confidence set of 73 genes 40 whose mutation causes increased PARP inhibitor sensitivity. In addition to an expected 41 42 enrichment for HR-related genes, we discovered that mutation in all three genes encoding 43 RNase H2 sensitized cells to PARP inhibition. We establish that the underlying cause of the 44 PARP inhibitor hypersensitivity of RNase H2-deficient cells is impaired ribonucleotide excision repair (RER)<sup>5</sup>. Embedded ribonucleotides, abundant in the genome of RER-45 46 deficient cells, are substrates for topoisomerase 1 cleavage, resulting in PARP-trapping 47 lesions that impede DNA replication and endanger genome integrity. We conclude that 48 genomic ribonucleotides are a hitherto unappreciated source of PARP-trapping DNA 49 lesions, and that the frequent deletion of RNASEH2B in metastatic prostate cancer and 50 chronic lymphocytic leukemia could provide an opportunity to exploit these findings 51 therapeutically.

52

We carried out dropout CRISPR screens with olaparib in three cell lines of diverse origins, representing both neoplastic and non-transformed cell types (Fig 1a and ED Fig 1a,b). The cell lines selected were HeLa, derived from a human papilloma virus-induced cervical adenocarcinoma; RPE1-hTERT, a telomerase-immortalized retinal pigment epithelium cell line; and SUM149PT, originating from a triple-negative breast cancer with a hemizygous *BRCA1* mutation<sup>6</sup>. SUM149PT cells express a partially defective BRCA1 protein (BRCA1- $\Delta$ 11q)<sup>7</sup> and thus provided a sensitized background to search for enhancers of PARP inhibition cytotoxicity in 60 HR-compromised cells. The screens were performed in technical triplicates, and a normalized 61 depletion score for each gene was computed using  $DrugZ^8$ . To identify high-confidence hits, we 62 used a stringent false discovery rate (FDR) threshold of 1%. To this initial list, we added genes 63 that were found at an FDR threshold of <10% in at least two cell lines. This analysis identified 64, 61 and 116 genes whose inactivation caused sensitization to olaparib in the HeLa, RPE1-65 hTERT and SUM149PT cell lines, respectively, giving a total of 155 different genes 66 (Supplementary Table 1).

67 Out of this list, 13 genes scored positive in all three cell lines and a further 60 genes were 68 common to two cell lines, which we combine to define a core set of 73 high-confidence PARP 69 inhibitor (PARPi)-resistance genes (Fig 1b and Supplementary Table 1). Gene ontology analysis 70 of the 73- and 155-gene sets (Fig 1c and ED Fig 1c, respectively) shows strong enrichment for 71 HR-related biological processes, providing unbiased confirmation that the screens identified 72 bona fide regulators of the response to PARP inhibition. Mapping the 73-gene set on the 73 HumanMine protein-protein interaction data (Fig 1d) generated a highly connected network 74 consisting of DNA damage response genes that include many HR regulators (such as BRCA1, 75 BARD1, BRCA2 and PALB2), components of the Fanconi anemia pathway, as well as the kinases 76 ATM and ATR. Outside or at the edge of the network, we noted the presence of genes encoding 77 the MUS81-EME1 nuclease, splicing and general transcription factors (such as SF3B3/5 and 78 CTDP1) and the three genes coding for the RNase H2 enzyme complex (RNASEH2A, 79 RNASEH2B and RNASEH2C). RNASEH2A/B/C were hits in all three cell lines, with RNASEH2A 80 and B being the two highest-scoring genes, as determined by the mean DrugZ value from the 81 three cell lines (Supplementary Table 1). A similar analysis of the 155-gene set generated an 82 even denser network, with additional genes lying at the periphery of an HR and Fanconi anemia 83 core (ED Fig 1d).

84 Next, we generated RNase H2-null HeLa, RPE1, SUM149PT and HCT116 clonal cell 85 lines using genome editing (denoted as KO; ED Fig 2a-d) and confirmed that RNase H2 86 deficiency caused hypersensitivity to both olaparib and a second clinical-stage PARPi, 87 talazoparib, in all cell lines tested (Fig 2a,b and ED Fig 2e-g, with EC50 values reported in ED 88 Fig. 2h). The RNASEH2A/B-KO cells also exhibited elevated levels of apoptosis after PARP 89 inhibition (ED Fig 2i-l), a phenotype that was particularly prominent with talazoparib treatment 90 (ED Fig 2i-l). Given the strength of the PARPi-induced phenotypes in RNase H2-deficient cells, 91 and since RNase H2 had not been previously linked to the response to PARP inhibition, we 92 sought to determine the mechanism of PARPi sensitization in RNase H2-deficient cells.

93 Since HR deficiency causes PARPi sensitivity, we first considered that RNase H2 might 94 promote HR. Consistent with this possibility, fission yeast cells that combine mutations in RNase H2 and RNase H1 are HR-defective<sup>9</sup>. However, in RNase H2-deficient cells, RAD51 readily 95 96 formed ionizing radiation-induced foci, suggesting efficient recombinase filament assembly (Fig 97 2c,d and ED Fig 3a,b). Furthermore, HR efficiency, as assessed by the direct repeat (DR)-GFP assay<sup>10</sup>, was at near wild-type levels in cells transduced with RNASEH2A and RNASEH2B 98 99 sgRNAs (Fig 2e and ED Fig 3c,d). Thirdly, rather than reduced HR, RNASEH2A-KO cells 100 displayed higher levels of sister chromatid exchanges, reminiscent of the 'hyper-rec' phenotype observed in RNase H2-deficient yeast<sup>11</sup> (Fig 2f). This phenotype was likely due to elevated 101 102 levels of replication-dependent DNA damage, as determined by  $\gamma$ -H2AX staining (Fig 2g and 103 ED Fig 3e-h) and marked poly(ADP-ribosylation) of PARP1 (Fig 2h and ED Fig 3i,j), 104 supporting previous observations of replication-associated genome instability in yeast and mammalian cells deficient in RNase H2<sup>12-14</sup>. 105

The increased levels of sister chromatid exchanges prompted us to test if RNase H2deficient cells required HR for survival. Indeed, we observed synthetic lethality when an sgRNA against *RNASEH2B* was delivered into engineered *BRCA1-KO* and *BRCA2-KO* cell lines in the RPE1-hTERT and DLD-1 backgrounds, respectively (Fig 2i and ED Fig 3k-o).

110 RNase H2 cleaves single ribonucleotides incorporated into DNA, as well as longer RNA:DNA hybrids<sup>15</sup>. To distinguish between these two functions, we carried out cellular 111 112 complementation experiments with variants of RNase H2. The sensitivity of RNASEH2A-KO 113 cells to olaparib was not rescued by either a catalytically-inactive RNase H2 enzyme 114 (RNASEH2A D34A/D169A), or by a separation-of-function mutant (RNASEH2A P40D/Y210A<sup>16</sup>) that retains activity against RNA:DNA hybrids, but not DNA-embedded 115 116 monoribonucleotides (Fig 2j and ED Fig 4). These data indicate that it is likely the removal of 117 genome-embedded ribonucleotides by RER, and not RNA:DNA hybrid cleavage by RNase H2, 118 which protects cells from PARPi-induced cytotoxicity.

119 To determine the genetic basis of the sensitivity of RNASEH2A-KO cells to PARPi, we 120 carried out CRISPR screens to identify mutations that restored resistance to talazoparib in RNase 121 H2-deficient HeLa and RPE1-hTERT cell lines (Fig 3a, ED Fig 5a and Supplementary Table 2). 122 The screens identified a single common gene, PARP1. The genetic dependency on PARP1 for 123 talazoparib- and olaparib-induced cytotoxicity was confirmed in double RNASEH2A-124 KO/PARP1-KO cells (Fig 3b and ED Fig 5b-e), providing evidence that the lethality associated with PARP inhibition requires formation of trapped PARP1-DNA adducts<sup>4</sup>. Consistent with this 125 126 finding, treatment with veliparib, a PARP inhibitor with poor trapping ability<sup>4</sup> induced much less 127 apoptosis than olaparib or talazoparib in RNASEH2A-KO cells (ED Fig 5f).

128 Analysis of DNA content by flow cytometry revealed that *RNASEH2A-KO* cells arrest in

S phase in a PARP1-dependent manner upon talazoparib treatment (Fig 3c and ED Fig 5g). *RNASEH2A-KO* cells also demonstrated elevated levels of talazoparib-induced  $\gamma$ -H2AX and these levels did not decline upon drug removal (Fig 3c and ED Fig 5h). These observations suggest that unresolved DNA lesions induced by PARP trapping are the likely cause of cell death in PARPi-treated *RNASEH2A-KO* cells.

Genome instability in RER-deficient yeast cells is dependent on an alternative, 134 topoisomerase 1 (TOP1)-mediated ribonucleotide excision pathway<sup>18-20</sup>. In this process, TOP1 135 136 enzymatic cleavage 3' of the embedded ribonucleotide results in DNA lesions predicted to engage PARP1, including nicks with difficult-to-ligate 2'-3' cyclic phosphate ends<sup>18,19,21</sup> and 137 covalent TOP1-DNA adducts (TOP1 cleavage complexes<sup>22</sup>) in conjunction with single-strand 138 DNA gaps or DSBs<sup>23</sup>. Given that the mechanisms promoting genome instability in mammalian 139 140 RNase H2-deficient cells remain poorly defined, we assessed whether TOP1 action on 141 misincorporated ribonucleotides contributed to the DNA damage observed in human RER-142 deficient cells. Short-term TOP1 depletion with short interfering RNAs (siRNAs) reduced the 143 number of  $\gamma$ -H2AX foci in RNase H2-deficient cells to nearly wild-type levels (Fig. 3d-f and ED 144 Fig 6a). Furthermore, TOP1-mediated ribonucleotide cleavage contributed to PARPi sensitivity, 145 as depletion of TOP1 with independent siRNAs in RNASEH2A-KO cells reduced the levels of talazoparib-induced apoptosis (Fig 3g and ED Fig 6b-e). TOP1 depletion also reduced 146 147 talazoparib-induced apoptosis in the RER-deficient RNASEH2A P40D/Y210A cells (ED Fig 6f-148 h) and ameliorated the talazoparib-induced S-phase arrest (ED Fig 6i). Together, these results 149 strongly suggest that the processing of genome-embedded ribonucleotides by TOP1 leads to 150 DNA lesions that engage PARP1, creating a vulnerability to PARP trapping.

151 The *RNASEH2B* gene resides on chromosome 13q14 in proximity to two tumour

suppressor loci. One of them, the *DLEU2-mir-15-16* microRNA cluster, is a target of 13q14 deletions observed in over 50% of chronic lymphocytic leukemia (CLL) cases<sup>24</sup>. As a result, collateral homozygous deletion of *RNASEH2B* can occur in CLL and other hematopoietic malignancies<sup>25</sup>. Additionally, in prostate cancer, frequent deletions at 13q14 involving the *RB1* but not the *BRCA2* locus<sup>26</sup> might also result in *RNASEH2B* loss. Such 13q14 deletions are late events associated with endocrine therapy resistance, luminal-to-basal phenotype transition and rapid disease progression<sup>27,28</sup>.

We determined *RNASEH2B* copy number by multiplex ligation-dependent probe amplification (MLPA) in 100 CLL patients. *RNASEH2B* deletions were present in 43% of CLL samples, with biallelic loss detected in 14%. Co-deletion of the *DLEU2* microRNA cluster was confirmed by CGH microarray (Fig 4a and ED Fig 7a,b), establishing that collateral *RNASEH2B* loss is frequent in CLL. Furthermore, analysis of whole-exome sequencing of metastatic castration-resistant prostate cancers<sup>29</sup> demonstrated frequent collateral loss of *RNASEH2B* with *RB1* gene deletion co-occurring in 34% of tumours (2% biallelic loss; ED Fig 7c).

166 The frequent collateral deletion of RNASEH2B prompted us to test whether RNASEH2B 167 loss in cancer cells could be an actionable vulnerability to PARP inhibition. To do so, we 168 performed ex-vivo analysis on primary CLL cells derived from 21 of the 100 patient samples 169 assayed above. Patient characteristics of selected samples were similar across groups (ED Table 170 1). RNase H2 status was confirmed by enzymatic assay of CLL lysates (Fig 4b) and short-term 171 CLL cultures were established from peripheral blood leukocyte samples by stimulating their 172 proliferation with IL21 and co-culture with CD40-ligand expressing MEFs (ED Fig 8a,b and 173 Supplementary Fig. 2). RNASEH2B-deficient cells were found to be significantly more sensitive 174 to PARPi and especially to talazoparib, with the degree of sensitivity correlating with number of 175 *RNASEH2B* alleles lost (Fig 4c and ED Fig 8c).

