1 WRN Inhibition Leads to its Chromatin-Associated Degradation Via the PIAS4-RNF4-p97/VCP Axis

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6 Abstract

7 Synthetic lethality, the concept in which the co-occurrence of two genetic events leads to cell death while 8 either single event alone does not, is an attractive strategy for targeted cancer therapies. A recent 9 example of synthetic lethality as a therapeutic paradigm is the observation that cancer cells with high 10 levels of microsatellite instability (MSI-H) are dependent on the Werner (WRN) RecQ helicase for survival. 11 However, the mechanisms that regulate WRN spatiotemporal dynamics are not fully understood. In this 12 study, we used our single molecule tracking (SMT) platform in combination with a recently disclosed WRN 13 inhibitor to gain insights into WRN's dynamic localization within the nuclei of live cancer cells. We observe 14 that WRN inhibition results in the helicase becoming trapped on chromatin, requiring p97/VCP for 15 extraction and shuttling to the proteasome for degradation. Interestingly, this sequence of events 16 resulting in WRN degradation appears to be MSI-H dependent. Using a phenotypic screen, we identify the 17 PIAS4-RNF4 axis as the pathway responsible for WRN degradation and show that co-inhibition of WRN 18 and SUMOylation has an additive toxic effect in MSI-H cells. Taken together, our work elucidates a novel 19 regulatory mechanism for WRN. Gaining a deeper understanding into this regulatory pathway for WRN 20 can aid in the identification of new high value targets for targeted cancer therapies.

21 Main

Werner Syndrome is a rare genetic condition caused by mutations in the *WRN* gene. It is marked by accelerated aging and a predisposition to a variety of cancers (Opresko et al., 2003). The *WRN* gene encodes a RecQ helicase that plays a critical role in genomic integrity and is unique amongst RecQ
helicases as it possesses an exonuclease domain in addition to its helicase domain (Croteau et al., 2014).
WRN typically resides in the nucleoli of cells but undergoes DNA damage-induced translocation to the
nucleoplasm to perform its repair functions, resolving a diverse array of DNA substrates including D-loops,
replication forks bubble structures, Holliday junctions and other secondary structures. (Constantinou et
al., 2000; von Kobbe and Bohr 2002; Bendtsen et al., 2014). Additionally, WRN is required for telomere
maintenance, a necessary process for maintaining stem cell cellular homeostasis (Shen and Loeb 2001).

31 Synthetic lethality has emerged as an appealing approach to target cancer cells while minimizing 32 collateral damage to otherwise healthy cells and tissue (O'Neil et al., 2017). Synthetic lethality occurs 33 when there is simultaneous disturbance of two essential biological events leading to cell death, which 34 would not occur in the presence of one genetic disruption alone. To highlight the importance of WRN in 35 DNA repair, WRN has been identified as a synthetic lethal target in microsatellite instability high (MSI-H) 36 tumor types that are deficient in mismatch repair (MMR) pathways (Chan et al., 2019; Kategaya et al., 37 2019; Lou et al., 2019; Picco et al., 2021). Microsatellites are short tandem repeats of repetitive 38 nucleotides that reside throughout the genome. Microsatellite stable (MSS) cells have two DNA repair mechanisms - (1) MMR machinery and (2) WRN to ensure genetic integrity. In MSS cells, disruption to 39 40 either MMR or WRN does not lead to cell death. However, in MSI-high cells, MMR processes are 41 compromised so the combination of inhibiting WRN leads to cell death due to this synthetic lethal 42 relationship (Chan et al., 2019; Lieb et al., 2019; Picco et al., 2021). This WRN synthetic modality can be exploited for therapeutic value and has recently entered the clinic for the treatment of MSI-H cancers 43 44 (NCT05838768 2018; NCT06004245 2023). A deeper understanding of DNA replication and repair 45 regulatory mechanisms may provide insights that can be exploited for therapeutic-based purposes. Here, 46 we present the identification of a new WRN degradation mechanism revealed by single molecule tracking 47 (SMT). Upon WRN inhibition, WRN becomes trapped on chromatin and is SUMOylated by PIAS4.

SUMOylated WRN is recognized by the ubiquitin E3 ligase RNF4, which then ubiquitinates WRN leading to
 p97/VCP-mediated chromatin extraction and ultimately proteasomal dependent degradation, providing
 new avenues for therapeutic targeting.

51 To investigate the consequence of WRN inhibition in WRN dynamics, we utilized a recently 52 published clinical WRN inhibitor HRO761 (WRNi, Fig. 1a) (NCT05838768 2018) (Bordas et al., 2022). We 53 first established a panel of MSI-H and MSS cells and tested for WRN sensitivity by depleting WRN using 54 siRNAs (Extended Data Fig. 1a-d). Testing this compound in cells showed a >100-fold induction of DNA 55 damage response in MSI-H cells compared to MSS cells, resulting in apoptosis and cell death (Fig. 1b-d, Extended Data Fig. 1g). Additionally, cellular toxicity was only observed in MSI-H, but not MSS cells, further 56 57 highlighting the specificity of WRNi (Fig. 1e, Extended Data Fig. 1h). To profile the specificity of this 58 compound in vitro, we purified WRN protein and its related homologue, BLM (Extended Data Fig. 1e-f). 59 Using this recombinant system, we observed a 1000-fold difference in ATPase and helicase inhibition with 60 WRN vs BLM (Fig. 1f-g, Extended Data Fig. 1i-j). Further highlighting the accumulation of DNA damage by 61 WRNi, we observed an increase in nuclear size, indicative of DNA damage accumulation and potential cell 62 senescence prior to apoptosis (Extended Data Fig. 2a-b) (Dedov et al., 2003; Rello-Varona et al., 2006; Kang et al., 2010). To enable imaging of proteins at the single molecule level we used CRISPR to generate 63 endogenous HaloTagTM WRN fusion protein (WRN^{Halo}) in MSS (U2OS) and MSI-H (HCT-116) cells 64 65 (McSwiggen et al., 2023). Prior to our imaging studies, we validated the successful tagging of WRN in these cells by depleting the HaloTag[™] WRN signal using a HaloTag[™] PROTAC (Extended Data Fig. 2c-d). 66 Furthermore, comparison between WT and HaloTag[™]-WRN HCT-116 cells showed no discernable 67 68 differences between the two upon WRN inhibition (Extended Data Fig. 2e).

69 WRN undergoes a sub-compartmental translocation in response to DNA damage using standard 70 immunofluorescence (Marciniak et al., 1998; von Kobbe and Bohr 2002). As previously reported, we 71 observed robust translocation of WRN from the nucleolus to the nucleoplasm upon induction of DNA 72 damage (Fig. 2a) (Kamath-Loeb et al., 2017; Veith et al., 2019; Zhu et al., 2021). We employed SMT to 73 understand the link between WRN enzymatic activity mobility in cells using WRNi. Our fully automated 74 SMT platform enables the measurement of thousands of experimental conditions and millions of cells per 75 day (McSwiggen 2023). Recently we introduced substantial improvements to the platform, utilizing a light-76 sheet-based illumination strategy that improves both the throughput and quality of the SMT platform 77 (Driouchi et al., submitted). Upon WRNi treatment, we observed a significant and dose-dependent 78 decrease in average WRN mobility in the HCT-116 background (Fig. 2b-d, Extended Data Fig. 2f). Strikingly, 79 no changes in protein diffusion coefficient were observed in U2OS, suggesting that the microsatellite state 80 of the cell influenced the effect of WRNi on WRN dynamics (Fig. 2c, Extended Data Fig. 2g). Due to the 81 virtue of SMT being a single molecule assay, we next sought to extract a more granular view of dynamic 82 states of WRN in cells. To this end, we plotted the proportion of WRN proteins moving at a particular 83 diffusion rate to generate a "state array" that provides a glimpse of the various states in which WRN is 84 found, ranging from an immobile chromatin-bound state to a freely diffusing state (McSwiggen et al., 85 2023). We observed a decrease in the fastest-diffusing states and more than a 2-fold increase in the 86 chromatin bound fraction of WRN in HCT-116 cells upon WRNi treatment (Fig. 2e-g). In contrast, we did 87 not observe these changes in microsatellite stable U2OS cells or with treatment of general DNA damaging 88 compounds (Fig. 2c and Extended Data Fig. 2g). The observed decrease in WRN diffusion coefficient was 89 subsequently followed by a decrease in SMT spot density, suggesting possible degradation of the WRN 90 protein (Fig. 2h).

91 We confirmed this observation by performing a WRNi time-course which resulted in a reduction 92 of WRN protein levels in HCT-116 cells but not U2OS cells (Fig. 3a-b, Extended Data Fig. 3c). WRN 93 degradation does not result from general DNA damage, but instead is specific to WRNi in a dose-94 dependent and proteasomal dependent way, as co-treatment of HCT-116 with the proteasome inhibitor 95 carfilzomib rescues this effect (Extended Data Fig. 3a-b). Using cycloheximide, we determined that WRN

96 inhibition reduced the half-life of WRN protein by over an order of magnitude, from 16.6h to 1.5h (Fig.
97 3b, and Extended Data Fig. 3d). The increased proportion of chromatin-bound ("bound") WRN molecules,
98 and the concomitant decrease in the freely diffusing ("fast") population, shown in the state array data,
99 suggests that inhibition of WRN by WRNi leads to its trapping on DNA (Fig. 2e-g). This mode of action has
100 been widely studied with PARP1 and its inhibitors (Helleday 2011; Lord and Ashworth 2017).

101 To investigate whether a similar mechanism of action was occurring upon WRN inhibition, we treated WRN^{Halo} cells with WRNi followed by permeabilization with a mild detergent to liberate soluble 102 103 contents of the cells followed by fixation for immunolocalization studies (Illuzzi et al., 2022). These 104 solubilization experiments revealed that treatment with WRNi led to the accumulation of WRN bound to 105 chromatin based on the retention of WRN signal detected in the nucleus after treatment with detergent 106 (Fig. 3d, 3k). Strikingly, we also observed increased colocalization of ubiquitin and WRN upon treatment 107 with WRNi, further fortifying the link between WRN inhibition and the ubiquitin-proteasome pathway 108 (Fig. 3d-f). This link was also validated by using tandem ubiquitin binding entities (TUBEs), which revealed 109 a clear enrichment of higher molecular weight WRN species after WRNi treatment (Fig. 3g). Taken 110 together, these data indicate that WRN is ubiquitylated upon its inhibition. Consistent with the lack of 111 yH2A.X induction observed in U2OS cells upon WRN inhibition, WRNi treatment in this MSS background 112 did not result in WRN trapping or degradation, suggesting that microsatellite instability plays a role in the 113 regulation of WRN upon its inhibition (Fig. 3c, 3h, 3l and Extended Data Fig. 3e). Replication machinery 114 has the potential to become stalled on DNA due to blockade by other bound proteins or DNA lesions (Edenberg et al., 2014; Liao et al., 2018; Le et al., 2023). To mitigate trapping, cells take advantage of 115 116 ubiquitin-mediated proteasomal degradation in which these stalled proteins are marked for degradation 117 by classical proteasome pathways after ubiquitin deposition (Vaz et al., 2013; Franz et al., 2016; Challa et 118 al., 2021). p97/VCP has been implicated in the extraction of ubiguitylated membrane-bound, and 119 chromatin-bound proteins. Therefore, we next investigated if the p97/VCP-proteasome axis was

