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# Experimental evolution of extremophile levels of radiation resistance in *Escherichia coli*

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#### 12 Abstract

Recent human development of high-level sources of ionizing radiation (IR) prompts a 13 corresponding need to understand the effects of IR on living systems. One approach has 14 15 focused on the capacity of some organisms to survive astonishing levels of IR exposure. Using experimental evolution, we have generated populations of Escherichia coli with IR resistance 16 comparable to the extremophile *Deinococcus radiodurans*. Every aspect of cell physiology is 17 affected. Cellular isolates exhibit approximately 1,000 base pair changes plus major genomic 18 and proteomic alterations. The IR resistance phenotype is stable without selection for at least 19 100 generations. Defined and probable contributions include alterations in cellular systems 20 involved in DNA repair, amelioration of reactive oxygen species, Fe metabolism and repair of 21 iron-sulfur centers, DNA packaging, and intermediary metabolism. A path to new mechanistic 22 discoveries, exemplified by an exploration of *rssB* function, is evident. Most important, there 23 is no single molecular mechanism underlying extreme IR resistance. 24

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# 26 Keywords

27 Ionizing radiation, experimental evolution, reactive oxygen species, double-strand breaks, 28 DNA ropain *Eccharichia coli*, *Deinococcus radiodurans* 

28 DNA repair, Escherichia coli, Deinococcus radiodurans

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# 30 Introduction

Ionizing radiation (IR), particularly from gamma rays, induces severe damage to all 31 cellular macromolecules. Gamma radiation works primarily via H2O radiolysis to produce 32 reactive oxygen species (ROS) (1, 2). Direct ionization and indirect oxidation of DNA through 33 ROS leads to the accumulation of double-strand breaks (DSBs) of genomic DNA, which are 34 lethal if left unrepaired (3-7). Cellular proteomes and lipidomes are also damaged (8-16). Some 35 organisms exhibit an extraordinary capacity to survive high doses of IR. Of seminal interest 36 has been the bacterium Deinococcus radiodurans, which was first isolated due to its ability to 37 survive IR-based sterilization of canned meat (17). Whereas a dose of 3-5 Gy is lethal to a 38 human, Deinococcus can survive 5,000 Gy without lethality (18-21). Research has revealed 39

several mechanisms utilized by *D. radiodurans* to survive high levels of IR exposure, including
specialized mechanisms of DNA end protection and DSB repair, and a notable ability to
accumulate Mn<sup>2+</sup> ions as a means of ameliorating the ROS generated by IR (22-30).

For most of the history of our planet, there have been no environments that would expose living organisms to high levels of ionizing radiation. That situation has changed with the advent of nuclear power, X-rays, the prospect of extended spaceflight, and more. Understanding how cells respond to ionizing radiation becomes more important as the potential for IR exposure increases.

Our understanding of IR resistance is rudimentary at best. The study of extremophiles 48 such as *D. radiodurans* has been illuminating but can never provide a complete view of IR 49 resistance mechanisms. The reason is simple. The evolution of *Deinococcus* was not driven by 50 ionizing radiation. Prior to its discovery in the 1950s (17), this organism was never exposed to 51 the requisite IR-laden environments (31). Instead, Deinococcus is a desert dweller and evolved 52 to survive desiccation (32-35). IR resistance is merely a byproduct of that parched origin. It is 53 thus unlikely that evolution has equipped *Deinococcus* with all possible biological strategies for 54 IR resistance. It is not clear that we know what to look for with respect to additional possible 55 mechanisms. It is not clear that we even understand all the strategies embodied in Deinococcus. 56 More broadly, no research has ever tested the limits. How resistant to ionizing radiation can 57 an organism become? 58

An alternative and unbiased approach to defining mechanisms of IR resistance, pioneered for this phenotype by Evelyn Witkin in 1946 (*36*, *37*), is experimental evolution. Take a species that is not IR resistant, convert it into something that is IR resistant, and see what changes. We are not the first to take this approach (*38-40*). However, previous work was carried out before the development of "omics" technologies that could efficiently identify cellular changes associated with the phenotype.

With a goal of elucidating new mechanisms of IR resistance, we are pursuing a long-65 term evolution experiment with the goal of generating IR resistance in four populations of *E*. 66 coli equivalent to and eventually surpassing D. radiodurans. After 150 cycles of selection carried 67 out over 5 years, our initial goal of IR resistance parity with *Deinococcus* has been achieved. 68 Earlier reports of results after 50 or 100 cycles of selection have described important 69 intermediate steps in this project (41, 42). Here, we present E. coli strains with documented 70 extremophile levels of IR resistance. Two of our four replicate populations of evolved E. coli 71 now match the survival of *D. radiodurans* out to a dose of 6,000 Gy of IR, whereas doses in 72 73 excess of 1,000 Gy effectively sterilize cultures of the Founder E. coli strain. Combined genomic and proteomic approaches have revealed that changes to systems involved in DNA repair, 74 ROS amelioration, Fe metabolism, and aerobic metabolism have partially driven acquired IR 75 76 resistance in at least one of the four evolved populations. A path to describing additional mechanisms is evident. In addition, we demonstrate that an elevated stress response 77 contributes to IR resistance. The overall effort highlights the lack of a single molecular 78 79 mechanism underlying biological resistance to ionizing radiation. Layers of contributing mechanisms are evident within these four evolved *E. coli* populations. 80