176 We then asked whether RNase H2 deficiency also confers PARPi sensitivity to tumours 177 in xenograft experiments, utilizing isogenic HCT116 cells with and without RNASEH2A deletion 178 (ED Fig 2a,g,l). Cells were implanted in the flanks of CD1 nude mice and, following 179 establishment of tumours, mice were treated with talazoparib given its higher trapping activity. 180 While talazoparib treatment did not lead to tumor regression, we observed significantly higher 181 sensitivity to talazoparib in tumours lacking RNase H2 (Fig 4d). Furthermore, a second 182 xenograft experiment confirmed this sensitivity to be specific to RNase H2 loss as 183 complementation with an *RNASEH2A* transgene abrogated PARPi sensitivity (ED Fig 8d). Taken 184 together, we conclude that collateral loss of RNase H2 enhances the vulnerability of cancer cells 185 to PARP-trapping drugs.

186 Finally, we note that genome-embedded ribonucleotides are by far the most abundant aberrant nucleotides in the genome of cycling cells<sup>13</sup> and may thus represent a major source of 187 188 the traps that mediate the cytotoxicity of PARPi alongside base excision repair (BER) 189 intermediates. In support of this possibility, RNASEH2A-KO cells are more sensitive to PARPi 190 than isogenic cell lines with homozygous mutations in the catalytic domain of DNA polymerase 191  $\beta$  (*POLB* $\Delta$ *188-190*), a key BER enzyme (ED Fig 9). We therefore propose a model whereby the 192 RER pathway and TOP1 compete for the processing of genome-embedded ribonucleotides (Fig 193 4e). Whereas RNase H2 cleavage initiates their problem-free removal, the action of TOP1 on 194 ribonucleotides create PARP-trapping DNA lesions that impair successful completion of DNA 195 replication and the resulting burden of genomic lesions ultimately causes cell death. We propose 196 that the manipulation of genomic ribonucleotide processing could be harnessed for therapeutic 197 purposes and this strategy may expand the use of PARP inhibitors to some HR-proficient tumors.

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218

#### 219 Author contributions

220 M.Z. performed the initial CRISPR screens with the help of M.A., A.M., M.C., S.A. and J.M;

221 T.H	analyzed the data	. M.Z. and O.M.	performed suppress	sor screens; A.M.	helped with	data
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- analysis. Unless otherwise stated, M.Z. and O.M, with input from M.A.M.R., performed all
- 223 additional experiments and data analysis. M.A.M.R. performed biochemical characterization of
- 224 RER-deficient RNase H2, and together with Ž.T. and A.F. contributed to the generation of HeLa
- and HCT116 RNASEH2A-KO cell lines. A.A., under the supervision of T.S., conducted ex-vivo
- 226 CLL studies and CGH arrays. S.P. and P.M. clinically characterized CLL patients and provided
- 227 CLL blood samples. R.C. performed MLPA assays. W.Y., M.C. and M.L., under the supervision
- of J.B., analysed CNA in the RB1-RNASEH2B region in CRPCs. M.M. and O.M., under the
- supervision of V.G.B., conducted xenograft experiments. A.P.J. and D.D. designed and directed
- the study. D.D. and A.P.J. wrote the manuscript with help of M.Z., O.M. and M.A.M.R. and all

authors reviewed it.

232

#### 233 Competing Financial Interests

- 234 DD and TH are advisors to Repare Therapeutics
- 235

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#### 310 FIGURE LEGENDS



334	replication-associated damage and increased PARP1 activation in RNASEH2A-KO cells. g,
335	Quantification of mean $\gamma$ -H2AX immunofluorescent foci number / nucleus in EdU positive (+)
336	and -negative (-) WT and RNASEH2A-KO cells. h, Representative poly(ADP-ribose) (PAR)
337	immunoblot of PARP1 immunoprecipitates (IP) from whole cell extracts (WCE). Mean fold-
338	increase in PARylation between WT and $RNASEH2A-KO$ indicated ( $n = 3$ biologically
339	independent experiments, normalized to immunoprecipitated PARP1 levels). Tubulin and IgG
340	heavy chain, loading controls. i, Synthetic lethality in combined absence of RNase H2 and
341	BRCA1. Quantification of colony formation of BRCA1-proficient (WT) and BRCA1-KO RPE1-
342	hTERT Cas9 TP53-KO cells transduced with sgLacZ or sgRNASEH2B constructs. Open circles,
343	individual values normalized to $sgLacZ$ ; red lines, mean ( $n = 3$ biologically independent
344	experiments). g, PARPi sensitivity is associated with ribonuclease excision repair (RER)
345	deficiency. Survival of olaparib-treated HeLa WT and RNASEH2A-KO cells transduced with
346	indicated FLAG-tagged constructs. Mean $\pm$ SD, normalized to untreated cells ( $n = 3$ biologically
347	independent experiments). Solid lines, nonlinear least squares fit to a three-parameter dose
348	response model. For <b>d</b> , <b>e</b> , and <b>g</b> : open circles, individual values; red lines, mean ( $n = 3$
349	biologically independent experiments; $\geq 100$ ( <b>d</b> , <b>g</b> ) and $\geq 1000$ ( <b>e</b> ) cells / sample / experiment
350	analyzed). P values in d-g and i, unpaired two-tailed t-test. See also ED Fig 2-4.
351	

#### 352 Figure 3. PARPi-induced PARP1 trapping occurs in RER-deficient cells as a result of

353 TOP1-mediated processing of genomic ribonucleotides. a,b, PARP1 is required for PARPi-

- 354 induced toxicity in RNASEH2A-KO cells. CRISPR screens for talazoparib sensitivity suppressors
- in RNASEH2A-deficient HeLa Cas9 and RPE1 Cas9 TP53-KO cell lines. MAGeCK positive 355
- 356 scores for each gene plotted. Colors indicate gene density in each hexagonal bin. b, Percentage

357	of cleaved caspase-3+ cells of indicated genotypes with or without talazoparib treatment
358	measured by flow cytometry (FACS). Open circles, individual experiments; red lines, mean ( $n =$
359	3 biologically independent experiments). c. DNA damage persists on withdrawal of PARPi in
360	RNASEH2A-KO cells. HeLa WT and RNASEH2A-KO cells were treated with talazoparib and
361	released into fresh medium for the indicated times before being processed for $\gamma$ -H2AX
362	immunofluorescence and propidium iodide (PI) staining. Representative ( $n = 3$ biologically
363	independent experiments) $\gamma$ -H2AX (pseudocolor plots) and cell cycle (histograms) IF/FACS
364	profiles shown. <b>d-f</b> , Increased $\gamma$ -H2AX foci formation in <i>RNASEH2A-KO</i> cells depends on TOP1
365	(images representative of $n = 5$ biologically independent experiments). <b>d</b> , HeLa WT and
366	RNASEH2A-KO cells were transfected with non-targeting (siCTRL) or TOP1-targeting (siTOP1)
367	siRNAs. Immunoblot of WCEs, probed for TOP1. Actin, loading control. e, Representative
368	micrographs of HeLa WT and RNASEH2A-KO cells transfected with siCTRL or siTOP1
369	immunostained for $\gamma$ -H2AX. Scale bars, 10 $\mu$ m. <b>f</b> , Quantification of experiments shown in <b>e</b> .
370	Mean number of foci / nucleus / experiment (open circles) with mean of $n = 5$ biologically
371	independent experiments (red lines). $\geq 100$ cells / sample / experiment analyzed. <b>g</b> , TOP1
372	depletion alleviates PARPi-induced apoptosis in RNASEH2A-KO cells. Quantification of cleaved
373	caspase-3+ WT and RNASEH2A-KO cells transfected with indicated siRNAs, with or without
374	talazoparib treatment. Mean $\pm$ SD normalized to untreated cells ( $n = 3$ biologically independent
375	experiments). $\geq$ 10,000 cells / sample / experiment. <i>P</i> values in <b>b</b> , <b>f</b> , <b>g</b> , unpaired two-tailed t-test.
376	See also <b>ED Fig 5</b> and <b>6</b> .
377	

378 Figure 4. Talazoparib selectively suppresses growth of RNase H2 deficient tumours. a-c,

379 PARP inhibitors selectively kill *RNASEH2B*-deficient chronic lymphocytic leukemia (CLL)

380	primary cancer cells. <b>a</b> , <i>RNASEH2B</i> deletion frequency in a panel of 100 primary CLL samples,
381	determined by multiplex ligation-dependent probe amplification (MLPA). <b>b</b> , Reduced RNase H2
382	activity in lysates from CLL samples with monoallelic and biallelic RNASEH2B deletions. Top,
383	substrate schematic. Individual data points, mean of technical duplicates for each sample. Red
384	lines, mean of individual genotypes ( $n = 8$ WT, 4 monoallelic and 9 biallelic deleted biologically
385	independent primary CLL samples). Data normalized to mean of RNASEH2B-WT samples. c,
386	Reduced survival of CLL cells with monoallelic and biallelic RNASEH2B loss following
387	treatment with talazoparib. Individual points, mean $\pm$ s.e.m. ( $n = 8, 4$ and 9 CLL samples as in
388	<b>b</b> ), each analysed in technical triplicates. <i>P</i> -values, unpaired two-tailed t-test ( <b>b</b> ) and two-way
389	ANOVA (c). d, Selective inhibition of <i>RNASEH2A-KO</i> xenograft tumour growth. HCT116
390	TP53-KO RNASEH2A-WT or -KO cells were injected subcutaneously into bilateral flanks of CD-
391	1 nude mice. Mice were randomized to either vehicle or talazoparib (0.333 mg/kg) treatment
392	groups ( $n = 8$ animals / group) and tumour volumes measured twice-weekly. Mean ± s.e.m. <i>P</i> -
393	value, two-way ANOVA. e, Model. Genome-embedded ribonucleotides (R) can be processed by
394	TOP1 as an alternative to RNase H2-dependent RER. DNA lesions that engage PARP1 (black
395	circles) are formed as a result, and PARP inhibitors induce PARP1 trapping on these TOP1-
396	dependent lesions, causing replication arrest, persistent DNA damage and cell death. See also
397	ED Fig 7, 8, ED Table 1 and Supplementary Table 3.

#### 399 EXTENDED DATA FIGURE LEGENDS

400

401 ED Figure 1 | Related to Figure 1. a, Cas9 immunoblot of whole cell extracts (WCEs) from
402 parental HeLa, RPE1-hTERT and SUM149PT cells and clones stably transduced with a

403	lentiviral FLAG-Cas9-2A-Blast construct (representative of $n \ge 2$ biologically independent
404	experiments). Tubulin, loading control. b, Validation of CRISPR/Cas9 gene editing efficiency in
405	Cas9-expressing HeLa, RPE1-hTERT and SUM149PT clones. Cell proliferation was monitored
406	after transduction with a control sgRNA construct (sgLacZ) or sgRNAs targeting essential genes
407	<i>PSMD1</i> , <i>PSMB2</i> and <i>EIF3D</i> <sup>30</sup> . Solid circles, individual values. Bars, mean $\pm$ SD (normalized to
408	sg <i>LacZ</i> , $n = 3$ technical replicates), <b>c</b> , Gene ontology (GO) terms significantly ( $P < 0.05$ ,
409	binomial test with Bonferroni correction) enriched among hits from olaparib screens common to
410	at least two cell lines. Enrichment was analyzed using PANTHER. d, esyN network analysis of
411	interactions between hits common to at least two cell lines. Node size corresponds to mean
412	DrugZ score across cell lines. 77/155 genes are mapped on the network.
413	
414	ED Figure 2   Related to Figure 2a,b. a, CRISPR-mediated inactivation of RNASEH2A or
415	RNASEH2B in the cell lines used in this manuscript. WCEs of indicated cell lines and genotypes
416	were processed for immunoblotting using antibodies against RNASEH2A, RNASEH2B or
417	RNASEH2C. Vinculin, tubulin and GAPDH, loading controls. Representative immunoblots (of
418	$n \ge 2$ biologically independent experiments). <b>b-d.</b> Abolished RNase H2 enzymatic activity and
419	increased levels of genome-embedded ribonucleotides in RNASEH2A-KO cells. b.
420	Representative ( $n = 3$ biologically independent experiments) analysis of total nucleic acids from
421	WT and RNASEH2A-KO HeLa cells treated with recombinant RNase H2 and separated by
422	alkaline agarose gel electrophoresis. Ribonucleotide-containing genomic DNA from
423	<i>RNASEH2A-KO</i> HeLa cells is nicked and therefore has increased electrophoretic mobility <sup>13</sup> . $c$ ,
424	Densitometric quantification of the alkaline gel shown in <b>b</b> . <b>d</b> , Cleavage of an RNase H2-specific
425	double-stranded DNA oligonucleotide with a single incorporated ribonucleotide (DRD:DNA;