120 responsible for degradation of WRN (Rape et al., 2001; Jarosch et al., 2002; Wojcik et al., 2004; Meyer et 121 al., 2012). Indeed, upon co-treatment with p97/VCP or proteasome inhibitors we were able to rescue the 122 WRN degradation phenotype induced by WRNi in HCT-116 cells, suggesting an endogenous ubiquitin-123 dependent regulatory paradigm for WRN (Fig. 3i, Extended Data Fig. 3f). This finding was further validated 124 using cellular SMT (Extended Data Fig. 3g) (Kim and Crews 2013; Anderson et al., 2015). Having 125 determined that WRN degradation by WRNi treatment is mediated by the p97/VCP-proteasome axis, we 126 next set out to identify the ubiquitin E3 ligase that activates this process. Depletion of E3 ligases, MIB1 127 and MDM2, that have been reported to regulate WRN did not prevent the degradation of WRN protein 128 upon WRN inhibition (Extended Data Fig. 4a) (Liu et al., 2019; Li et al., 2020). Therefore, we set out to 129 perform a ubiguitome-focused phenotypic siRNA screen and identified six potential genes that upon 130 knock-down rescue WRNi-mediated degradation of WRN or regulate WRN protein levels in other ways 131 (Fig. 4a and Extended Data Fig. 4b). Further validation of these siRNA hits identified RNF4 as the ligase 132 responsible for WRN degradation upon WRNi treatment (Fig. 4a-b). Knock-down of RNF4 showed the most 133 significant rescue of the phenotype, with all RNF4 siRNA oligos rescuing WRN protein levels to at least 134 75% of control treated cells (Extended Data Fig. 4c). We speculate that the partial rescues from the other 135 ubiquitin modulators indicate putative genetic interactors, but the lack of a full rescue effect suggests 136 these are not direct modulators, such as UBE2D3, which is a known E2 conjugating enzyme for RNF4 (Fig. 137 4c and Extended Data Fig. 4c) (DiBello et al., 2016; Roman-Trufero and Dillon 2022). We also observed a 138 rescue of the WRN degradation phenotype using SMT by measuring the density of WRN molecules after 139 RNF4 siRNA knock-down (Extended Data Fig. 4d). Depletion of RNF4 abolished the ubiquitylation of WRN 140 after WRNi treatment, further establishing that RNF4 is responsible for WRN degradation (Fig. 4d-f and 141 Extended Data Fig. 4e). Surprisingly, depletion of RNF4 only slightly rescued the change in WRN diffusion 142 coefficient after inhibition by WRNi, suggesting that WRN remains trapped on chromatin in the absence 143 of ubiquitylation (Fig. 4g). Trapped WRN is detrimental for genome integrity, and failure to remove and

degrade trapped WRN results in further DNA damage, as evidenced by increased yH2A.X accumulation 144 145 following knockdown of RNF4 (Ext data fig 4f). Co-treatment with the E1 ubiquitin activating enzyme 146 inhibitor TAK-234 (E1i) and WRNi led to a slight decrease in the diffusion coefficient of WRN, which could 147 suggest that inhibited WRN remains bound to chromatin if it cannot be ubiquitylated (Hyer et al., 2018) 148 (Fig. 4h). State array analysis revealed that the global inhibition of ubiquitylation by E1i alone led to an 149 increase in the slow diffusing fraction of WRN protein (Fig. 4i, j). This indicates that ubiquitylation is indeed 150 required for the regulation of WRN beyond its function to drive protein degradation as has been shown 151 for various cellular processes (Manford et al., 2020; Rodriguez-Perez et al., 2021; Padovani et al., 2022). 152 However, the state array analysis also revealed that co-treatment with E1i and WRNi lead to a slight but 153 significant increase in the chromatin bound fraction of WRN (Fig. 4i, k). Taken together, these data suggest 154 that WRN inhibition in MSI-H cells leads to an increase in chromatin binding, followed by ubiquitylation 155 and subsequent degradation. This ubiquitylation is necessary to remove WRN from chromatin, as 156 exemplified by the increase in WRN bound to chromatin in the presence of an E1 inhibitor. RNF4 is known 157 as a SUMO-targeted ubiquitin ligase (STUBL) and has been reported to regulate trapped chromatin 158 proteins such as PARP1, DNMT1, and TOP1A (Sun et al., 2020; Liu et al., 2021; Krastev et al., 2022). To 159 investigate if SUMOylation is required for the degradation of inhibited WRN, we depleted cells of the 160 SUMO ligase PIAS4. The depletion of PIAS4 provided a modest, yet significant rescue of WRN degradation 161 upon WRNi treatment (Fig. 5a, 5c). We posit that this failure to completely rescue the degradation 162 phenotype is due to redundancies among the different PIAS proteins. To investigate this, we co-treated 163 cells with the SUMO E1 activating enzyme inhibitor ML-792 (SUMOi) and WRNi, leading to a full rescue of 164 the degradation phenotype by various means of detection, including SMT (Fig. 5b-f and Extended Data 165 Fig. 6a-b) (He et al., 2017). Furthermore, upon SUMOylation inhibition, we failed to pull down 166 ubiquitylated WRN even in the presence of WRNi (Fig. 5g), indicating that the SUMO cascade is necessary 167 to initiate ubiquitylation of WRN. This is evident by the appearance of higher molecular weight WRN,

168 which corresponds to ubiquitylated and SUMOylated WRN, as they disappear upon co-treatment with 169 SUMOi (Fig. 5h). Co-treatment with SUMOi and WRNi lead to a further decrease in WRN diffusion 170 coefficient (Fig. 5i). State array analysis indicated that this decrease was due to an increase in the "slow" 171 fraction of WRN, further suggesting that the SUMO/ubiquitin cascade is important for the regulation of 172 inhibited WRN (Fig. 5j). Indeed, inhibition of the SUMO-Ubiquitin-p97/VCP cascade lead to an increase in 173 chromatin-bound WRN following inhibition by WRNi (Extended Fig. 5a-b). Taken together, these data 174 clearly show that small molecule inhibition of WRN leads to its trapping onto chromatin, leading to its 175 SUMOylation and subsequent ubiquitylation by RNF4 which ultimately results in proteasomal degradation 176 (Fig. 5m). This work has elucidated a novel molecular pathway that regulates WRN activity upon its 177 inhibition. The identification of WRN as a synthetic lethal target in MSI-H cancers has opened new avenues 178 for therapeutic intervention of these malignancies. Clinical medicine has been successful in the selective 179 killing of cancer cells mediated by DNA binding protein inhibition [etoposide citation]. For example, the 180 enzyme, Poly ADP-ribose polymerase (PARP) can induce specific cell toxicity by PARP trapping. PARP 181 carries out its enzymatic function by repairing single-stranded DNA breaks, but inhibition leads to trapping 182 on DNA. As a result, PARP function is attenuated leading to an accumulation of DNA damage, stalled replication forks and cytotoxicity (Shen et al., 2015; Hopkins et al., 2019; Rose et al., 2020). Understanding 183 184 the mechanistic underpinnings of WRN regulation has implications for treating MSI-H cancers in the clinic. 185 WRN that cannot be released from chromatin after trapping is detrimental and leads to persistent DNA 186 damage, even after removal of WRN inhibition (Fig. 5I and Extended Data Fig. 6c-d). This regulatory 187 cascade has various points in which potential therapeutic interventions are possible, such as co-inhibition of the SUMOylation cascade or depletion of RNF4, which results in synergistic shift in the EC₅₀ of WRNi 188 189 (Fig. 5k and Extended Data Fig. 6e-h). Many other areas of potential intervention exist, such as disruption 190 of the translocation of WRN from the nucleolus to the nucleoplasm, or perturbation of the molecular 191 players that recognize chromatin-trapped WRN protein. This work provides deeper insights into the

- 192 regulation of WRN protein activity upon small molecule inhibition that was gained in part through cutting-
- 193 edge super resolution microscopy techniques like cellular SMT. Such detailed understanding will likely be
- 194 critical in defining optimal clinical strategies as first-in-class WRN inhibitors undergo clinical testing.

195 Acknowledgements

- 196 The authors extend their deepest gratitude to all the employees and consultants of Eikon, past and
- 197 present, especially, Pratir Doshi, Nick Vaquera, Jeff Dove, Tiffany Cheng, Roma Moore, Bruno da Rocha
- 198 Azevedo, Madhu Mena, Emily Kirkeby, Puneet Kumar, Dave Piotrowski, Melissa Dumble, Geeta Sharma,
- and Alex Therian. Their tireless work enabled the experiments described here. We thank Roger
- 200 Perlmutter, Robert Tjian for helpful discussions and critical feedback on the direction of our investigation
- and on the resulting manuscript. Eikon Therapeutics provided all funding.

202 Declaration of interests

203 The authors are employees and/or shareholders of Eikon Therapeutics.

204

205 Materials and Methods:

206 Tissue Culture

207 HCT-116 (MSI genotype), U2OS, RKO, SW48, DLD-1 and HT-29 (MSS genotype) cells were used. 208 Cells were cultured in Gibco McCoy's 5A Medium (Thermo, 16600082) containing 10% fetal bovine serum 209 (FBS), 1X GlutaMAX supplement (Thermo, 35050061), 1X MEM Non-Essential Amino Acids solution 210 (Thermo, 11140076) and 50,000 Units/µg of Penicillin/Streptomycin (Thermo, 15140122) at 37°C and 5% 211 CO₂. Cells were seeded on Greiner 384-Well Black Microplate with Optical-Bottom (Greiner, 781906) using 212 a Thermo Multidrop Combi. HCT-116 cells were seeded at 2,500 cells per well and HT-29 cells were seeded 213 at 3,000 cells per well and incubated at 37°C and 5% CO₂ for 24 hours before compound treatment. 214 Compounds were added to the plates using a Beckman Echo 655 acoustic liquid handler. Compound 215 treated plates were incubated for 24 h at 37°C and 5% CO₂.

216 Cell Line Engineering

217 WRN^{Halo} HCT-116 and U2OS were generated by nucleofection of ribonucleoprotein (RNP) complexes (DeWitt et al., 2017) using a guide (5' – AAAGATGAGTGAAAAAAAAT – 3') targeting the N-218 219 terminus of the WRN gene locus and a megamer coding for the Halo tag as a donor template (5' -220 221 222 ACATTGTTTTTTGGACTCTGCAAATACCACATTTCAAAGATGGCAGAAATCGGTACTGGCTTTCCATTCGACCCCCA 223 TTATGTGGAAGTCCTGGGCGAGCGCATGCACTACGTCGATGTTGGTCCGCGCGATGGCACCCCTGTGCTGTTCCTG 224 CACGGTAACCCGACCTCCTCCTACGTGTGGCGCAACATCATCCCGCATGTTGCACCGACCCATCGCTGCATTGCTCC 225 AGACCTGATCGGTATGGGCAAATCCGACAAACCAGACCTGGGTTATTTCTTCGACGACCACGTCCGCTTCATGGAT 226 GCCTTCATCGAAGCCCTGGGTCTGGAAGAGGTCGTCCTGGTCATTCACGACTGGGGCTCCGCTCTGGGTTTCCACT 227 GGGCCAAGCGCAATCCAGAGCGCGTCAAAGGTATTGCATTTATGGAGTTCATCCGCCCTATCCCGACCTGGGACG 228 AATGGCCAGAATTTGCCCGCGAGACCTTCCAGGCCTTCCGCACCACCGACGTCGGCCGCAAGCTGATCATCGATCA

229 230 CGAGCCGTTCCTGAATCCTGTTGACCGCGAGCCACTGTGGCGCTTCCCAAACGAGCTGCCAATCGCCGGTGAGCC 231 AGCGAACATCGTCGCGCTGGTCGAAGAATACATGGACTGGCTGCACCAGTCCCCTGTCCCGAAGCTGCTGTTCTG 232 233 234 TGTCGACGCTCGAGATTTCCGGCGAAAACCTGTATTTTCAGAGCAGTGAAAAAAATTGGAAACAACTGCACAGC 235 236 TGACTATTCTTTTGGGTGAGAAATTTAATTTAATTTGACTGTGCAAAGAGTCAGTTGTTACTTGTAAACTTCAAGT 237 CATTGTTTAGGTCAGAG – 3'). RNPs were assembled with Alt-R[™] S.p. HiFi Cas9 Nuclease V3 (IDT, 1081060) 238 in a 10 µl reaction of 1 µM of Cas9, 120 pmol sgRNA, and 100 pmol ssODN in Cas9 buffer (20 mM HEPES 239 7.5, 150 mM KCl, 10% glycerol, 1 mM TCEP). Reactions were gently mixed for 30s and incubated for 10min 240 at room temperature. RNP complexes and 200k cells resuspended in 20 µl buffer SE (Lonza) were added 241 to a nucleofection strip and the mixture pulsed with program EH-100 (Lonza 4D-Nucleofector). Cells were 242 plated into 96-well plates for recovery with a fresh media change after 1 day. Two days after RNP 243 electroporation, pooled cells then were single-cell seeded into 384-well plates. After single-cell derived 244 clones emerged, they were split into 2 plates, one of which was used to detect positive clones containing the desired HaloTag[™] sequence through Sanger sequencing. 245

246 Antibodies and Reagents

Goat anti-Rabbit IgG Alexa Fluor Plus 647 (Thermo, A32733), anti-phospho-Histone H2A.X
(Ser139)-clone JBW301 (γH2A.X, Millipore, 05-636-I), anti-WRN clone EPR6392 (abcam, ab124673), Goat
anti-Rabbit IgG (H+L)-HRP (Invitrogen, 31460), anti-GAPDH-HRP (CST, 31460), anti-Ubiquitin, clone P4D1
(for WB; CST, 3936), anti-Ubiquitin, clone FK2 (For IF; Millipore sigma, 04-263), anti-SUMO-2/3, clone 18H8
(CST, 4971S), anti-phospho-CHK1 (Ser345; CST, 2341), phospho-Chk2 (Thr68; clone C13C1; CST, 2661),

cleaved PARP1 (Asp214; clone D64E10; CST, 5625), anti-RNF4 (R&D Systems, AF7964), anti-MDM2 (R&D
Systems, D1V2Z), anti-MIB1 (CST, 4400).