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#### 82 **Results and Discussion**

#### 83 Ionizing radiation resistance has matched that of Deinococcus radiodurans

To summarize our experimental evolution protocol (detailed in the Materials and 84 *Methods*), at each cycle of selection, each of the four replicate populations is initially grown to 85 early exponential phase and cultures are washed with PBS to remove all nutrients or potential 86 ROS-ameliorating agents present in growth media. The washed cultures are treated with 87 sufficient IR to kill 99% of each of the four replicate populations at each cycle of selection (thus 88 the dose used increases as IR resistance increases). The irradiation utilizes a clinical linear 89 accelerator (Linac) dosing at 70 Gy/min. Following irradiation, a portion of each culture is used 90 to determine percent survival and the rest is allowed to recover in fresh growth medium 91 overnight before storage of the new population at -80 °C the following day. This protocol has 92 resulted in four evolved lineages (IR9, IR10, IR11, and IR12), with distinct populations at each 93 cycle of selection (i.e. IR9-150 is the population from the IR9 lineage at round 150 of selection), 94 and 10 isolates from each population (i.e. IR9-150-1 and IR9-150-2 are two separate isolates 95 from population IR9-150). We note that population IR9 has two sub-populations which have 96 competed in clonal interference for about 80 selection cycles through round 150 (42). Isolates 97 IR9-150-1 and IR9-150-2 are taken from different sub-populations and exhibit many genetic 98 and proteomic distinctions in spite of their presence in the same population. 99

After beginning with a dose of 500 – 750 Gy, at round 150 of selection a dose of 3700-4000 Gy is required to achieve 99% killing **(Figure S1)**. Furthermore, two populations, IR9-150 and IR10-150, now exhibit survival curves equivalent to *D. radiodurans*, where cultures of each population and *D. radiodurans* have approximately 0.1% survival at a dose of 6000 Gy **(Figure 1**).

105 Radioresistance in these four populations is a stable phenotype. After growth for ~100 generations in the absence of IR exposure, isolates from each of these populations maintains 106 the same level of IR resistance (Figure S2). However, these isolates are not free of fitness 107 tradeoffs. Each isolate has clear growth rate defects (Figure S3) as well as readily observable 108 changes in cell morphology (Figure S4). Such differences are expected, as after 150 cycles of 109 selection isolates contain about 1000 single-nucleotide polymorphisms (SNPs) and other small 110 111 changes on average (Table S1). Larger genomic alterations are evident in the populations as 112 well. Some were reported previously (42). A new genomic deletion present in IR11 is added in 113 Figure S5.

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#### 115 *Evolved isolates exhibit enhanced DNA repair and resistance to protein oxidation*

DNA double strand breaks accumulate with increasing doses of IR and must be repaired in order for the exposed cell to survive. Thus, we assayed IR-mediated damage to genomic DNA and the dynamics of DNA repair in evolved isolates from each population after a dose of 1000 or 4000 Gy, utilizing Pulsed-Field Gel Electrophoresis (PFGE). All evolved isolates demonstrated a clear capacity to repair shattered genomic DNA at a dose of 4000 Gy, attesting to enhanced capacity for DNA repair. In contrast, the Founder strain did not recover from this

dose (Figure 2). These results align with our previous observations that by rounds 50 and 100 122 of selection, numerous mutations in DNA repair proteins (RecA, RecD, RecN, and RecJ) were 123 drivers of evolved IR resistance (41, 42). Many of these mutants affect proteins involved in 124 double strand break repair. The evolved isolates exhibited different rates of DNA repair. 125 126 Repaired genomic DNA is apparent in isolate IR12-150-1 only 3 hr post-irradiation with 4000 Gy. Furthermore, IR11-150-1 and IR12-150-1 appear to better protect their genomic DNA from 127 IR-mediated DSBs, where cells harvested immediately post-irradiation with 1000 Gy contain 128 at least some more or less intact genomes. Thus, it appears that the evolved lineages may have 129 developed differing mechanisms of maintaining genome stability despite equivalent IR 130 exposure. 131

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#### 133 Proteomics and genomics provide a new entrée to mechanisms of evolved IR resistance

The four evolved populations were subjected to deep sequencing to identify all 134 mutations present in more than 2% of the cells (Supplementary Dataset S1). Genes subject to 135 mutation in multiple populations are of particular interest as potential candidates for 136 phenotype contributions, and these are listed in **Table S2**. In parallel, using mass spectrometry 137 approaches we have described previously (10, 11), we surveyed the proteomes of the Founder 138 wild type strain and two evolved isolates from the current experiment. Analysis of the 139 quantified IR9-150-1 and IR9-150-2 proteomes at early exponential phase growth under normal 140 growth conditions (no IR) revealed significant changes to the composition of their proteomes 141 compared to MG1655 (Figure 3). With a significance threshold of at least a 2-fold increase or 142 decrease (adjusted p-value < 0.05), each isolate had over 300 proteins with altered abundance 143 144 (IR9-150-1: 156 proteins increased and 165 decreased; IR9-150-2: 150 proteins increased and 317 145 proteins decreased). Such changes potentially reveal what pathways are beneficial or expendable for IR resistance. 146