426	ribonucleotide position is shown in red) by WT and RNASEH2A-KO WCEs of the indicated cell
427	types was measured using a fluorescence quenching-based assay <sup>31</sup> . Individual values (open
428	circles) with mean (red lines, $n = 3$ biologically independent experiments). e-l, RNase H2
429	deficiency leads to PARPi sensitivity in multiple cell types. e-h, Clonogenic survival assays of
430	the indicated cell lines treated with the indicated PARPi. Mean ±SD, normalized to untreated
431	cells ( $n = 3$ biologically independent experiments). Solid lines, nonlinear least squares fit of the
432	data to a three-parameter dose response model. h. EC50 values for olaparib (left) and talazoparib
433	(right) in the indicated cell lines as determined by nonlinear least squares fitting of data in <b>e</b> , <b>f</b> , <b>g</b>
434	and Fig 2a,b. Bars, EC50 value ± 95% confidence interval. i-l, Increased apoptosis in HeLa
435	RNASEH2A-KO, SUM149PT Cas9 RNASEH2B-KO and HCT116 RNASEH2A-KO cells
436	following PARPi treatment. <b>i</b> , Representative ( $n = 3$ biologically independent experiments)
437	cleaved caspase-3 immunofluorescence / flow cytometry (IF/FACS) profiles of untreated and
438	talazoparib-treated HeLa WT and <i>RNASEH2A-KO</i> cells. FSC = forward scatter. <b>j-l</b> , Percentages
439	of cleaved caspase-3-positive (caspase-3+) cells of the indicated genotypes treated with the
440	indicated PARPi. Individual values (coloured symbols) with mean (solid lines, $n = 3$ biologically
441	independent experiments). Inset: Levels of cleaved caspase-3+ cells without PARPi treatment.
442	Red lines, mean ( $n = 3$ biologically independent experiments). $P$ values, unpaired two-tailed t-
443	test. In <b>a</b> , <b>d</b> , <b>g</b> and <b>l</b> , HCT116 <i>RNASEH2A-KO</i> cells were transduced either with an empty vector
444	(+EV) or a full-length RNASEH2A expression construct (+WT), where indicated.
445	
446	ED Figure 3   Related to Figure 2. a-d, HR is not affected by inactivation of RNase H2. a,

447 Representative micrographs (*n* = 3 biologically independent experiments) of RPE1-hTERT Cas9

*TP53-KO* (WT) and *RNASEH2A-KO* cells exposed to 3 Gy of X-rays (IR) and processed for γ-

449	H2AX and RAD51 immunofluorescence (IF) 4 h later. <b>b.</b> Quantification of the experiment in <b>a</b>
450	at the indicated time points after IR, plotted as percentage of cells with $>5 \gamma$ -H2AX and RAD51
451	colocalizing foci. Individual values (open circles) with mean (red lines, $n = 3$ biologically
452	independent experiments). $P$ values, unpaired two-tailed t-test. <b>c</b> , Representative ( $n = 3$
453	biologically independent experiments) quantitative image-based cytometry (QIBC) plots of DR-
454	GFP experiments in Fig 2e. Each point shows the mean GFP and RNASEH2A IF intensities per
455	nucleus of mock- or I-SceI-transfected HeLa DR-GFP cells transduced with indicated
456	Cas9/sgRNA constructs (EV = empty vector). Dashed lines separate RNASEH2A+/- and GFP+/-
457	cell populations. <b>d</b> , Quantification of RNASEH2A+ cells in DR-GFP experiments shown in $\mathbf{c}$
458	and <b>Fig 2e</b> as determined by QIBC. Individual values (open circles) with mean (red lines; $n = 3$
459	biologically independent experiments). e-h, Replication-dependent endogenous DNA damage in
460	RNase H2-deficient cells. <b>e</b> , Representative ( $n = 3$ biologically independent experiments)
461	micrographs for experiments quantified in Fig 2g. $\gamma$ -H2AX immunofluorescence (IF) in EdU
462	positive (EdU+) and negative (EdU-) WT and RNASEH2A-KO HeLa cells. Scale bars, 5 $\mu$ m. <b>f</b> ,
463	Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>e</b> and <b>Fig 2g</b> . Dots, foci
464	number in individual nuclei. Red lines, mean ( $n = 3$ biologically independent experiments). <b>g,h</b> .
465	HeLa WT and RNASEH2A-KO cells were treated with aphidicolin and EdU as indicated in the
466	schematic (top), and immunostained with antibodies to $\gamma$ -H2AX. Mean number of foci per EdU-
467	positive (EdU+) nucleus in each experiment (g, open circles) or the number of foci in individual
468	EdU+ nuclei ( <b>h</b> , dots). Red lines, mean ( $n = 3$ biologically independent experiments, $\geq 100$ cells /
469	sample / experiment analyzed). P value, unpaired two-tailed t-test. i, j, Increased poly(ADP-
470	ribosylation) of PARP1 in G1 as well as in S/G2/M phases in RNASEH2A-KO cells. i,
471	Representative ( $n = 2$ biologically independent experiments) FACS plots of HeLa WT and

472	RNASEH2A KO cells expressing the FUCCI cell cycle reporters mKO2-Cdt1 and mAG-
473	Geminin <sup>32</sup> . <b>j</b> , PARP1 immunoprecipitates from WCEs of FUCCI-sorted G1 or S/G2/M HeLa
474	WT and RNASEH2A-KO cells, probed with the indicated antibodies in immunoblotting
475	(representative of $n = 2$ biologically independent experiments). Tubulin, loading control.
476	Densitometric quantification of PAR signals normalized to immunoprecipitated PARP1 is shown
477	as fold changes from WT to RNASEH2A-KO cells. <b>k-o</b> , Inactivation of RNase H2 in BRCA1- or
478	BRCA2-deficient backgrounds results in synthetic lethality. k, BRCA1 and BRCA2 expression,
479	respectively, in RPE1-hTERT TP53-KO WT and BRCA1-KO (top) or DLD-1 WT and BRCA2-
480	KO (bottom) cells. WCEs were processed for immunoblotting with the indicated antibodies.
481	Tubulin and KAP1, loading controls. Representative of $n \ge 2$ biologically independent
482	experiments. <b>l</b> , RNase H2 levels in cells used in m, <b>n</b> , <b>o</b> (bottom) and <b>Fig 2i</b> . Cells were
483	transduced with the indicated sgRNA- (top) or Cas9/sgRNA vectors (bottom; EV = empty
484	vector) and processed for RNASEH2A IF. Each point represents mean RNASEH2A intensity per
485	nucleus as measured by QIBC ( $n = 1$ experiment). $\geq 2000$ cells analyzed per sample. Percentages
486	of RNASEH2A+ cells in individual samples are shown above each plot. <b>m</b> , Representative
487	images ( $n = 3$ biologically independent experiments) of clonogenic survival assays quantified in
488	Fig 2i. n, o, Synthetic lethality after inactivation of RNASEH2A or RNASEH2B in BRCA2-
489	deficient cells. Clonogenic survival of DLD-1 WT and BRCA2-KO cells was assessed after
490	transduction with indicated Cas9/sgRNA vectors. <b>n</b> , Representative images of $n = 3$ biologically
491	independent experiments. o, Quantification of the experiment in n. Individual values (open
492	circles) with mean (red lines; $n = 3$ biologically independent experiments). <i>P</i> values, unpaired
493	two-tailed t-test.

495	ED Figure 4   Related to Figure 2j. RNASEH2A P40D/Y210A is a separation-of-function
496	mutant that cannot excise single DNA-embedded ribonucleotides, but cleaves RNA:DNA
497	heteroduplexes (similar to the yeast <i>rnh201-P45D-Y219A</i> mutant <sup>16</sup> ). <b>a</b> , Schematic depicting
498	enzymatic activity against two different RNase H2 substrates (DRD:DNA, dsDNA with
499	embedded ribonucleotide, or RNA:DNA hybrids) in cell lines used in <b>b-d</b> and <b>Fig 2j</b> . WT and
500	RNASEH2A-KO cells were transduced with either an empty vector (EV) or the indicated
501	RNASEH2A constructs. b, Complementation of HeLa RNASEH2A-KO cells with FLAG-tagged
502	RNASEH2A variants restores RNase H2 complex protein levels. WCEs from HeLa WT and
503	RNASEH2A-KO cells stably expressing indicated lentiviral constructs were processed for
504	immunoblotting with the indicated antibodies. Vinculin, loading control. Asterisk indicates a
505	non-specific band. Representative of $n = 3$ biologically independent experiments. c,d,
506	Complementation of HeLa RNASEH2A-KO cells with WT RNASEH2A, but not with the
507	D34A/D169A (catalytic-dead) or P40D/Y210A (separation-of-function) mutants, rescues
508	increased levels of genome-embedded ribonucleotides. $c$ , Total nucleic acids from the cell lines
509	shown in <b>a</b> , <b>b</b> were treated with recombinant RNase H2 and separated by alkaline agarose gel
510	electrophoresis (representative of $n = 4$ experiments). <b>d</b> , Densitometric quantification of alkaline
511	gel shown in c. e, Purified human RNase H2 complexes consisting of RNASEH2B, RNASEH2C
512	and either RNASEH2A WT, P40D/Y210A or D34A/D169A subunits separated by SDS-PAGE
513	and stained with Coomassie Blue ( $n = 1$ ). <b>f-k</b> , RNase H2 activity assays with fluorescein-labeled
514	RNA:DNA substrate (f) or double-stranded DNA with a single incorporated ribonucleotide
515	(DRD:DNA) (g) and increasing amounts of recombinant WT, P40D/Y210A or D34A/D169A
516	RNase H2. Products were separated by polyacrylamide gel electrophoresis and detected by
517	fluorescence imaging. Representative of $n = 3$ biologically independent experiments. <b>h</b> , <b>k</b> ,

518 Quantification of **f**, **g**. Product signal plotted relative to substrate signal per lane. Mean  $\pm$ SD (n =519 3 biologically independent experiments).

520

#### 521 ED Figure 5 | Related to Fig 3a-c. PARP1 trapping is the underlying cause of PARPi 522 sensitivity in RNase H2-deficient cells. a, Schematic representation of CRISPR screens for 523 suppressors of talazoparib sensitivity in RNase H2-deficient cells. Cas9-expressing cells were 524 transduced with the TKOv1 library, talazoparib was added on day 6 (t6; HeLa: 20 nM, RPE1-525 hTERT: 50 nM) and cells were cultured in its presence until day 18 (t18). Cells were subcultured 526 once at day 12 (RPE1) or 13 (HeLa). sgRNA representations in the initial (t6) and final (t18) 527 populations were quantified by next-generation sequencing. Gene knockouts that were enriched at t18 over t6 were identified by MAGeCK<sup>33</sup>. **b**, CRISPR-mediated inactivation of *RNASEH2A* 528 529 and/or *PARP1* in cell lines used in **c-e** and **Fig 3b**. WCEs were processed for immunoblotting 530 with the indicated antibodies. KAP1, loading control. Representative of n = 2 biologically 531 independent experiments. c-e, Loss of PARP1 restores PARPi-resistance in RNASEH2A-KO 532 cells. c, Percentage of cleaved caspase-3+ HeLa cells of indicated genotypes with or without 533 olaparib treatment measured by flow cytometry (FACS). Individual values (open circles) with 534 mean (red lines, n = 3 biologically independent experiments; *P*-value, unpaired two-tailed t-test). 535 d,e. Clonogenic survival assays with HeLa (d) and RPE1-hTERT (e) cells of the indicated 536 genotypes treated with olaparib (left) or talazoparib (right). Mean $\pm$ SD (n = 3 biologically 537 independent experiments). Solid lines, nonlinear least squares fit to a three-parameter dose 538 response model. **f**. Trapping activity of PARPi correlates with the ability to induce apoptosis in 539 RNASEH2A-KO cells. Quantification of cleaved caspase-3-positive HeLa WT and RNASEH2B-540 KO cells without treatment or treated with the indicated PARPi. Individual values with mean