JF549 (Promega, GA1110), etoposide (CST, 2200), 5-FU (5-fluorouracil; Selleckckem, S1209), doxorubicin (CST, 5927), HaloPROTAC3 (Promega, GA3110), HaloPROTAC-E (MedChem Express, HY-145752), MLN-4924 (CST, 85923), carfilzomib (CST, 15022), TAK-243 (MedChem Express, HY-100487), ML-792 (MedChem Express, HY-108702).

258 Immunofluorescence and DNA Damage Quantification

259 Cell culture medium was removed using the Blue Cat Bio Blue®Washer. Paraformaldehyde 260 solution (8% PFA (Electron Microscopy Sciences, 15710-S), 1X phosphate buffered solution (PBS) (Teknova 261 P0191)) was added to the cells via Thermo Multidrop® Combi and treated for 7 min at room temperature 262 to fixate the cells. The paraformaldehyde solution was removed with the Blue Cat Bio BlueWasher. Triton 263 solution (0.1% TritonX-100 (Sigma, T8787), 1X PBS) was added to the cells via Thermo Multidrop Combi 264 and treated for 15 min at room temperature to permeabilize the cells. The triton solution was removed 265 with the Blue Cat Bio BlueWasher. Serum solution (10% Goat Serum (Gibco, 16210), 1X PBS) was added 266 to the cells via Thermo Multidrop Combi and treated for 15 min at room temperature to block the cells. 267 The serum solution was removed with the Blue Cat Bio BlueWasher. Primary antibody solution (1:1000 268 primary antibody, 10% Goat Serum, 1X PBS) was added to the cells via a Thermo Multidrop Combi and 269 treated for 3 hours at room temperature with shaking. The primary antibody solution was removed with 270 the Blue Cat Bio BlueWasher. Secondary antibody solution (secondary antibody (1:1000), Hoechst 33342 271 (AnaSpec, AS-83210) (1:3000), 10% Goat Serum, 1X PBS) was added to the cells via Thermo Multidrop 272 Combi and treated for 30 min at room temperature, covered from light, with shaking. Cells were washed 273 with PBS 3 times using the Blue Cat Bio BlueWasher with a final dispense of PBS. Sealed plates were 274 imaged using an ImageXpress Micro slit confocal microscope (Molecular Devices) using a 40x water 275 immersion objective, and 6 fields of view per well. Exposure parameters were optimized to prevent pixel 276 saturation for each channel. Images were analyzed using MetaXpress Custom Module Editor, by using a 277 Hoechst mask to identify nuclei, then measuring the integrated AlexaFluor 648 intensity across all nuclei 278 in the field of view (FOV) and then averaged. Percent yH2A.X signal was calculated by using the following equation: %S = (T - C_{pos})/(C_{pos} - C_{neg}) * 100, where %S is percent γ H2A.X signal and T is the measured 279 280 yH2A.X fluorescence of the wells treated with test compound. The effective compound concentration 281 leading to a 50% induction of yH2A.X signal (EC_{50}), and the resulting cell yH2A.X signal measured at the 282 highest tested compound tested (c_{max}) was carried out by fitting a 4-parameter non-linear regression using 283 GraphPad Prism. At least three biological replicates were done per compound tested.

284 Western Blotting

Samples were lysed in 1x NuPAGE LDS sample buffer (Invitrogen, NP0008), sonicated, and heated
 at 75 °C for 10 min. Samples were normalized to protein concentration and volume using a Pierce 660
 reagent kit (Invitrogen, 22660). SDS-Page was performed using standard protocols (ref). Western blot
 transfers were performed using an iBlot2 nitrocellulose membrane system (Invitrogen, IB23001).
 Membranes were blocked with Pierce™ Protein-Free Blocking Buffer (Thermo Scientific, 37572) for 30 min
 at room temperature, and subsequently probed with desired antibodies.

291 Cell Viability Assay

292 To seed cells, cells were trypsinized and resuspended in complete culture media to the desired 293 concentration (HCT-116: 10,000 cells/mL; RKO: 15,000 cells/mL; SW-48, HT-29, and SW-480: 25,000 294 cells/mL; U2OS: 7,000 cells/mL). Cell suspensions were seeded in 50 μ L of complete culture media and 295 plated onto 384-well white clear-bottom plates (ThermoFisher Cat# 164610) using a Multidrop[™] Combi 296 liquid dispenser in the slowest setting in triplicate. After 24 h, cells were treated with compounds in a 10-297 pt, 3.16 step serial dilution using an Echo acoustic liquid dispenser (Beckman). DMSO was backfilled to a 298 final concentration of 0.1%. 10 μ M etoposide (C_{pos}) and DMSO (C_{neg}) were used as reference compounds. 299 96 h post compound addition, plates were evacuated to decant all media. To measure viability, cellular

ATP concentrations were measured by adding 20 μL of 1x CellTiterGlo 2.0 (CTG) solution (1 part PBS, 1
 part CellTiterGlo2.0 stock solution; Promega Cat# G9241) using a MultidropTM Combi liquid dispenser
 and measuring the luminescence on a SpectraMax iD3 (Molecular Devices).

Percent viability was calculated by the following equation: $%V = (T - C_{pos})/(C_{pos} - C_{neg}) * 100$, where %V is percent viability and T is the measured luminescence of the wells treated with test compound. The effective compound concentration leading to a 50% reduction in viability (EC₅₀), and the resulting cell viability measured at the highest tested compound tested (c_{max}) was carried out by fitting a 4-parameter non-linear regression using GraphPad Prism. At least three biological replicates were done per compound tested.

309 Tandem Ubiquitin Binding Entities (TUBE) Pull Down

HCT-116 cells were grown as described above. 10⁶ cells were seeded on 10 cm tissue culture 310 311 dishes (ThermoFisher, 150464), and allowed to attach for 48h. Cells were subsequently treated with 312 compounds for 8 h ($[ML-792] = 1 \mu M$, [Carfilzomib] = 10 nM, [TAK-243] = 10 nM, $[WRNi] = 10 \mu M$), 313 harvested by trypsinization, centrifuged (300 x g), and flash frozen in liquid nitrogen. Cell pellets were 314 lysed in 500 µL of TUBE lysis buffer (150 mM NaCl, 2 mM MgCl₂, 25 mM HEPES pH 7.0, 0.03% SDS, 1 M 315 urea, 10 µM PR-619, 1x protease inhibitor cocktail) for 10 min on ice, with frequent mixing. While cell lysis 316 was occurring, 100 µL of slurry/reaction of TUBE magnetic beads (UM501M, LifeSensors) were washed in 317 1x PBS three times, and once with TUBE lysis buffer. After cell lysis, lysates were cleared (20,000 x g, 10 318 min), inputs were taken, and lysates were subsequently added to TUBE magnetic beads. Lysate-TUBE bead 319 mixtures were incubated at 4 °C on a nutator for 2 h. Samples were subsequently washed five times with 320 TUBE wash buffer (150 mM NaCl, 2 mM MgCl₂, 25 mM HEPES pH 7.0, 0.5% Triton X-100, 1 M urea), and 321 resuspended in 1X LDS buffer to prepare for downstream Western blot analysis.

322 HaloTag[™] Protein labeling

Cells were seeded in black 384-well plates using a combidrop multidrop dispenser and seeded for at least 24 h in complete growth media at either 45K cells/mL or 100K cells/mL, depending on the experiment. To label HaloTagged proteins, JF549 (Promega, GA1110) was added to cells using an Echo acoustic liquid dispenser to a final concentration of 25 pM. Cells were incubated at 37 °C for 2 h, and subsequently washed 5 times in PBS, with the final wash leaving the wells empty. After washing, cells were either fixed in 4% paraformaldehyde (Electron Microscopy Sciences, 15710), or growth media was replenished to perform subsequent compound treatments.

330 WRN Imaging Degradation Assays

331 Cell lines and growth conditions are identical to the above conditions. To seed cells for the assay, cells were trypsinized and resuspended in complete culture media to the desired concentration (WRN^{Halo} 332 HCT-116: 50,000 cells, WRN^{Halo} U2OS: 30,000 cells/mL). Cell suspensions were seeded in 50 µL of complete 333 334 culture media and onto 384-well black clear-bottom optical plastic plates (Greiner Bio-One Cat# 781097) 335 using a MultidropTM Combi liquid dispenser in the slowest setting in triplicate. After 24 h, cells were labeled 336 with Halo dye as described above and subsequently treated with compounds in a 10-pt, 3.16 step serial 337 dilution using an Echo acoustic liquid dispenser and incubated at 37 °C in a humidified 5% CO₂ incubator. 10 μM Halo-PROTAC3 (C_{pos:} Promega, GA3110) and DMSO (C_{neg}) were used as reference compounds.24 h 338 339 after compound treatments, cells were fixed in 4% paraformaldehyde for 10 min, washed three times with 340 PBS, then blocked and permeabilized in PBS containing 10% goat serum and 0.1% triton X-100 containing the DNA counterstain Hoechst for 30 min. Plates were washed 3 times in PBS, and sealed with thermal foil 341 342 seals.

343 Sealed plates were imaged using an ImageXpress Micro slit confocal microscope (Molecular 344 Devices) using a 40x water immersion objective, and 6 fields of views per well. Exposure parameters were 345 optimized to prevent pixel saturation for each channel. Images were analyzed using MetaXpress Custom

346 Module Editor, by using a Hoechst mask to identify nuclei, then measuring the average Halo dye intensity 347 across all nuclei in the FOV, averaged, and then background subtracted.

Percent WRN^{Halo} signal was calculated by using the following equation: %S = (T - C_{pos})/(C_{pos} - C_{neg}) * 100, where %S is percent WRN^{Halo} signal and T is the measured WRN^{Halo} fluorescence of the wells treated with test compound. The effective compound concentration leading to a 50% induction of WRN^{Halo} signal (EC₅₀), and the resulting cell WRN^{Halo} signal measured at the highest tested compound tested (C_{max}) was carried out by fitting a 4-parameter non-linear regression using GraphPad Prism. At least three biological replicates were done per compound tested.

354 Phenotypic Screening and Imaging

355 An arrayed human ON-TARGETplus ubiquitome siRNA SMARTpool library (Horizon Discovery, 356 106205-E2-01) was resuspended to a final concentration of 20 uM in 1x siRNA Buffer (Horizon Discovery, 357 B-002000-UB-100). siRNA oligos were transferred onto black 384 μClear plates using an Echo acoustic 358 dispenser to yield a final oligo concentration of 20 nM. 5 µL of opti-MEM (Gibco, 31985062) was added to 359 resuspend transferred oligos. RNA oligo:lipid complexes were formed by adding 0.3 µL of Lipofectamine 360 RNAiMAX (Invitrogen, 13778075) in 5 µL of opti-MEM. Complexes were incubated for 5 min before dispensing 50 µL of HCT-116 cells (100K cells/mL) and incubating in growth conditions described above. 361 After 24 h, growth media was exchanged, and labeled with HaloTag[™] dye as described above. WRNi, or 362 363 DMSO, was added at a final concentration of 10 µM, treated for 24 h, and subsequently fixed with 4% 364 PFA. Cells were permeabilized with 0.1% triton-x 100 for 20 min to remove non-specific dye staining and 365 imaged as described above.

366 siRNA Depletions

367 Cells were processed as described for quantifying WRN levels after siRNA knock-down. For 368 Western blot analysis, HCT-116 (90K cells/well) or U2OS (60K cells/well) cells were seeded in 6-well TC 369 treated plates (Corning, 3516). siRNA depletions were performed following the recommended protocol

for Lipofectamine RNAiMax. Samples were harvested in 1x LDS sample buffer and prepared for Western
blot analysis as described.