Across both isolates, large changes in abundance of greater than 10-fold (observed in 18 147 proteins increasing and 52 proteins decreasing in IR9-150-1, and 24 proteins increasing and 42 148 decreasing in IR9-150-2) are largely explained through genomic sequencing data. Large 149 increases in abundance are often due to mutations in their cognate regulatory proteins. For 150 example, the pentose phosphate pathway protein RpiB is increased due to a frameshift in its 151 regulator, RpiR (also called AlsR) (43, 44); pilus proteins from the fim operon are likely 152 increased due to mutations in the FimE site-specific recombinase, locking the *fim* promoter in 153 the 'on' orientation (45-47); and RpoS is increased in IR9-150-1 due to an early frameshift in its 154 regulator, RssB, that effectively eliminates RssB function (48-50). Some increases in protein 155 abundance may be linked to intergenic mutations affecting transcription or translation (such 156 as two mutations at –150 and +21 relative to the transcription start site of *uvrB* in both IR9-150 157 158 isolates), or due to regulatory changes (i.e. Fe-starvation, explained below). Large decreases in protein abundance are typically due to introduced stop codons or frameshift mutations, or 159 160 deletion of the gene encoding the protein in question (particularly due to a > 100 kb deletion in 161 IR9 (42)).

In prior work characterizing the evolved lineages after 50 (41) and 100 (42) cycles of 162 selection, we found that evolved IR resistance appeared to proceed through at least two distinct 163 phases: (1) adaptation of DNA repair pathways in all four lineages, followed by (2) lineage-164 specific adaptations, potentially focusing at least in part on adaptations to respiration, Fe-S 165 166 cluster repair, and polyamine metabolism in lineage IR9 (42). The proteomics and additional genomic sequencing conducted now at round 150 of selection largely confirm the trends first 167 discovered through genomics datasets, as well as highlight new potential players in evolved 168 radioresistance. In the following analysis, we highlight mutations confirmed to be contributors 169 to IR resistance as well as proteomic changes that suggest additional mechanisms to guide 170 additional investigation. 171

DNA repair. The proteomics effort highlights adaptations of the DNA repair systems 172 that were not documented earlier. Each of the two round 150 isolates have increased levels of 173 proteins involved in nucleotide excision repair (UvrB, LigA) as well as proteins involved in 174 remodeling DNA (RuvA, GyrA, GyrB, ParC), suggesting new ways to bolster a cells capacity 175 to process damaged DNA substrates (Figure S6). In conjunction with increased levels of GyrA 176 (IR9-150-2) and GyrB (IR9-150-1), a decrease in the gyrase inhibitor SbmC in both isolates 177 further underlines the potential importance of increased gyrase activity in radioresistance. 178 Additional decreases in DNA repair proteins focus on exo and endonucleases (RecD, RecJ, 179 SbcD, Nth). As IR-induced double strand breaks increase, the capacity of exonucleases to 180 wreak genomic havoc also increases. Hence, a reduction in cellular nuclease activity may 181 contribute to the maintenance of genome integrity in a high IR environment. Mutations in 182 proteins involved in double strand break repair (RecD, RecN, RecJ, RecA) make clear 183 184 contributions to IR resistance, as documented previously (41), and the IR resistance conferred 185 by the changes in RecD and RecJ may reflect declines in nuclease function.

Response to ROS. ROS generated by IR or respiration have the capacity to not only 186 generate DNA damage, but also oxidize proteins side chains and destroy protein-coordinated 187 Fe-S clusters (51, 52). IR9-150-1 and IR9-150-2 exhibit extensive proteomic changes which are 188 suggestive of suppressing ROS formation linked to respiration by reducing the level of 189 associated proteins (Figure S6). Furthermore, both isolates exhibit significant increases in 190 proteins involved in repair of damaged Fe-S clusters, in particular members of the suf operon 191 (Figure S6). These changes build on previous observations that variants of the ATP synthase 192 component AtpA and Fe-S repair protein SufD were partially responsible for the radioresistant 193 phenotype of lineage IR9 after 100 cycles of selection (42). 194

The new proteomics data show that isolate IR9-150-1 has increased levels of the ROS 195 amelioration enzymes Hmp, SodA, and AhpF in addition to a large increase in the stress 196 response sigma factor RpoS (which in turn controls transcription of the ROS amelioration 197 198 proteins PqiA, PqiB, PqiC, and KatE). All of these changes have the potential to reduce the oxidative damage to cellular constituents of all kinds. In contrast, numerous ROS-responsive 199 proteins are observed at decreased levels in IR9-150-2 (KatG, PqiB, PqiC, OxyR, Dps) 200 201 suggesting that the two sub-populations of IR9 are pursuing quite distinct paths to IR 202 resistance.

