1 Supplementary Information

3 **Predictable and precise template-free CRISPR editing of pathogenic variants**

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48 Supplementary Discussion

49

50 Cellular repair of double-stranded DNA breaks and inDelphi

51 DNA double-strand breaks are detrimental to genomic stability, and as such the detection 52 and faithful repair of genomic lesions is crucial to cellular integrity. A large number of 53 genes have evolved to respond to and repair DNA double-strand breaks, and these genes 54 can be broadly grouped into a set of DNA repair pathways¹, each of which differs in the 55 biochemical steps it takes to repair DNA double-strand breaks. Accordingly, these 56 pathways tend to produce characteristically distinguishable non-wildtype genotypic 57 outcomes.

58

59 The goal of our machine learning algorithm, inDelphi, is to accurately predict the identities 60 and relative frequencies of non-wildtype genotypic outcomes produced following a 61 CRISPR/Cas9-mediated DNA double-strand break. To accomplish this goal, we 62 developed parameters to classify three distinct categories of genotypic outcomes, 63 microhomology deletions, microhomology-less deletions, and insertions, informed by the 64 biochemical mechanisms underlying the DNA repair pathways that typically give rise to 65 them.

66

67 Double strand breaks are thought to be repaired via four major pathways: classical non-68 homologous end-joining (c-NHEJ), alternative-NHEJ (alt-NHEJ), microhomologymediated end-joining (MMEJ), and homology-directed repair (HDR)¹. To create inDelphi, 69 70 we developed three machine learning modules to model genotypic outcomes assuming 71 characteristic of the c-NHEJ, microhomology mediated alt-NHEJ, and MMEJ pathways. 72 While template-free CRISPR/Cas9 DNA double-strand break may lead to HDR repair via 73 endogenous homology templates that exist in *trans*², we do not explicitly model HDR-74 characteristic outcomes using our algorithm.

75

Before proceeding, it is important to note that while specific DNA repair pathways are 76 77 characteristically associated with distinct genotypic outcomes, the proteins involved in the 78 various pathways and the resulting repair products may at times overlap. This fact has 79 several implications. First, we cannot make conclusive statements about the role of 80 specific proteins or pathways in specific genotypic outcomes without perturbation experiments (e.g. our comparison of wildtype and *Prkdc^{-/-}Lig4^{-/-}* mESCs can illuminate the 81 82 roles of these proteins, specifically). Second, because assigning genotypic outcomes to 83 biochemical mechanisms is likely imperfect, we use machine learning methods to identify 84 trends and patterns in genotype frequencies that refine this crude binning process.

85

In the first step of the inDelphi method, we separate genotypic outcomes into three classes: microhomology deletions (MH deletions), microhomology-less deletions (MH-less deletions), and single-base insertions (1-bp insertions) (Figure 1e). Below we outline the algorithmic definitions of each genotypic outcome class, the pathways associated with each class, and the DNA sequence parameters included in inDelphi training of each class. For more detailed technical algorithmic definitions of the genotypic outcome classes, see Supplementary Methods

- 92 Supplementary Methods.
- 93

94	MH deletions are predicted from MH length, MH GC content, and deletion length
95	The majority of Cas9-mediated double-strand break repair genotypes we observe in our
96	datasets are what we classify as MH deletions (53-58% in mESC, K562, HCT116, and
97	HEK293). We hypothesize that these deletions occur through MMEJ-like processes and
98	use known features of this pathways to inform a machine learning module to predict MH
99	deletion outcomes. Following 5'-end resection as occurs in MMEJ, alt-NHEJ, and HDR ¹ ,
100	microhomologous basepairing of single-stranded DNA (ssDNA) sequences occurs
101	across the border of the double strand breakpoint ^{3,4} . To restore a contiguous double-
102	strand DNA chain, the 5'-overhangs not participating in the microhomology are removed
103	up until the paired microhomology region, and the unpaired ssDNA sequences are
104	extended by DNA polymerase using the opposing strand as a template (Figure 1d,
105	Extended Data Fig. 2).
106	
107	Assuming these same processes, inDelphi calculates the set of all MH deletions available
108	given a specific sequence context and cleavage site.
109	
110	As an example workflow, given the following sequence and its cleavage site:
111	
112	ACGTG CATGA
113	TGCAC GTACT
114	
115	for every possible deletion length from 1-bp to 60-bp deletions, we overlap the 3'-
116	overhang downstream of the cut site under the upstream 3'-overhang and determine if
117	there is any microhomologous basepairing. As an example, given the 4-bp deletion
118	length:
119	
120	ACGTG
121	
122	GTACT
123	
124	we see that there are three microhomologous basepairing events.
125	
126	we then choose a particular micronomology (here, the highlighted C:G):
127	A C C C C
120	
129	
121	GIACI
121	then generate its unique repair genetice by following left to right along the ten strend and
132	impring down to the complement of the bettern strend to simulate DNA polymerose fill
133	jumping down to the complement of the bottom strand to simulate DNA polymerase mi-
134	11.
135	Here this vields:
130	
137	ΑΓΑΤΩΑ
139	ТСТАСТ
140	
110	

141 This can also be displayed as an alignment. We note that by "jumping down" after the first 142 base in the top strand, we can also describe this outcome using the delta-position 1. (See 143 section on delta-positions). A deletion at delta-position 0 yields the same genotype.

- 144 145 **Deletion a:** AC---ATGA
- 146 Wt: ACGTGCATGA
- 147

This same sequence context and cleavage site could produce a distinct 4-bp MH deletion genotypic outcome through use of the TG:AC microhomology. This single outcome can be described as using delta-positions 2, 3, or 4. inDelphi uses only the single maximum delta-position (here, 4) to described a unique MH deletion.

- 152

 153
 Deletion b:
 ACGTG----A

 154
 Wt:
 ACGTGCATGA
- 155

Thus, there may be multiple MH deletion outcome genotypes for a given deletion length, and there is always a 1:1 mapping between the microhomologous basepairing used in that MH deletion and the resultant genotypic outcome. The set of MH deletions thus includes all 1-bp to 60-bp deletions that can be derived from the steps above that simulate the MMEJ mechanism.

161

162 MMEJ efficiency has been reported to depend on the thermodynamic favorability and stability of a candidate microhomology^{3,4}. To parameterize MH deletions using the 163 biochemical sequence features that influence this form of DNA repair, inDelphi calculates 164 165 the MH length, MH GC content, and resulting deletion length for each possible MH 166 deletion. These features are input into a machine learning module referred to in the Supplementary Methods as the microhomology neural network (MH-NN) to learn the 167 168 relationship between these features and the frequency of an MH deletion outcome in a 169 training CRISPR/Cas9 genotypic outcome dataset. While we predict and empirically find 170 that favored MH deletions have long MH lengths relative to total deletion length and high 171 MH GC-contents, we do not provide any explicit direction or comparative weighting to 172 these parameters at the outset. inDelphi then outputs a phi-score for any MH deletion 173 genotype (whether it was in the training data or not) that represents the favorability of that 174 outcome as predicted by MH-NN.

175

176 It is important to emphasize that the phi-score of a particular MH deletion does not itself 177 represent the likelihood of that MH deletion occurring in the context of all MH deletions at a given site. Some CRISPR/Cas9 target sites may have many possible favorable MH 178 179 deletion outcomes while other sites have few, and thus phi-score must be normalized for 180 a given target site to generate the fractional likelihood of that genotypic outcome at that site. Total unnormalized MH deletion phi-score is one factor that is further used to predict 181 182 the relative frequency of the different repair classes: MH deletions, MH-less deletions, 183 and insertions.

- 184
- 185

186 MH-less deletions are predicted from their length

We define MH-less deletions as all possible deletions that have not been accounted for by the workflow described above for MH deletions. Mechanistically, our data analysis suggests that MH deletions are associated with repair genotypes produced by c-NHEJ and microhomology-mediated alt-NHEJ pathways.

191

192 Following a double-strand break, c-NHEJ-associated proteins rapidly bind the DNA 193 strands flanking the double-strand DNA breakpoint and recruit ligases, exonucleases, and 194 polymerases to process and re-anneal the breakpoint in the absence of 5'-end resection (Extended Data Fig. 2)^{1,5}. Commonly, c-NHEJ repair is error-free; however, in the context 195 196 of Cas9-mediated cutting, faithful repair leads to repeated cutting, thereby increasing the 197 eventual likelihood of mutagenic repair. Erroneous c-NHEJ repair products are mainly 198 thought to consist of small insertions or deletions or combinations thereof that most frequently occur in the direct vicinity of the DNA break point⁵⁻⁷. The resulting deletions, 199 200 which we refer to as medial end-joining MH-less deletions, have often lost bases both 201 upstream and downstream of the cleavage site.

202

203 Microhomology-mediated alt-NHEJ is a distinct pathway that produces MH-less deletion 204 products. In contrast to c-NHEJ, which is microhomology independent, this form of alt-205 NHEJ repair occurs following 5'-end resection and is mediated by microhomology in the sequence surrounding the double-strand break-point¹. Microhomologous basepairing 206 207 stabilizes the 3'-ssDNA overhangs following 5'-end resection, similarly to in MMEJ, 208 allowing DNA ligases to join the break across one of the strands of this temporarily 209 configured complex. The opposing un-annealed flap is then removed, and newly 210 synthesized DNA templated off of the remaining strand is annealed to repair the lesion 211 (Extended Data Fig. 2).

212

213 While alt-NHEJ uses microhomology, the repair products it produces do not follow the 214 predictable genotypic patterns induced by MMEJ and are thus grouped into MH-less 215 deletion genotypes. MH deletions are a direct merger of both annealed strands, in which 216 the outcome genotype switches from top to bottom strand at the exact end-point of a 217 microhomology. In contrast, while alt-NHEJ employs microhomology in its repair 218 mechanism, the deletion outcomes it generates comprise bases exclusively derived from 219 either the top or bottom strand. Mechanistically, this occurs because ligation of a 3'-220 overhang to its downstream ligation partner results in removal of the entire opposing 221 ssDNA overhang up until the point of ligation. This process prevents any deletion from 222 occurring in the 3'-overhang strand that is first attached to the DNA backbone, while 223 inducing loss of an indeterminant length of sequence on the opposing strand. The 224 resulting deletion genotypes, which we refer to as unilateral end-joining MH-less 225 deletions, do not retain information on the exact microhomology causal to their 226 occurrence, and are thus also referred to as MH-less.

227

228 Consequently, the various mechanisms that give rise to MH-less deletions are capable of 229 generating a vast number of genotypic outcomes for any given deletion length. Having 230 less information on the biochemical mechanisms that impact the relative frequency of NHEJ deletion products, inDelphi models these deletions without assuming any particular
 mechanism.

232 mecl 233

inDelphi detects MH-less deletions from training data as the set of all deletions that are
not MH deletions and parameterizes them solely by the length of the resulting deletion.
This is based on the simple assumption that c-NHEJ and alt-NHEJ processes are most
likely to produce short deletions, supported by our empirical observation. As with MH
deletions, this assumption is not explicitly coded into the inDelphi MH-less deletion
prediction module, instead allowing it to be "learned" by a neural network called MHlessNN.

241

MHIess-NN optimizes a phi-score for a given MH-less deletion length, grounded in the frequency of MH-less deletion outcomes of that length observed in the training data. We observe that MHIess-NN learns a near-exponential decaying phi-score for increasing deletion length, that reflects the sum total frequency of all MH-less deletion genotypes. The total unnormalized MH-less deletion phi-score for a given target and cut site is also

- 247 employed to inform the relative frequency of different repair classes.
- 248

249 **1-bp insertions are predicted from sequence context and deletion phi-scores**

250 Lastly, inDelphi predicts 1-bp insertions from both the broader sequence context and the 251 immediate vicinity of the cleavage site. We empirically find that 1-bp insertions are far 252 more common than longer insertions, so we focus on their prediction. It is classically assumed that short sequence insertions are the result of c-NHEJ^{6,7}, however, little else is 253 known about their biochemical mechanism as it pertains to local sequence context to help 254 255 inform prediction. Nonetheless, we find powerful correlations between the identities of the 256 bases surrounding the Cas9 cleavage site and the frequency and identity of the inserted 257 base (see main text). Motivated by these empirical observations, inDelphi is fed with training data on 1-bp insertion frequencies and identities at each training site 258 259 parameterized with the identities of the -3, -4, and -5 bases upstream of the NGG PAMsequence (when the training set is sufficiently large, and the -4 base alone when training 260 data is limited) as features. Also added as features are the precision score of the deletion 261 262 length distribution and the total deletion phi-score at that site. These features are combined into a k-nearest neighbor algorithm that predicts the relative frequencies and 263 identities of 1-bp insertion products at any target site. 264

265

The combination of the MH, MH-less, and insertion model predict genotype fractions

Altogether, informed by known paradigms of DNA repair, we build 2 neural networks and 268 269 a k-nearest neighbor model to predict genotypic outcomes following Cas9 cutting. These 270 models compete and collaborate in inDelphi to generate predictions of the relative frequencies of these products. This competition within inDelphi among repair types 271 reflects empirical evidence from Lib-A and Lib-B that sequence contexts do influence 272 273 classes of repair outcomes. Sequence contexts with high phi scores (high 274 microhomology) have higher efficiencies of MH deletions among all editing outcomes 275 (Figure 2d, Extended Data Fig. 3), and sequence contexts with low phi scores (low microhomology) have higher efficiencies of 1-bp insertions among all editing outcomes 276

277 (Figure 2d, Extended Data Fig. 3). While it is tempting to generalize that the competition 278 and collaboration among outcome classes modeled by inDelphi reflects interactions 279 among components of distinct DNA repair pathways, the classes of outcomes considered 280 by inDelphi do not necessarily arise from distinct DNA repair pathways as they are described above. inDelphi is trained on the repair outcomes only and cannot distinguish 281 282 between the nature of genotypes when they may occur through MH-mediated and MH-283 less mechanisms, and it is imaginable that some repair products result through more than 284 one repair pathway.