372 WRN In-Situ Trapping Assay

WRN^{Halo} Cells (HCT-116 or U2OS) were seeded in black 384-well plates using a combidrop 373 374 multidrop dispenser and seeded for 48 h in complete media at 100K cells/mL (HCT-116) or 30 K cells/mL (U2OS). Cells were labeled with HaloTag[™] Dye and subsequently treated with desired compounds as 375 described above for 8 h ([ML-792] = 1 μ M, [Carfilzomib] = 10 nM, [TAK-243] = 10 nM, [WRNi] = 10 μ M). 376 377 Following compound treatments, samples were decanted to remove all media and treated with ice cold 378 CSK buffer (25 mM PIPES pH 7.0, 300 mM NaCl, 2 mM MgCl₂, 0.3 % TritonX-100, 200 mM sucrose) for 2 379 min, on ice. Without removing buffer in wells, cells were fixed in 4% PFA supplemented with Hoechst for 380 15 min and washed with 1x PBS 5 times using an AquaMax Plate washer (Molecular Devices). Samples 381 were then imaged as described above. Quantifications were done by measuring the total intensity of the 382 Halo dye signal and using the Hoechst channel as a nuclear mask.

383 **Protein Purification**

384 Human WRN residues 480–1251 were cloned into pFastBac vector containing an 8xHis N-terminal tag and expressed in Sf9 insect cells. Harvested cells were resuspended in lysis buffer (50 mM HEPES, pH 385 7.5, 500 mM NaCl, 25 mM imidazole, 1mM TCEP, 5 U/mL benzonase (Millipore Sigma), EDTA-free 386 387 cOmplete protease inhibitor cocktail tablet (Roche)) and lysed by addition of Insect PopCulture reagent 388 (Millipore Sigma). Lysate was loaded on to a HiTrap TALON Crude column (Cytiva) and protein was eluted 389 using 500mM Imidazole. Fractions containing WRN protein were collected, pooled, then diluted to in 390 buffer to drop the NaCl concentration to ~100mM. The pooled and diluted sample was then loaded onto 391 a heparin column (Cytiva) and eluted using a step-wise gradient from 200mM to 1000mM NaCl. Fractions 392 containing purified WRN protein were pooled, and buffer exchanged into 50mM HEPES, 150mM NaCl, 393 10% glycerol, 1mM TCEP using a HiPrep desalting column (Cytiva).

394 Human BLM residues 636-1298 were cloned into pFastBac vector containing an 8xHis N-terminal 395 tag and expressed in Sf9 insect cells. Harvested cells were resuspended in lysis buffer (50 mM HEPES, pH 396 8, 200 mM NaCl, 0.5mM TCEP, 5 U/mL benzonase (Millipore Sigma), EDTA-free cOmplete protease 397 inhibitor cocktail tablet (Roche)) and lysed by addition of Insect PopCulture reagent (Millipore Sigma). 398 Lysate was loaded onto a Ni-NTA column (Thermo Fisher) and protein was eluted using 300mM Imidazole. 399 Fractions containing BLM protein were collected, pooled, and diluted in buffer to drop the NaCl 400 concentration to ~50mM. The pooled and diluted sample was then loaded on a HiTrap Heparin column 401 (Cytiva) and eluted with a linear gradient from 50mM to 1M NaCl. The pure fractions were pooled and 402 concentrated and purified by size exclusion chromatography using S200 Increase (Cytiva) in 50mM HEPES 403 pH 8, 200mM NaCl, 5% glycerol, 0.5mM TCEP.

404 Helicase Unwinding and ATPase Assay

- 405 The helicase unwinding and ATPase were carried out in a multiplexed fashion based on previously
- 406 published BLM assay (Chen et al., 2021). Single-stranded DNA was purchased from IDT:

- 409 B-dark: 5'-CGTACCCGATGTGTTCGTTCY-3'
- 410 Strands A and B were annealed in TE + 50mM NaCl in a slowly-cooling thermocycler.

First, compounds in DMSO were dispensed to a 384-well white ProxiPlate (Perkin Elmer) using an Echo acoustic liquid handler. WRN or BLM Protein was diluted into assay buffer and 2.5 µL was dispensed into each well. Protein and compound then incubated at room temperature for 15 min. Following preincubation, 2.5µL of DNA and ATP in assay buffer was dispensed into each well to initiate the reaction and incubated for 20 min at room temperature. At the 20 minute time point, Cy3 fluorescence was read on an Envision plate reader to measure unwinding activity. ATP hydrolysis was then measured using ADP-Glo kit (Promega) by adding 5uL ADP-Glo reagent for 40 minutes followed by 10uL Kinase Detection Reagent for

418 1hour. Luminescence was measured using Envision plate reader. Data was normalized to DMSO (100%
419 activity). The final reaction conditions are: 1mM ATP, 15nM dsDNA substrate, 1.5uM B-dark in reaction
420 buffer composed of 50mM Tris-HCl pH 8.0, 50mM NaCl, 2mM MgCl2, 0.01% Tween-20, 2.5µg/mL poly(dl421 dC), 1mM DTT, 1% DMSO, and 12.5nM WRN or 2.5nM BLM.

422 Cellular SMT Sample preparation

423 For SMT experiments, WRN Halo-tagged HCT-116 cells were seeded in FluoroBrite DMEM (Thermo Fisher, cat. no. 1896701) supplemented with 10% FBS (Corning), 1% Penicillin/Streptomycin 424 425 (Gibco, 1510-122), and 1% GlutaMAX (Gibco, 35050-061) on plasma-coated 384-well glass-bottom plates 426 (Cellvis, P384-1.5H-N) at 1.5x10⁴ cells per well. WRN Halo-tagged U2OS cells were seeded in GlutaMAX-427 supplemented DMEM (Gibco, 10566-016) with 10% FBS and 1% Penicillin/Streptomycin at 6x10³ cells per 428 well. Prior to treatment and imaging, HCT-116 cells were incubated at 37 °C and 5% CO₂ for ~48 hours to 429 allow for cell adherence to the plates, while U2OS cells were incubated under the same conditions for 24 430 hours. For all SMT experiments, cells were treated with 10-40 pM of JF549-HTL (synthesized in-house) and 431 200 nM Hoechst 33342 (Thermo Fisher, cat. No. 62249) for 1 hour in complete medium at 37 °C and 5% 432 CO₂. Cells were then washed three times in PBS and twice in FluoroBrite DMEM supplemented with 2% 433 FBS, 1% Penicillin/Streptomycin, and 1% GlutaMAX. All compounds were prepared on Echo Qualified 384-434 Well Low Dead Volume Source Microplates (Labcyte, cat. No. LP-0200) in DMSO and administered onto 435 cells at a final 1:500 dilution in cell culture medium. Unless otherwise specified, cells were incubated with 436 compounds for 4 hours at 37 °C prior to image acquisition. When possible, well replicate conditions were 437 randomized across each plate. For all experiments, control conditions included vehicle (DMSO) treatment 438 and wells lacking JF549-HTL to assess possible effects from detection of non-dye signal.

439 Cellular SMT Image Acquisition

440 Unless otherwise stated, all image acquisition for cSMT was performed on a customized Nikon
441 Eclipse Ti2-E inverted fluorescence microscope with a motorized stage. The microscope system was

outfitted with a stage top environmental chamber with temperature and CO₂ control (OKO labs), Nikon 442 443 objective water dispenser, an Oblique Line Scanning (OLS) illumination module (Driouchi et al., submitted) with laser launch containing 405 nm, 560 nm, and 642 nm lasers, three-band emission filter set (ET 444 445 445/58m, FF01-585/40-25, FF01-676/37-25, Chroma), motorized filter wheel (Lambda 10-B; Sutter 446 Instruments), and a high-speed sCMOS camera equipped with light-sheet mode capability (ORCA-Fusion 447 BT, Hamamatsu). Images were acquired with a 60X 1.27 NA water immersion objective (CFI SR Plan Apo IR 448 60XC WI, Nikon, Japan). The environmental chamber was maintained at 37° C, 95% humidity, and 5% CO₂. 449 For each field of view, 150 SMT frames were collected at a frame rate of 100 Hz with a 407-microsecond 450 stroboscopic laser pulse, and 1 frame in the Hoechst channel was subsequently collected for downstream 451 registration of trajectories to nuclei. Each frame captured an FOV that was 1728 x 2304 pixels (187.14 x 452 249.52 microns) in size. Automated microscope control and image acquisition was performed using 453 customized scripts in MicroManager.

For all experiments, conditions were tested by acquiring JF549 movies and Hoechst images at 4 FOVs per well, a minimum of 2 well replicates per plate, and a minimum of 2 plate replicates. Unless otherwise stated, reported averages for each condition are the mean value of all FOVs collected. For time course experiments, reported averages at each time point are the mean value of all FOVs collected across 6 consecutive wells per plate replicate.

459 Cellular SMT Image Processing

Image acquisition yielded one JF549 movie and one Hoechst movie per FOV. The JF549 movie was
used to track the motion of individual JF549 molecules, while the Hoechst movie was used for nuclear
segmentation.

For tracking, we used a custom pipeline that operates in three sequential steps. First, dye molecules are detected using a generalized log likelihood ratio detector (Serge et al., 2008). The position of each detected emitter is then estimated using a Levenberg-Marquardt fitting routine (Levenberg 1944; 466 Marguardt 1963; Laurence and Chromy 2010) with an integrated 2D Gaussian spot model (Smith et al., 467 2010) starting from an initial guess afforded by the radial symmetry method (Parthasarathy 2012). 468 Detected emitters were then linked into trajectories using a custom algorithm. Briefly, this method first 469 estimates the marginal probabilities of each potential link between particles using the graphical softmax 470 operator (Cuturi 2013; Mena et al., 2018) applied to a Brownian motion model, then attempts to find the 471 set of trajectories with maximum marginal log probability using a modification of Sbalzerini's hill-climbing 472 algorithm (Sbalzarini and Koumoutsakos 2005). Identical tracking settings were used for all movies in this 473 manuscript.

474 For nuclear segmentation, all frames of the Hoechst movie were averaged to generate a mean 475 projection. This mean projection was then segmented with a UNET-based convolutional neural network 476 trained on human-labeled nuclei (Ronneberger et al., 2015). Each spot was then assigned to at most one 477 nucleus using its subpixel coordinates. To recover dynamical information from trajectories, we used state 478 arrays (Heckert et al., 2022), a Bayesian inference approach, with the "RBME" likelihood function and a 479 grid of 100 diffusion coefficients from 0.01 to 100.0 μ m²/sec and 31 localization error magnitudes from 480 0.02 µm to 0.08 µm (1D RMSD). After inference, localization error was marginalized out to yield a one-481 dimensional distribution over the diffusion coefficient for each FOV.

482 Synthesis of HRO761

The WRN inhibitor HRO761 was synthesized following an established synthesis route for Compound 42,
as described in WO2022/249060. (Bordas et al., 2022).