Fe metabolism. In addition to proteomic confirmation of adaptations previously 203 identified through genomics, the proteome of IR9-150-2 exhibits a strong indication of altered 204 Fe metabolism. Pathways focused on the production and uptake of Fe-scavenging siderophore 205 enterobactin are all increased in abundance, while Fe-storage proteins Dps and Bfr are both 206 207 decreased. The changes to Fe metabolism may be related to a frameshift in the ExbB protein (ExbB S67fs), which would additionally knock ExbD out of frame. The ExbBD protein complex 208 (in conjunction with TonB) provides the mechanical energy necessary to facilitate enterobactin 209 uptake across the outer membrane. Without these proteins cells can no longer utilize 210 enterobactin as a means of Fe acquisition, leading to Fe starvation (53, 54). Interestingly, metals 211 analysis through ICP-MS (Figure S7) did not reveal a decreased concentration of Fe in IR9-150-212 2 cells, suggesting that while an inability to uptake enterobactin may trigger the Fe starvation 213 response, the cells are able to acquire Fe by other means. However, the Mn/Fe ratio was 214 somewhat increased in several of the populations, perhaps telegraphing some evolutionary 215 movement towards an ROS amelioration mechanism characterized by Daly and coworkers (22-216 30). 217

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An altered stress response through RpoS highlights evolutionary divergence of IR resistance
 mechanisms

The new proteomic and genomic data highlight many potential new contributions to 221 the IR resistance phenotype, but each requires testing. To illustrate how these evolved 222 223 populations can be exploited for mechanistic discovery, we further explored the frameshift in 224 gene *rssB*, resulting in increased expression of RpoS. The frameshift occurs early in the gene (at codon 3), essentially deleting *rssB* function. This mutation is present in one sub-population 225 226 of IR9, exemplified by isolate IR9-150-1, but not in the sub-population from which isolate IR9-150-2 is derived or in population IR10. The *rssB* mutation contributes to IR resistance in a 227 context-dependent manner. When a wild type *rssB* gene is restored in IR9-150-1, while 228 retaining all other mutations in this evolved isolate, there is no evident effect on IR resistance 229 at 3,000 Gy. However, at 4,000 Gy, IR resistance is strongly suppressed (Figure 4). Thus, the 230 *rssB* mutation is important in this genetic background primarily at high doses of IR. When an 231 rssB deletion was engineered into isolate IR9-150-2, no increase in IR resistance was observed. 232 Instead, the change proved to be slightly deleterious (Figure 4). A similar deletion engineered 233 into an isolate from population IR10 (IR-150-1) had no significant effect on IR resistance. An 234 rssB deletion introduced into a wild type background produced a significant increase in IR 235 resistance at 1,000 Gy (Figure S8), providing additional confirmation that this alteration can 236 contribute to the IR resistance phenotype. Notably, elevated expression of RpoS in a wild type 237 background can slow growth under at least some conditions (55), an effect that may have 238 239 constrained the emergence of similar mutations earlier in the experimental evolution trial. The mutation does not measurably add to the IR resistance of IR9-150-2 or IR10-150-1, indicating 240 that its effects depend on genomic context. 241

Whereas we have identified quite a number of mutations that contribute to IR resistance 243 244 in the present study and previously (41, 42), we cannot yet account for the complete phenotype in any one population or sub-population. Some obvious candidates for additional study are 245 revealed here and more will be identified. As exemplified by the *rssB* mutation, these 246 247 experimentally evolved populations provide a robust platform for the discovery of new mechanisms of IR resistance, although complexity is evident. This study also makes several 248 more general and important points. First, we have not yet reached a dose of ionizing radiation 249 250 that would preclude survival if appropriate genomic changes are present. The potential for amelioration of IR-mediated damage in living organisms is great and may be greater than that 251 seen in any extant extremophile studied to date. Second, there is not just one way to survive 252 253 the damage inflicted by IR. Instead, survival may depend on layers of mechanisms that both prevent IR-inflicted damage and facilitate repair of any damage that occurs. The existence of 254 multiple paths to IR resistance may eventually offer multiple points of intervention for cells 255 from bacteria to human that may require this phenotype. Finally, it is likely that at least some 256 of the mechanisms that may contribute to IR resistance involve processes never or rarely 257 discussed in the context of this phenotype. Contributions by alterations in genes involved in 258 DNA packaging (cadA) and repair of Fe-S clusters (sufD)(42), as well as the rssB mutation 259 described here, provide examples. Experimental evolution provides an unbiased path to their 260 discovery. 261

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# 276 **Dedication**

277 This paper is dedicated to Evelyn Witkin on the occasion of her 100<sup>th</sup> birthday.

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#### 279 Data and Resource Availability Statement:

280 Mutations detected via deep sequencing at round 150 are listed in Supplementary Dataset 1. MS

281 datasets for each strain are available online at the Proteomics Identification Database

282 (<u>https://www.ebi.ac.uk/pride/</u>; accession number: PXD024784) and are included as