285

As an additional note, while NHEJ is generally assumed to dominate double-strand break repair from environmentally induced damage⁵, we find in the context of Cas9 cutting that MH deletion genotypes are more common than MH-less deletions and insertions. It is possible that error-free c-NHEJ is occurring frequently in response to Cas9 cutting but that its perfect repair allows for recurring Cas9 cutting that goes undetected by our workflow, thus skewing the observed relative frequency profile of mutagenic outcomes toward MMEJ-type repair.

293

294 Rarer CRISPR-Cas9 outcomes

295 Our library assay and workflow involved data processing of high-throughput sequencing 296 data using sequence alignments and a designed procedure for categorizing sequence 297 alignments into categories of CRISPR-related outcomes. Beyond simple deletions and 298 insertions, we identified other rarer outcomes that were explained as indels caused by 299 CRISPR, such as combination insertion/deletions involving and/or near the cleavage site 300 (0.5-2% of all products) and indels near but not immediately at the cleavage site (3-5% of 301 all products), which occurred more often on the PAM-distal side of the double-strand 302 break (data not shown). Our library assay is unable to observe events that occur outside 303 of our high-throughput sequencing window.

304

305 Default sequence alignment procedures can generate sequence alignments involving simple CRISPR-caused deletions and insertions that do not occur immediately at the 306 cleavage site, but that can be transformed into an equal-scoring sequence alignment 307 308 where the indel does occur immediately at the cleavage site. This straightforward 309 processing step is not performed by the most common bioinformatic tools for sequence alignment, since they were not expressly designed for CRISPR. We note here that our 310 311 sequence alignment procedure takes this into account (see Supplementary Methods for 312 more detailed description). This attention to detail enables us to accurately identify simple 313 indels that occur near but not immediately at the cleavage site. We observe that the 314 frequency of these indels across target sites correlates significantly with the total on-target 315 editing efficiency (measured by the frequency of non-wild-type outcomes out of all nonnoise outcomes) at these target sites in HEK293 and mES cells. We also observe 316 significantly higher frequencies in postCas9 treatment conditions than preCas9 control 317 318 conditions. Together, these observations suggest that these indels are caused by 319 CRISPR editing.

320

321 *Prkdc^{-/-}Lig4^{-/-}* mutants have distinct and predictable DNA repair product 322 distributions While it is generally true that our work cannot establish roles for specific DNA repair pathways in specific types of Cas9-mediated outcomes, we have performed an experiment in which we measure Cas9-mediated genotypic outcomes from mESCs that are lacking *Prkdc* and *Lig4*, two proteins known to be key in c-NHEJ⁵. We find an increase in relative frequency of MH deletions as compared to MH-less deletions in *Prkdc*-/-*Lig4*-/- mESCs as compared to wild-type mESCs (see main text), which is suggestive of an increase in MMEJ outcomes at the expense of NHEJ outcomes.

330

Intriguingly, we also find that *Prkdc*^{-/-}*Lig4*^{-/-} mESCs are impaired in unilateral deletions, 331 332 where only bases from one side of the cutsite are removed, but not medial MH-less 333 deletion outcomes that have loss of bases on both sides of the breakpoint. (Extended 334 Data Fig. 6). As discussed earlier, microhomology-mediated alt-NHEJ, which we 335 hypothesize may give rise to unilateral MH-less deletions, proceeds through a mechanism 336 in which DNA repair intermediates that mimic MMEJ-mediated repair are formed initially 337 (Extended Data Fig. 2), as microhomology base-pairing temporarily stabilizes 3'-338 overhangs following 5'-end resection. Subsequently, ligation joins one 3' overhang with the sequence on the other side of the DNA double-strand break, giving rise to a unilateral 339 340 deletion. If the unilateral joining products we observe in our experiments indeed arise through similar mechanisms as those described by this form of alt-NHEJ, it is conceivable 341 342 that the MMEJ pathway may overtake 3'-end ligation at this microhomology-containing 343 intermediate step when ligation is impaired through loss of Lig4. Thus, cross-talk of 344 microhomology-mediated repair pathways could account for loss of unilateral end-joining 345 MH-less outcomes and concomitant increase in MH deletion outcomes. Medial joining outcomes are not hypothesized to originate from intermediates that overlap with 346 347 microhomology-mediated deletion products (Extended Data Fig. 2). Therefore, the repair 348 genotypes generated via this orthogonal pathway may be afforded more time to be completed by ligases other than Lig4, thus explaining why these outcomes appear 349 350 unaffected by NHEJ impairment.

351

While DNA repair products in Prkdc-/-Lig4-/- mESCs differ substantially from those in wild-352 type cells, we find that these DNA repair products are also highly predictable. In particular, 353 354 inDelphi performed well on held-out Prkdc-/-Lig4-/- data when trained on Prkdc-/-Lig4-/- data (indel genotype prediction median Pearson correlation = 0.84, indel length frequency 355 prediction Pearson correlation = 0.80), showing that our modeling approach is robustly 356 357 capable of learning accurate predictions for Cas9 editing data in not just wild-type 358 experimental settings but also settings with significant biochemical perturbation. As such, 359 we suggest here that inDelphi's modeling approach can be useful on additional tasks 360 unexplored here provided that inDelphi is supplied with appropriate training data.

361

362 NU7041, DPKi3, and MLN4924 induce a distinct DNA repair product distribution

We further investigated the role of DNA repair pathways by three separate experiments involving HTS characterization of Lib-B in mESCs treated with three separate small molecules: NU7041, a DNA dependent protein kinase (DNA-PK) inhibitor; DPKi3, another DNA-PK inhibitor, and MLN4924, a NEDD8-activating enzyme (NAE) inhibitor. DNA-PK and NAE are proteins involved in c-NHEJ^{5,8}.

- 369 MLN4924 is thought to inhibit the release of the Ku70/Ku80 heterodimer following proper 370 c-NHEJ repair, potentially disrupting downstream processes such as transcription and 371 replication, which may lead to decreased cell survival and a depletion of Ku70/Ku80-372 dependent DNA repair genotypes in a population.⁸
- 373
- DNA-PK is commonly recruited to DSBs during c-NHEJ and is known to phosphorylate *in vitro* many c-NHEJ-related factors including Ku70/80, XRCC4, DNA Ligase IV, Artemis, H2AX, p53, and itself. Inhibition of DNA-PK leads to DNA repair defects⁵. The catalytic subunit of DNA-PK is encoded by the Prkdc gene, which was knocked out in Prkdc^{-/-}Lig4⁻/- cells.
- 379

380 From HTS data, we observed that the frequency of MH deletions among all deletions 381 clustered into three approximate groups: wild-type (median 77%) and MLN4924, then DPKi3 and NU7041 (median 81%), and lastly Prkdc^{-/-}Lig4^{-/-} (median 90%) (Extended Data 382 383 Fig. 6). These data suggest that impairing DNA-PK (via DPKi3, NU7041 and Prkdc^{-/-}) 384 yields a moderate 17% reduction in the frequency of MH-less deletions (23% to 19%). This reduction appears to be non-redundant with knockout of Lig4 evidenced in Prkdc⁻ 385 ¹Lig4⁻¹ cells with a 57% reduction (23% to 10%) in MH-less frequency. Lastly, impairing 386 NAE did not have a significant impact on the frequency of MH-less deletions. 387

388

We observed an overall increased frequency of repair to wild-type at pathogenic microduplication alleles after treatment with DPKi3, MLN4924, and NU7041 (Extended Data Fig. 6). Along with Prkdc^{-/-}Lig4^{-/-} cells, the change in repair efficiency was associated with deletion length ($p < 2.2x10^{-3}$), with decreased efficiency compared to wild-type at short deletion lengths and increased efficiency at longer deletion lengths.

394

The change in repair efficiency caused by separate treatments of DPKi3, MLN4924, and NU7041 was highly consistent across different target sites (r = 0.73, 0.77, and 0.81,Extended Data Fig. 6). This is surprising since MLN4924 inhibits a different target than DPKi3 and NU7041. We observed a similar but weaker relationship between the three small molecules and *Prkdc^{-/-}Lig4^{-/-}*, with Pearson correlations of 0.09, 0.16, and 0.18. Taken together, these observations suggest a relationship between DNA sequence and the propensity of DNA repair outcomes through c-NHEJ.

402

In DPKi3, MLN4924, and NU7041 treated cells, the decrease in MH-less deletions primarily occurs medial joining products (Extended Data Fig. 6), suggesting that DNA-PK is a strong contributor to medial joining products. However, when both DNA-PK and Lig4 are knocked out in *Prkdc^{-/-}Lig4^{-/-}* cells, the average frequency of medial joining products is not significantly changed, and instead the primary decrease occurs in unilateral joining products.

409

410 Interestingly, MLN4924 increases the average frequency of unilateral joining events.

411 Combined with its effect of decreasing medial joining products, the overall net effect of

412 MMLN4924 is an absence of significant change to the frequency of MH-less deletions.

The frequency distribution of medial joining products in *Prkdc^{-/-}Lig4^{-/-}* reveals a decrease in median frequency in combination with an inflation in high frequency outliers (target sites where >80% of all deletion products are MHless medial products) which skews the distribution's average to be above the median. Taken together, these data confirm that both medial and unilateral products are both generally depleted in *Prkdc^{-/-}Lig4^{-/-}* cells, and suggest that knocking out DNA-PK depletes medial MHless products while knocking out Lig4 depletes unilateral MHless products.

- 422 Supplementary Methods
- 423
- 3
- 424 **Library cloning protocol** 425
- 426 Synthesized oligo library sequence
- 427 GATGGGTGCGACGCGTCAT [55bpTarget] AGATCGGAAGAGCACACGTCTG**AATA<u>TT</u>GTGGA**
- 428 <u>AAGGACGAAACACCG[19/20-nt_PROTOSPACER</u> depending on whether it
- 429 naturally starts with a G]GTTTAAGAGCTATGCTGGAAACAGC
- 430
- 431 Linker region / Oligo library amplification primer anneal region
- 432 Read 2 sequencing primer stub
- 433 Sspl restriction site
- 434 <u>U6-promoter stub</u>
- 435 sgRNA-hairpin stub
- 436
- 437
- 438 1. Oligo library QPCR to determine number of amplification cycles for Oligo
 439 Library PCR
- 440 Notes: Amplification of oligos with relatively low GC-content is less efficient than GC-rich
 441 sequences. We found NEBNext polymerase to be the least biased in amplification of our
 442 library. Increasing the elongation time to 1 min per cycle for all cloning and sequencing
 443 library prep PCRs eliminates GC-skewing of library sequences and reduces the rate of
 444 PCR-recombination.
- 445 446
- Set up the following reaction:
- 447

0.4 ng	Synthesized Oligo Library
10 ul	NEBNext 2x Master Mix
0.5 ul	20uM OligoLib_Fw
0.5 ul	20uM OligoLib_Rv
0.2 ul	SybrGreen Dye (100x)
to 20 ul	H ₂ O

448

- 449 67°C annealing temperature
- 450 451
- Check 246bp amplicon size on 2.5% agarose gel.
- 452 Determine the point that signal amplification has plateaued.453

454 **2. Oligo Library PCR amplification**

- Set up the following reaction:
- 455 456

4 ng	Synthesized Oligo Library
50 ul	NEBNext 2x Master Mix
2.5 ul	20uM OligoLib_Fw
2.5 ul	20uM OligoLib_Rv
to 100 ul	H ₂ O

- 458 67°C annealing temperature, 1 minute extension time.

459 Cycle number is half the number of cycles needed to reach signal amplification plateau 460 in the QPCR in step 1, reduced by 1 cycle to scale for DNA input.

- PCR purify amplified sequence.

463464 3. Donor template amplification

- 465 Set up the following reaction:

5 ng	spCas9 sgRNA plasmid (71485)
50 ul	NEBNext 2x Master Mix
2.5 ul	20uM CircDonor_Fw
2.5 ul	20uM CircDonor_Rv
to 100 ul	H ₂ O

- 468 62°C annealing temperature
- **20 cycles**

471 - Gel purify 167bp band from 2.5% agarose gel.

4. Circular assembly and restriction digest linearization

474 Note: We use a molar ratio of donor template to amplified oligo library of 3:1. An increase
475 in amplified oligo library compounds cross-over within library members resulting in
476 mismatch of protospacer and target sequences.
477

- Set up the following reaction:

429 ng	Donor template
239 ng	Amplified Oligo Library
30 ul	Gibson Assembly 2x Master Mix
to 60 ul	H ₂ O

50°C incubation for 1 hour.

- Exonuclease treatment

60 ul	Circular assembly reaction
9 ul	ATP (25mM)
9 ul	10x Plasmid Safe Buffer
3 ul	Plasmid Safe Nuclease
9 ul	H ₂ O

37°C incubation for 1 hour.

488 - PCR purify and elute in 50 ul.

- 489 Digest to linearize library
- 490

50 ul	Purified assemblies
10 ul	10x CutSmart Buffer
3 ul	SspI-HF
37 ul	H ₂ O

491

- 37° C incubation for ≥ 3 hours. 492
- 493 Gel purify 273bp band from 2.5% agarose gel. _
- 494

495 Note: Band is sometimes fuzzy and poorly visible. If not clearly discernible, proceed with 496 gel isolation between 200-300bp.