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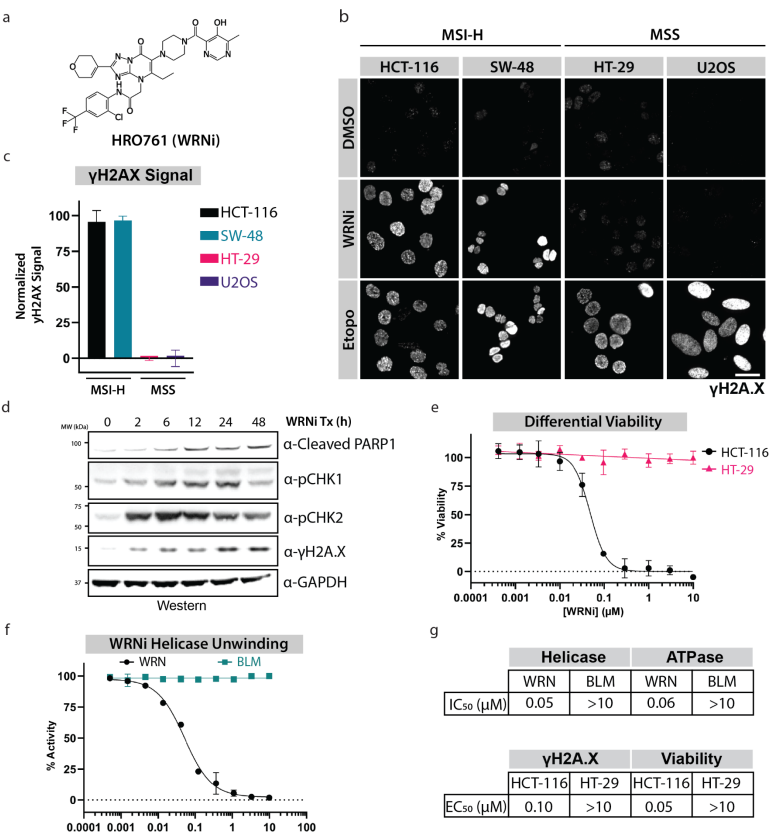
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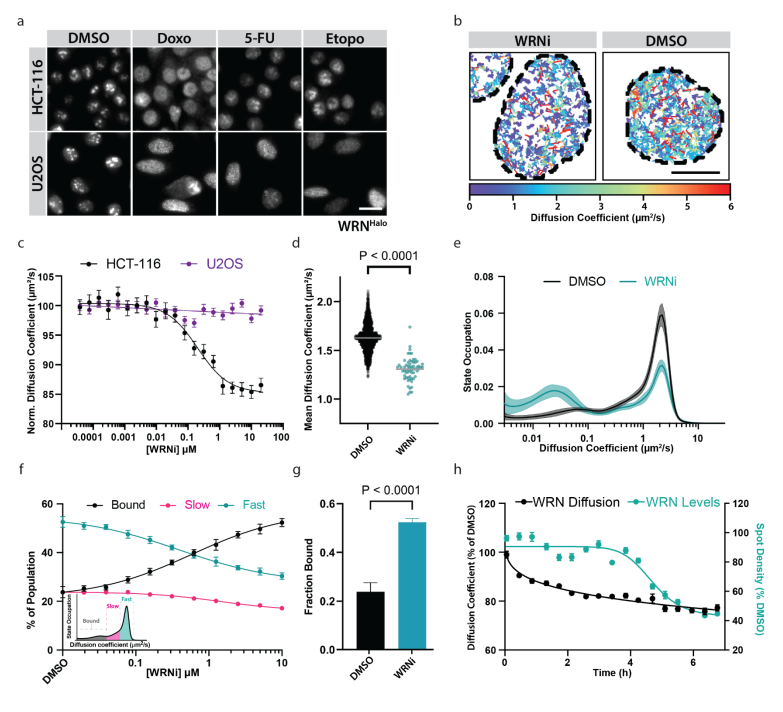
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[WRNi] (µM)

Fig. 1: HRO761 is a specific and potent WRN inhibitor.

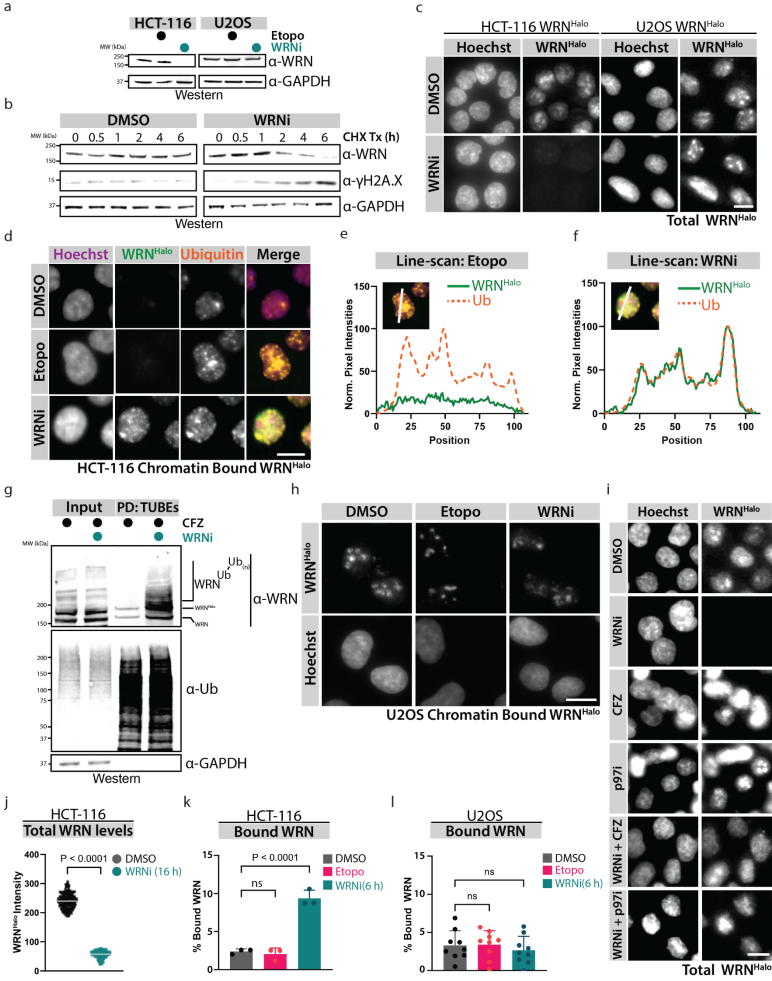
665 a. Chemical structure of HRO761 (WRNi). b. WRN inhibition leads to induction of the DNA damage 666 response in MSI-H cells. Phospho-histone H2A.X (Ser139) (yH2A.X) staining was used to visualize DNA 667 damage in MSI-H (HCT-116, SW-48) and MSS (HT-29, U2OS) cells after 24 h treatment with indicated 668 compounds. Scale bar = 20 μ m. c. Quantification of yH2A.X signal levels in cells in b, normalized to DMSO 669 and Etoposide. Graphs represent averages of at least 3 plate replicates, consisting of at least 6 fields of 670 view (FOVs). d. WRN inhibition by WRNi leads to an induction of the DNA damage response resulting in 671 apoptosis. HCT-116 cells treated with 10 µM WRNi and analyzed via Western to assess DNA damage 672 response markers. e. Dose response curves measuring viability of HCT-116 cells or HT-29 cells after 673 treatment with WRNi for 4 days. Graphs represent averages from n = 6 plates. f. Dose response curves measuring in vitro WRN or BLM unwinding activity after treatment with WRNi. Graphs represent averages 674 675 from n = 4 replicates. WRN unwinding activity is normalized to DMSO and ATP-y-S; BLM activity is normalized to DMSO and 10 µM BLMi. g. Summary of IC₅₀ or EC₅₀ values of WRNi in the indicated 676 biochemical and cellular assays. All curve fits were done by fitting a 4-parameter logarithmic regression 677 678 curve. All error bars represent standard deviation (s.d.). DMSO is dimethyl sulfoxide; BLMi is BLM inhibitor 679 Compound 2; Etopo is etoposide. MW is molecular weight.



681 Fig. 2: Single molecule tracking shows a change in WRN cellular dynamics in an MSI-H

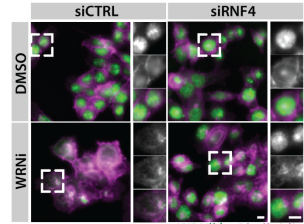
682 dependent manner.

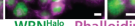
a. WRN^{HALO} cell lines, HCT-116 and U2OS, were validated by visualizing the subcellular localization of WRN 683 in the presence or absence of DNA damaging compounds. WRN^{Halo} successfully translocates across 684 compartments, suggesting a functional protein. Scale bar = 20 μ m. **b**. The decrease in WRN diffusion 685 686 coefficient upon inhibition can be captured by SMT, showing a dramatic slowdown of WRN protein. WRN 687 inhibition leads to a decrease in molecule diffusion coefficient. Representative SMT track overlays over 688 Hoechst nuclear stain outlines, in the presence or absence of 10 µM WRNi. Tracks are colored according 689 to the diffusion coefficient of the molecules. c. Inhibition of WRN only affects its mean diffusion coefficient 690 in MSI-H cells. Dose response curves with WRNi measuring the diffusion coefficient of WRN^{HALO} after 4 h treatments in HCT-116^{WRN-HALO} or U2OS^{WRN-HALO}. Graph represents the average from n = 4 plates per 691 692 condition. Error bars represent standard error of the mean (s.e.m.). All curve fits were done by fitting a 4-693 parameter logarithmic regression curve. d. Dot plot quantification of WRN diffusion coefficient from SMT 694 movies after treatment with 10 µM WRNi. Each point represents the average WRN diffusion coefficient 695 within all the nuclei in an FOV. n = 20 plates. Lines represent sample medians. e. WRNi shifts a large fraction of molecules from the free-diffusing state ("fast") to the chromatin-bound ("bound") state. 696 Distribution of diffusive states in HCT-116^{WRN-Halo} cells showing the relative proportion of WRN molecules 697 698 as a function of diffusion coefficient occupation, in the presence and absence of 10 μ M WRNi. Shaded 699 area represents s.d. f. Treatment with WRNi shows a dose-dependent increase in the bound fraction of 700 WRN, suggesting binding of WRN onto chromatin. Dose response curves with WRNi measuring the 701 different diffusive states of WRN protein. Inset is representation of how diffusive states are classified. Error bars represent s.d., g. Quantification from f of the chromatin bound fraction of WRN^{Halo} protein in 702 the presence or absence of 10 µM WRNi. Error bars represent s.d.. h. SMT can capture changes in protein 703 diffusion before degradation is observed. Overlay of WRN^{HALO} diffusion coefficient and molecule spot 704 705 densities after treatment with 10 µM WRNi over the indicated time points. Values are normalized to 706 DMSO by dividing all FOV-level measurements at each time point by the median DMSO value at that time 707 point, then multiplying by 100. n = 2 plates. Error bars represent s.e.m.. Curve fits were done by fitting a 708 4-parameter logarithmic regression curve. P-values were calculated using a two-tailed, unpaired t-test. 709 DMSO is dimethyl sulfoxide; WRNi is HRO761; 5-FU is 5-fluorouracil, Doxo is doxorubicin.

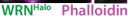


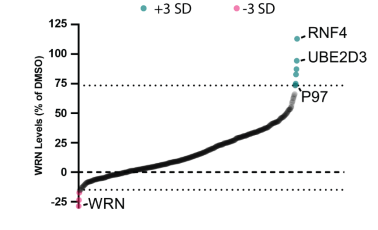
711 Fig. 3: WRN Inhibition leads to its chromatin associated degradation.

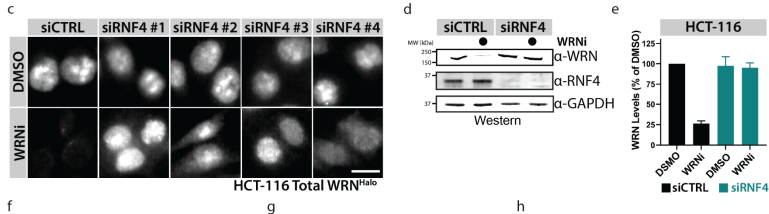
a. Inhibition of WRN leads to its degradation in an MSI-H dependent manner. HCT-116^{WRN-Halo} or U2OS^{WRN-} 712 713 Halo were treated with 10 μ M WRNi or etoposide for 16 h and analyzed by Western blot. **b**. Inhibition of WRN in MSI-H leads to a change in the half-life of WRN protein. HCT-116^{WRN-Halo} were treated with 100 714 μ g/mL of cycloheximide (CHX) to inhibit protein synthesis in the presence or absence of 10 μ M WRNi, 715 716 then harvested at the indicated time points for Western blot analysis. Quantifications of CHX chase are in 717 **Extended Data Fig. 3d. c.** Loss of WRN protein after treatment with WRNi can be visualized by microscopy. HCT-116^{WRN-Halo} or U2OS^{WRN-Halo} were treated with 10 µM WRNi for 16 h then imaged to visualize total WRN 718 protein levels. Scale bar = 10 μ m. d. WRN inhibition leads to WRN trapping on chromatin and its 719 720 ubiquitylation. HCT-116^{WRN-Halo} cells were treated with 10 μ M WRNi or etoposide for 6 h, at which point 721 cells were permeabilized by treating with a mild detergent, then fixed and imaged. WRN signal retention 722 is only seen when WRN is inhibited, and not with general DNA damaging compounds. Ubiquitin signal was also observed to colocalize with WRN^{Halo} when probing with a ubiquitin antibody. Scale bar = 10 μ m. e. 723 724 Line-scan guantification of etoposide treated cells from d, showing a lack of co-localization of the ubiquitin 725 signal channel and the WRN signal channel. f. Line-scan quantification of WRNi treated cells from d, 726 showing clear colocalization of the ubiquitin and WRN signal channels. g. WRN inhibition leads to its 727 ubiquitylation. Tandem ubiquitin binding entities (TUBEs) pulldown (PD) of HCT-116 cells after treatment 728 with 10 μ M WRNi for 6 h in the presence of 1 μ M carfilzomib (CFZ). Blotting for endogenous WRN using 729 an α -WRN antibody shows higher molecular weight species in the TUBEs pull down only in the presence 730 of WRNi. h. WRN chromatin trapping upon its inhibition is MSI-H dependent. Cells were treated with WRNi as in **d** but using the MSS cell line U2OS^{WRN-Halo}. **i**. Degradation of WRN is dependent on the p97/VCP-731 proteasome axis. HCT-116^{WRN-Halo} were treated with 10 µM of WRNi and 1 µM of either CB-5083 (p97i) or 732 733 CFZ for 6 h, then imaged. WRN protein degradation by WRNi is rescued upon co-treatment with CFZ or 734 p97i. Quantifications are in Extended Data Fig. 3f. j-l. Quantifications of c, d, and h. Graph in j represents 735 averages from n = 3 plates, with each individual point representing one well. Graph in k represents averages from n = 3 plates, each individual point is the average of 6 wells. Graph in I represents averages 736 737 from n = 9 plates, each individual point is the average of 6 wells. Error bars represent s.d. DMSO is 738 dimethyl sulfoxide; WRNi is HRO761; CHX is cycloheximide; CFZ is carfilzomib; Etopo is etoposide. P-values 739 were calculated using a two-tailed, unpaired Student's t-test. ns = not significant. MW is molecular weight.