- 283 Supplementary Dataset 2. All data utilized here are incorporated within this publication or in
- one of these datasets. In addition, all "omics" data, as well as all bacterial populations,
- isolates, and cellular constructs associated with this project, <u>both published and unpublished</u>,
- are available to any interested researcher at any time. For a current listing, please inquire.
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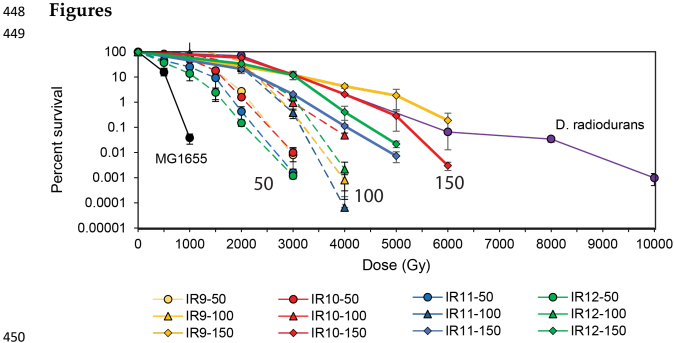
# 289 References290

- J. A. Reisz, N. Bansal, J. Qian, W. L. Zhao, C. M. Furdui, Effects of ionizing radiation on biological molecules-mechanisms of damage and emerging methods of detection. *Antiox. Redox. Signal.* 21, 260-292 (2014).
- P. A. Riley, Free radicals in biology oxidative stress and the effects of ionizing radiation. *Int. J. Rad. Biol.* 65, 27-33 (1994).
- 3. G. Iliakis, The role of DNA double strand breaks in ionizing radiation-induced killing
  of eukaryotic cells. *Bioessays* 13, 641-648 (1991).
- R. E. Krisch, F. Krasin, C. J. Sauri, DNA breakage, repair, and lethality accompanying
   125I decay in microorganisms. *Current Topics Rad. Res.h Quarterly*. 12, 355-368 (1978).
- 3005.T. Haaf *et al.*, Sequestration of mammalian Rad51-recombination protein into301micronuclei. J. Cell Biol. 144, 11-20 (1999).
- M. Toma, T. Skorski, T. Sliwinski, DNA double strand break repair related synthetic
   lethality. *Curr. Med. Chem.* 26, 1446-1482 (2019).
- 304 7. C. Shee *et al.*, Engineered proteins detect spontaneous DNA breakage in human and
  305 bacterial cells. *eLife* 2, e01222 (2013).
- G. Z. Cao *et al.*, Effects of X-ray and carbon ion beam irradiation on membrane
   permeability and integrity in Saccharomyces cerevisiae cells. *J. Rad. Res.* 56, 294-304
   (2015).
- 309 9. A. Krisko, M. Radman, Protein damage and death by radiation in *Escherichia coli* and
  310 *Deinococcus radiodurans*. *Proc. Natl. Acad. Sci. U.S.A.* 107, 14373-14377 (2010).
- S. T. Bruckbauer *et al.*, Ionizing radiation-induced proteomic oxidation in *Escherichia coli*. *Mol. Cell. Proteom.* 19, 1375-1395 (2020).
- B. B. Minkoff, S. T. Bruckbauer, G. Sabat, M. M. Cox, M. R. Sussman, Covalent
  modification of amino acids and peptides Induced by ionizing radiation from an
  electron beam linear accelerator used in radiotherapy. *Rad. Res.* 191, 447-459 (2019).
- G. H. Xu, M. R. Chance, Hydroxyl radical-mediated modification of proteins as probes
  for structural proteomics. *Chem. Rev.* 107, 3514-3543 (2007).
- A. F. Fragopoulou *et al.*, Hippocampal lipidome and transcriptome profile alterations
  triggered by acute exposure of mice to GSM 1800 MHz mobile phone radiation: An
  exploratory study. *Brain Behav.* 8, (2018).
- 14. E. L. Pannkuk *et al.*, A lipidomic and metabolomic serum signature from nonhuman
   primates exposed to ionizing radiation. *Metabolomics* 12, (2016).
- M. Upadhyay *et al.*, Identification of plasma lipidome changes associated with low
   dose space-type radiation exposure in a murine model. *Metabolites* 10, (2020).
- R. L. Chang *et al.*, Protein structure, amino acid composition and sequence determine
   proteome vulnerability to oxidation-induced damage. *EMBO J.* 39, (2020).
- A. W. Anderson, H. C. Nordon, R. F. Cain, G. Parrish, D. Duggan, Studies on a radioresistant micrococcus. I. Isolation, morphology, cultural characteristics, and resistance
  to gamma radiation. *Food Technol.* 10, 575-578 (1956).