541	(black lines, $n = 3$ biologically independent experiments). Note that PARP-trapping activity
542	decreases as follows: talazoparib > olaparib > veliparib $^{4,17}$ . <b>g</b> , PARPi-induced S-phase arrest in
543	RNASEH2A-KO cells is alleviated in the absence of PARP1. Top, schematic of talazoparib and
544	EdU treatment. Bottom, representative ( $n = 3$ biologically independent experiments) EdU
545	(pseudocolor plots) and DNA content (histograms) FACS profiles of untreated and talazoparib-
546	treated HeLa WT, PARP1-KO, RNASEH2A-KO and PARP1-KO/RNASEH2A-KO cells. DNA
547	content was determined by propidium iodide (PI) staining. <b>h</b> Quantification of mean $\gamma$ -H2AX
548	intensities in experiments shown in Fig 3c. Individual values (open circles) with mean (red lines,
549	$n = 3$ biologically independent experiments, $\geq 10,000$ cells / sample / experiment analyzed).
550	
551	ED Figure 6   Related to Figure 3d-g. TOP1-mediated cleavage at genome-embedded
552	ribonucleotides leads to PARPi sensitivity in RER-deficient cells. a, Reduced endogenous
	i v v
553	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per
553 554	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines,
553 554 555	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi-
553 554 555 556	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi-induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A
553 554 555 556 557	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi- induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b</b> , Representative ( $n = 3$ biologically
553 554 555 556 557 558	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi- induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b</b> , Representative ( $n = 3$ biologically independent experiments) cleaved caspase-3 FACS plots for experiments quantified in <b>Fig 3g</b> .
553 554 555 556 557 558 559	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi- induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b</b> , Representative ( $n = 3$ biologically independent experiments) cleaved caspase-3 FACS plots for experiments quantified in <b>Fig 3g</b> . FSC, forward scatter. <b>c</b> , HeLa WT and <i>RNASEH2A-KO</i> cells were transfected with non-targeting
553 554 555 556 557 558 559 560	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi- induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b</b> , Representative ( $n = 3$ biologically independent experiments) cleaved caspase-3 FACS plots for experiments quantified in <b>Fig 3g</b> . FSC, forward scatter. <b>c</b> , HeLa WT and <i>RNASEH2A-KO</i> cells were transfected with non-targeting (siCTRL-SP) or TOP1-targeting (siTOP1-SP) SMARTpool siRNAs. WCEs analyzed by
553 554 555 556 557 558 559 560 561	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i</b> , TOP1 depletion alleviates PARPi-induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b</b> , Representative ( $n = 3$ biologically independent experiments) cleaved caspase-3 FACS plots for experiments quantified in <b>Fig 3g</b> . FSC, forward scatter. <b>c</b> , HeLa WT and <i>RNASEH2A-KO</i> cells were transfected with non-targeting (siCTRL-SP) or TOP1-targeting (siTOP1-SP) SMARTpool siRNAs. WCEs analyzed by immunoblotting with antibodies to TOP1 and actin (loading control). Representative of $n = 3$
553 554 555 556 557 558 559 560 561 562	DNA damage in TOP1-depleted <i>RNASEH2A-KO</i> cells. Quantification of $\gamma$ -H2AX foci per nucleus in experiments shown in <b>Fig 3e,f</b> . Dots, focus number in individual nuclei. Red lines, mean ( $n = 5$ biologically independent experiments). <b>b-i,</b> TOP1 depletion alleviates PARPi-induced apoptosis and S-phase arrest in HeLa <i>RNASEH2A-KO</i> cells ( <b>b-e</b> ) and in RNASEH2A P40D/Y210A separation-of-function mutant cells ( <b>f-h</b> ). <b>b,</b> Representative ( $n = 3$ biologically independent experiments) cleaved caspase-3 FACS plots for experiments quantified in <b>Fig 3g</b> . FSC, forward scatter. <b>c,</b> HeLa WT and <i>RNASEH2A-KO</i> cells were transfected with non-targeting (siCTRL-SP) or TOP1-targeting (siTOP1-SP) SMARTpool siRNAs. WCEs analyzed by immunoblotting with antibodies to TOP1 and actin (loading control). Representative of $n = 3$ biologically independent experiments. <b>d,</b> Representative ( $n = 3$ biologically independent

564	RNASEH2A-KO HeLa cells after talazoparib treatment. FSC, forward scatter. e, Quantification
565	of the experiment shown in <b>d</b> . <b>f</b> , HeLa <i>RNASEH2A-KO</i> cells stably expressing the indicated
566	FLAG-tagged constructs were transfected with non-targeting (siCTRL) or TOP1-targeting
567	(siTOP1) siRNAs. WCEs were analyzed by immunoblotting with antibodies to TOP1, FLAG
568	and actin (loading control). Representative of $n = 3$ biologically independent experiments. <b>g</b> ,
569	Representative ( $n = 3$ biologically independent experiments) FACS plots of cleaved caspase-3 in
570	siCTRL- or siTOP1-transfected HeLa RNASEH2A-KO cells expressing RNASEH2A-WT or
571	P40D/Y210A mutant. <b>h</b> , Quantification of the experiment shown in <b>g</b> . Data in <b>e,h</b> , mean $\pm$ SD
572	normalized to untreated cells ( $n = 3$ biologically independent experiments, $\geq 10,000$ cells /
573	sample / experiment analyzed; $P$ values, unpaired two-tailed t-test). <b>i</b> , Representative ( $n = 3$
574	biologically independent experiments) cell cycle profiles, prior and post talazoparib treatment, of
575	HeLa WT and RNASEH2A-KO cells transfected with the indicated siRNAs. DNA content was
576	assessed by PI staining and FACS.
577	
578	ED Table 1   Related to Figure 4a-c. Clinical and molecular characteristics of primary CLL
579	samples used in Fig. 4b,c, ED Fig. 7a,b and ED Fig. 8a,b. See table for details.
580	
581	ED Figure 7   Related to Figure 4a-c. Collateral loss of RNASEH2B in CLL and metastatic
582	castration-resistant prostate cancer (CRPC). a, b, Multiplex ligation-dependent probe
583	amplification (MLPA) analysis (a) and comparative genomic hybridization (CGH) array profiles
584	for chromosome 13q (b) of representative CLL samples carrying two wild-type (WT)
585	RNASEH2B alleles (top), a monoallelic RNASEH2B deletion (middle) or biallelic deletion
586	(bottom). a, For MLPA analysis, genomic DNA from reference and experimental samples was

587 analyzed using probes targeting control loci and individual RNASEH2B exons (Exon 1-11). 588 MLPA ratio calculated per probe and normalised to control probes and reference samples. Error 589 bars indicate SD of the mean from 8 control probes for each sample. Dashed lines indicate the 590 threshold set for diploid copy number. **b**, For each CGH array profile the y-axes of the top and 591 bottom plots indicate copy number probe intensity (log R ratio) and the x axes mark the position 592 on chromosome 13 represented by the ideogram (middle). An enlargement of the frequently 593 deleted 13q14.2-14.3 region, including the miRNA-15A/16-1 gene cluster and the RNASEH2B 594 gene, is shown in the bottom plot. n = 1 experiment. c, RNASEH2B is frequently co-deleted with 595 *RB1* in CRPC. Copy number alterations (CNA) in the *RB1-RNASEH2B* region in CRPC (n = 226 596 cases) are shown. Horizontal lines represent the CNA profile for individual CRPC samples (dark 597 blue, homozygous loss; light blue, heterozygous loss; grey, no change; pink, copy number gain 598 (CNA 3-4); red, copy number amplification (CNA > 4); white, insufficient data to determine 599 CNA). Samples are clustered based on RNASEH2B gene status. CNA frequencies for 600 *RNASEH2B* and the *RB1-RNASEH2B* region without a copy number breakpoint are shown on 601 the right.

602

603 **ED Figure 8** | **Related to Figure 4. a,b,** Proliferating cells, and not quiescent cells, are the major 604 population of viable cells in *ex-vivo* cultured primary CLL patient samples irrespective of 605 treatment group. Quantification of absolute (**a**) and relative (**b**) quiescent and proliferating cell 606 numbers as determined by FACS analysis of the primary CLL samples used in **Fig 4b,c**. 607 (*RNASEH2B* WT, n = 8 individual samples; monoallelic deletion, n = 4 individual samples; 608 biallelic deletion, n = 9 individual samples). Mean  $\pm$  SD (n = 3 technical replicate). FACS gating 609 strategy for stimulated peripheral blood lymphocytes (PBLs) from CLL patients is shown in

610	Supplementary Fig 2. c, RNase H2-deficient primary CLL cells have reduced survival when
611	cultured with olaparib. Mean of individual samples $\pm$ s.e.m. ( $n = 3$ biologically independent CLL
612	samples / group, each analyzed in technical triplicates). P-value, two-way ANOVA. d,
613	Talazoparib selectively inhibits the growth of RNASEH2A-KO xenograft tumours. RNASEH2A-
614	KO cells complemented either with empty vector (EV) or RNASEH2A-WT were injected
615	subcutaneously into bilateral flanks of CD-1 nude mice. Mice were randomized to either vehicle
616	or talazoparib (0.333 mg/kg) treatment groups (n = 8 animals / group) and tumour volumes were
617	measured twice-weekly. Data plotted as mean ± s.e.m. P-values. two-way ANOVA under the
618	null hypothesis that talazoparib does not supress the tumour growth.
619	
620	ED Figure 9. RNase H2-deficient cells are more sensitive to PARPi than DNA polymerase $\beta$
621	<b>mutants. a,</b> Schematic representation of the <i>POLB</i> $\Delta$ 188-190 CRISPR mutation. The Mg <sup>2+</sup> -
622	coordinating aspartate residues (D190, D192 and D256, red triangles) are highlighted in the
623	domain structure of the human $Pol\beta$ protein. The sgRNA target site and antibody epitope are
624	indicated by black lines. <b>b</b> , WCEs from parental RPE1-hTERT Cas9 <i>TP53-KO</i> cells and two
625	<i>POLB</i> $\Delta$ <i>188-190</i> clones were processed for immunoblotting with antibodies to Pol $\beta$ and tubulin
626	(loading control). Representative of $n = 2$ biologically independent experiments. <b>c</b> , The
627	POLB 2188-190 mutation impairs base excision repair. RPE1-hTERT Cas9 TP53-KO WT or
628	POLB  2188-190 cells were exposed to different concentrations of methyl-methanesulfonate
629	(MMS) for 24 h, followed by growth in drug-free media for an additional 48 h. Cell viability was
630	determined by the Cell Titer Glo assay. d, Sensitivity of RPE1-hTERT Cas9 TP53-KO WT,
631	RNASEH2A-KO and POLBA188-190 cells to indicated talazoparib concentrations in clonogenic
632	survival assays. Data in <b>c</b> and <b>d</b> represent mean $\pm$ SD, normalized to untreated cells ( $n = 3$

biologically independent experiments). Solid lines denote a nonlinear least-squares fit to a three-parameter dose response model.

635

#### 636 METHODS

#### 637 Cell culture

- 638 HeLa, RPE1-hTERT and 293T cells were purchased from ATCC and grown in Dulbecco's
- 639 Modified Eagle Medium (DMEM; Gibco/Thermo Fisher) supplemented with 10% fetal bovine
- 640 serum (FBS; Wisent), 200 mM GlutaMAX, 1x non-essential amino acids (both Gibco/Thermo
- Fisher), 100U/ml penicillin and 100µg/ml streptomycin (Pen/Strep; Wisent). HCT116 TP53-KO
- 642 cells<sup>34</sup>, a kind gift from B. Vogelstein (Johns Hopkins University School of Medicine), were
- 643 maintained in modified McCoy's 5A medium (Gibco/Thermo Fisher) supplemented with 10%
- 644 FBS and Pen/Strep. SUM149PT cells were purchased from Asterand BioScience and grown in a
- 645 DMEM:F12 medium mixture (Gibco/Thermo Fisher) supplemented with 5% FBS, Pen/Strep, 1
- 646 μg/ml hydrocortisone and 5 μg/ml insulin (both Sigma). DLD-1 WT and BRCA2-KO cells were
- 647 purchased from Horizon and maintained in RPMI media (Gibco/Thermo Fisher) supplemented
- 648 with 10% FBS and Pen/Strep. All cell lines were grown at 37°C and 5% CO<sub>2.</sub> HeLa, RPE1-
- 649 hTERT (with the exception of *BRCA1-KO* and *POLBΔ188-190* clones) and HCT116 cells were
- 650 grown at atmospheric O<sub>2</sub>. RPE1-hTERT BRCA1-KO and POLB Δ188-190 clones, as well as
- 651 DLD-1 and SUM149PT cell lines were maintained at  $3\% O_2$ .
- 652

#### 653 Lentiviral and retroviral transduction

- To produce lentivirus,  $4.5 \times 10^6$  293T cells in a 10-cm dish were transfected with packaging
- 655 plasmids (5 μg pVSVg, 3 μg pMDLg/pRRE and 2.5 μg pRSV-Rev) along with 10 μg of transfer

656	plasmid using calcium phosphate. Medium was refreshed 12-16 h later. Virus-containing
657	supernatant was collected ~36-40 h post transfection, cleared through a 0.4 $\mu$ m filter,
658	supplemented with 4 $\mu$ g/ml polybrene (Sigma) and used for infection of target cells. The TKOv1
659	library virus was prepared as previously described <sup>30</sup> . The following antibiotics were used for
660	selection of transductants: puromycin (HeLa, SUM149PT: 2 $\mu$ g/ml; RPE1-hTERT 15-20 $\mu$ g/ml;
661	each for 48-72 h unless indicated otherwise) and blasticidin (5 $\mu$ g/ml, 4-5 d for all cell lines).
662	Cells stably expressing FLAG-Cas9-2A-Blast were maintained in the presence of 2 $\mu$ g/ml
663	blasticidin.
664	To complement the HCT116 TP53-KO RNASEH2A-KO cell line, cells were infected with
665	retroviral supernatant produced in amphotropic Phoenix packaging cells <sup>35</sup> using either
666	pMSCVpuro empty vector (EV) or pMSCVpuro-RNASEH2A-WT in the presence of 4 $\mu$ g/ml
667	polybrene (Sigma) and 48 h later selected for stable integration using 2 $\mu$ g/ml puromycin.
668	

#### 669 RNASEH2A expression plasmids

670 A FLAG-tagged human RNASEH2A cDNA (NM\_006397.2; encoding amino acids 2-299) and

671 the D34A/D169A mutant<sup>31</sup> were cloned into the pCW57.1 vector (a gift from David Root;

672 Addgene #41393) using the Gateway system (Life Technologies/Thermo Fisher) according to

673 the manufacturer's protocol. The P40D/Y210A mutations were generated by site-directed