497

Linearized library QPCR to determine number of amplification cycles for PCR 498 5. 499 amplification

- 500
- Set up the following reaction: -

501

0.5 %	Purified linearized library
10 ul	NEBNext 2x Master Mix
0.5 ul	20uM PlasmidIns_Fw
0.5 ul	20uM PlasmidIns_Rv
0.2 ul	SybrGreen Dye (100x)
to 20 ul	H ₂ O

502

503 65°C annealing temperature

504 505

Determine the point that signal amplification has plateaued.

506 507 6. Linearized Library PCR amplification

- Set up the following reaction:
- 508 509

50 %	Purified linearized library
50 ul	NEBNext 2x Master Mix
2.5 ul	20uM PlasmidIns_Fw
2.5 ul	20uM PlasmidIns_Rv
to 100 ul	H ₂ O

510

- 511 65°C annealing temperature, 1 minute extension time.
- Cycle number is number of cycles needed to reach signal amplification plateau in the 512
- QPCR in step 5, reduced by 4 cycles to scale for increased DNA input. 513
- 514
- 515 Gel purify 375bp band from 2.5% agarose gel. -

516 517 Vector backbone digest 7.

- 518 Set up the following reaction:
- 519

2 ug	spCas9 sgRNA plasmid (71485)
10 ul	10x Buffer 2.1
3 ul	BbsI
2 ul	Xbal
to 100 ul	H ₂ O

520

522

523

- 521 37° C incubation for ≥ 3 hours.
 - Gel purify 5.9 kb band from 1% agarose gel.

524 8. Vector assembly and cleanup

525 Note: Include a ligation with water for insert as a control.

526 527

- Set up the following reaction:

528

300	ng	Digested vector backbone
42 ng		Amplified Oligo Library
30 ul		Gibson Assembly 2x Master Mix
to 60 ul		H ₂ O

529

530 50°C incubation for 1 hour.

531 532

Isopropanol precipitation

533

40 ul	Vector assembly reaction
0.4 ul	GlycoBlue Coprecipitant
0.8 ul	50mM NaCl
38.8 ul	Isopropanol

534

537

- 535 Vortex and incubate at room temperature for 15 minutes.
- 536 Spin down at \geq 15.000g for 15 minutes, and carefully remove supernatant.
 - Wash pellet with 300ul 80% EtOH and repeat spin at ≥15.000g for 5 minutes.
- 538 Carefully remove all liquid without disturbing pellet, and let air dry for 1-3 minutes.
- 539 Dissolve dried pellet in 10 ul H₂O at 55°C for 10 minutes.

540

541 **9.** Transformation

Note: Electroporation competent cells give a higher transformation efficiency than chemically competent cells. We use NEB10beta electro-competent cells, however these can be substituted for other lines and transformed according to the manufacturer's instructions.

546

547 Note: We use DRM as recovery and culture medium to enhance yield. If substituting for 548 a less rich medium such as LB, we recommend scaling up the culture volume to obtain 549 similar plasmid DNA quantities.

551 Note: Antibiotic-free recovery time should be limited to 15 minutes to prevent shedding of 552 transformed plasmids from replicating bacteria.

- 553
- 554 Note: Also transform water ligation as control. 555
- Pre-warm 3.5mL recovery medium per electroporation reaction, at 37°C for 1 hour.
- 557 Pre-warm LB-agar plates containing appropriate antibiotic.
- 558 Per reaction, add 1 ul purified vector assembly to 25ul competent cells on ice.
 559 Perform 8 replicate reactions.
- 560 Electroporate according to the manufacturer's instructions.
- 561 Immediately add 100 ul pre-warmed recovery media per cuvette and pool all 562 replicates into culture flask.
- Add 1 mL recovery media per replicate reaction to culture flask and shake at
 200rpm 37°C for 10 15 minutes.
- Plate a dilution series from 1:10⁴ 1:10⁶ on LB-agar plates containing antibiotic
 and grow overnight at 37°C
- 567 Add 2 mL media per replicate reaction and admix appropriate antibiotic.
- 568 Grow overnight in shaking incubator at 200rpm 37°C
- Assess transformation efficiency from serial dilution LB-agar plates. Expect ~10⁶
 clones.
- 572

573 The development of this cloning protocol was guided by work described in Videgal et al. 574 2015.

576 Sequence alignment and data processing

577 For library data, each sequenced pair of gRNA fragment and target was associated with 578 a set of designed sequence contexts G by finding the designed sequence contexts for all 579 gRNAs whose beginning section perfectly matches the gRNA fragment (read 1 in general 580 does not fully sequence the gRNA), and by using locality sensitive hashing (LSH) with 7-581 mers on the sequenced target to search for similar designed targets. An LSH score on 7-582 mers between a reference and a sequenced context reflects the number of shared 7-583 mers between the two. If the best reference candidate scored, through LSH, greater than 584 5 higher than the best LSH score of the reference candidates obtained from the gRNA-585 fragment, the LSH candidate is also added to G. LSH was used due to extensive (~33%) rate) PCR recombination between read1 and read2 which in sequenced data appears as 586 587 mismatched read1 and read2 pairs. The sequenced target was aligned to each candidate 588 in G and the alignment with the highest number of matches is kept. Sequence alignment 589 was performed using the Needleman-Wunsch algorithm using the parameters: +1 match, 590 -1 mismatch, -5 gap open, -0 gap extend. For library data, starting gaps cost 0. For all 591 other data, starting and ending gaps cost 0. For VO data, sequence alignments were 592 derived from SAM files from SRA.

593

594 Alignments with low-accuracy or short matching sections flanked by long (10 bp+) 595 insertions and deletions were filtered out as PCR recombination products (observed 596 frequency of ~5%). These PCR recombination products are different than that occurring 597 between read1 and read2; these occur strictly in read2. Alignments with low matching 598 rates were removed. Deletions and insertions were shifted towards the expected 599 cleavage site while preserving total alignment score. CRISPR-associated DNA repair 600 events were defined as any alignment with deletions or insertions occurring within a 4 bp 601 window centered at the expected cut site and any alignment with both deletions and insertions (combination indel) occurring with a 10 bp window centered at the expected 602 603 cut site. All CRISPR-associated DNA repair events observed in control data had their 604 frequencies subtracted from treatment data to a minimum of 0.

605

606 We carried out replicate experiments for library data in each cell type. For each cell-type, 607 each target site not fulfilling the following data quality criteria was filtered: in each 608 replicate, data at this target site must have a total of at least 1,000 reads for all CRISPR editing outcomes at that target site (see section on "Calling CRISPR editing outcomes 609 610 with high confidence" below for a discussion on the 1,000 reads threshold), and a Pearson 611 correlation of at least 0.85 in the frequency of microhomology-based deletion events. The class of microhomology-based deletion events was used for this criterion since it is a 612 613 major repair class with the highest average replicability across experiments.

614

615 **Details on alignment processing**

All alignments with gaps were shifted as much as possible towards the cleavage site while preserving the overall alignment score. Then, the following criteria were used to categorize the alignments into noise, not-noise but not CRISPR-associated (for example, wildtype); as well as primary and secondary CRISPR activity. All data used in modeling and analysis derive solely from outcomes binned into primary CRISPR activity.

- 622 The following criteria was used to filter library alignments into "noise" categories.
- 623
- 624 Homopolymer: Entire read is homopolymer of a single nucleotide. Not considered a 625 CRISPR repair product.
- Has N: Read contains at least one N. Discarded as noise, not considered a CRISPR repair product.
- 628 PCR Recombination: Contains recombination alignment signature: (1) if a long indel (10
- bp+) followed by chance overlap followed by long indel (10 bp+) of the opposite type, e.g.,
- 630 insertion-randommatch-deletion and deletion-randommatch-insertion. OR, if one of these
- two indels is 30 bp+, the other can be arbitrarily short. If either criteria is true, and if the chance overlap is length 5 or less, or any length with less than 80% match rate, then it
- satisfies the recombination signature. In addition, if both indels are 30 bp+, regardless of
- the middle match region, it satisfies the recombination signature. Finally, if randommatch
- 635 is length 0, then indel is allowed to be any length. Not considered a CRISPR repair 636 product.
- ⁶³⁷ Poor-Matches: 55bp designed sequence context has less than 5 bp representation (could
- 638 occur from 50 bp+ deletions or severe recombination) or less than 80% match rate. Not 639 considered a CRISPR repair product.
- 640 Cutsite-Not-Sequenced: The read does not contain the expected cleavage site.
- 641 Other: An alignment with multiple indels where at least one non-gap region has lower
- than an 80% match rate. Or generally, any alignment not matching any defined category
- above or below. In practice, can include near-homopolymers. Not considered a CRISPR
- 644 repair product.
- 645
- 646 The following criteria was used to filter library alignments into "main" categories.
- 647 Wildtype: No indels in all of alignment. Not considered a CRISPR repair product.
- 648
- 649 Deletion: An alignment with only a single deletion event. Subdivided into:
- 650 Deletion Not CRISPR: Single deletion occurs outside of 4 bp window centered around 651 cleavage site. Not considered a CRISPR repair product.
- 652 Deletion Not at cut: Single deletion occurring within 4 bp window centered around
- 653 cleavage site, but not immediately at cleavage site. Considered a CRISPR repair product.
- Deletion: Single deletion occurring immediately at cleavage site. Considered a CRISPR
- 655 repair product.
- 656
- 657 Insertion: An alignment with only a single insertion event. Subdivided into:
- Insertion Not CRISPR: Single insertion occurs outside of 10 bp window around cleavagesite. Not considered a CRISPR repair product.
- 660 Insertion Not at cut: Single insertion occurring within 4 bp window centered around
- 661 cleavage site, but not immediately at cleavage site. Considered a CRISPR repair product.
- Insertion: Single insertion occurring immediately at cleavage site. Considered a CRISPR
 repair product.
- 664
- 665 Combination indel: An alignment with multiple indels where all non-gap regions have at 666 least 80% match rate. Subdivided into:

667 Combination Indel: All indels are within a 10 bp window around the cleavage site. 668 Considered a primary CRISPR repair product.

669 Forgiven Combination Indel: At least two indels, but not all, are within a 10 bp window

- around the cleavage site. Considered a rarer secondary CRISPR repair product, ignored.
- 671 Forgiven Single Indel: Exactly one indel is within a 10 bp window around the cleavage 672 site. Considered a rarer secondary CRISPR repair product, ignored.
- 673 Combination Indel Not CRISPR: No indels are within a 10 bp window around the 674 cleavage site. Not considered a CRISPR repair product.
- 675
- We note that deletion and insertion events, even those spanning many bases, are defined to occur at a single location between bases. As such, events occurring up to 5 bp away from the cleavage site are defined as events where there are five or fewer matched/mismatched alignment positions between the event and the cleavage site, irrespective of the number of gap dashes in the alignment.
- 681

682 Calling CRISPR editing outcomes with high confidence

- Following the processing steps above, we performed the following further analysis and processing steps to call high-confidence CRISPR editing outcomes. These steps largely follow heuristics, and we believe that a thorough and unbiased methodological standardization in counting CRISPR editing outcomes will be valuable future work.
- 687

688 DNA repair at Cas9-mediated double-strand breaks is known to result in a large diversity 689 of outcomes, with indels of varying length and positions around the cleavage site. The 690 frequencies of many of these editing outcomes, though enriched in Cas9-treatment data 691 over control data, are rare (<0.5% of all edited products) and can be challenging to assign 692 as a CRISPR editing outcome due to a lack of foundational biological or computational 693 models on the exact mechanisms of DNA repair. In addition, rare outcomes can 694 sometimes be attributed to sequencing errors.

695

In this work, we focus on CRISPR editing outcomes that are enriched in treatment data over control data and agree with a relatively conservative and strict model of DNA repair, in order to ensure a high degree of confidence in the editing outcomes that we call. As a result, we underestimate the total number of unique CRISPR editing outcomes, though we believe this underestimation is not by an order of magnitude, though it may be by a factor of 2x or so.

702

703 We define high-confidence CRISPR editing outcomes as bins of alignments categorized 704 by the previously described pipeline into CRISPR-associated categories that have no 705 mismatches. Each unique deletion genotype consistent with microhomology is treated as 706 a single unique outcome, though we note that microhomology deletions may arise by 707 noise or chance though we expect this to be a rare event. Each unique insertion genotype 708 is also treated as a single unique outcome, and as with MH deletions, we note that some 709 insertions may arise by noise or chance though we anticipate this to be rare. In sum, we 710 likely overestimate by a slight amount the true number of unique microhomology deletion 711 and insertion events.

713 All microhomology-less deletion genotypes are binned together for a particular deletion 714 length, which almost always will bin together multiple unique MH-less deletion genotypes. 715 However, the class of MH-less deletions, in general, has lower replicate consistency and 716 higher stochasticity than MH deletions, and the space of all possible MH-less deletions is 717 orders of magnitude larger than that of MH deletions. The class of MH-less deletions is 718 also less frequent than MH deletions in all five human and mouse cell types we examined. 719 In sum, we characterize MH-less deletions as comprising a large number of rare 720 genotypes that lack high replicate consistency. As such, we conservatively count all 721 binned MH-less deletions for a particular deletion length as a single unique outcome. In sum, we likely underestimate by a moderate amount the true number of unique 722 723 microhomology-less deletion events.