- J. R. Battista, Against all odds the survival strategies of deinococcus radiodurans.
   *Ann. Rev. Microbiol.* 51, 203-224 (1997).
- M. M. Cox, J. R. Battista, *Deinococcus radiodurans* The consummate survivor. *Nature Rev. Microbiol.* 3, 882-892 (2005).
- 334 20. M. J. Daly, OPINION A new perspective on radiation resistance based on *Deinococcus* 335 *radiodurans*. *Nature Rev. Microbiol*. 7, 237-245 (2009).
- D. Slade, M. Radman, Oxidative Stress Resistance in *Deinococcus radiodurans*. *Microbiol. Mol. Biol. Rev.* 75, 133-191 (2011).
- E. Bentchikou, P. Servant, G. Coste, S. Sommer, A major role of the RecFOR pathway
   in DNA double-strand-break repair through ESDSA in *Deinococcus radiodurans*. *PLoS Genet* 6, e1000774 (2009).
- 341 23. D. A. Bernstein *et al.*, Crystal structure of the *D. radiodurans* single-stranded DNA
  342 binding protein suggests a novel mechanism for coping with DNA damage. *Proc. Natl.*343 *Acad. Sci. U.S.A.* 101, 8575-8580 (2004).
- M. J. Daly, L. Ouyang, P. Fuchs, K. W. Minton, In vivo damage and *recA*-dependent
  repair of plasmid and chromosomal DNA in the radiation-resistant bacterium *Deinococcus radiodurans. J. Bacteriol.* **176**, 3508-3517 (1994).
- M. J. Daly, K. W. Minton, An alternative pathway of recombination of chromosomal fragments precedes *recA*-dependent recombination in the radioresistant bacterium *Deinococcus radiodurans*. J. Bacteriol. **178**, 4461-4471 (1996).
- M. J. Daly *et al.*, Accumulation of Mn(II) in, *Deinococcus radiodurans* facilitates gamma radiation resistance. *Science* 306, 1025-1028 (2004).
- 27. D. R. Harris *et al.*, Preserving Genome Integrity: The DdrA Protein of *Deinococcus radiodurans* R1. *PLoS Biology* 2, e304 (2004).
- D. R. Harris, K. V. Ngo, M. M. Cox, The stable, functional core of DdrA from
   *Deinococcus radiodurans* R1 does not restore radioresistance in vivo. *J Bacteriol* 190,
   6475-6482 (2008).
- J.-I. Kim, M. M. Cox, The RecA proteins of *Deinococcus radiodurans* and *Escherichia coli*promote DNA strand exchange via inverse pathways. *Proc. Natl. Acad. Sci. U.S.A.* 99,
  7917-7921 (2002).
- 360 30. C. Norais *et al.*, The *Deinococcus radiodurans* DR1245 Protein, a DdrB Partner
  361 Homologous to YbjN Proteins and Reminiscent of Type III Secretion System
  362 Chaperones. *PLoS One* 8, (2013).
- 363 31. P. A. Karam, S. A. Leslie, Calculations of background beta-gamma radiation dose
  364 through geologic time. *Health Physics* 77, 662-667 (1999).
- 365 32. J. K. Fredrickson *et al.*, Protein oxidation: key to bacterial desiccation resistance? *Isme J.*366 2, 393-403 (2008).
- 367 33. V. Mattimore, J. R. Battista, Radioresistance of *Deinococcus radiodurans*: functions
   368 necessary to survive ionizing radiation are also necessary to survive prolonged
   369 desiccation. J. Bacteriol. 178, 633-637 (1996).
- 370 34. M. Tanaka *et al.*, Analysis of Deinococcus radiodurans's transcriptional response to
   ionizing radiation and desiccation reveals novel proteins that contribute to extreme
   radioresistance. *Genetics* 168, 21-33 (2004).
- 373 35. F. A. Rainey *et al.*, Extensive diversity of ionizing-radiation-resistant bacteria
  374 recovered from Sonoran Desert soil and description of nine new species of the genus
  375 Deinococcus obtained from a single soil sample. *Appl. Environ. Microbiol.* 71, 5225-5235
  376 (2005).
- 377 36. E. M. Witkin, A case of inherited resistance to radiation in bacteria. *Genetics* 31, 236-236
  378 (1946).