674 mutagenesis using the following primers (5'-3'): P40D –

675 GGCCCAGCACGTCGCCCTGCCCG (Forward - F),

676 CGGGCAGGGGCGACGTGCTGGGCC (Reverse - R); Y210A -

677 GTCTTGGGATCATTGGGGGGGCGCCTGAGCCATAATCAGT (F),

678 ACTGATTATGGCTCAGGCGCCCCCAATGATCCCAAGA (R). Expression constructs were

679	introduced into HeLa RNASEH2A-KO cells by lentiviral transduction and expression was
680	induced by the addition of 1 $\mu$ g/ml doxycycline (Clontech) 24 h prior to starting experiments.
681	The pMSCVpuro-RNASEH2A-WT plasmid was generated by cloning the coding sequence of
682	human RNASEH2A into pMSCVpuro-Dest, a Gateway-compatible version of pMSCVpuro
683	(Clontech), and introduced into HCT116 TP53-KO RNASEH2A-KO cells by retroviral
684	transduction.
685	
686	sgRNA target sequences
687	sgRNAs targeting the following sequences (5' to 3') were used to generate CRISPR knockouts.
688	RNASEH2A: TGCCCGCCTCATCGACGCCC and CCCGTGCTGGGTGCGCCCCT (for Hela
689	RNASEH2A-KO), GACCCTATTGGAGAGCGAGC (for HeLa Cas9, RPE1-hTERT, HeLa DR-
690	GFP); RNASEH2B: TCCACCACAACTTGATCAAG; PARP1:
691	TAACGATGTCCACCAGGCCA; BRCA1: AAGGGTAGCTGTTAGAAGGC; POLB:
692	GAGAACATCCATGTCACCAC; LacZ: CCCGAATCTCTATCGTGCGG; PSMD1-1:
693	TGTGCGCTACGGAGCTGCAA; PSMD1-2: ACCAGAGCCACAATAAGCCA
694	PSMB2-1: ATGTTCTTGTCGCCTCCGAC; PSMB2-2: AATATTGTCCAGATGAAGGA;
695	<i>EIF3D-1</i> : TGTAGGTTGCCTCCATGGCC; <i>EIF3D-2</i> : AGACGACCCTGTCATCCGCA; <i>TP53</i> :
696	CAGAATGCAAGAAGCCCAGA.
697	Vectors expressing the Cas9n D10A nickase together with guide RNAs designed against
698	exon 1 and intron 1 of human RNASEH2A were generated by cloning annealed DNA
699	oligonucleotides into pSpCas9n(BB)-2A-GFP and pSpCas9n(BB)-2A-Puro vectors (Addgene
700	plasmid #48140 and #48141, respectively; gifts from Feng Zhang) as previously described <sup>36</sup> . All
701	other sgRNA-expressing constructs were generated by cloning annealed DNA oligonucleotides

- into lentiGuide-Puro or lentiCRISPR v2 vectors (Addgene #52963 and 52961, gifts from Feng
  Zhang) as previously described<sup>37</sup>.
- 704

#### 705 **RNA interference**

- TOP1 was targeted with 40 nM of either a custom siRNA (siTOP1, target site sequence
- 707 AAGGACTCCATCAGATACTAT, Sigma) previously described<sup>38</sup> or an ON-TARGETplus
- 708 SMARTpool siRNA (siTOP1-SP, L-005278-00, Dharmacon/BD Technologies), previously used
- to knock down TOP1<sup>39-41</sup>. A custom siRNA targeting Luciferase (siCTRL,
- 710 CTTACGCTGAGTACTTCGA, Sigma) or an ON-TARGETplus non-targeting pool (siCTRL-
- 711 SP, D-001810-10-05, Dharmacon/BD Technologies) were used as controls<sup>42</sup>. siRNA oligos were
- 712 transfected in Opti-MEM reduced serum medium using Oligofectamine (Life
- 713 Technologies/Thermo Fisher). To improve knockdown efficiency for the ON-TARGETplus
- siRNA, a second round of transfection was conducted after 24 h. Following siRNA transfection,
- 715 cells were seeded either for cell cycle analysis (24 h post last transfection) or for
- 716 immunofluorescence analysis (48 h post transfection) as described below. Knockdown was
- 717 optimised to minimize cell death, while maintaining efficient TOP1 depletion (apoptosis levels  $\leq$
- 718 14% of control transfected cells).
- 719

#### 720 **DNA damaging drugs**

- 721 PARP inhibitors olaparib, talazoparib and veliparib were purchased from Selleck Chemicals.
- 722 Talazoparib for the xenograft experiments was a kind gift of T. Heffernan and N. Feng (The
- 723 University of Texas MD Anderson Cancer Center). Methyl methanesulfonate (MMS) and

aphidicolin were obtained from Sigma. Concentrations and durations of treatment are indicatedin the sections below and in the respective figures.

726

#### 727 Generation of Cas9-expressing cells

728 Cells were transduced with the Lenti-FLAG-Cas9-2A-Blast vector<sup>30</sup> and transductants were

selected with blasticidin. Cells were then seeded at low densities (500-1,000 cells, depending on

cell line) on 15-cm dishes and single colonies were isolated using glass cylinders. Cas9

731 expression was confirmed by immunoblotting and gene editing efficiency was tested as follows:

732 Cells were transduced at a low ( $\sim 0.3$ ) multiplicity of infection (MOI) with either a control

733 LacZ sgRNA construct or sgRNA constructs targeting essential genes PSMD1, PSMB2 and

 $EIF3D^{30}$  followed by puromycin selection. 2.5 x  $10^4$  cells were subsequently seeded in 6-well

plates, medium was exchanged 3 days later and the experiment was terminated at day 6. Cells

were trypsinized, resuspended in media and the live cell count was determined by trypan blue
exclusion on a ViCELL instrument (Beckman Coulter). Cell numbers were plotted relative to

738 sgLacZ-transduced samples.

739

#### 740 Generation of CRISPR knockout cell lines

To establish HeLa and HCT116 *TP53-KO RNASEH2A-KO* cell lines,  $0.5 \times 10^6$  cells were

seeded in 6-well plates and transfected with two vectors encoding both Cas9n and sgRNAs

targeting *RNASEH2A* (derivatives of pSpCas9n(BB)-2A-GFP and pSpCas9n(BB)-2A-Puro)

- via using Lipofectamine 2000 (Life Technologies/Thermo Fisher). 48 h after transfection, single
- 745 GFP-positive cells were sorted into 96-well plates on a BD FACSJazz instrument (BD
- 746 Biosciences) and grown until colonies formed. RNASEH2A-KO clones were selected on the basis

of the size of PCR amplicons from the targeted region to detect clones that underwent editing,

which was subsequently confirmed by Sanger sequencing. Oligonucleotides (5' to 3') used for

749 PCR amplification and sequencing of targeted *RNASEH2A* loci were as follows:

750 ACCCGCTCCTGCAGTATTAG and TCCCTTGGTGCAGTGCAATC. The absence of

functional RNASEH2A was confirmed by immunoblotting, an RNase H2 activity assay and

alkaline gel electrophoresis as described below. Functionally wild-type *RNASEH2A* clones wereidentified in parallel and used as controls.

754 To generate the remaining CRISPR-edited HeLa and RPE1-hTERT cell lines, cells were 755 electroporated with 5 µg of vectors encoding either the sgRNA (lentiGuide-Puro, for cells stably 756 expressing Cas9) or encoding both the sgRNA and Cas9 (lentiCRISPR v2) using an Amaxa Nucleofector II instrument (Lonza). 0.7 x 10<sup>6</sup> RPE1-hTERT cells in a buffer containing 100 mM 757 758 Na<sub>2</sub>HPO<sub>4</sub> (pH 7.75), 10 mM KCl and 11 mM MgCl<sub>2</sub> were electroporated using program T-23. 759 For HeLa cells, the Amaxa Cell Line Nucleofector Kit R (Lonza) was used with program I-13 760 according to the manufacturer's instructions. Cells were re-plated into antibiotic-free McCoy's 761 5A media supplemented with 10% FBS and allowed to recover for 24 h. Puromycin was 762 subsequently added to growth media to enrich for transfectants and removed 24 h later. Cells 763 were then cultured for additional 3-5 days to provide time for gene editing and eventually seeded 764 at low densities (400-1,000 cells, depending on cell line) on 15-cm dishes. Single colonies were 765 isolated using glass cylinders two to three weeks later. SUM149PT Cas9 RNASEH2B-KO cells 766 were generated by transient transfection of parental SUM149PT Cas9 cells with a lentiGuide-767 puro-sgRNASEH2B construct using Lipofectamine 2000 (Thermo Fisher) as per the 768 manufacturers protocol (2  $\mu$ g plasmid DNA and 2  $\mu$ l of Lipofectamine 2000 was used for 1x 10<sup>5</sup>

cells in a 6-well plate). Transfected cells were selected with puromycin for 24 h, grown foradditional 4 days and single clones were isolated as above.

- 771 Targeted clones were identified by immunofluorescence and/or immunoblotting and
- successful gene editing was confirmed by PCR and TIDE analysis (https://tide-

calculator.nki.nl)<sup>43</sup>. The following PCR primers (5' to 3') were used for amplification of targeted

- 174 loci in RNASEH2A: AGATCTGGAGGCGCTGAAAGTGG (F),
- 775 AGTGGCTGTATCATGTGACAGGG (R); RNASEH2B: TAGATGGTGTGTGTGTGG (F),
- 776 TGCTCAGCTTGTCATTGACC (R); BRCA1: TCTCAAAGTATTTCATTTTCTTGGTGCC
- 777 (F), TGAGCAAGGATCATAAAATGTTGG (R); PARP1: AAGCAAACAGGACTGCCAGC
- 778 (F), TACGCCACTGCACTCCAGC (R); POLB: TTACTGTTGTCATCACAGATTCTGC (F),
- 779 AGCAACTCATGGAAGAATAATAGG (R); TP53: GCATTGAAGTCTCATGGAAGC (F);
- 780 TCACTGCCATGGAGGAGC (R).
- 781

#### 782 Generation of HeLa FUCCI WT and RNASEH2A KO cells

783 To establish HeLa WT and RNASEH2A-KO cells expressing the FUCCI cell cycle reporters

784 mKO2-Cdt1 and mAG-Geminin.<sup>32</sup> HeLa WT and *RNASEH2A-KO* cells were transduced at a low

- 785 MOI with pLenti6-mKO2-Cdt1 and pLenti6-mAG-Geminin vectors and transductants were
- selected with 2 µg/ml blasticidin. Subsequently, cells positive for both mKO2-Cdt1 and mAG-
- 787 Geminin fluorescence were collected by sorting on a BD Biosciences FACS Aria II instrument,
- expanded and used for further experiments. Expression of mKO2-Cdt1 and mAG-Geminin was
- 789 confirmed by immunofluorescence and FACS analysis.
- 790

#### 791 CRISPR/Cas9 screening

CRISPR screens were performed as described<sup>30</sup>. Cas9-expressing cells were transduced with the 792 793 lentiviral TKOv1 library at a low MOI (~0.2-0.3) and puromycin-containing media was added 794 the next day to select for transductants. Selection was continued until 72 h post transduction, 795 which was considered the initial time point, t0. At this point the transduced cells were split into 796 technical triplicates. During negative-selection screens (for PARPi sensitizers), cells were 797 subcultured at day 3 (t3) and at day 6 (t6) each of the three replicates was divided into two 798 populations. One was left untreated and an to the other an LD20 dose of olaparib (HeLa:  $2 \mu M$ , 799 RPE1-hTERT: 0.5 µM, SUM149PT: 0.2 µM) was added. Cells were grown with or without 800 olaparib until t15 and subcultured every three days. Sample cell pellets were frozen at each time 801 point for genomic DNA (gDNA) isolation. A library coverage of ≥200 cells/sgRNA was 802 maintained at every step. Positive-selection screens (for suppressors of sensitivity) were carried 803 out in a similar way, but the untreated control was left out, an LD80 dose of talazoparib was used 804 (20 and 50 nM for HeLa and RPE1-hTERT, respectively), cells were subcultured only once after 805 drug addition (t12 – t13) and screens were terminated at t18. Library coverage was  $\geq$ 100 cells / 806 sgRNA.

807 gDNA from cell pellets was isolated using the QIA amp Blood Maxi Kit (Qiagen) and 808 genome-integrated sgRNA sequences were amplified by PCR using the KAPA HiFi HotStart 809 ReadyMix (Kapa Biosystems). i5 and i7 multiplexing barcodes were added in a second round of 810 PCR and final gel-purified products were sequenced on Illumina HiSeq2500 or NextSeq500 811 systems to determine sgRNA representation in each sample. DrugZ (see Related Manuscript 812 *File*) was used to identify gene knockouts, which were depleted from olaparib-treated t15 813 populations but not depleted from untreated cells. Gene knockouts enriched at t18 as compared 814 to t6 in positive-selection screens were identified using MAGeCK<sup>33</sup>.

#### 816 Gene ontology and interaction network analyses

PANTHER (http://pantherdb.org)<sup>44</sup> was used to identify gene ontology (GO) biological 817 818 processes enriched in datasets of screen hits as compared to genome-wide representation. Hits 819 [false discovery rate (FDR)  $\leq 0.01$  in at least one cell line + FDR  $\leq 0.1$  in at least 2 cell lines] 820 from individual cell lines or hits common to at least two cell lines were analyzed with the 821 "statistical overrepresentation test" (GO ontology database released 2017-02-28; annotation data 822 set 'GO biological process complete') with Bonferonni correction for multiple testing. Mapping 823 of the hits on the HumanMine protein interaction network was done through the esyN interface 824 (http://www.esyn.org/). The network was then exported and visualized in Cytoscape version 825 3.4.0 (http://www.cytoscape.org/) and the node sizes adjusted to be proportional to the averaged 826 DrugZ score over the three cell lines.