724

As MH-less deletions represents the larger space of possible unique genotypes, in total, we are likely underestimating the total number of unique outcomes in our procedure that calls unique outcomes with high confidence. We provide statistics on the total number of high-confidence unique outcomes in the manuscript and in Extended Data Fig. 1 and 5.

729

730 Based on a computational simulation of subsampling the data, we empirically set 1,000 731 reads per target site as a minimum quality threshold. The diversity of editing outcomes 732 requires some minimum read count to consider the data as representative of editing 733 outcomes at that target site. In Lib-A and Lib-B data in U2OS and mESCs, we empirically 734 observe that 1,000 reads per target site lies above the "elbow" in the curve plotting the 735 number of unique high-confidence outcomes and subsampled read count. We 736 recommend this quality filtering methodology in general for future work studying CRISPR 737 editing outcomes, and based on our data, empirically suggest that 1,000 reads per target 738 site may be a useful guideline for future experimental design.

739

Controlling for cell-type specific 1-bp insertion frequencies when measuring replicability of indel frequencies across cell-types

All indels not belonging to "all major repair outcomes" were filtered out. To adjust the frequencies of all 1-bp insertion genotypes in a target site in two cell-types, the average of the total 1-bp insertion frequency among all major repair outcomes was calculated between the two cell-types, then frequencies of each 1-bp insertion genotype was adjusted proportionally such that the resulting total 1-bp insertion frequency in that is equal to the aforementioned average, and thereby equal to the adjusted 1-bp insertion frequency in the data from the other cell-type.

749

750 Selection of variants from disease databases

Disease variants were selected from the NCBI ClinVar database (downloaded September
 9, 2017)⁹ and the Human Gene Mutation Database (publicly available variant data from
 before 2014.3)¹⁰ for computational screening and subsequent experimental correction.

754

A total of 4,935 unique variants were selected from Clinvar submissions where the functional consequence is described as complete insertions, deletions, or duplications where the reference or alternate allele is of length less than or equal to 30 nucleotides. Variants were included where at least one submitting lab designated the clinical significance as 'pathogenic' or 'likely pathogenic' and no submitting lab had designated
the variant as 'benign' or 'likely benign', including variants will all disease associations.
More complex indels and somatic variants were included. A total of 18,083 unique
insertion variants were selected from HGMD which were between 2 to 30 nucleotides in
length. Variants were included with any disease association with the HGMD classification
of 'DM' or disease-causing mutation.

765

766 SpCas9 gRNAs and their cleavage sites were enumerated for each disease allele. Using 767 a previous version of inDelphi, genotype frequency and indel length distributions were 768 predicted for each tuple of disease variant and unique cleavage site. Among each unique disease, the single best gRNA was identified as the gRNA inducing the highest predicted 769 770 frequency of repair to wildtype genotype, and if this was impossible (due to, for example, 771 a disease allele with 2+ bp deletion), then the single best gRNA was identified as the 772 gRNA inducing the highest predicted frameshift repair rate. 1327 sequence contexts were 773 designed in this manner for Lib-B. An additional 265 sequence contexts were designed 774 by taking the 265 sequence contexts in any disease in decreasing order of predicted wildtype repair rate, starting with Clinvar, stopping at 45% wildtype repair rate, then 775 continuing with HGMD. This yielded 1592 total sequences derived from Clinvar and 776 777 HGMD. 778

780 **Definition of Delta-Positions**

781 Using the MMEJ mechanism, deletion events can be predicted at single-base resolution. 782 For computational convenience, we use the tuple (deletion length, delta-position) to 783 construct a unique identifier for deletion genotypes. A delta-position associated with a 784 deletion length N is an integer between 0 and N inclusive (Extended Data Fig. 2). In a 785 sequence alignment, a delta-position describes the starting position of the deletion gap in 786 the read with respect to the reference sequence relative to the cleavage site. For a 787 deletion length N and a cleavage site at position C such that seq[:C] and seq[C:] yield the 788 expected DSB products where the vector slicing operation vector[index1:index2] is 789 inclusive on the first index and exclusive on the second index (python style), a delta-790 position of 0 corresponds to a deletion gap at seq[C-N+0 : C+0], and generally with a 791 delta-position of D, the deletion gap occurs at seq[C-N+D : C+D]. Microhomologies can 792 be described with multiple delta-positions. To uniquely identify microhomology-based 793 deletion genotypes, the single maximum delta-position in the redundant set is used. 794 Microhomology-less deletion genotypes are associated with only a single delta position 795 and deletion length tuple; we use this as its unique identifier.

796

Another way to define delta-positions can be motivated by the example workflow in the Supplementary Discussion on MH deletions describing how each microhomology is associated with a deletion genotype. In that workflow, the delta-position is the number of bases included on the top strand before "jumping down" to the bottom strand.

801

MH-less medial end-joining products correspond to all MH-less genotypes with deltaposition between 1 and N-1 where N is the deletion length. MH-less unilateral end-joining products correspond to MH-less genotypes with delta-position 0 or N. We note that a deletion genotype with delta position N does not immediately imply that it is a microhomology-less unilateral end-joining product since it may contain microhomology (it's possible that delta-positions N-j, N-j+1, ..., N all correspond to the same MH deletion.)

809 **Definition of Precision Score**

810 For a distribution X, where |X| indicates its cardinality (or length when represented as a 811 vector):

812 $PrecisionScore(X) = 1 - \frac{-\sum_{i=1}^{n} P(x_i) \log(P(x_i))}{\log(|X|)}$

813 This precision score ranges between zero (minimally precise, or highest entropy) to one 814 (maximally precise, or lowest entropy).

815

816 inDelphi Deletion Modeling: Neural network input and output

inDelphi receives as input a sequence context and a cleavage site location, and outputs
 two objects: a frequency distribution on deletion genotypes, and a frequency distribution

- 819 on deletion lengths.
- 820

To model deletions, inDelphi trains two neural networks: MH-NN and MHless-NN. MH-NN receives as input a microhomology that is described by two features: microhomology

823 length and GC fraction in the microhomology. Using these features, MH-NN outputs a

number (psi). MHless-NN receives as input the deletion length. Using this feature,
MHless-NN outputs a number (psi).

826

A phi score is obtained from a psi score using: phi_i = exp(psi_i - 0.25*deletion_length), where 0.25 is a "redundant" hyperparameter that serves to increase training speed by helpful scaling. This relationship between psi and phi is differentiable and encodes the assumption that the frequency of an event exponentially increases with neural network output psi (which empirically appears to reflect MH strength) and exponentially decreases with its minimum necessary resection length (deletion length).

833

The architecture of the MH-NN and MHless-NN networks are *input-dimension* -> 16 -> 16 -> 1 for a total of two hidden layers where all nodes are fully connected. Sigmoidal activations are used in all layers except the output layer. All neural network parameters are initialized with Gaussian noise centered around 0. No regularization or dropout was used.

839

840 inDelphi Deletion Modeling: Making predictions

Given a sequence context and cleavage site, inDelphi enumerates all unique deletion
genotypes as a tuple of its deletion length and its delta-position for deletion lengths from
1 bp to 60 bp. For each microhomology enumerated, an MH-phi score is obtained using
MH-NN. In addition, for each deletion length from 1 bp to 60 bp, an MHindep-phi score is
obtained using MHless-NN.

846

inDelphi combines all MH-phi and MHindep-phi scores for a particular sequence context
into two objects – a frequency distribution on deletion genotypes, and a frequency
distribution on deletion lengths – which are both compared to observations for training.
The model is designed to output two separate objects because both are of biological
interest, and separate but intertwined modeling approaches are useful for generating
both. By learning to generate both objects, inDelphi jointly learns about microhomologybased deletion repair and microhomology-less deletion repair.

854

To generate a frequency distribution on deletion genotypes, inDelphi assigns a score for each microhomology. Score assignment considers the concept of "full" microhomology and treats full and not full MHs differently.

858

859 A microhomology is "full" if the length of the microhomology is equal to its deletion length. 860 The biological significance of full microhomologies is that there is only a single deletion genotype possible for the entire deletion length, while in general, a single deletion length 861 862 is consistent with multiple genotypes. In addition, this single genotype can be generated through not just the MH-dependent MMEJ mechanism but also through MH-less end-863 864 joining, for example as mediated by Lig4. Therefore, we model full microhomologies as receiving contributions from both MH-containing and MH-less mechanisms by scoring full 865 microhomologies as MH-phi[i] + MHindep-phi[j] for deletion length *j* and microhomology 866 867 index *i*.

- 868
- 869 Microhomologies that are not "full" are assigned a score of MH-phi[i] for MH index *i*.

870

871 Scores for all deletion genotypes assigned this way are normalized to sum to 1 to produce 872 a predicted frequency distribution on deletion genotypes.

873

To generate a frequency distribution on deletion lengths, inDelphi assigns a score for each deletion length. Score assignment integrates contributions from both MH-dependent and MH-independent mechanisms via the following procedure: For each deletion length *j*, its score is assigned as MHindep-phi[j] plus the sum of MH-phi for each microhomology with that deletion length. Scores for all deletion lengths are normalized to sum to 1 to produce a frequency distribution.

880

inDelphi trains its parameters using a single sequence context by producing both a
 predicted frequency distribution on deletion genotypes and deletion lengths and
 minimizing the negative of the sum of two values: the mean squared Pearson correlation
 for the deletion genotype frequency distribution at each target site in the training set plus
 the mean squared Pearson correlation for the deletion length frequency distribution at
 each target site in the training set. This represents a multitask learning framework.

887

888 In practice, deletion genotype frequency distributions are formed from observations for 889 deletion lengths 1-60, and deletion length frequency distributions are formed from 890 observations for deletion lengths 1-28. Both neural networks are trained jointly and 891 simultaneously on both tasks. inDelphi is trained with stochastic gradient descent with 892 batched training sets. inDelphi is implemented in Python using the autograd library. We 893 used a batch size of 200, an initial weight scaling factor of 0.10, an initial step size of 0.10, 894 and an exponential decaying factor for the step size of 0.999 per step. We observed 895 performance convergence within about 50 epochs.

896

897 inDelphi Deletion Modeling: Multitask learning improves performance

898 Over the course of developing our model, at an intermediate stage we considered a 899 simpler model for predicting the frequencies of MH deletions. This model featurizes all 900 sequence microhomologies at a target site using MH length and GC content and uses a 901 2x16x16x1 neural network with sigmoidal activations except at the output layer to output 902 psi. This psi value is adjusted using exp(psi – 0.25 * deletion length) to obtain phi for a 903 particular microhomology, which are normalized across all microhomologies to sum to 1 904 to achieve a predicted distribution of frequencies. Altogether, this model is identical to the 905 MH module used in inDelphi, with the notable difference of not including contributions of 906 MH-less phi at "full" microhomologies.

907

This simple model, henceforth known as the baseline model, does not recognize the possibility that a MH genotype may arise from both MH and MH-independent repair pathways. We compared the performance of the baseline model to inDelphi's MH module and observed a statistically significant relative improvement of 10% in model performance as measured on test set data (p ~ 0.02). These measurements were performed using Lib-A target sites in mESCs.

- We note that the multitask model used in inDelphi also jointly trains MHless-NN and, in addition to predicting MH deletion frequencies more accurately than the baseline, also
- 917 provides strong performance on deletion length frequency prediction.
- 918

919 Using random seed A, the baseline model mean Pearson r on held-out data was .905, 920 while the multitask model mean Pearson r on the same held-out data was 0.913, for a 921 8.5% relative improvement (p = 0.009, one-sided *t*-test). Using random seed B, the 922 baseline model mean Pearson r on held-out data was .924, while the multitask model 923 mean Pearson r on the same held-out data was 0.928, for a 5.3% relative improvement 924 (p = 0.02, one-sided t-test). Using random seed C, the baseline model mean Pearson r 925 on held-out data was .912, while the multitask model mean Pearson r on the same held-926 out data was 0.917, for a 5.7% relative improvement (p = 0.03, one-sided *t*-test).

927

928 inDelphi Deletion Modeling: Summary and Revisiting Assumptions

In summary, inDelphi trains MH-NN, which uses as input (microhomology length, microhomology GC content) to output a psi score which is translated into a phi score using deletion length. This phi score represents the "strength" of the microhomology corresponding to a particular MH deletion genotype. It also trains MHless-NN which uses as input (deletion length) to directly output a phi score representing the "total strength" of all MH-independent activity for a particular deletion length.

935

936 While the model assumes that microhomology and microhomology-less repair can 937 overlap in contributions to a single repair genotype, this assumption is made 938 conservatively by assuming that their contributions overlap only when there is no 939 alternative. Specifically, in the context of a deletion length with full microhomology, the 940 model assumes that they must overlap, while in the context of a deletion length without full microhomology, inDelphi allows MHindep-phi to represent all MH-less repair 941 942 genotypes and none of the MH-dependent repair genotypes which are represented solely 943 using their MH-phi scores. This can be seen by noting that at a deletion length j without 944 full microhomology, MH genotypes are scored using their MH-phi scores, while the length j is scored by MHindep-phi[j] plus the sum of MH-phi for each microhomology. Therefore, 945 946 the subset of MH-less genotypes at this deletion length have a score MHindep-phi[j].

947

When the subset of MH-less genotypes includes only one MH-less genotype, this single genotype's score is equal to MHindep-phi[j]. In general, multiple MH-less genotypes are possible, in which case the total score of all of the MH-less genotypes is equal to MHindep-phi[j].