379 37. E. M. Witkin, inherited differences in sensitivity to radiation in *Escherichia coli*. Proc. Natl. Acad. Sci. U.S.A. 32, 59-68 (1946). 380 38. R. Davies, A. J. Sinskey, Radiation-resistant mutants of Salmonella typhimurium LT2: 381 382 development and characterization. J Bacteriol **113**, 133-144 (1973). 383 39. I. E. Erdman, F. S. Thatcher, K. F. Macqueen, Studies on the irradiation of microorganisms in relation to food preservation. II. Irradiation resistant mutants. Can J 384 *Microbiol* 7, 207-215 (1961). 385 40. A. Parisi, A. D. Antoine, Increased radiation resistance of vegetative Bacillus pumilus. 386 387 Appl Microbiol 28, 41-46 (1974). S. T. Bruckbauer *et al.*, Experimental evolution of extreme resistance to ionizing 41. 388 radiation in *Escherichia coli* after 50 cycles of selection. J. Bacteriol. **201**, e00784 (2019). 389 42. S. T. Bruckbauer et al., Physiology of highly radioresistant Escherichia coli after 390 experimental evolution for 100 cycles of selection. *Frontiers Microbiol.* **11**, 582590 (2020). 391 392 43. K. I. Sorensen, B. HoveJensen, Ribose catabolism of Escherichia coli: Characterization 393 of the *rpiB* gene encoding ribose phosphate isomerase B and of the *rpiR* gene, which is 394 involved in regulation of *rpiB* expression. J. Bacteriol. **178**, 1003-1011 (1996). 44. G. A. Sprenger, Genetics of pentose-phosphate pathway enzymes of Escherichia coli K-395 396 12. Archives Microbiol. 164, 324-330 (1995). 397 45. I. C. Blomfield, D. H. Kulasekara, B. I. Eisenstein, Integration host factor stimulates 398 both FimB- and FimE-mediated site-specific DNA inversion that controls phase variation of type 1 fimbriae expression in *Escherichia coli*. Molecular Microbiology 23, 399 400 705-717 (1997). 46. S. A. Joyce, C. J. Dorman, A Rho-dependent phase-variable transcription terminator 401 402 controls expression of the FimE recombinase in *Escherichia coli*. Mol. Microbiol. 45, 1107-1117 (2002). 403 404 47. M. S. McClain, I. C. Blomfield, B. I. Eisenstein, Roles of FimB and FimE in site-specific DNA inversin associated with phase variation of type-1 fimbriae in *Escherichia coli*. J. 405 Bacteriol. 173, 5308-5314 (1991). 406 G. Becker, E. Klauck, R. Hengge-Aronis, Regulation of RpoS proteolysis in Escherichia 407 48. *coli*: The response regulator RssB is a recognition factor that interacts with the turnover 408 element in RpoS. Proc. Natl. Acad. Sci. U.S.A. 96, 6439-6444 (1999). 409 49. D. Micevski, J. E. Zammit, K. N. Truscott, D. A. Dougan, Anti-adaptors use distinct 410 modes of binding to inhibit the RssB-dependent turnover of RpoS (sigma(S)) by 411 412 ClpXP. Frontiers Mol. Biosci. 2, (2015). F. Mika, R. Hengge, A two-component phosphotransfer network involving ArcB, 50. 413 ArcA, and RssB coordinates synthesis and proteolysis of sigma(S) (RpoS) in E-coli. 414 Genes Develop. 19, 2770-2781 (2005). 415 J. A. Imlay, The mismetallation of enzymes during oxidative stress. J. Biological 51. 416 417 *Chemistry* **289**, 28121-28128 (2014). J. A. Imlay, R. Sethu, S. K. Rohaun, Evolutionary adaptations that enable enzymes to 52. 418 419 tolerate oxidative stress. Free Rad. Biol. Med. 140, 4-13 (2019). 53. B. M. M. Ahmer, M. G. Thomas, R. A. Larsen, K. Postle, Characterization of the exbBD 420 421 operon of *Escherichia coli* and the role of ExbB and ExbD in TonB function and 422 stabillity. J. Bacteriol. 177, 4742-4747 (1995). S. Maki-Yonekura *et al.*, Hexameric and pentameric complexes of the ExbBD energizer 423 54. in the Ton system. *Elife* 7, (2018). 424 O. Patange *et al.*, Escherichia coli can survive stress by noisy growth modulation. 425 55. *Nature Commun.* 9, (2018). 426

- 427 56. J. H. Miller, A Short Course in Bacterial Genetics: A Laboratory Manual and Handbook for
  428 Escherichia coli and Related Bacteria. (Cold Spring Harbor Laboratory, Cold Spring
  429 Harbor, NY, 1992).
- 430 57. F. R. Blattner *et al.*, The complete genome sequence of *Escherichia coli* K-12. *Science* 277, 1453-1474 (1997).
- 432 58. S. Warming, N. Costantino, D. L. Court, N. A. Jenkins, N. G. Copeland, Simple and
  433 highly efficient BAC recombineering using gaIK selection. *Nuc. Acids Res.* 33, e36
  434 (2005).
- 435 59. K. A. Datsenko, B. L. Wanner, One-step inactivation of chromosomal genes in
  436 *Escherichia coli* K-12 using PCR products. *Proc. Natl. Acad. Sci. U.S.A.* 97, 6640-6645
  437 (2000).
- 438 60. P. R. Almond *et al.*, AAPM's TG-51 protocol for clinical reference dosimetry of high-439 energy photon and electron beams. *Med. Phys.* **26**, 1847-1870 (1999).
- 440 61. D. R. Harris *et al.*, Directed evolution of radiation resistance in *Escherichia coli*. J.
  441 Bacteriol. 191, 5240-5252 (2009).
- 442 62. H. Li, R. Durbin, Fast and accurate short read alignment with Burrows-Wheeler transform. *Bioinformatics* 25, 1754-1760 (2009).
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452 Figure 1. After 150 cycles of selection experimentally evolved *E. coli* populations exhibit IR resistance comparable to *D. radiodurans*. Survival curves of evolved populations compared 453 to previously evolved Escherichia coli populations and Deinococcus radiodurans. MG1655 is the 454 Founder strain used to begin the evolution experiment. IR'X'-50 and IR'X'-100 are populations 455 of the indicated lineages after 50 or 100 cycles of selection with IR. Early exponential phase 456 cultures of the indicated strains were exposed to electron beam IR as described in the Materials 457 458 and methods. Error bars represent the standard deviation of CFU/mL calculations from a single experiment performed in biological triplicate. Survival data for MG1655, populations after 50 459 cycles of selection, and D. radiodurans strain R1 at 2000 and 6000 Gy are as previously reported 460 (41, 42). 461