827

#### 828 Clonogenic survival assays

829 To determine PARPi sensitivity cells were seeded on 10-cm dishes (500-3,000 cells/plate, 830 depending on cell line and genotype) into drug-free media or media containing a range of PARPi 831 concentrations. Cells were either treated either for 2 days with talazoparib followed by additional 832 9-12 days of growth in drug-free media (HeLa, SUM149PT), treated for 7 days with talazoparib 833 followed by 5-6 days in drug-free media (RPE1-hTERT, HCT116), or treated continuously for 834 12-13 days with olaparib. The cultures were incubated at 3% O<sub>2</sub>, with the exception of the 835 experiment in Fig 3g, which was carried out at atmospheric O<sub>2</sub>. Medium (with or without 836 PARPi) was refreshed every 4-7 days in all cases. At the end of the experiment medium was 837 removed, cells were rinsed with PBS and stained with 0.4% (w/v) crystal violet in 20% (v/v)

838 methanol for 30 mins. The stain was aspirated and plates were rinsed 2x in ddH<sub>2</sub>O and air-dried. 839 Colonies were manually counted and data were plotted as surviving fractions relative to 840 untreated cells. To calculate EC50 values the data were fit to a three-parameter dose response 841 model ['log(inhibitor) vs. normalized response'] using the non-linear regression function in 842 Graphpad PRISM v6.0. 843 To analyze the synthetic lethality of combined BRCA1 and RNASEH2B knockouts, 844 RPE1-hTERT Cas9 TP53-KO WT and BRCA1-KO cells were transduced at a high (>1.0) MOI 845 with lentiGuide sgRNA constructs targeting either RNASEH2B or LacZ (control) and seeded for 846 clonal growth 48 h later. WT and BRCA1-KO colonies were grown at 3% O<sub>2</sub> for 12 and 20 days 847 (due to slower growth of BRCA1-deficient cells), respectively. Synthetic lethality between 848 RNase H2 and BRCA2 was assessed by transducing DLD-1 WT and BRCA2-KO cells with 849 either an empty lentiCRISPR v2 vector or lentiCRISPR v2 constructs carrying sgRNASEH2A or 850 sgRNASEH2B. Cells were selected with puromycin and seeded for clonogenic assays 7 days post 851 infection. Clones were grown at 3% O2 for 11 (WT) or 14 days (BRCA2-KO). 852

#### 853 Immunofluorescence microscopy

854 To analyze  $\gamma$ -H2AX focus formation, cells were grown on coverslips for 24 h, incubated in

855 media containing 10 µM EdU for 20 min to label cells undergoing DNA replication, then pre-

extracted for 5 min on ice with ice-cold buffer (25 mM HEPES, pH 7.4, 50 mM NaCl, 1 mM

EDTA, 3 mM MgCl<sub>2</sub>, 300 mM sucrose and 0.5% Triton X-100) and fixed with 4%

- 858 paraformaldehyde (PFA) for 15 min at room temperature (RT). After fixation, cells were washed
- with PBS and blocked in 3% fetal cals serum (FCS) in PBS for 30 min at RT. EdU
- 860 immunolabeling was performed using the Click-iT EdU Imaging Kit (Invitrogen, C10337).

861 Afterwards cells were incubated with a primary mouse antibody against y-H2AX (Millipore 05-862 636; 1:800) for 1.5 h at RT and then stained with anti-mouse secondary antibodies conjugated to 863 Alexa Fluor-568 (Life Technologies) for 1 h at RT. Coverslips were mounted using Vectashield 864 antifade mounting medium with 4,6-diamidino-2-phenylindole (DAPI; Vector Laboratories). For 865 quantification of  $\gamma$ -H2AX foci images were visualized on a Zeiss Axioplan 2 microscope with a 866 40× Plan-neofluar objective, captured using Micro-Manager (http://open-imaging.com/) and analyzed using an ImageJ-based script as previously described<sup>45</sup>. Nuclei were defined on the 867 868 basis of DAPI staining, and  $\gamma$ -H2AX foci were detected using the "Find maxima" function of 869 ImageJ within each nuclear region. Exposure time, binning, microscope settings, light source 870 intensity and the noise level in the "Find maxima" function were kept constant for all the 871 samples within each individual experiment. More than 100 cells were analyzed per condition in 872 each experiment.

For combined  $\gamma$ -H2AX / RAD51 immunofluorescence, 0.25x 10<sup>6</sup> cells were seeded on 873 874 coverslips and ~24 h later were subjected to 3 Gy X-irradiation. 2, 4 or 6 h post irradiation cells 875 were incubated with nuclear extraction buffer (20 mM HEPES pH 7.4, 20 mM NaCl, 5 mM 876 MgCl<sub>2</sub>, 0.5% NP-40, 1 mM DTT and protease inhibitors) for 10 min on ice, rinsed with ice-cold 877 PBS and subsequently fixed with 4% PFA for 10 min at RT. Cells were blocked in IF blocking 878 buffer (10% goat serum, 0.5% NP-40, 0.5% saponin in PBS) for 30 min and incubated with 879 primary antibodies diluted in blocking buffer (Santa-Cruz Biotechnologies rabbit anti-RAD51 / 880 Millipore mouse anti-y-H2AX; 1:150 and 1:2,000, respectively) for 2 h at RT. Cells were then 881 washed with PBS (3x 5 min) and stained with fluorescent secondary antibodies (Alexa Fluor 882 488-conjugated goat anti-rabbit IgG and Alexa Fluor 555-conjugated goat anti-mouse IgG, Life 883 Technologies/Thermo Fisher: 1:1.000 in blocking buffer) and 0.5 ug/ml DAPI for 1 h at RT.

884 Cells were washed as above, mounted in ProLong Gold mounting media (Life

885 Technologies/Thermo Fisher) and imaged used a Zeiss LSM780 laser-scanning microscope.

886 Cells with more than 5 colocalizing γ-H2AX and RAD51 foci were quantified by manual

887 counting. At least 100 cells per condition were analyzed in each experiment.

888

#### 889 Immunofluorescence/flow cytometry (IF/FACS)

For detection of apoptotic cells by cleaved caspase-3 IF/FACS,  $0.5 \times 10^6$  cells were plated on 6-890 891 cm dishes and either left untreated or treated with PARPi for 48 h (PARPi doses are indicated in respective figures). For analysis of apoptotic cells following TOP1 depletion, 0.25 x 10<sup>6</sup> cells 892 893 were plated on 6-cm dishes, transfected with siCTRL or siTOP1 the next day and 24 h post-894 transfection were either left untreated or treated with PARPi for 48 h. Medium was removed and 895 stored in a conical tube, cells were harvested by trypsinization, resuspended in the original 896 conditioned media and centrifuged at 1,500RPM for 5 min at 4°C. Pellets were washed in PBS 897 and fixed in 1 ml 4% PFA for 10 min at RT. Cells were pelleted as above, resuspended in 100 µl 898 PBS and chilled on ice. 900 µl of -20°C methanol was then added drop-wise while gently 899 vortexing. Fixed cells were stored at -20°C overnight or longer.

Before staining, cells were spun down as above, washed in PBS and blocked in IF
blocking buffer (see 'Immunofluorescence' above). Cells were then centrifuged and resuspended
in 200 µl of diluted rabbit anti-cleaved caspase-3 antibody (Cell Signaling #9661; 1:800 in IF
blocking buffer). After 2 h incubation the antibody was diluted with 2 ml PBS, cells were spun
down, and incubated for 1 h in 200 µl Alexa Fluor 488-conjugated goat anti-rabbit secondary
antibody (Molecular Probes/Thermo Fisher, 1:1,000 in IF blocking buffer). The antibody was
diluted with 2 ml PBS, cells were centrifuged, resuspended in 1 ml PBS and cleaved caspase-3

907 signal was analyzed on a BD FACSCalibur or BD LSRFortessa X-20 instruments. Data were
908 analyzed using FlowJo software (Tree Star). See Supplementary Figure 2 for examples of gating
909 strategies.

910 For analysis of recovery from talazoparib-induced replication blockage, cells were treated
911 with 1 μM talazoparib for 24 h, washed extensively with PBS and grown in drug-free media for
912 additional 10, 24 or 34 h. Cells were then harvested, fixed, stained as described above using an
913 anti-γH2AX primary antibody (JBW301, Millipore #05-636, 1:1,000 in blocking buffer) and
914 finally DNA was labeled with propidium iodide (see below).
915
916 Cell cycle analysis by FACS

917  $0.5 \times 10^6$  cells were seeded on 6-cm dishes into media with or without PARPi (doses and

918 durations are indicated in respective figures). Cells were then harvested by trypsinization,

919 resuspended in media and centrifuged (1,000 RPM, 5 min, 4°C). Pellets were resuspended in

920 PBS, centrifuged again and resuspended in 1 ml propidium iodide (PI) staining buffer (20 µg/ml

921 PI, 0.02% Triton X-100, 0.2 mg/ml RNase A in PBS). Cells were stained for 15 min at 37°C and

922 analyzed on a BD FACSCalibur or BD LSR Fortessa X-20 instruments.

For combined PI/EdU staining, cells were treated and harvested as above and fixed in
70% ethanol (added dropwise while gently vortexing) overnight at -20°C. Cells were then
centrifuged as above, washed in PBS and incubated with 10μM Alexa Fluor 488 azide
(Molecular Probes/Thermo Fisher) in a buffer containing 100 mM Tris-HCl pH 8.5, 1 mM
CuSO<sub>4</sub> and 100 mM ascorbic acid for 30 min before centrifugation, washing in PBS and PI
staining. See Supplementary Figure 2 for examples of gating strategies.

#### 930 Sister chromatid exchange assay

931  $0.5 \times 10^{6}$  HeLa cells were seeded in 10-cm dishes and BrdU (final concentration 10  $\mu$ M) was 932 added the next day. BrdU containing-medium was refreshed 24 h later and cells were grown for 933 another 22 h (46 h BrdU incubation in total). 100 ng/ml KaryoMAX colcemid (Gibco/Thermo 934 Fisher) was added for the final 2 h and cells were harvested as follows:

935 Growth medium was removed and stored in a conical tube. Cells were gently washed 936 with 1 ml of trypsin (the trypsin wash was combined with the original media), trypsinized, 937 resuspended in the original conditioned media (+trypsin wash) and centrifuged (1000 RPM, 5 938 min, 4°C). Cells were then washed with PBS, spun down, resuspended in pre-warmed 75 mM 939 KCl and incubated for 30 min at 37°C. Cells were centrifuged again, the supernatant was 940 removed and cells were fixed by drop-wise addition of 1 ml fixative (ice-cold methanol : acetic 941 acid, 3:1) while gently vortexing. An additional 10 ml of fixative was then added and cells were 942 fixed at 4°C for at least 16 h. Once fixed, metaphases were dropped on glass slides, rinsed with 943 fixative and air-dried overnight (protected from light). 944 To visualize sister chromatid exchanges (SCE) slides were rehydrated in PBS for 5 min 945 and stained with 2 ug/ml Hoechst 33342 (Molecular Probes/Thermo Fisher) in 2xSSC (final 300 946 mM NaCl, 30 mM sodium citrate, pH 7.0) for 15 min. Stained slides were placed in a plastic 947 tray, covered with a thin layer of 2xSSC and irradiated with 254 nM UV light ( $\sim$ 5400 J/m<sup>2</sup>). 948 Slides were subsequently dehydrated in a 70%, 95% and 100% ethanol series (5 min each), air-

949 dried and mounted in DAPI-containing ProLong Gold mounting medium (Molecular

950 Probes/Thermo Fisher). Images were captured on a Zeiss LSM780 laser-scanning microscope.

951

#### 952 DR-GFP assay and quantitative image-based cytometry (QIBC)

953 HeLa DR-GFP cells (a gift from R. Greenberg) were transduced with either a lentiCRISPR v2 954 empty vector or sgRNA-expressing constructs targeting RNASEH2A or RNASEH2B. 7 days after transductions 4-5 x  $10^3$  cells were plated per well of 96-well imaging plates (Corning 3603) and 955 956 next day either mock transfected or transfected with either 100 ng of a plasmid expressing I-SceI 957 or a GFP-expressing plasmid (to assess transfection-efficiency) using Lipofectamine 2000. 958 Medium was exchanged 6-8 h later and at 48 h post-transfection cells were fixed in 4% 959 paraformaldehyde. Immunofluorescence for RNASEH2A was performed as described above and 960 plates were imaged on an InCell Analyzer 6000 automated microscope (GE Life Sciences) with a 961 20x objective. Image analysis was performed using Columbus (PerkinElmer). Nuclei were 962 segmented and a sum of DAPI intensity, mean RNASEH2A intensity and mean GFP intensity 963 was quantified for each nucleus. Cells showing a DNA content between 1N and 2N were 964 selected based on DAPI intensity, RNASEH2A-positive and -negative populations were 965 separated and percentages of GFP-positive cells were calculated. Only RNASEH2A+ cells were 966 analyzed in vector-infected samples, whereas only RNASEH2A- cells were considered in 967 sgRNA-transduced samples. Percentages of GFP+ cells in each sample were normalized to the 968 transfection efficiency of a control GFP plasmid.