952

The relative frequency of MH deletions and MH-less deletions is learned implicitly by the balancing between the sum of all MH-phi and MHindep-phi. Since MHindep-phi does not vary by sequence context while MH-phi does, the model assumes that variation in the fraction of deletions that use MH can at least partially be explained by varying sequence microhomology as represented by MH-NN.

958

959 inDelphi Insertion Modeling

960 Once inDelphi is trained on both deletion tasks, it predicts insertions from a sequence 961 context and cleavage site by using the precision score of the predicted deletion length distribution and total deletion phi (from all MH-phi and MHindep-phi). inDelphi also uses 962 963 one-hot-encoded binary vectors encoding nucleotides -4 and -3. In a training set, these 964 features are collected and normalized to zero mean and unit variance, and the fraction of 965 1-bp insertions over the sum counts of 1-bp insertions and all deletions are tabulated as 966 the prediction goal. A k-nearest neighbor model is built using the training data. inDelphi 967 uses the default parameter k = 5.

968

969 On test data, the above procedure is used to predict the frequency of 1-bp insertions out 970 of 1-bp insertions and all deletions for a particular sequence context. Once this frequency 971 is predicted, it is used to make frequency predictions for each of the 4 possible insertion 972 genotypes, which are predicted by deriving from the training set the average insertion 973 frequency for each base given its local sequence context. When the training set is small, 974 only the -4 nucleotide is used. When the training set is relatively large, nucleotides -5, -4, 975 and -3 are used.

976

To produce a frequency distribution on 1-bp insertions and 1-60 bp deletion genotypes,
scores for all deletion genotypes and all 1-bp insertions are normalized to sum to 1. To

produce a frequency distribution on indel lengths (+1 to -60), scores for all deletion lengths
 and 1-bp insertions are normalized to sum to 1.

980 981

982 inDelphi: Repair classes predicted at varying resolution

inDelphi predicts MH-deletions and 1-bp insertions at single base resolution. Measuring
 performance on the task of genotype frequency prediction considers this subset of repair
 outcomes only (about 60-70% of all outcomes).

986

inDelphi predicts MH-less deletions to the resolution of deletion length. That is, inDelphi
 predicts a single frequency corresponding to the sum total frequency of all unique MH less deletion genotypes possible for a particular deletion length. This modeling choice
 was made because genotype frequency replicability among MH-less deletions is
 substantially lower than among MH deletions.

992

Measuring performance on the task of indel length frequency considers MH deletions,
 MH-less deletions, and 1-bp insertions (90% of all outcomes).

995

In practice, if end-users desire, they can extend inDelphi predictions to frequency
 predictions for specific MH-less deletion genotypes by noting that MH-less deletions are
 distributed uniformly between 0 delta-position genotypes, medial genotypes, and N delta position genotypes.

1000

1001 **Comparison with a linear baseline model**

We compared inDelphi to a baseline model with the same model structure but replacing the deep neural networks with linear models. We compared using Lib-A mESC data. While inDelphi achieves a mean held-out Pearson correlation of 0.851 on deletion genotype frequency prediction and 0.837 on deletion length frequency prediction, the linear baseline model achieves a mean held-out Pearson correlation of 0.816 on deletion genotype frequency prediction and 0.796 on deletion length frequency prediction. When including the third model component for 1-bp insertion modeling and testing on genotype frequency prediction for 1-bp insertions and all deletions, inDelphi achieves a median held-out Pearson correlation of 0.937 and 0.910 on the task of indel length frequency prediction. The linear baseline model achieves a median held-out Pearson correlation of 0.919 and 0.900 on the two tasks respectively.

1013

From these results, we can see that much of the model's power is derived from its designed structure which is independent of the choice of linear or non-linear modeling. While the baseline does not significantly cripple the model, the use of deep nonlinear neural networks offers a substantial performance improvement (10-24%) above linear modeling. In addition, the strong performance of the linear baseline model highlights that the prediction task, given the model structure, is relatively straightforward. This suggests that our model should be able to generalize well to unseen data.

1021

1022 The deep neural network version of MH-NN learns that microhomology length is more 1023 important than % GC (Extended Data Fig. 2). The linear version learns the same 1024 concept, with a weight of 1.1585 for MH length and 0.332 for % GC.

1025

1026 **Comparison with a baseline model lacking microhomology length as a feature**

1027 Microhomology length is an important feature for MH-NN (Extended Data Fig. 2). We

trained a model that uses only % GC as input to MH-NN while keeping the rest of the

model structure identical. On held-out data at Lib-A target sites in mESCs, this baseline

1030 model at convergence achieves to a mean Pearson correlation of 0.59 on the task of

1031 predicting deletion genotype frequencies, and a mean Pearson correlation of 0.58 on 1032 the task of predicting deletion length frequencies. Notably, a model at iteration 0 with

1032 randomly initialized weights achieves mean Pearson correlations of 0.55 and 0.54 on

1034 the two respective tasks on held-out data. This basal Pearson correlation is relatively

1035 high due to the model structure, in particular, the exponential penalty on deletion length.

1036 In sum, removing MH length as a feature severely impacts model performance,

1037 restricting it to predictive performance not appreciably better than random chance.

1038

1039 inDelphi training and testing on data from varying cell-types

1040 For predicting genotype and indel length frequencies in any particular cell-type C where data D is available, we first trained inDelphi's deletion component on a subset of Lib-A 1041 1042 mESC data. Then, we apply k-fold cross-validation on D where D is iteratively split into 1043 training and test datasets. For each cross-validation iteration, the training set is used to 1044 train the insertion frequency model (k-nearest neighbors) and insertion genotype model (matrix of observed probabilities of each inserted base given local sequence context, 1045 which is just the -4 nucleotide when the training dataset is small, and -5, -4 and -3 1046 nucleotides when the training dataset is large). For each cross-validation iteration, 1047 predictions are made at each sequence context in the test set which are compared to 1048 1049 observations for each sequence context to yield a Pearson correlation. For any particular 1050 sequence context, the median test-time Pearson correlation across all cross-validation 1051 iterations is used as a single number summary of the overall performance of inDelphi. For all reported results, we used 100-fold cross-validation with 80%/20% training and testing
 splits. Empirically, we observed small variance in test-time Pearson correlation,
 highlighting the stability of inDelphi's modeling approach.

1055

1056 inDelphi testing on endogenous VO data

On this task, the deletion component of inDelphi was trained on a subset of the Lib-A 1057 1058 mESC data. For each cell type in HCT116, K562, and HEK293T, all VO sequence contexts (about 100) were randomly split into training and test datasets 100 times. During 1059 each split, the training set was used for *k*-nearest neighbor modeling of 1-bp insertion 1060 1061 frequencies. Feature normalization to zero mean and unit variance was not performed. The average frequency of each 1-bp insertion genotype was derived from the training set 1062 as well. For each of the ~100 sequence contexts, the median test-time Pearson 1063 1064 correlation was used for plotting in Figure 3. Due to the small size of the training set, only the -4 nucleotide was used for modeling both the insertion frequency and insertion 1065 1066 genotype frequencies.

1067

1068 inDelphi testing on library data

1069 On this task, the deletion component of inDelphi was trained on a subset of the Lib-A mESC data. The remaining test set was used for measuring test-time prediction 1070 performance on Lib-A. Nucleotides -5, -4, and -3 were used for the insertion genotype 1071 model. For testing on Lib-B, Lib-B was split into training and test datasets in the same 1072 1073 manner as with VO data. Nucleotide -4 was used for the insertion genotype model. The 1074 median test-time Pearson correlation is used as a single number summary of the overall performance of inDelphi on any particular sequence context. For reporting predictive 1075 1076 results in Figure 4, sequence contexts with low replicability (less than 0.85 Pearson 1077 correlation) in observed editing outcome frequencies were first removed.

1078

1079 inDelphi training and testing on *Prkdc^{-/-}Lig4^{-/-}* data

inDelphi was trained on data from 946 Lib-A sequence contexts and tested on 168 held out Lib-A sequence contexts. Nucleotide -4 was used for insertion rate modeling, all other
 modeling choices were standard as described above. On held-out data, this version of
 inDelphi achieved a median Pearson correlation of 0.84 on predicting indel genotype
 frequencies, and 0.80 on predicting indel length frequencies.

1085

1086 Training the online public version of inDelphi and its expected properties

For general-use on arbitrary cell types, we trained a version of inDelphi using additional 1087 1088 data from diverse types of cells. Deletion modeling was trained using data from 2,464 1089 sequence contexts from high-replicability Lib-A and Lib-B data (including clinical variants and microduplications, fourbp, and longdup) in mES and data from VO sequence contexts 1090 in HEK293 and K562. Insertion frequency modeling is implemented as above. Insertion 1091 1092 genotype modeling uses nucleotides -5, -4, and -3. The insertion frequency model and insertion genotype model are trained on VO endogenous data in K562 and HEK293T, 1093 1094 Lib-A data in mESC, and Lib-B data (including clinical variants and microduplications, 1095 fourbp, and longdup) in mESC and U2OS.

1097 Though MHless-NN, as trained on library data, never receives information on deletion 1098 lengths beyond 28, we allow it to generalize its learned function and make predictions on 1099 deletion lengths up to 60 bp to match the supported range of MH-NN.

1100

inDelphi makes predictions on 1-bp insertions and 1-60-bp deletions, which we 1101 empirically show to consist of higher than 90% of all Cas9 editing outcomes in data from 1102 1103 multiple human and mouse cell lines. Nevertheless, there is a subset of repair (about 8% 1104 on average) that inDelphi does not attempt to predict. We suggest that end-users, 1105 depending on what predictive quantities are of interest, take this into account when using 1106 inDelphi. For example, if inDelphi predicts that 60% of 1-bp insertions and 1-60-bp deletions at a disease allele correspond to repair to wildtype genotype, a quantity of 1107 1108 interest may be the rate of wildtype repair in all Cas9 editing outcomes (including the 8% 1109 not predicted by inDelphi). In such a situation, this quantity can be calculated as 1110 (92%*60%) = 55.2%.

1111

By the design of 1872 sequence contexts in Lib-A, our training dataset has rich and uniform representation across all quintiles of several major axes of variation including GC content, precision, and number of bases participating in microhomology as measured empirically in the human genome. This design strategy enables inDelphi to generalize well to arbitrary sequence contexts from the human genome.

1117

1118 These training data further include data in the outlier range of statistics of interest, including extremely high and low precision repair distributions, and extremely weak and 1119 1120 strong microhomology (minimal microhomology and extensive microduplication 1121 microhomology sequences). The availability of such sequences in our training data enables inDelphi to generalize well to sequence contexts of clinical interest and sequence 1122 contexts supporting unusually high frequencies of precision repair. In particular, by 1123 1124 training on more than 1000 examples of repair at clinical microduplications, inDelphi has 1125 received strong preparation for accurate prediction on other clinical microduplications. 1126

- 1127 By training on data from many cell-types, we enable inDelphi to make predictions that are generally applicable to many human cell-types. We note that the HCT116 human colon 1128 1129 cancer cell line experiences a markedly higher frequency of single base insertions 1130 compared to all other cell lines we studied, possibly due to the MLH1 deficiency of this 1131 cell line leading to impaired DNA mismatch repair. For this reason, we excluded HCT116 data from our training dataset. For best results, we suggest end-users keep in mind that 1132 repair class frequencies can be cell type-dependent, and this issue has not been well-1133 1134 characterized thus far.
- 1135

We note that inDelphi's main error tendency is on the side of overestimating rather than underestimating the precision of repair (Figure 4). In general, this tendency can be explained by noting that inDelphi only considers sequence microhomology as a factor, while it's plausible and likely in biological experimental settings that even sequence contexts with very strong sequence microhomology may not yield precise results due to noise factors that are not considered by inDelphi. For best results, we recommend endusers take this tendency into account when using inDelphi predictions for further experiments. In particular, if gRNAs are designed by using a minimum precision threshold, end-users should recognize that observed repair outcomes may have empirical precision under this threshold. However, conversely, it is unlikely that a gRNA will have precision higher than what inDelphi predicts.

1147

1148 Lib-A design

All designed sequence contexts were 55 bp in length with cutting between the 27th and 28th base.

1151

1152 1872 sequence contexts were designed by empirically determining the distribution of four statistics in sequence contexts from the human genome. These four statistics are GC 1153 1154 content, total sum of bases participating in microhomology for 3-27-bp deletions, Azimuth 1155 predicted on-target efficiency score, and the statistical entropy of the predicted 3-27-bp 1156 deletion length distribution from a previous version of inDelphi. For each of these 1157 statistics, empirical quintiles were derived by calculating these statistics in a large number 1158 of sequence contexts from the human genome. For the library, sequence contexts were designed by randomly generated DNA that categorized into each combination of quintiles 1159 across each of the four statistics. For example, a sequence context falling into the 1st 1160 quintile in GC, 2nd quintile of total MH, 1st quintile of Azimuth score, and 5th quintile of 1161 entropy, was found by random search. With four statistics and five bins each (due to 1162 quintiles), there are $5^4 = 625$ possible combinations. For each combination, we attempted 1163 1164 to design three sequence contexts for a total of 1875; 3 sequences could not be designed 1165 (for a total of 1872) though each bin was filled. 90 sequence contexts were designed from VO sequence contexts. Other sequence contexts were also designed for a total of 2000 1166 1167 sequence contexts in Lib-A. Lib-A sequence names, gRNAs, and sequence contexts are 1168 listed in Supplementary Table 2.

1169

1170 Lib-B design

All designed sequence contexts were 55 bp in length with cutting between the 27th and 28th base.