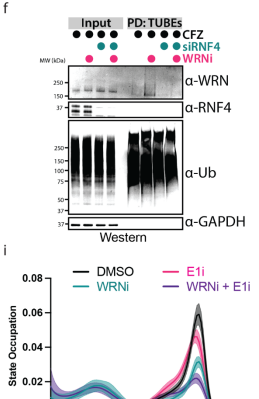












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Diffusion coefficient (µm²/s)

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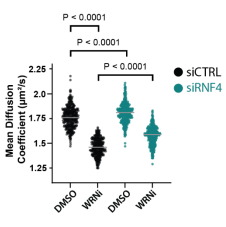
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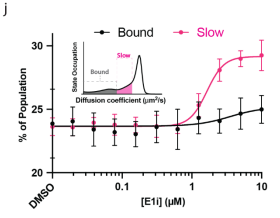
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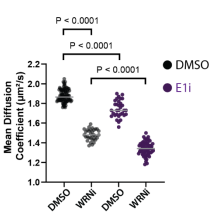
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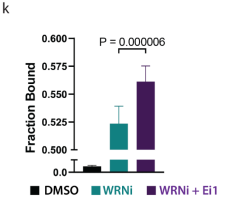
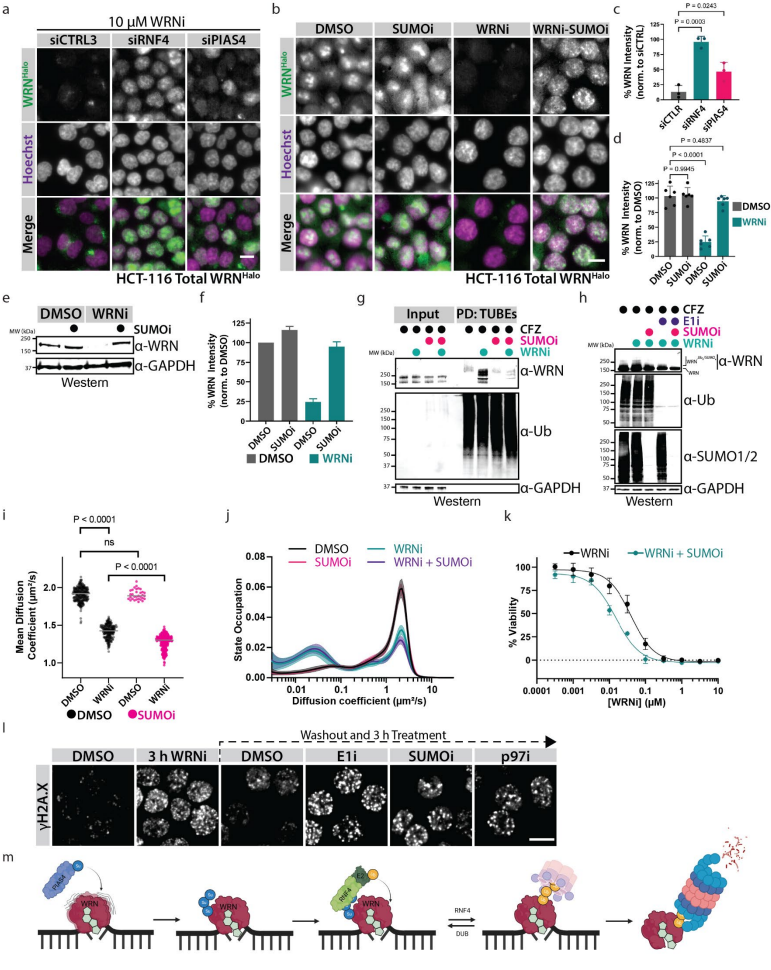


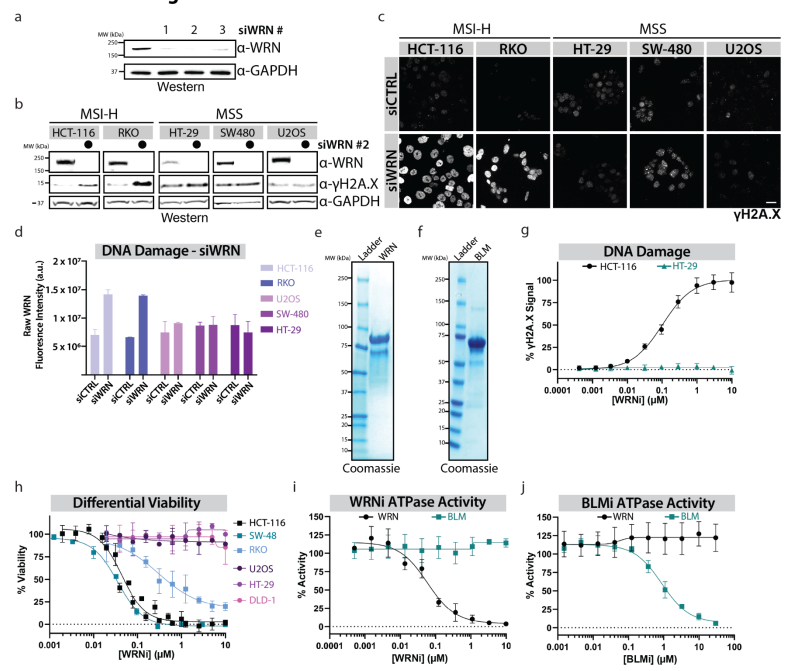
Fig. 4: Phenotypic siRNA screen identified RNF4 as the ubiquitin E3 ligase targeting WRN for degradation.

743 a. A phenotypic siRNA screen identified genes involved in the degradation of WRN after WRNi treatment. 744 Colored circles indicate hits that are 3 standard deviations (SD) from the mean. Quantification of WRN 745 protein levels after treatment with SMARTpool siRNAs in the presence or absence of 10 μ M WRNi for 24 746 h. Depletion of RNF4 leads to stabilization of WRN protein in the presence of WRNi. UBE2D3 is an E2 known to interact with RNF4. b. Representative images of HCT-116^{WRN-Halo} cells the siRNA screen 747 748 performed in a, showing a rescue of WRNi-induced degradation of WRN when RNF4 is depleted. Scale 749 bars = 10 μ m. c. Decomplexifying the siRNA SMARTpool validates RNF4 as the ligase responsible for WRN degradation by WRNi treatment. Representative images of HCT-116^{WRN-Halo} cells show all individual RNF4 750 751 siRNA oligos rescue the degradation phenotype induced by WRNi. Scale bar = 10 μ m. Quantifications are 752 in Extended Data Fig. 4c. d. Western blot analysis validates RNF4 as the ligase responsible for degradation 753 of WRN induced by WRNi. HCT-116 cells were treated with siRNF4 oligos for 24 h, then treated with or 754 without 10 µM WRNi for an additional 24 h, at which point cells were lysed and analyzed by Western blot 755 with indicated antibodies. **e**. Quantifications of **d**. Graphs represent the mean value of n = 2 Western blot runs. Error bars represent s.d.. f. Depletion of RNF4 directly prevents ubiquitylation of WRN after WRNi 756 757 treatment. HCT-116 cells were treated with siRNF4 oligos for 24 h, then cotreated with or without 10 μM 758 of WRNi for 6 h. All samples were treated with CFZ to stabilize ubiquitin-modified proteins. TUBEs 759 pulldowns (PD) and Western blot analysis were performed, probing with indicated antibodies. High 760 molecular weight WRN species are detected in the TUBEs PD when treated with WRNi, which are not 761 observed when treated with siRNF4 oligos, suggesting ubiquitylation of WRN is dependent on RNF4. g. 762 RNF4 depletion leads to a slight but significant increase in the mean diffusion coefficient of WRN. Dot plots of WRN diffusion coefficient via SMT after co-treatment with siRNF4 and either DMSO or WRNi. Each 763 764 point represents the average WRN diffusion coefficient within all the nuclei in an FOV. n = 4 plates. Lines 765 represent sample medians. h. The ubiquitin pathway is involved in regulating WRN dynamics. Dot plots of 766 WRN diffusion coefficient via SMT after co-treatment with the ubiquitin-activating enzyme (E1) inhibitor 767 (E1i) and either DMSO or WRNi. E1i treatment caused a reduction in the diffusion coefficient of WRN. This 768 decrease in diffusion was exacerbated by cotreatment with WRNi. Each point represents the average WRN 769 diffusion coefficient within all the nuclei in an FOV. n = 4 plates. i. Distribution of diffusive states in HCT-116^{WRN-Halo} cells showing the relative proportion of WRN molecules as a function of diffusion coefficient 770 771 occupation after treatment with 1 μ M E1i in the presence or absence of 10 μ M WRNi. E1 inhibition leads 772 to a shift from the "fast" to the "slow" state of WRN molecules. This shift is exacerbated by the 773 cotreatment with WRNi. Shaded area represents s.d., j. Treatment with E1i shows a dose-dependent 774 increase in the slow fraction of WRN, suggesting that the ubiquitin pathway regulates WRN dynamics. 775 Dose response curves with E1i measuring the "bound" and "slow" fractions of WRN protein. Inset is 776 representation of how diffusive states are classified. Error bars represent s.d.. k. Bar graph quantification of the fraction bound of WRN^{Halo} protein after WRNi treatment in the presence or absence of E1i. Error 777 778 bars represent s.d. DMSO is dimethyl sulfoxide; WRNi is HRO761; E1i is TAK-243; CFZ is carfilzomib. P-779 values were calculated using a two-tailed, unpaired Student's t-test. ns = not significant. MW is molecular 780 weight.