969

#### 970 Immunoblotting

971 Cell pellets were resuspended in hot 2x sample buffer (166.7 mM Tris-HCl pH 6.8, 2% SDS, 20

972 mM DTT, 10% glycerol, 0.01% bromophenol blue) at a concentration of 5 x  $10^6$  cells/ml and

973 denatured at 95°C for 5 min. An equivalent of 0.25-1 x 10<sup>5</sup> cells was separated by SDS-PAGE

and transferred to a nitrocellulose or PVDF (for RNASEH2B) membrane. Membranes were

blocked with 5% milk / TBST (TBS + 0.1% Tween-20) for at least 1 h at RT and incubated with

976 primary antibodies diluted in 5% milk/TBST overnight at 4°C. Membranes were then washed 3 977 times with TBST, incubated with horseradish peroxidase-conjugated secondary antibodies for 1 978 h at RT, washed again and protein bands were detected using the SuperSignal West Pico 979 enhanced chemiluminescence reagent (Thermo Fisher). 980 To assess the efficiency of TOP1 depletion, whole cell extracts were obtained by lysis 981 and sonication of cells in UTB buffer (8 M urea, 50 mM Tris-HCl, pH 7.5, 150 mM β-982 mercaptoethanol, protease inhibitor cocktail (Roche)). Whole cell extracts for RNase H activity 983 assays and for determining protein levels of the RNase H2 subunits were prepared by lysing cells 984 in 50 mM Tris-HCl pH 8.0, 280 mM NaCl, 0.5% NP-40, 0.2 mM EDTA, 0.2 mM EGTA, 10% 985 glycerol (vol/vol), 1 mM DTT and 1 mM PMSF for 10 min on ice, and subsequent addition of an 986 equal volume of 20 mM HEPES pH 7.9, 10 mM KCl, 1 mM EDTA, 10% glycerol (vol/vol), 1 987 mM DTT and 1 mM PMSF for an additional 10 min. Whole cell extracts were cleared by 988 centrifugation (17,000 g for 10 min at 4°C) and protein concentration was determined by 989 Bradford assay (Protein Assay Kit, BioRad). Protein samples (35 µg total protein) were run on 990 NuPAGE 4-12% Bis-Tris Protein Gels (Thermo Fisher Scientific) and transferred to 991 nitrocellulose or PVDF membranes. Membranes were blocked in 5% milk / TBST (TBS + 0.2% 992 Tween-20) and immunoblotting was performed as described above.

993

#### 994 Immunoprecipitation

995 Cells were collected by trypsinization, washed once with PBS supplemented with 1 µM ADP-

HPD (PARG inhibitor; Enzo) and  $4 \times 10^6$  cells were snap-frozen in liquid nitrogen and then

997 lysed in 1 ml of lysis buffer [50 mM HEPES pH 8.0, 100 mM KCl, 2 mM EDTA, 0.5% NP-40,

998 10% glycerol, 1 mM DTT, complete protease inhibitor cocktail (Roche), 1 μM ADP-HPD].

999 Lysates were incubated with gentle rotation at 4°C for 15 min and then centrifuged at 15,000x g 1000 for 10 min. 50 µl of total cell lysates were used as "input" and 950 µl were incubated with 5 µl of 1001 mouse anti-PARP1 antibody [Enzo (F1-23) ALX-804-211-R050] for 5 h at 4 °C. Protein G-1002 agarose beads (60 µl slurry; Pierce) were added for an additional hour. Beads were collected by 1003 centrifugation, washed four times with lysis buffer and eluted by boiling in 60 µl 2x sample 1004 buffer. Samples were run on an 8% SDS-PAGE gel and immunoblotting was performed as 1005 described above (see 'Immunobloting').

1006 To analyze PARP1 poly(ADP-ribosylation) in a specific phase of the cell cycle, HeLa 1007 FUCCI WT or RNASEH2A-KO were trypsinized, washed once with PBS, collected in tubes with 1008 PBS supplemented with 3% FCS and 1 µM ADP-HPD, and sorted based on mKO2-Cdt1 (G1 1009 phase) and mAG-Geminin (S/G2/M phases) fluorescence on a BD Biosciences FACS Aria II instrument. See Supplementary Figure 2 for examples of gating strategies. 4 x 10<sup>6</sup> FACS-sorted 1010 1011 cells were snap-frozen and lysed as described above. Equivalent amounts of proteins ( $\sim 0.5-1$  mg) 1012 were incubated with 25 µl of PARP1-Trap A pre-equilibrated bead slurry (ChromoTek) for 2.5 1013 h at 4 °C, washed four times with lysis buffer and eluted by boiling in 2x sample loading buffer 1014 (31.25 mM Tris pH 6.8, 25% glycerol, 1% SDS, 0.01% bromophenol blue, β-mercaptoethanol) 1015 prior to immunoblotting. Samples were run on a NuPAGE 4–12% Bis-Tris Protein Gel (Thermo 1016 Fisher Scientific) and immunoblotting was performed as described above.

1017

#### 1018 Antibodies

1019 The following antibodies were used for immunofluorescence (IF) and immunoblotting (IB) at

1020 indicated dilutions: sheep anti-pan-RNase H2 (raised against human recombinant RNase H2<sup>13</sup>,

1021 IB 1:1,000, IP 5 μl / 1 ml lysate); rabbit anti-RNASEH2C (Proteintech 16518-1-AP; IB 1:1,000);

1022	rabbit anti-RNASEH2A	(Origene TA306706.	IB 1:1.000): mouse	anti-RNASEH2A	Abcam
		(			

- 1023 ab92876; IF 1:500); mouse anti-RNASEH2A G-10 (Santa Cruz Biotechnologies sc-515475; WB
- 1024 1:1000); mouse anti-γH2AX JBW301 (Millipore 05-636, IF 1:800 1:2,000); rabbit anti-RAD51
- 1025 H-92 (Santa Cruz Biotechnologies sc-8349, IF 1:150); rabbit anti-BRCA1<sup>46</sup> (IB 1:1,000); mouse
- 1026 anti-Cas9 (Diagenode C15200203, IB 1:1,000); rabbit anti-PARP1 H-250 (Santa Cruz
- 1027 Biotechnologies sc-7150, IB 1:1,000); mouse anti-PAR 10H (Enzo ALX-804-220-R100, IB
- 1028 1:1,000); rabbit anti-Topoisomerase I (Abcam ab109374; IB 1:5,000); rabbit anti-DYKDDDDK
- 1029 (Cell Signaling Technologies 2368, IB 1:1,000); rabbit anti-actin (Sigma A2066, IB 1:5,000);
- 1030 mouse anti-α-tubulin DM1A (Millipore CP06, IB 1:5,000); mouse anti-α-tubulin B512 (Sigma
- 1031 T6074, IB 1:5,000); rabbit anti-GAPDH (Sigma G9545, IB 1:20,000); mouse anti-vinculin
- 1032 (Sigma V9264, IB 1:1,000); rabbit anti-DNA polymerase beta (Abcam ab26343, IB 1:1,000);
- 1033 rabbit anti-cleaved caspase-3 (Cell Signaling Technologies 9661S, IF 1:800).
- 1034

#### 1035 Cell Titer Glo assay

200 cells per condition were plated on 96-well assay plates in technical triplicates either in drugfree media or in a range of MMS concentrations. MMS was washed out 24 h later and cells were
grown in drug-free media for another 48 h. Cell viability was analyzed using the Cell Titer Glo

- 1039 assay kit (Promega) according to the manufacturer's instructions. Luminescence was read on an
- 1040 Envision 2104 plate reader (Perkin Elmer).
- 1041

#### 1042 Detection of ribonucleotides in genomic DNA

1043 Total nucleic acids were isolated from 10<sup>6</sup> cells by lysis in ice-cold buffer (20 mM Tris-HCl pH

1044 7.5, 75 mM NaCl, 50 mM EDTA) and subsequent incubation with 200 µg/ml proteinase K

1045 (Roche) for 10 min on ice followed by addition of sarkosyl (Sigma) to a final concentration of
1046 1%. Nucleic acids were sequentially extracted with TE-equilibrated phenol,

1047 phenol:chloroform:isoamyl alcohol (25:24:1), and chloroform, and then precipitated with

1048 isopropanol. Nucleic acids were collected by centrifugation, washed with 75% ethanol, air-dried

- 1049 and dissolved in nuclease-free water.
- 1050 For alkaline gel electrophoresis, 500 ng of total nucleic acids were incubated with 1 pmol

1051 of purified recombinant human RNase H2<sup>31</sup> and 0.25 µg of DNase-free RNase (Roche) for 30

1052 min at 37°C in 100 μl reaction buffer (60 mM KCl, 50 mM Tris-HCl pH 8.0, 10 mM MgCl<sub>2</sub>,

1053 0.01% BSA, 0.01% Triton X-100). Nucleic acids were ethanol-precipitated, dissolved in

1054 nuclease-free water and separated on a 0.7% agarose gel in 50 mM NaOH, 1 mM EDTA. After

loss electrophoresis, the gel was neutralised in 0.7 M Tris-HCl pH 8.0, 1.5 M NaCl and stained with

1056 SYBR Gold (Invitrogen). Imaging was performed on a FLA-5100 imaging system (Fujifilm),

1057 and densitometry plots generated using an AIDA Image Analyzer (Raytest).

1058

#### 1059 RNase H2 activity assay

1060 Recombinant RNase H2 was expressed in Rosetta-2 *Escherichia coli* cells using a polycistronic
 1061 construct based on pGEX6P1 (pMAR22) and purified as previously described<sup>31</sup>. Site-directed

1062 mutagenesis to introduce the D34A/D169A or P40D/Y210A mutations was performed using the

1063 Quikchange method (Agilent). To measure enzyme activity, a range of RNase H2 concentrations

1064 (0.06 - 2 nM) was incubated with 2  $\mu$ M substrate in 5  $\mu$ l reactions (in a buffer containing 60 mM

1065 KCl, 50 mM Tris-HCl pH 8.0, 10 mM MgCl<sub>2</sub>, 0.01% BSA and 0.01% Triton X-100) at 37°C for

1066 30 min or 1 h. Substrate was formed by annealing a 3'-fluorescein-labelled oligonucleotide

1067 (GATCTGAGCCTGGGaGCT, DRD:DNA, or gaucugagccugggagcu, RNA:DNA; uppercase

1068 DNA, lowercase RNA) to a complementary 5' DABCYL-labelled DNA oligonucleotide

1069 (Eurogentec). Reactions were stopped by adding an equal volume of 96% formamide, 20 mM

1070 EDTA, and heating at 95°C. Products were resolved by denaturing PAGE (20%, 1x TBE),

1071 visualized on a FLA-5100 imaging system (Fujifilm) and quantified using ImageQuant TL (GE

1072 Healthcare).

1073 To assess RNase H2 activity in whole cell extracts a FRET-based fluorescent substrate

1074 release assay was performed as previously described<sup>31</sup>. RNase H2 specific activity was

1075 determined against a DRD:DNA substrate (described above). Activity against a double-stranded

1076 DNA substrate of the same sequence was measured and used to correct for non-RNase H2

1077 activity against the DRD:DNA substrate. Reactions were performed in 100 µl of buffer with 250

1078 nM substrate in 96-well flat-bottomed plates at 25°C. Whole cell lysates were prepared as

1079 described above, and the final protein concentration used per reaction was 100 ng/µl.

1080 Fluorescence was read for 100 ms using a VICTOR2 1420 multilabel counter (Perkin Elmer),

1081 with a 480-nm excitation filter and a 535-nm emission filter.

1082

#### 1083 Ex-vivo CLL studies

Peripheral blood mononuclear cells were isolated from blood samples collected from patients with a new or existing diagnosis of CLL, irrespective of the stage of disease or time/type of treatment from two Birmingham hospitals (Heartlands and Queen Elizabeth). This study was approved by the South Birmingham Ethics Committee (REC number 10/H1206/58), performed according to institutional guidelines and written consent was obtained from all participants.

1089 Primary chronic lymphocytic leukemia (CLL) cells ( $5 \times 10^4$ ) and CD40L-expressing mouse

1090 embryonic fibroblasts (5  $\times$  10<sup>3</sup>) were seeded in each well of a 96-well plate (Corning) in 100  $\mu$ l of

1091 RPMI media supplemented with 10% FBS (Sigma-Aldrich, UK) and 25 ng/ml IL-21 1092 (eBioscience)<sup>47</sup>. After 24 h, 200 µl more media was gently added and cells were incubated for 1093 another 48 h. The media was then aspirated, replaced with 200 ul of media containing 1094 talazoparib and cells were incubated for a further 72 h. The cytotoxic effect of PARPi was 1095 determined by propidium iodide exclusion as measured by flow cytometry with an Accuri C6 1096 flow cytometer (Applied Biosystems). Only cells, which entered the cell cycle upon stimulation 1097 (as determined by forward- and side-scatter FACS profiles), were considered for analysis. Data 1098 was expressed as a surviving fraction relative to untreated cells, for gating strategies see 1099 Supplementary Fig 2.