1173

1174 1592 sequence contexts were designed from Clinvar and HGMD (see section on Selection of variants from disease databases). Some disease sequence contexts were 1175 designed that such that the corrected wildtype or frameshift allele supports further cutting 1176 1177 by the original gRNA; data from such sequence contexts were ignored during analysis. 57 "longdup" sequence contexts were designed by repeating the following procedure 1178 three times: for N = 7 to 25, an N-mer was randomly generated, then duplicating and 1179 1180 surrounded by randomly generated sequences, while ensuring that SpCas9 NGG was included and appropriately positioned for cutting between positions 27 and 28. 90 1181 sequence contexts were designed from VO sequence contexts. 228 "fourbp" sequence 1182 contexts were designed at 3 contexts with random sequences (with total phi score on 1183 average lower than VO sequence contexts) while varying positions -5 to -2; for each of 1184 the 3 "low-microhomology" contexts,76 four bases were randomly designed while 1185 ensuring representation from all possible 2 bp microhomology patterns including no 1186 microhomology, one base of microhomology at either position, and full two bases of 1187 microhomology. Other sequence contexts were also designed for a total of 2000 1188

sequence contexts in Lib-B. Lib-B sequence names, gRNAs, and sequence contexts arelisted in Supplementary Table 3.

1191

1192 **1bpInsDisLib design**

1193 12 sequence contexts were designed from Clinvar and HGMD. Pathogenic alleles were

selected for a high predicted frequency of correction to the wild-type genotype via a Cas9mediated 1-bp insertion. Sequence names, gRNAs, and sequence contexts are listed in

- 1196 Supplementary Table 4.
- 1197

1198 PHG design

1199 18 sequence contexts were designed using inDelphi to select SpCas9 gRNAs targeting 1200 the coding regions of genes including VEGFA, VEGFR2, PDCD1, APOB, CCR5, CD274, 1201 CXCR4, PCSK9, and APOBEC3B, such that a frameshift would be induced with higher 1202 frequency than typical SpCas9 gRNAs. Of these 18 frameshift designs, 10 were designed 1203 to induce a single deletion genotype with high precision, and 8 were designed to induce 1204 a single 1-bp insertion genotype with high precision. 6 sequence contexts were designed using inDelphi from Clinvar and HGMD where pathogenic 1-bp insertion alleles were 1205 selected based on a high predicted frequency of induction from Cas9 editing of the wild-1206 type allele. Sequence names, gRNAs, and sequence contexts are listed in 1207 1208 Supplementary Table 5.

1209

1210 Generating a DNA motif for 1-bp insertion frequencies

1211 Nucleotides from positions -7 to 0 were one-hot-encoded and used in ridge regression to predict the observed frequency of 1-bp insertions out of all Cas9 editing events in 1996 1212 1213 sequence contexts from Lib-A mESC data. The data were split into training and testing sets (80/20 split) 10,000 times to calculate a bootstrapped estimate of linear regression 1214 weights and test-set predictive Pearson correlation. The median test-set Pearson 1215 1216 correlation was found to be 0.62. To generate a DNA motif, any features that included 0 1217 within the bootstrapped weight range were excluded (probability that the weight is zero > 1218 1e-4). The average bootstrapped weight estimate was used as the "logo height" for all 1219 remaining features. Each feature is independent; vertical stacking of features follows the 1220 published tradition of DNA motifs.

1221

1222 Predicting precision repair of genomic SpCas9 gRNAs

1223 In this work, we determined the distributions of the most frequent deletion and

insertion outcomes among major editing outcomes at SpCas9 gRNAs targeting human 1224 exons and introns as predicted by inDelphi trained on data from Lib-A target sites in 1225 1226 mESCs and U2OS cells separately (Fig. 3f, Extended Data Table 1). A combination of computational constraint (the inability to make predictions at ~350 million target sites 1227 comprising all SpCas9 gRNAs in the human genome), uncertainty in the exact predictions 1228 of the model and a preference for avoiding overfitting our training data, and lack of 1229 sufficient held-out data to verify our predictions and identify potential bias, motivated us 1230 to smooth the exact predictions made by the model. We resampled each predicted value 1231 from a Gaussian centered at the predicted value with a specified standard deviation. For 1232 mESCs, we set the standard deviation as the predicted value divided by 4, up to a 1233 1234 maximum of 3% for insertions, while for deletions we used the predicted value divided by 4 with a minimum of 6%. For U2OS cells, we set the standard deviation as the predicted value divided by 4 for insertions, and the predicted value divided by 4 with a minimum of 6% for deletions. The scaling of standard deviation at higher predicted values reflects the abundance of data and therefore higher relative confidence at lower predicted values. The use of symmetrical noise reflects our prior belief that our predictions are equally likely to underestimate and overestimate the true value.

1242	
1243	
1244	Plasmid and insert sequences
1245	
1246	P2T-CAG-MCS-P2A-GFP-PuroR complete plasmid sequence
1247	CCACCTAAATTGTAAGCGTTAATATTTTGTTAAAATTCGCGTTAAATTTTTGTTAAAT
1248	CAGCTCATTTTTTAACCAATAGGCCGAAATCGGCAAAATCCCTTATAAATCAAAAGA
1249	ATAGACCGAGATAGGGTTGAGTGTTGTTCCAGTTTGGAACAAGAGTCCACTATTAA
1250	AGAACGTGGACTCCAACGTCAAAGGGCGAAAAACCGTCTATCAGGGCGATGGCCC
1251	ACTACGTGAACCATCACCCTAATCAAGTTTTTTGGGGTCGAGGTGCCGTAAAGCAC
1252	TAAATCGGAACCCTAAAGGGAGCCCCCGATTTAGAGCTTGACGGGGAAAGCCGGC
1253	GAACGTGGCGAGAAAGGAAGGGAAGAAAGCGAAAGGAGCGGGCGCTAGGGCGC
1254	TGGCAAGTGTAGCGGTCACGCTGCGCGTAACCACCACCCGCCGCGCGTTAATGC
1255	GCCGCTACAGGGCGCGTCCCATTCGCCATTCAGGCTGCGCAACTGTTGGGAAGG
1256	GCGATCGGTGCGGGCCTCTTCGCTATTACGCCAGCTGGCGAAAGGGGGGATGTGC
1257	TGCAAGGCGATTAAGTTGGGTAACGCCAGGGTTTTCCCAGTCACGACGTTGTAAAA
1258	CGACGGCCAGTGAGCGCGCGTAATACGACTCACTATAGGGCGAATTGGGTACCG
1259	GCATATGGTTCTTGACAGAGGTGTAAAAAGTACTCAAAAATTTTACTCAAGTGAAAG
1260	TACAAGTACTTAGGGAAAATTTTACTCAATTAAAAGTAAAAGTATCTGGCTAGAATC
1261	TTACTTGAGTAAAAGTAAAAAGTACTCCATTAAAATTGTACTTGAGTATTAAGGAA
1262	GTAAAAGTAAAAGCAAGAAAGATCGATCTCGAAGGATCTGGAGGCCACCATGGTG
1263	TCGATAACTTCGTATAGCATACATTATACGAAGTTATCGTGCTCGACATTGATTATT
1264	GACTAGTTATTAATAGTAATCAATTACGGGGTCATTAGTTCATAGCCCATATATGGA
1265	GTTCCGCGTTACATAACTTACGGTAAATGGCCCGCCTGGCTGACCGCCCAACGAC
1266	CCCCGCCCATTGACGTCAATAATGACGTATGTTCCCATAGTAACGCCAATAGGGAC
1267	TTTCCATTGACGTCAATGGGTGGAGTATTTACGGTAAACTGCCCACTTGGCAGTAC
1268	ATCAAGTGTATCATATGCCAAGTACGCCCCCTATTGACGTCAATGACGGTAAATGG
1269	CCCGCCTGGCATTATGCCCAGTACATGACCTTATGGGACTTTCCTACTTGGCAGTA
1270	CATCTACGTATTAGTCATCGCTATTACCATGGTCGAGGTGAGCCCCACGTTCTGCT
1271	TCACTCTCCCCATCTCCCCCCCCCCCCCCCCCCCCCCCC
1272	ATTATTTTGTGCAGCGATGGGGGGGGGGGGGGGGGGGGG
1273	GGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
1274	CAGCCAATCAGAGCGGCGCGCGCCCGAAAGTTTCCTTTTATGGCGAGGCGGCGGC
1275	GGCGGCGGCCCTATAAAAAGCGAAGCGCGCGGGGGGGGGG
1276	GCTGCCTTCGCCCCGTGCCCGCTCCGCCGCCGCCGCCGCC
1277	CTCTGACTGACCGCGTTACTCCCACAGGTGAGCGGGCGGG
1278	CGGGCTGTAATTAGCGCTTGGTTTAATGACGGCTTGTTTCTTTC
1279	GAAAGCCTTGAGGGGCTCCGGGAGGGCCCTTTGTGCGGGGGGGG
1280	GGGTGCGTGCGTGTGTGTGCGTGGGGGAGCGCCGCGTGCGGCTCCGCGCTGC
1281	CCGGCGGCTGTGAGCGCTGCGGGGCGCGCGGGGGCTTTGTGCGCTCCGCAGT
1282	GTGCGCGAGGGGAGCGCGGCCGGGGGGGGGGGGGGGGGG

1283 1284 GCTGAGCACGGCCCGGCTTCGGGTGCGGGGCTCCGTACGGGGCGTGGCGCGGG 1285 1286 1287 1288 GCGCCGGCGGCTGTCGAGGCGCGCGGCGAGCCGCAGCCATTGCCTTTTATGGTAAT 1289 CGTGCGAGAGGGCGCAGGGACTTCCTTTGTCCCAAATCTGTGCGGAGCCGAAATC 1290 TGGGAGGCGCCGCCGCACCCCCTCTAGCGGGCGCGGGGGCGAAGCGGTGCGGCG CCGGCAGGAAGGAAATGGGCGGGGGGGGGGCCTTCGTGCGTCGCCGCGCCGCCGT 1291 1292 1293 GGGGGACGGGGCAGGGCGGGGTTCGGCTTCTGGCGTGTGACCGGCGGCTCTAG AGCCTCTGCTAACCATGTTCATGCCTTCTTCTTTTTCCTACAGCTCCTGGGCAACGT 1294 1295 GCTGGTTATTGTGCTGTCTCATCATTTTGGCAAAGAATTCCTCGAGCGGCCGCCAG 1296 TGTGATGGATATCGGATCCGCTAGCGCTACTAACTTCAGCCTGCTGAAGCAGGCT GGAGACGTGGAGGAGAACCCTGGACCTGGACCGGTCGCCACCATGGTGAGCAAG 1297 GGCGAGGAGCTGTTCACCGGGGTGGTGCCCATCCTGGTCGAGCTGGACGGCGAC 1298 GTAAACGGCCACAAGTTCAGCGTGTCCGGCGAGGGCGAGGGCGATGCCACCTAC 1299 GGCAAGCTGACCCTGAAGTTCATCTGCACCACCGGCAAGCTGCCCGTGCCCTGG 1300 1301 CCCACCCTCGTGACCACCCTGACCTACGGCGTGCAGTGCTTCAGCCGCTACCCCG ACCACATGAAGCAGCACGACTTCTTCAAGTCCGCCATGCCCGAAGGCTACGTCCA 1302 GGAGCGCACCATCTTCTTCAAGGACGACGGCAACTACAAGACCCGCGCGAGGT 1303 1304 GAAGTTCGAGGGCGACACCCTGGTGAACCGCATCGAGCTGAAGGGCATCGACTT 1305 CAAGGAGGACGGCAACATCCTGGGGCACAAGCTGGAGTACAACTACAACAGCCAC AACGTCTATATCATGGCCGACAAGCAGAAGAACGGCATCAAGGTGAACTTCAAGAT 1306 CCGCCACAACATCGAGGACGGCAGCGTGCAGCTCGCCGACCACTACCAGCAGAA 1307 CACCCCCATCGGCGACGGCCCCGTGCTGCTGCCCGACAACCACTACCTGAGCAC 1308 1309 CCAGTCCGCCCTGAGCAAAGACCCCCAACGAGAAGCGCGATCACATGGTCCTGCTG GAGTTCGTGACCGCCGCGGGATCACTCTCGGCATGGACGAGCTGTACAAGTAAA 1310 GCGGCCGCCACCGCGGTGGAGCTCGAATTAATTCATCGATGATGATCCAGACATG 1311 ATAAGATACATTGATGAGTTTGGACAAACCACAACTAGAATGCAGTGAAAAAAATG 1312 1313 CTTTATTTGTGAAATTTGTGATGCTATTGCTTTATTTGTAACCATTATAAGCTGCAAT 1314 TGGGAGGTTTTTTAAAGCAAGTAAAACCTCTACAAATGTGGTATGGCTGATTATGAT 1315 1316 1317 1318 GGGTCGTGGGGCGGGGCGTCAGGCACCGGGCTTGCGGGTCATGCACCAGGTGCG CGGTCCTTCGGGCACCTCGACGTCGGCGGTGACGGTGAAGCCGAGCCGCTCGTA 1319 GAAGGGGAGGTTGCGGGGGCGCGGAGGTCTCCAGGAAGGCGGGCACCCCGGCGC 1320 1321 GCTCGGCCGCCTCCACTCCGGGGGAGCACGACGGCGCTGCCCAGACCCTTGCCCT 1322 GGTGGTCGGGCGAGACGCCGACGGTGGCCAGGAACCACGCGGGCTCCTTGGGC