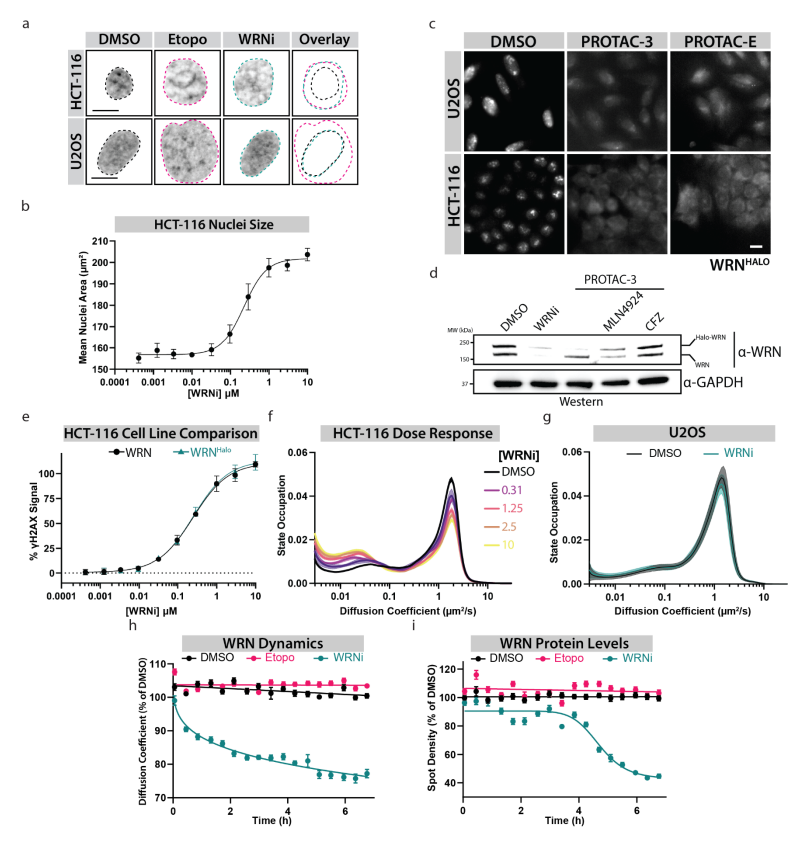
782 Fig. 5: The PIAS4-RNF4 axis is responsible for the chromatin-associated degradation of WRN.

a. WRN degradation is dependent on the SUMO E3 ligase PIAS4. HCT-116^{WRN-Halo} cells were treated with 783 784 the indicated siRNA oligos for 24 h, and subsequently treated with 10 µM of WRNi for 24 h then imaged. 785 WRN degradation is rescued by the depletion of RNF4, and partially rescued by PIAS4 depletion. b. SUMOylation is required for WRN degradation induced by WRNi. HCT-116^{WRN-Halo} cells were treated with 786 787 1 μ M of the SUMO-activating enzyme (SAE) inhibitor ML792 (SUMOi) in the presence or absence of 10 788 µM WRNi, for 6 h and then fixed for imaging. Treatment with SUMOi results in a full rescue of the WRN 789 degradation phenotype. c and d. Quantifications of a and b, respectively. Graphs represent the average 790 of n = 3 plates for siRNA experiments and n = 6 plates for the small molecule experiments. Each dot 791 represents the average of 6 wells, and 6 FOVs per well. Error bars represent s.d.. e. Whole cell lysate 792 analysis shows SUMOylation is necessary for the degradation of WRN induced by WRNi. HCT-116 cells 793 were treated with 1 μ M SUMOi in the presence or absence 10 μ M WRNi 16 h, at which point cells were 794 lysed and analyzed by Western blot with indicated antibodies. Western blot analysis of cells showing WRN 795 protein levels are stabilized when treated with SUMOi. f. Quantification of e. Graphs represent the mean 796 value of n = 2 Western blot runs. Error bars represent s.d., g. Inhibition of the SUMO pathway directly 797 prevents ubiquitylation of WRN after WRNi treatment. HCT-116 cells were treated with 1 μ M SUMOi in 798 the presence or absence of 10 µM of WRNi for 6 h. All samples were treated with CFZ to stabilize ubiquitin-799 modified proteins. TUBEs pulldowns (PD) and Western blot analysis were performed, probing with 800 indicated antibodies. High molecular weight WRN species are detected in the TUBEs PD when treated with 801 WRNi, which are not observed when treated with SUMOi, suggesting that the SUMO pathway is required 802 for the ubiquitylation of WRN upon its inhibition by WRNi. h. Whole cell lysate analysis by Western blot 803 showing that higher molecular weight WRN species are dependent on the SUMO and ubiquitin pathways. 804 Co-treatment with WRNi in the presence or absence of E1i and SUMOi show the disappearance of higher 805 molecular weight species when co-treated with SUMOi or E1i, and a full disappearance of these species 806 when co-treated with both SUMOi and E1i. i. Dot plots of WRN diffusion coefficient via SMT after co-807 treatment with SUMOi and either DMSO or WRNi. Each point represents the average diffusion coefficient 808 of all the nuclei in an FOV. n = 4 plates. Lines represent sample medians; P-value was calculated using a 809 two-tailed, unpaired t-test. Co-treatment with both compounds leads to a further decrease in the diffusion coefficient of WRN. j. Distribution of diffusive states for WRN^{HALO} in HCT-116^{WRN-Halo} after 810 treatment with 1 µM of SUMOi in the presence or absence of 10 µM WRNi, showing that SUMOi leads to 811 812 a decrease of the "fast" moving WRN molecules. k. Inhibition of the SUMO pathway synergizes with WRN 813 inhibition. Dose response of WRNi in HCT-116 cells treated in the absence or presence of 10 nM SUMOi 814 showing a shift in the EC₅₀ viability of WRNi. I. Failure to extract trapped WRN leads to persistent DNA 815 damage. Treating HCT-116 cells with WRNi for 3 h, then washing and replenishing with fresh growth media 816 containing 1 µM of the indicated inhibitors show a retention in DNA damage as measured by yH2A.X when 817 the SUMO-ubiquitin axis is perturbed. Scale bar = 10 μ m. m. Model for targeting trapped WRN to 818 proteasomal degradation by WRNi. WRN that is bound to chromatin surveying DNA damage in MSI-H cells 819 becomes trapped upon inhibition by WRNi. This stalled WRN is SUMOylated by the SUMO ligase PIAS4. 820 SUMOylated WRN recruits the STUBL RNF4, leading to its ubiquitylation. Ubiquitylated WRN is extracted 821 from chromatin by p97/VCP, leading to its degradation by the proteasome. DMSO is dimethyl sulfoxide; 822 WRNi is HRO761; E1i is TAK-243; SUMOi is ML-792; CFZ is carfilzomib. P-values were calculated using a 823 two-tailed, unpaired Student's t-test. ns = not significant. MW is molecular weight.



825 Extended Data Fig. 1:

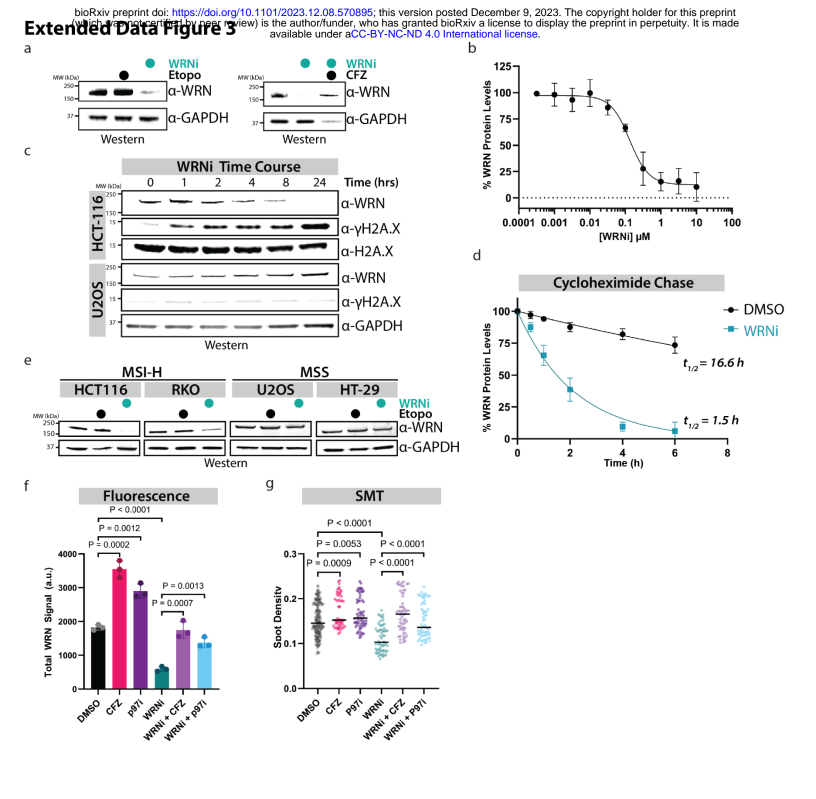
826 a. Whole cell lysates of HCT-116 after siRNA depletions of WRN for 24 h and probing with a WRN antibody 827 show a robust loss of WRN protein. **b**. WRN depletion is synthetic lethal in MSI-H cells but not MSS cells. 828 Whole cell lysates in the indicated cell lines after treatment with siWRN or siCTRL oligos for 48 h, and 829 subsequently analyzed for DNA damage induction by Western blot. c. As in b, but cells were fixed in 830 paraformaldehyde after siRNA treatments, and DNA damage was measured by measuring yH2A.X levels 831 via immunofluorescence. Scale bar = 20 μ m. **d**. Quantifications of **c**. Each graph represents the mean of *n* 832 = 3 plates. e. Purification of WRN protein from SF9 insect cells. Coomassie gel staining shows a product of 833 the expected protein molecular weight after purification. f. Protein purification of BLM protein from E. 834 *coli*. Coomassie gel staining shows a product of the expected protein molecular weight after purification. g. DNA damage induction in MSI-H cells after WRN inhibition is dose-dependent. Dose response curves 835 836 measuring DNA damage response via yH2A.X levels in HCT-116 cells or HT-29 cells after treatment with 837 WRNi for 24 h. Graphs represent averages from n = 6 plates. h. Cell viability panel of MSI-H and MSS cells showing the differential viability effect of WRN inhibition towards MSI-H cells. Dose response curves 838 839 measuring the viability of the indicated cell lines after WRNi treatment for 4 days. HCT-116, SW-48, and 840 RKO are MSI-H cells; U2OS, HT-29, and DLD-1 are MSS cells. Graphs represent averages from n = 3 plates. 841 All error bars represent standard deviation (s.d.). i. Dose response curves measuring the *in vitro* ATPase 842 activity of WRN or BLM after WRNi treatment. Graphs represent averages from n = 6 plates. All curve fits 843 were done by fitting a 4-parameter logarithmic regression curve. j. Purified BLM protein is active. 844 Benchmarking of BLM protein by treatment with BLMi. Dose responses measuring ATPase and helicase 845 inhibition by BLMi. All curve fits were done by fitting a 4-parameter logarithmic regression curve. All error 846 bars represent standard deviation (s.d.). DMSO is dimethyl sulfoxide; WRNi is HRO761; BLMi is the BLM 847 inhibitor Compound 2. MW is molecular weight.



849 Extended Data Fig. 2:

850 a. WRNi causes morphological changes to nuclei due to DNA damage accumulation in MSI-H cells. Images 851 of Hoechst-stained nuclei of HCT-116 or U2OS cells treated in the presence or absence of 10 µM WRNi or 852 etoposide for 24 h. Nuclei outlines are overlaid on top of each other, showing the large change in area 853 after WRNi treatment in HCT-116 cells. This increase in nuclei size is observed in HCT-116 and U2OS cells after etoposide treatment. Scale bar = 10 μ m. **b**. Nuclear morphology changes induced by DNA damage 854 are dose-dependent. Dose response curve measuring nuclei area in HCT-116 cells after treatment with 855 WRNi for 24 h. Error bars represent s.d.. c. Validation of endogenous WRN Halo tagging of HCT-116 and 856 U2OS. WRN protein levels in HCT-116^{WRN-Halo} and U2OS^{WRN-Halo} visualized by staining with JF549 dye after 857 858 treatment with 10 μ M degraders of HaloTag, PROTAC-E or PROTAC-3, for 24 h. Scale bar = 10 μ m. **d**. 859 Treatment with Halo-PROTAC-3 leads to proteasomal dependent degradation of WRN^{Halo}. Further validation of the WRN^{Halo} tag, showing Western blot analysis of HCT-116-WRN^{Halo} cells after treatment 860 with 10 µM PROTAC-3 in the presence or absence of 2 nM CFZ or 5 nM MLN-4924 for 24 h. PROTAC-3 861 uses CUL2^{VHL} as a ligase, therefore CUL2 inhibition via MLN-4924 leads to a rescue of the WRN^{Halo} 862 degradation phenotype. e. Wild type HCT-116 cells and HCT-116-WRN^{Halo} cells have identical responses to 863 WRNi, suggesting WRN^{Halo} is functional. Dose response curves of HCT-116^{WT} and HCT-116-WRN^{Halo} 864 measuring DNA damage induction. Error bars represent s.d.. f. WRNi leads to a dose-dependent increase 865 of WRN molecules bound to chromatin. Distribution of diffusive states for WRN^{Halo} in increasing 866 867 concentrations of WRNi. WRNi leads to a decrease of free-disusing WRN molecules with a concomitant 868 increase in chromatin-bound WRN molecules. g. WRNi does not lead to chromatin trapping in MSS cells. SMT measurements show that the distribution of WRN^{Halo} diffusive states in U2OS^{WRN-Halo} cells remains 869 870 unchanged in the presence or absence of 10 µM WRNi. Shaded area represent s.d.. h and i. WRN diffusion 871 coefficient (h) and protein levels (i) remain unchanged upon DNA damage induction. Kinetic SMT of HCT-116-WRN^{Halo} after treatment with 10 μ M WRNi or Etopo over the indicated time points. Error bars 872 873 represent s.e.m.. CFZ is carfilzomib; DMSO is dimethyl sulfoxide; WRNi is HRO761; Etopo is etoposide. 874 MW is molecular weight.