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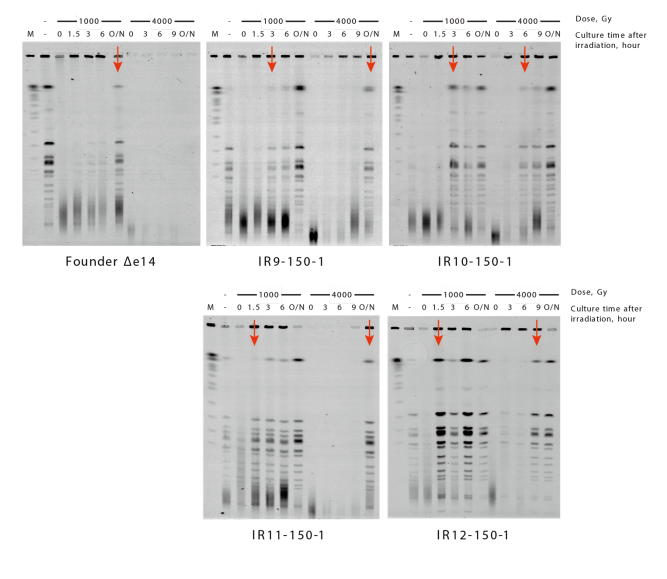
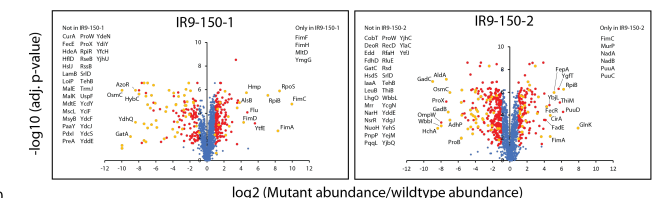


Figure 2. Evolved isolates experience and repair extensive IR-induced DNA damage. Pulsed-field gel electrophoresis (PFGE) was used to assay the extent and repair of DNA damage induced by IR exposure at 1000 and 4000 Gy. Genomic DNA is digested with restriction enzyme NotI to facilitate observation of intact versus degraded genomes. At 4000 Gy, cultures of MG1655 are completely killed and no repair of genomic DNA is observed. Red arrows indicate the first appearance of a banding pattern suggesting intact genome. Results are representative of two independent experiments. PFGE was performed as described previously (61), and as in *Materials and methods*. 



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Figure 3. The proteome composition of IR9-150-1 and IR9-150-2 have diverged from E. coli 483 484 MG1655. Volcano plots depict each protein detected with a circle, with the fold difference in abundance compared to the same protein in MG1655 on the x-axis and the p-value of that 485 change on the y-axis. Proteins with a significant fold change are colored in red. Numerous 486 proteins were not detected, or only detected in the evolved isolates. Such proteins are listed 487 next to their respective volcano plots, as they cannot be plotted due to no actual fold increase 488 or decrease. The proteome composition of MG1655, IR9-150-1, and IR9-150-2 was determined 489 utilizing label free quantification mass spectrometry (LFQ-MS). Significant changes in protein 490 abundance were defined as an increase or decrease greater than two-fold, with adjusted p-491 values less than 0.05 (calculated using Benjamini-Hochberg correction). Supplementaary 492 Datasets 1 and 2 contain the genomics and proteomics data used for these analyses, 493 494 respectively.

	10 <sup>0</sup>	10 <sup>-1</sup>	10-2	<sup>2</sup> 10	<sup>3</sup> 10	<sup>4</sup> 10 <sup>-5</sup>	10 <sup>0</sup>	10 <sup>-1</sup>	10 <sup>-2</sup>	10 <sup>-3</sup>	10 <sup>-4</sup>	10 <sup>-5</sup>	10 <sup>0</sup>	10 <sup>-1</sup>	10 <sup>-2</sup>	10-3	10 <sup>-4</sup>	10 <sup>-5</sup>
IR9-150-1				•	<b>E</b>	-		۲	۲	1.1.4 1.5.4	1			۲	菊	٨,	•	
IR9-150-2				۲	dig.	***		۲		1	•			۲	٢	$\hat{s}^{*}_{i}$	•	•
IR10-150-1	•		۲	۲	1.22	****		۲	۲	16.	·.**	•		۲		ŝ.	•	÷.
IR9-150-1 + rssB wt	•	•		•	-	-		۲	٢	14	•		60		•			
IR9-150-2 ∆rssB			۲	0		4		۲	S.	-25				۲	15		in.	-:
IR10-150-1 ∆rssB			۲	۲	¢	• <sup>8</sup> •		۲	۲	索	Ÿ.			۲	۲	1.5	· · · ·	
	0 Gy						3000 Gy						4000 Gy					

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500 Figure 4. Loss of *rssB* enhances IR resistance in a context-dependent manner. When *rssB* is

converted to the wildtype sequence in IR9-150-2, a loss of IR-resistance is observed at 4000 Gy.
However, at 3000 Gy no effect is observed from loss of the frameshifted *rssB*, highlighting the
dose-dependent context of some beneficial mutations. Furthermore, when *rssB* is deleted from
IR9-150-2, or IR10-150-1 we see no effect (IR10-150-1) or deleterious phenotypes (IR9-150-2) at
the doses tested. Therefore, beneficial effects from the loss of *rssB* is also dependent on genomic
context (see Figure S8 for the effects of an *rssB* deletion in a wild type background.