CGGTGCGGCGCCAGGAGGCCTTCCATCTGTTGCTGCGCGGCCAGCCGGGAACCG 1323 1324 CTCAACTCGGCCATGCGCGGGCCGATCTCGGCGAACACCGCCCCGCTTCGACG CTCTCCGGCGTGGTCCAGACCGCCACCGCGGCGCCGTCGTCCGCGACCCACACC 1325 1326 TTGCCGATGTCGAGCCCGACGCGCGTGAGGAAGAGTTCTTGCAGCTCGGTGACC 1327 1328 1329 GGCGCACCGTGGGCTTGTACTCGGTCATGGAAGGTCGTCTCCTTGTGAGGGGTCA 1330 GGGGCGTGGGTCAGGGGGATGGTGGCGGCACCGGTCGTGGCGGCCGACCTGCAG GCATGCAAGCTTTTTGCAAAAGCCTAGGCCTCCAAAAAAGCCTCCTCACTACTTCT 1331 1332 TCAGCCATGGGGCGGAGAATGGGCGGAACTGGGCGGAGTTAGGGGCGGGATGG 1333 1334 1335 ACTTCTGCCTGCTGGGGAGCCTGGGGACTTTCCACACCTGGTTGCTGACTAATTG 1336 AGATGCATGCTTTGCATACTTCTGCCTGCTGGGGAGCCTGGGGACTTTCCACACC CTAACTGACACACATTCCACAGAATTCAAGTGATCTCCAAAAAATAAGTACTTTTG 1337 ACTGTAAATAAAATTGTAAGGAGTAAAAAGTACTTTTTTTCTAAAAAAATGTAATTA 1338 AGTAAAAGTAAAAGTATTGATTTTTAATTGTACTCAAGTAAAGTAAAAATCCCCAAAA 1339 ATAATACTTAAGTACAGTAATCAAGTAAAATTACTCAAGTACTTTACACCTCTGGTTC 1340 1341 TTGACCCCCTACCTTCAGCAAGCCCAGCAGATCCGAGCTCCAGCTTTTGTTCCCTT TAGTGAGGGTTAATTGCGCGCGTTGGCGTAATCATGGTCATAGCTGTTTCCTGTGTG 1342 AAATTGTTATCCGCTCACAATTCCACACAACATACGAGCCGGAAGCATAAAGTGTA 1343 AAGCCTGGGGTGCCTAATGAGTGAGCTAACTCACATTAATTGCGTTGCGCTCACTG 1344 1345 CCCGCTTTCCAGTCGGGAAACCTGTCGTGCCAGCTGCATTAATGAATCGGCCAAC GCGCGGGGAGAGGCGGTTTGCGTATTGGGCGCTCTTCCGCTTCCTCGCTCACTGA 1346 1347 GGTAATACGGTTATCCACAGAATCAGGGGGATAACGCAGGAAAGAACATGTGAGCA 1348 1349 AAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGTTGCTGGCGTTTTTCC ATAGGCTCCGCCCCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTG 1350 GCGAAACCCGACAGGACTATAAAGATACCAGGCGTTTCCCCCTGGAAGCTCCCTC 1351 GTGCGCTCTCCTGTTCCGACCCTGCCGCTTACCGGATACCTGTCCGCCTTTCTCC 1352 1353 CTTCGGGAAGCGTGGCGCTTTCTCATAGCTCACGCTGTAGGTATCTCAGTTCGGT GTAGGTCGTTCGCTCCAAGCTGGGCTGTGTGCACGAACCCCCCGTTCAGCCCGAC 1354 CGCTGCGCCTTATCCGGTAACTATCGTCTTGAGTCCAACCCGGTAAGACACGACTT 1355 ATCGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGC 1356 GGTGCTACAGAGTTCTTGAAGTGGTGGCCTAACTACGGCTACACTAGAAGGACAG 1357 1358 TATTTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGGAAAAAGAGTTGGTAGC 1359 GCAGATTACGCGCAGAAAAAAGGATCTCAAGAAGATCCTTTGATCTTTCTACGG 1360 1361 GGTCTGACGCTCAGTGGAACGAAAACTCACGTTAAGGGATTTTGGTCATGAGATTA 1362

1363 AAAGTATATATGAGTAAACTTGGTCTGACAGTTACCAATGCTTAATCAGTGAGGCAC 1364 CTATCTCAGCGATCTGTCTATTTCGTTCATCCATAGTTGCCTGACTCCCCGTCGTGT AGATAACTACGATACGGGAGGGCTTACCATCTGGCCCCAGTGCTGCAATGATACC 1365 1366 1367 ATTGTTGCCGGGAAGCTAGAGTAAGTAGTTCGCCAGTTAATAGTTTGCGCAACGTT 1368 GTTGCCATTGCTACAGGCATCGTGGTGTCACGCTCGTCGTTTGGTATGGCTTCATT 1369 1370 CAGCTCCGGTTCCCAACGATCAAGGCGAGTTACATGATCCCCCATGTTGTGCAAAA 1371 1372 GTTATCACTCATGGTTATGGCAGCACTGCATAATTCTCTTACTGTCATGCCATCCGT AAGATGCTTTTCTGTGACTGGTGAGTACTCAACCAAGTCATTCTGAGAATAGTGTAT 1373 GCGGCGACCGAGTTGCTCTTGCCCGGCGTCAATACGGGATAATACCGCGCCACAT 1374 1375 AGCAGAACTTTAAAAGTGCTCATCATTGGAAAACGTTCTTCGGGGCGAAAACTCTC 1376 AAGGATCTTACCGCTGTTGAGATCCAGTTCGATGTAACCCACTCGTGCACCCAACT GATCTTCAGCATCTTTTACTTTCACCAGCGTTTCTGGGTGAGCAAAAACAGGAAGG 1377 CAAAATGCCGCAAAAAAGGGAATAAGGGCGACACGGAAATGTTGAATACTCATACT 1378 CTTCCTTTTTCAATATTATTGAAGCATTTATCAGGGTTATTGTCTCATGAGCGGATAC 1379 1380 ATATTTGAATGTATTTAGAAAAATAAACAAATAGGGGTTCCGCGCACATTTCCCCGA AAAGTG 1381 1382

1384 LDLRwt

1385 ATGGGGCCCTGGGGCTGGAAATTGCGCTGGACCGTCGCCTTGCTCCTCGCCGCG GCGGGGACTGCAGTGGGCGACAGATGCGAAAGAAACGAGTTCCAGTGCCAAGAC 1386 GGGAAATGCATCTCCTACAAGTGGGTCTGCGATGGCAGCGCTGAGTGCCAGGATG 1387 1388 GCTCTGATGAGTCCCAGGAGACGTGCTTGTCTGTCACCTGCAAATCCGGGGACTT 1389 CAGCTGTGGGGGGCCGTGTCAACCGCTGCATTCCTCAGTTCTGGAGGTGCGATGGC CAAGTGGACTGCGACAACGGCTCAGACGAGCAAGGCTGTCCCCCCAAGACGTGC 1390 1391 TCCCAGGACGAGTTTCGCTGCCACGATGGGAAGTGCATCTCTCGGCAGTTCGTCT GTGACTCAGACCGGGACTGCTTGGACGGCTCAGACGAGGCCTCCTGCCCGGTGC 1392 1393 TCACCTGTGGTCCCGCCAGCTTCCAGTGCAACAGCTCCACCTGCATCCCCCAGCT GTGGGCCTGCGACAACGACCCCGACTGCGAAGATGGCTCGGATGAGTGGCCGCA 1394 1395 GCGCTGTAGGGGTCTTTACGTGTTCCAAGGGGACAGTAGCCCCTGCTCGGCCTTC 1396 GAGTTCCACTGCCTAAGTGGCGAGTGCATCCACTCCAGCTGGCGCTGTGATGGTG GCCCCGACTGCAAGGACAAATCTGACGAGGAAAACTGCGCTGTGGCCACCTGTCG 1397 CCCTGACGAATTCCAGTGCTCTGATGGAAACTGCATCCATGGCAGCCGGCAGTGT 1398 GACCGGGAATATGACTGCAAGGACATGAGCGATGAAGTTGGCTGCGTTAATGTGA 1399 1400 CACTCTGCGAGGGACCCAACAAGTTCAAGTGTCACAGCGGCGAATGCATCACCCT GGACAAAGTCTGCAACATGGCTAGAGACTGCCGGGACTGGTCAGATGAACCCATC 1401 1402 AAAGAGTGCGGGACCAACGAATGCTTGGACAACAACGGCGGCTGTTCCCACGTCT 1403 GCAATGACCTTAAGATCGGCTACGAGTGCCTGTGCCCCGACGGCTTCCAGCTGGT 1404 GGCCCAGCGAAGATGCGAAGATATCGATGAGTGTCAGGATCCCGACACCTGCAGC 1405 CAGCTCTGCGTGAACCTGGAGGGTGGCTACAAGTGCCAGTGTGAGGAAGGCTTC CAGCTGGACCCCCACACGAAGGCCTGCAAGGCTGTGGGCTCCATCGCCTACCTCT 1406 1407 TCTTCACCAACCGGCACGAGGTCAGGAAGATGACGCTGGACCGGAGCGAGTACA 1408 CCAGCCTCATCCCCAACCTGAGGAACGTGGTCGCTCTGGACACGGAGGTGGCCA 1409 GCAATAGAATCTACTGGTCTGACCTGTCCCAGAGAATGATCTGCAGCACCCAGCTT GACAGAGCCCACGGCGTCTCTTCCTATGACACCGTCATCAGCAGAGACATCCAGG 1410 1411 CCCCCGACGGGCTGGCTGTGGACTGGATCCACAGCAACATCTACTGGACCGACTC 1412 TGTCCTGGGCACTGTCTCTGTTGCGGATACCAAGGGCGTGAAGAGGAAAACGTTA TTCAGGGAGAACGGCTCCAAGCCAAGGGCCATCGTGGTGGATCCTGTTCATGGCT 1413 TCATGTACTGGACTGACTGGGGAACTCCCGCCAAGATCAAGAAAGGGGGGCCTGAA 1414 1415 TGGTGTGGACATCTACTCGCTGGTGACTGAAAACATTCAGTGGCCCAATGGCATCA CCCTAGATCTCCTCAGTGGCCGCCTCTACTGGGTTGACTCCAAACTTCACTCCATC 1416 1417 TCAAGCATCGATGTCAATGGGGGGCAACCGGAAGACCATCTTGGAGGATGAAAAGA 1418 GGCTGGCCCACCCCTTCTCCTTGGCCGTCTTTGAGGACAAAGTATTTTGGACAGAT 1419 ATCATCAACGAAGCCATTTTCAGTGCCAACCGCCTCACAGGTTCCGATGTCAACTT GTTGGCTGAAAACCTACTGTCCCCAGAGGATATGGTCCTCTTCCACAACCTCACCC 1420 1421 AGCCAAGAGGAGTGAACTGGTGTGAGAGGACCACCCTGAGCAATGGCGGCTGCC AGTATCTGTGCCTCCCTGCCCGCAGATCAACCCCCACTCGCCCAAGTTTACCTG 1422 1423 CGCCTGCCCGGACGGCATGCTGCTGGCCAGGGACATGAGGAGCTGCCTCACAGA 1424 GGCTGAGGCTGCAGTGGCCACCCAGGAGACATCCACCGTCAGGCTAAAGGTCAG

1425	CTCCACAGCCGTAAGGACACAGCACAACCACCCGGCCTGTTCCCGACACCTCC
1426	CGGCTGCCTGGGGCCACCCCTGGGCTCACCACGGTGGAGATAGTGACAATGTCT
1427	CACCAAGCTCTGGGCGACGTTGCTGGCAGAGGAAATGAGAAGAAGCCCAGTAGC
1428	GTGAGGGCTCTGTCCATTGTCCTCCCCATCGTGCTCCTCGTCTTCCTTTGCCTGGG
1429	GGTCTTCCTTCTATGGAAGAACTGGCGGCTTAAGAACATCAACAGCATCAACTTTG
1430	ACAACCCCGTCTATCAGAAGACCACAGAGGATGAGGTCCACATTTGCCACAACCA
1431	GGACGGCTACAGCTACCCCTCGAGACAGATGGTCAGTCTGGAGGATGACGTGGC
1432	G
1433	
1434	LDLRDup252 with surrounding region
1435	CCCCCAAGACGTGCTCCCAGGACGAGTTTCGCTGCCACGATGGGAAGTGCATCTC
1436	TCGGCAGTTCGTCTGTGACTCAGACCGGGACTGCTTGGACGGCTCAGACGAGGC
1437	CTCCTGCCCGGTGCTCACCTGTGGTCCCGCCAGCTTCCAGTGCAACAGCTCCACC
1438	TGCATCCCCCAGCTGTGGGCCTGCGACAACGACCCCGACTGCGAAGATGGCTCG
1439	GAGGCTCGGATGAGTGGCCGCAGCGCTGTAGGGGTCTTTACGTGTTCCAAGGGG
1440	ACAGTAGCCCCTGCTCGGCCTTCGAGTTCCACTGCCTAAGTGGCGAGTGCATCCA
1441	CTCCAGCTGGCGCTGTGATGGTGGCCCCGACTGCAAGGACAAATCTGACGAGGA
1442	AAACTGCG
1443	
1444	LDLRDup254/255 with surrounding region
1445	CCCCCAAGACGTGCTCCCAGGACGAGTTTCGCTGCCACGATGGGAAGTGCATCTC
1446	TCGGCAGTTCGTCTGTGACTCAGACCGGGACTGCTTGGACGGCTCAGACGAGGC
1447	CTCCTGCCCGGTGCTCACCTGTGGTCCCGCCAGCTTCCAGTGCAACAGCTCCACC
1448	TGCATCCCCCAGCTGTGGGCCTGCGACAACGACCCCGACTGCGAAGATGGCTCG
1449	GATGAGTGGCCGCAGCGCTGTAGGGGTCTTTACGTGTTCCAAGGGGACAGTAGC
1450	CCCTGCTCGGCCTTCGAGTTCCACTGCCTAAGTGGCGAGTGCATCCACTCCAGCT
1451	GGCGCTGTGATGGTGGCCCCGACTGCAAGGACAAATCTGACAGGACAAATCTGAC
1452	GAGGAAAACTGCGCTGTGGCCACCTGTCGCCCTGACGAATTCCAGTGCTCTGATG
1453	GAAACTGCATCCATG
1454	
1455	LDLRDup258 with surrounding region
1456	CCCCCAAGACGTGCTCCCAGGACGAGTTTCGCTGCCACGATGGGAAGTGCATCTC
1457	TCGGCAGTTCGTCTGTGACTCAGACCGGGACTGCTTGGACGGCTCAGACGAGGC
1458	CTCCTGCCCGGTGCTCACCTGTGGTCCCGCCAGCTTCCAGTGCAACAGCTCCACC
1459	TGCATCCCCCAGCTGTGGGCCTGCGACAACGACCCCGACTGCGAAGATGGCTCG
1460	GATGAGTGGCCGCAGCGCTGTAGGGGTCTTTACGTGTTCCAAGGGGACAGTAGC
1461	CCCTGCTCGGCCTTCGAGTTCCACTGCCTAAGTGGCGAGTGCATCCACTCCAGCT
1462	GGCGCTGTGATGGTGGCCCCGACTGCAAGGACAAATCTGAGGACAAATCTGACGA
1463	GGAAAACTGCGCTGTGGCCACCTGTCGCCCTGACGAATTCCAGTGCTCTGATGGA
1464	AACTGCATCCATG