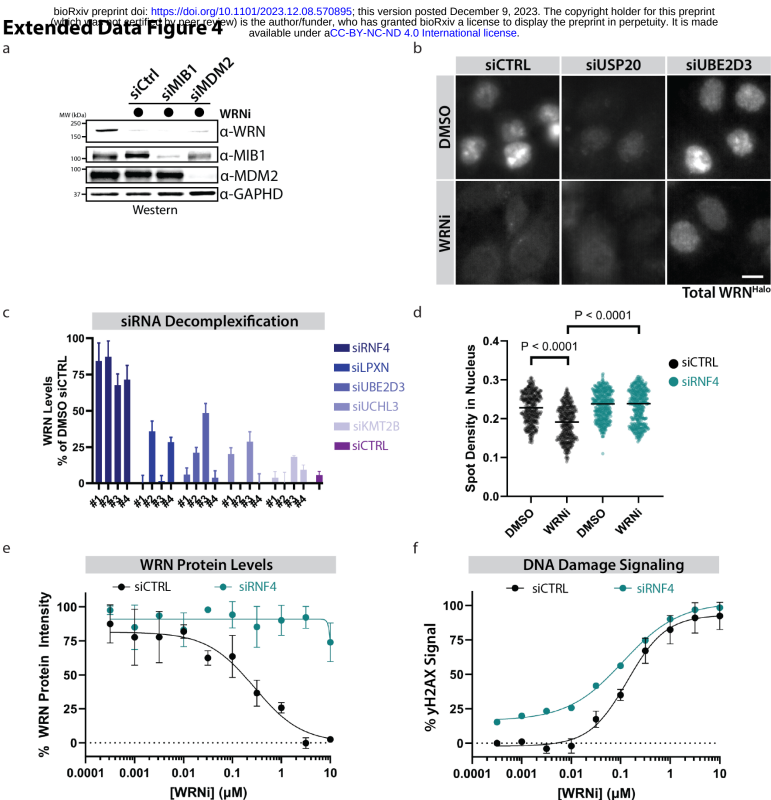
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877 Extended Data Fig. 3:

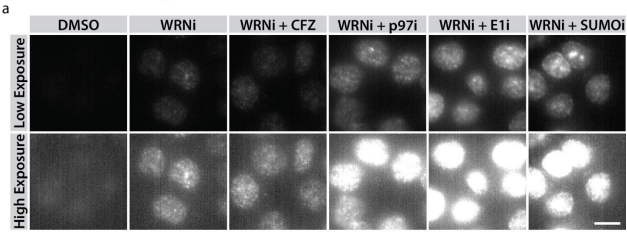
878 a. Degradation of WRN is induced by WRNi, but not by general DNA damage. Western blot analysis of 879 HCT-116 cells treated with 10 µM etoposide or WRNi. This degradation is rescued by the addition of CFZ. b. Dose response curves measuring WRN protein levels by staining HCT-116^{WRN-Halo} cells with JF549 after 880 881 treatment with WRNi for 24 h. Graphs represent averages from n = 3 plates, measuring 3 well per plate 882 and 6 FOVs per well. The curve fit was done by fitting a 4-parameter logarithmic regression curve. c. 883 Inhibition of WRN induces WRN degradation in a time-dependent manner in MSI-H cells. Steady state 884 Western blot analysis of HCT-116 or U2OS cells treated with 10 µM WRNi over the indicated time points. 885 d. Quantification of Fig. 3b. WRN inhibition leads to a decrease in the half-life of WRN protein. 886 Cycloheximide (CHX) chase experiments in HCT-116 cells in the presence or absence of 10 µM WRNi show 887 a dramatic decrease in the half-life of WRN protein upon its inhibition. Graphs represent n = 2 replicates. 888 The curve fit was done by fitting a half-life decay regression curve. e. WRN degradation upon its inhibition 889 is MSI-H dependent. The indicated MSI-H or MSS cell lines were treated as in a and analyzed by Western 890 blot with the indicated antibodies. f. Quantifications of Fig. 3i. Bar graphs are the mean of n = 3 replicates, 891 each point represents an the average of a well. Error bars represent s.d. g. SMT can be used to measure 892 protein degradation. SMT was used to measure WRN molecules after inhibition of the p97/VCP-893 proteasome pathway, showing a rescue in protein degradation. Each point represents the average WRN 894 spot density within all the nuclei in an FOV. n = 4 plates. Lines represent sample medians. DMSO is 895 dimethyl sulfoxide; CFZ is carfilzomib; p97i is CB-5083; WRNi is HRO761. P-values were calculated using a

two-tailed, unpaired Student's t-test. MW is molecular weight.

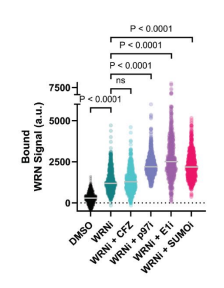


898 Extended Data Fig. 4:

899 a. Previously reported E3 ligases of WRN are not responsible for the WRNi-dependent degradation 900 phenotype. HCT-116 were treated with indicated siRNA oligos for 24 h, and subsequently treated with 10 901 µM WRNi for 16 h and analyzed by Western showing that depletion of the indicated E3 ligases does not 902 rescue the WRN degradation phenotype. b. Identification of additional ubiquitin pathway regulators of 903 WRN regulation. Representative images of HCT-116 cells after depletion of the indicated ubiquitin 904 pathway regulators. Scale bar = 10 μ m. c. Quantification of decomplexified siRNA screen hits from Fig. 4a after treatment of HCT-116^{WRN-Halo} with indicated siRNA oligos for 24 h and then subsequent treatment 905 906 with 10 µM WRNi for 24 h. Bar graphs are the average quantified WRN protein levels for each indicated 907 siRNA oligo of n = 3 plates. Error bars represent s.d.. **d**. SMT can be used to quantify protein degradation. 908 Depletion of RNF4 rescues WRN protein levels after WRN inhibition. WRNi dot plots showing the WRN 909 nuclear spot density from SMT experiments after co-treatment with siRNF4 and either DMSO or WRNi. 910 Each point represents the average WRN spot density within all the nuclei in an FOV. n = 4 plates. Lines 911 represent sample medians. e. RNF4 depletion rescues the WRN degradation phenotype. Dose response 912 curves measuring WRN protein levels by imaging HCT-116^{WRN-Halo} treated with the indicated siRNAs for 24 913 h, and subsequent treatment with WRNi for 24 h. Graphs represent averages from n = 3 plates, measuring 914 3 wells per plate and 6 FOVs per well. Error bars represent s.d., f. Depletion of RNF4 exacerbates DNA 915 damage induced by WRNi. HCT-116 cells were depleted with indicated siRNAs for 24 h, and subsequently 916 subjected to a dose response of WRNi for 16 h. DNA damage was assessed by measuring yH2A.X staining. 917 Graphs represent averages from n = 3 plates, measuring 3 wells per plate and 6 FOVs per well. Error bars 918 represent s.d.. All curve fits were done by fitting a 4-parameter logarithmic regression curve. DMSO is 919 dimethyl sulfoxide; WRNi is HRO761. P-values were calculated using a two-tailed, unpaired Student's t-920 test. MW is molecular weight.



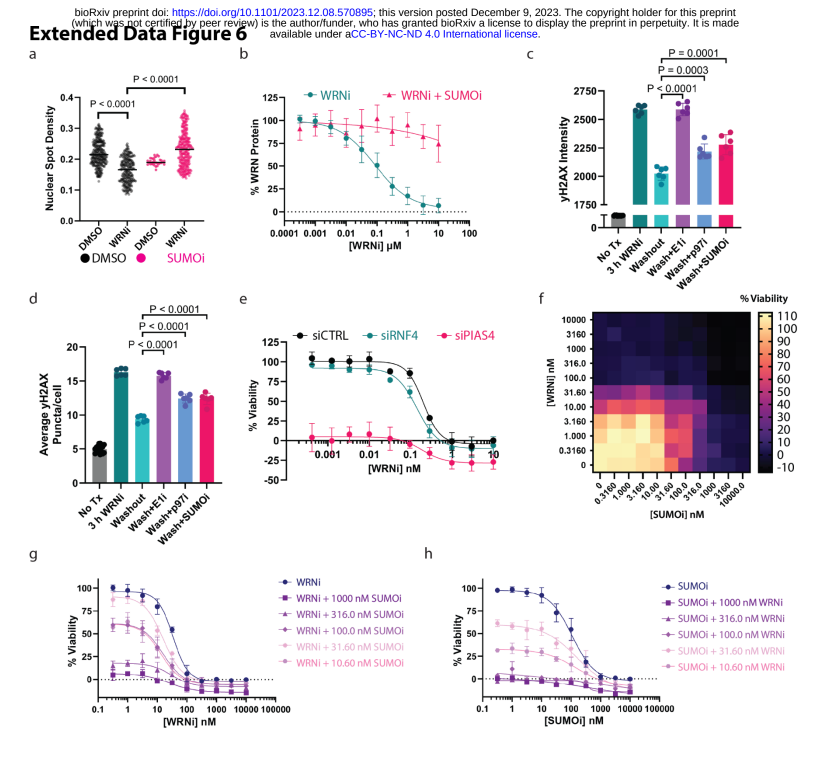
Chromatin-Bound WRN^{Halo}



b

922 Extended Data Fig. 5:

- 923 **a.** The SUMO-Ubiquitin-p97/VCP axis is required to remove trapped WRN from chromatin. Treatment of
- 924 HCT-116-WRN^{Halo} cells with 10 μM WRNi in the presence or absence of 1 μM of CFZ, p97i, E1i, or SUMOi,
- followed by detergent extraction and imaging. **b**. Dot plot quantification of **a**. Each point represents an
- 926 individual cell, measuring the nuclear intensity of WRN. Bars represent the means of the populations.
- 927 DMSO is dimethyl sulfoxide, WRNi is HRO761; CFZ is carfilzomib; p97i is CB-5083; E1i is TAK-243; SUMOi
- 928 is ML-792. P-values were calculated using a two-tailed, unpaired Student's t-test. ns = not significant.



930 **Extended Data Fig. 6:**

931 a. SMT can be used to elucidate molecular regulatory pathways, such as protein degradation. Inhibition 932 of SUMOylation rescues WRN protein levels after WRN inhibition. WRNi Dot plots showing the WRN 933 nuclear spot density from SMT experiments after co-treatment with SUMOi and either DMSO or WRNi. 934 Each point represents the average spot density within all the nuclei in an FOV. n = 4 plates. Lines represent 935 sample medians. b. Inhibition of SUMOylation prevents WRN degradation by WRNi. Dose response curves measuring WRN protein levels by in HCT-116^{WRN-Halo} treated with WRNi in the presence or absence of 1 936 937 μ M SUMOi, and subsequent treatment with WRNi for 24 h. Graphs represent averages from n = 3 plates, 938 measuring 3 well per plate and 6 FOVs per well. Error bars represent s.d. c and d. Quantification of 939 washout experiment in Fig. 5I. d quantifies the average vH2A.X nuclear intensities of an FOV, and c 940 quantifies the average number of vH2A.X puncta per nuclei in an FOV. Bar graphs in both c and d represent 941 the average of n = 3 plates. Each data point represents the average of one well, containing 6 FOVs. Error 942 bars represent s.d.. e. WRNi compound efficacy is independent of WRN degradation but shows potential 943 sensitization to inhibition of the PIAS4-RNF4 axis. HCT-116 cells were treated with the indicated siRNAs 944 for 24 h, followed by WRNi treatment at the indicated doses for 48 h. Cell viability was measured using a 945 CTG2 kit. Graphs represent averages from n = 3 plates. Error bars represent s.d. f. Co-treatment of WRNi 946 with SUMOi has potential synergy. Dose response matrix of both WRNi and SUMOi with indicated 947 concentrations of compound. HCT-116 cells were treated with indicated dose combinations for 48 h. 948 Viability was measured via CTG2. g and h. Quantifications of f, showing dose response plots for the 949 indicated concentration combinations. Graphs represent averages from n = 3 plates. Error bars represent 950 s.d.. All curve fits were done by fitting a 4-parameter logarithmic regression curve. DMSO is dimethyl sulfoxide; WRNi is HRO761; E1 is TAK-243; SUMOi is ML-792; p97i is CB-5083. P-values were calculated 951

952 using a two-tailed, unpaired Student's t-test.