1100

#### 1101 Multiplex Ligation-dependent Probe Amplification (MLPA) assay

1102 Genomic DNA was isolated from primary CLL cells using the Flexigene kit (Qiagen). To 1103 identify deletions in RNASEH2B gene the MLPA assay was performed on approximately 100 ng 1104 of genomic DNA (gDNA) per sample using the P388-A2 SALSA MLPA kit (MRC-Holland) 1105 according to the manufacturer's protocol. 2 µl of amplified products were separated by capillary 1106 electrophoresis on an ABI PRISM 3130xl Genetic Analyzer (Applied Biosystems) with a 1107 GeneScan 600 LIZ Size Standard (Thermo Fisher). Data were analyzed using GeneMaker 1108 software v2.4.0 (SoftGenetics). Data were normalized using gDNA from 4 control reference 1109 samples. Copy number changes represented as a MLPA ratio were detected by comparing 1110 normalized peak intensities between the reference and the CLL samples. The MPLA ratio 1111 thresholds (X) were set as follows:  $0.75 \ge X \le 1.25$ , diploid sample;  $0.4 \ge X < 0.75$ , monoallelic 1112 deletion; X < 0.4, biallelic deletion. Samples showing either a standard deviation (SD) of control

- 1113 probes above 0.15, or samples with large Q fragment peaks and with more than 4 control
- 1114 probes having MLPA ratios out of diploid range were excluded from the analysis.
- 1115

#### 1116 Comparative Genomic Hybridization (CGH) array

- 1117 Genotyping of CLL samples was accomplished using HumanCoreExome BeadChip arrays
- 1118 (Illumina Inc., San Diego, USA) by UCL Genomics (UCL Great Ormond Street Institute of
- 1119 Child Health, London) in accordance with the Infinium HTS Assay protocol (15045736\_A,
- 1120 Illumina Inc, San Diego, USA). Genotypes were called by GenomeStudio software Genotyping
- 1121 Module v.3.1 (Illumina). A call rate of 98% was accepted as the primary quality control for each
- 1122 sample. Log R Ratio and B Allele Frequency values generated by the GenomeStudio software
- 1123 were used to assess allelic losses in chromosome 13q.
- 1124

#### 1125 Analysis of copy number alterations (CNA) in the RB1-RNASEH2B region in castration-

#### 1126 resistant prostate cancer (CRPC)

- 1127 CRPC (n=226) whole exome sequencing data generated by the International Stand Up To
- 1128 Cancer/Prostate Cancer Foundation Prostate Cancer Dream Team were downloaded and re-
- analysed<sup>29,48</sup>. Paired-end sequencing reads were aligned to the human reference genome
- 1130 (GRCh37/hg19) using BWA (0.5.9), with default settings and re-aligned using stampy (1.0.2).
- 1131 ASCAT (version 2.3) was used to estimate CNA, tumour purity and ploidy.

1132

#### 1133 Xenograft experiments

- 1134 Female athymic CD-1 nude mice (5–7 weeks old, Charles River Laboratories) were used for *in*
- 1135 vivo xenograft studies and quarantined for at least 1 week before experiments. Exponentially

1136	growing HCT116 TP53-KO RNASEH2A-WT or RNASEH2A-KO cells were injected
1137	subcutaneously into the bilateral flanks of each animal $(2x10^6 \text{ cells per flank})$ . Tumours were
1138	measured by caliper every $3 - 4$ days and tumour volume was determined by the formula (length
1139	$\times$ width <sup>2</sup> )/2. When the tumour volumes reached approximately 100 mm <sup>3</sup> (10 days after injection),
1140	mice were randomized into treatment and control groups (8 animals per group, 32 animals in
1141	total; sample size was determined based on previous relevant studies). Talazoparib [BMN673,
1142	0.333 mg/kg, pharmacological grade, a kind gift of T. Heffernan and N. Feng (The University of
1143	Texas MD Anderson Cancer Center)] or vehicle [10% N,N-Dimethylacetamide (ACROS
1144	Organics), 5% Solutol HS 15 (Sigma-Aldrich) in PBS (Gibco)] was administered once daily by
1145	oral gavage (0.1 ml per 10 g of body weight) for the indicated length of time, or until the tumour
1146	reached the maximum size (15 mm in any direction) or ulcerated, or a body conditioning score of
1147	2 was reached, as determined by UK Home Office regulations. The data reported is the average
1148	tumor volume per mouse. Individual flanks that showed no evidence of tumour growth before
1149	initiation of treatment were excluded from subsequent measurements and analysis.
1150	A subsequent experiment was performed by injecting exponentially growing HCT116
1151	TP53-KO RNASEH2A-KO cells complemented either with an empty vector (EV) or a vector
1152	encoding RNASEH2A-WT ( $2x10^6$ cells per flank). To increase the potential treatment window,
1153	mice were randomized into treatment and control groups (8 animals per group, 32 animals in
1154	total), and treatment started 3 days after injection when palpable tumours were formed. The
1155	treatment was administered as described above. Animals that showed no evidence of tumour
1156	growth on both flanks within the first 11 days of treatment were excluded from analysis.
1157	The technician performing tumour measurements was blinded to the experimental
1158	design/identity of cells injected. All animal studies were carried out under Project Licence PPL

1159 70/8897 approved by the UK Home Office and by the University of Edinburgh Animal Welfare 1160 and Ethical Review Body.

1161

#### 1162 **Statistical analysis**

1163 Data were analyzed using a two-tailed Student's t-test and a two-way ANOVA under the 1164 assumption of normal distribution for biological parameters. No corrections for multiple testing 1165

were made. Test used are indicated in respective figure legends. The number of samples (n) in

1166 figure legends represents independent biological replicates, unless stated otherwise. No statistical

1167 methods were used to determine the sample size prior to starting experiments. Cell biology

1168 experiments were not randomized and the investigators were not blinded with regards to sample

1169 allocation and evaluation of the experimental outcome. For xenograft experiments blinding and

1170 randomisation were performed.

1171

#### 1172 Data availability statement

1173 The results of the PARP inhibitor CRISPR screens and source data for mouse xenograft 1174 experiments are included in the on-line version of the manuscript as Supplementary Tables 1, 2 1175 and 3. Unprocessed images of all immunoblots are presented in Supplementary Fig 1. 1176 Supplementary Fig 2 contains examples of gating strategies for FACS experiments. All other 1177 datasets generated during the study are available from the corresponding authors upon reasonable 1178 request.

- 1180 **Methods References**
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GO Biological Process Complete (p < 0.05)



![](_page_59_Figure_2.jpeg)

![](_page_59_Figure_3.jpeg)

![](_page_59_Figure_4.jpeg)

![](_page_59_Figure_5.jpeg)

![](_page_59_Picture_6.jpeg)

![](_page_60_Figure_0.jpeg)

![](_page_60_Figure_1.jpeg)

![](_page_61_Picture_0.jpeg)

f

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-

- -

\_

pan-RNas

ie H2

![](_page_61_Figure_4.jpeg)

k

Products

Product -

![](_page_61_Picture_8.jpeg)

![](_page_61_Figure_9.jpeg)

100-Substrate converted [%] 80 **RNASEH2A WT** P40D/Y210A -• 60-40 20-0 0.5 2.0 0.0 1.0 1.5 RNase H2 concentration [nM]

![](_page_62_Figure_0.jpeg)

С

а

![](_page_62_Figure_2.jpeg)

![](_page_62_Figure_4.jpeg)

f

![](_page_62_Figure_6.jpeg)

![](_page_62_Figure_7.jpeg)

b

![](_page_63_Figure_0.jpeg)

![](_page_63_Figure_1.jpeg)

С

![](_page_63_Figure_2.jpeg)

![](_page_63_Figure_3.jpeg)

d

![](_page_64_Figure_0.jpeg)

а

2.0-

С

![](_page_64_Figure_3.jpeg)

RNASEH2B biallelic deletion

![](_page_64_Figure_4.jpeg)

![](_page_64_Figure_5.jpeg)

![](_page_64_Picture_6.jpeg)

![](_page_64_Figure_7.jpeg)

![](_page_64_Figure_8.jpeg)

![](_page_64_Figure_9.jpeg)

![](_page_64_Picture_10.jpeg)

# 35.4% (n=80/226) RNASEH2B loss

33.6% (n=76/226) RB1-RNASEH2B co-deleted

## 54.9% (n=124/226) RNASEH2B no change

9.7% (n=22/226) RNASEH2B gain

![](_page_64_Picture_15.jpeg)

![](_page_64_Picture_16.jpeg)

![](_page_64_Picture_17.jpeg)

![](_page_65_Figure_0.jpeg)

![](_page_65_Figure_1.jpeg)

![](_page_65_Figure_3.jpeg)

![](_page_65_Figure_5.jpeg)

# Proliferating CLL cells

![](_page_66_Figure_0.jpeg)

![](_page_66_Figure_1.jpeg)

![](_page_66_Picture_2.jpeg)

Polβ (short exposure)  $Pol\beta$  (long exposure) Tubulin

Clinical characteristics							Molecular characteristics					
Sample	Age	Sex	Binet stage	Time from diagnosis (Months)	Treatment	Time on treatment (Days)	Response to treatment	Cytogenetics (FISH)	RNASEH2B status <sup>1</sup>	ATM status <sup>2</sup>	TP53 status <sup>3</sup>	<i>IgVH</i> status <sup>4</sup>
CLL1	67	F	А	35	Pre-treatment	0	-	Trisomy 12	WT	WT	WT	М
CLL2	74	F	А	24	Pre-treatment	0	-	Normal	WT	WT	c.658_663del, c.849_850insC#	UM
CLL3	67	М	А	176	Ibrutinib	0	PRL	Normal	WT	WT	WT	UM
CLL4	68	М	А	49	Pre-treatment	0	-	Normal	WT	WT	WT	М
CLL5	76	М	А	49	Pre-treatment	0	-	N/A	WT	WT	WT	UM
CLL6	65	F	А	153	Pre-treatment	0	-	N/A	WT	WT	WT	М
CLL7	63	F	А	199	Fludarabine+Cyclophosphamide+ Rituximab	37	CR	Trisomy 12	WT	WT	WT	UM
CLL8	39	М	В	80	<b>Pre-treatment</b>	0	-	Normal	WT	WT	WT	М
CLL9	80	F	А	33	Chlorambucil	83	PR	del(13q)	Monoallelic del	WT	WT	М
CLL10	57	F	А	136	<b>Pre-treatment</b>	0	-	del(13q)	Monoallelic del	WT	WT	М
CLL11	79	F	А	70	Bendamustine + rituximab	251	CR	N/A	Monoallelic del	WT	WT	М
CLL12	48	М	В	159	Ibrutinib	486	PR	N/A	Monoallelic del	WT	WT	UM
CLL13	62	F	А	203	Pre-treatment	0	-	N/A	Biallelic del	WT	WT	М
CLL14	63	М	А	27	Pre-treatment	0	-	del(13q)	Biallelic del	WT	WT	UM
CLL15	42	F	А	414	Bendamustine + rituximab +/- ibrutinib	120	SD	del(13q)	Biallelic del	WT	c.561A>G *	М
CLL16	84	F	А	19	<b>Pre-treatment</b>	0	-	N/A	Biallelic del	WT	WT	М
CLL17	72	F	А	153	Chlorambucil	63	PR	Trisomy 12, del(13q)	Biallelic del	WT	c.743G>A*	М
CLL18	79	F	А	36	<b>Pre-treatment</b>	0	-	del(13q)	Biallelic del	WT	WT	М
CLL19	48	F	В	8	Pre-treatment	0	-	del(17p), del(13q)	Biallelic del	WT	c.753_754insCC#	М
CLL20	70	F	В	10	Pre-treatment	0	-	del(13q)	Biallelic del	WT	WT	UM
CLL21	67	М	В	56	Pre-treatment	0	-	del(13q)	Biallelic del	WT	WT	UM

Extended Data Table 1: Clinical and molecular characteristics of primary CLL samples.

CLL samples grouped by *RNASEH2B* status. F, female; M, male; PRL, partial response with lymphocytosis; SD, stable disease; PR, partial response; CR, complete response; "-", not applicable; N/A, not available; WT, <sup>1</sup> Based on MLPA and CGH array, WT, wild-type; del, deleted; <sup>2</sup> WT, intact ATM status confirmed by next-generation sequencing (NGS) and/or functional assays; <sup>3</sup> *TP53* status determined by sequencing, WT, no mutations in coding sequencing; otherwise mutation(s) shown, \* monoallelic *TP53* alteration, # biallelic *TP53* alteration. <sup>4</sup> Maturational status of CLL assessed by detection of hypermutation in Immunoglobulin Heavy chain variable region (IgVH); UM, unmutated >98% sequence homology with germline sequence; M, mutated <98% sequence homology with germline sequence.