1465

1466 LDLRDup261 with surrounding region

CCCCCAAGACGTGCTCCCAGGACGAGTTTCGCTGCCACGATGGGAAGTGCATCTC 1467 TCGGCAGTTCGTCTGTGACTCAGACCGGGACTGCTTGGACGGCTCAGACGAGGC 1468 CTCCTGCCCGGTGCTCACCTGTGGTCCCGCCAGCTTCCAGTGCAACAGCTCCACC 1469 TGCATCCCCCAGCTGTGGGCCTGCGACAACGACCCCGACTGCGAAGATGGCTCG 1470 GATGAGTGGCCGCAGCGCTGTAGGGGTCTTTACGTGTTCCAAGGGGACAGTAGC 1471 CCCTGCTCGGCCTTCGAGTTCCACTGCCTAAGTGGCGAGTGCATCCACTCCAGCT 1472 GGCGCTGTGATGGTGGCCCCGACTGCAAGGACAAATCTGACGACAAATCTGACGA 1473 GGAAAACTGCGCTGTGGCCACCTGTCGCCCTGACGAATTCCAGTGCTCTGATGGA 1474 1475 AACTGCATCCATG 1476 1477 LDLRDup264 with surrounding region CTTCATGTACTGGACTGACTGGGGGAACTCCCGCCAAGATCAAGAAAGGGGGGCCTG 1478 AATGGTGTGGACATCTACTCGCTGGTGAGCTGGTGACTGAAAACATTCAGTGGCC 1479 CAATGGCATCACCCTAG 1480

1481

1483 **GAAwt**

1484 ATGGGAGTGAGGCACCCGCCCTGCTCCCACCGGCTCCTGGCCGTCTGCGCCCTC GTGTCCTTGGCAACCGCTGCACTCCTGGGGGCACATCCTACTCCATGATTTCCTGCT 1485 1486 GGTTCCCCGAGAGCTGAGTGGCTCCTCCCCAGTCCTGGAGGAGACTCACCCAGCT 1487 CACCAGCAGGGAGCCAGCAGACCAGGGCCCCGGGATGCCCAGGCACACCCCGG CCGTCCCAGAGCAGTGCCCACACAGTGCGACGTCCCCCCCAACAGCCGCTTCGA 1488 1489 TTGCGCCCCTGACAAGGCCATCACCCAGGAACAGTGCGAGGCCCGCGGCTGTTG 1490 CTACATCCCTGCAAAGCAGGGGCTGCAGGGGGGCCCAGATGGGGGCAGCCCTGGTG CTTCTTCCCACCCAGCTACCCCAGCTACAAGCTGGAGAACCTGAGCTCCTCTGAAA 1491 1492 TGGGCTACACGGCCACCCTGACCCGTACCACCCCCACCTTCTTCCCCAAGGACAT 1493 CCTGACCCTGCGGCTGGACGTGATGATGGAGACTGAGAACCGCCTCCACTTCACG ATCAAAGATCCAGCTAACAGGCGCTACGAGGTGCCCTTGGAGACCCCGCATGTCC 1494 1495 ACAGCCGGGCACCGTCCCCACTCTACAGCGTGGAGTTCTCCGAGGAGCCCTTCG 1496 GGGTGATCGTGCGCCGGCAGCTGGACGGCCGCGTGCTGCTGAACACGACGGTG 1497 GCGCCCCTGTTCTTTGCGGACCAGTTCCTTCAGCTGTCCACCTCGCTGCCCTCGC AGTATATCACAGGCCTCGCCGAGCACCTCAGTCCCCTGATGCTCAGCACCAGCTG 1498 GACCAGGATCACCCTGTGGAACCGGGACCTTGCGCCCACGCCCGGTGCGAACCT 1499 CTACGGGTCTCACCCTTTCTACCTGGCGCTGGAGGACGGCGGGTCGGCACACGG 1500 1501 GGTGTTCCTGCTAAACAGCAATGCCATGGATGTGGTCCTGCAGCCGAGCCCTGCC CTTAGCTGGAGGTCGACAGGTGGGATCCTGGATGTCTACATCTTCCTGGGCCCAG 1502 AGCCCAAGAGCGTGGTGCAGCAGTACCTGGACGTTGTGGGATACCCGTTCATGCC 1503 1504 GCCATACTGGGGCCTGGGCTTCCACCTGTGCCGCTGGGGCTACTCCTCCACCGCT 1505 ATCACCCGCCAGGTGGTGGAGAACATGACCAGGGCCCACTTCCCCCTGGACGTC CAGTGGAACGACCTGGACTACATGGACTCCCGGAGGGACTTCACGTTCAACAAGG 1506 1507 GCTACATGATGATCGTGGATCCTGCCATCAGCAGCTCGGGCCCTGCCGGGAGCTA 1508 1509 CAGGCCCTACGACGAGGGTCTGCGGAGGGGGGGTTTTCATCACCAACGAGACCGG CCAGCCGCTGATTGGGAAGGTATGGCCCGGGTCCACTGCCTTCCCCGACTTCACC 1510 AACCCCACAGCCCTGGCCTGGTGGGAGGACATGGTGGCTGAGTTCCATGACCAG 1511 GTGCCCTTCGACGGCATGTGGATTGACATGAACGAGCCTTCCAACTTCATCAGGG 1512 1513 GGGTGGTTGGGGGGGGCCCCCCCAGGCGGCCACCATCTGTGCCTCCAGCCACCAGT 1514 TTCTCTCCACACACTACAACCTGCACAACCTCTACGGCCTGACCGAAGCCATCGCC 1515 TCCCACAGGGCGCTGGTGAAGGCTCGGGGGGACACGCCCATTTGTGATCTCCCGC 1516 TCGACCTTTGCTGGCCACGGCCGATACGCCGGCCACTGGACGGGGGGGCGTGTGG 1517 1518 AGCTCCTGGGAGCAGCTCGCCTCCTCCGTGCCAGAAATCCTGCAGTTTAACCTGC TGGGGGTGCCTCTGGTCGGGGCCGACGTCTGCGGCTTCCTGGGCAACACCTCAG 1519 AGGAGCTGTGTGTGCGCTGGACCCAGCTGGGGGGCCTTCTACCCCTTCATGCGGAA 1520 1521 CCACAACAGCCTGCTCAGTCTGCCCCAGGAGCCGTACAGCTTCAGCGAGCCGGC 1522 CCAGCAGGCCATGAGGAAGGCCCTCACCCTGCGCTACGCACTCCTCCCCCACCT

1523 1524 CTTCCTGGAGTTCCCCAAGGACTCTAGCACCTGGACTGTGGACCACCAGCTCCTG 1525 TGGGGGGGGGCCCTGCTCATCACCCCAGTGCTCCAGGCCGGGAAGGCCGAAGTG 1526 ACTGGCTACTTCCCCTTGGGCACATGGTACGACCTGCAGACGGTGCCAGTAGAGG 1527 CGAGGGGCAGTGGGTGACGCTGCCGGCCCCCCTGGACACCATCAACGTCCACCT 1528 1529 CCGGGCTGGGTACATCATCCCCCTGCAGGGCCCTGGCCTCACAACCACAGAGTC 1530 CCGCCAGCAGCCCATGGCCCTGGCTGTGGCCCTGACCAAGGGTGGGGGAGGCCC GAGGGGAGCTGTTCTGGGACGATGGAGAGAGCCTGGAAGTGCTGGAGCGAGGG 1531 1532 GCCTACACACAGGTCATCTTCCTGGCCAGGAATAACACGATCGTGAATGAGCTGG 1533 TACGTGTGACCAGTGAGGGAGCTGGCCTGCAGCTGCAGAAGGTGACTGTCCTGG GCGTGGCCACGGCGCCCCAGCAGGTCCTCTCCAACGGTGTCCCTGTCTCCAACTT 1534 1535 CACCTACAGCCCCGACACCAAGGTCCTGGACATCTGTGTCTCGCTGTTGATGGGA 1536 GAGCAGTTTCTCGTCAGCTGGTGT

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1538 GAADup327/328

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1594 **GLB1wt**

1595 ATGCCGGGGTTCCTGGTTCGCATCCTCCCTCTGTTGCTGGTTCTGCTGCTTCTGG GCCCTACGCGCGGCTTGCGCAATGCCACCCAGAGGATGTTTGAAATTGACTATAG 1596 CCGGGACTCCTTCCTCAAGGATGGCCAGCCATTTCGCTACATCTCAGGAAGCATTC 1597 1598 ACTACTCCCGTGTGCCCCGCTTCTACTGGAAGGACCGGCTGCTGAAGATGAAGAT GGCTGGGCTGAACGCCATCCAGACGTATGTGCCCTGGAACTTTCATGAGCCCTGG 1599 CCAGGACAGTACCAGTTTTCTGAGGACCATGATGTGGAATATTTTCTTCGGCTGGC 1600 1601 TCATGAGCTGGGACTGCTGGTTATCCTGAGGCCCGGGCCCTACATCTGTGCAGAG 1602 TGGGAAATGGGAGGATTACCTGCTTGGCTGCTAGAGAAAGAGTCTATTCTTCTCCG

CTCCTCCGACCCAGATTACCTGGCAGCTGTGGACAAGTGGTTGGGAGTCCTTCTG 1603 1604 CCCAAGATGAAGCCTCTCCTCTATCAGAATGGAGGGCCAGTTATAACAGTGCAGG TTGAAAATGAATATGGCAGCTACTTTGCCTGTGATTTTGACTACCTGCGCTTCCTGC 1605 AGAAGCGCTTTCGCCACCATCTGGGGGGATGATGTGGTTCTGTTTACCACTGATGGA 1606 1607 GCACATAAAACATTCCTGAAATGTGGGGCCCTGCAGGGCCTCTACACCACGGTGG ACTTTGGAACAGGCAGCAACATCACAGATGCTTTCCTAAGCCAGAGGAAGTGTGA 1608 1609 1610 CTTGCCCGTGGGGCGAGTGTGAACTTGTACATGTTTATAGGTGGGACCAATTTTGC 1611 1612 CTATTGGAATGGGGCCAACTCACCCTATGCAGCACAGCCCACCAGCTACGACTAT GATGCCCCACTGAGTGAGGCTGGGGGACCTCACTGAGAAGTATTTTGCTCTGCGAA 1613 ACATCATCCAGAAGTTTGAAAAAGTACCAGAAGGTCCTATCCCTCCATCTACACCA 1614 1615 AAGTTTGCATATGGAAAGGTCACTTTGGAAAAGTTAAAGACAGTGGGAGCAGCTCT 1616 GGACATTCTGTGTCCCTCTGGGCCCATCAAAAGCCTTTATCCCTTGACATTTATCCA GGTGAAACAGCATTATGGGTTTGTGCTGTACCGGACAACACTTCCTCAAGATTGCA 1617 GCAACCCAGCACCTCTCTCTCCACCCCTCAATGGAGTCCACGATCGAGCATATGTT 1618 GCTGTGGATGGGATCCCCCAGGGAGTCCTTGAGCGAAACAATGTGATCACTCTGA 1619 ACATAACAGGGAAAGCTGGAGCCACTCTGGACCTTCTGGTAGAGAACATGGGACG 1620 1621 TGTGAACTATGGTGCATATATCAACGATTTTAAGGGTTTGGTTTCTAACCTGACTCT CAGTTCCAATATCCTCACGGACTGGACGATCTTTCCACTGGACACTGAGGATGCAG 1622 TGTGCAGCCACCTGGGGGGCTGGGGACACCGTGACAGTGGCCACCATGATGAAG 1623 CCTGGGCCCACAACTCATCCAACTACACGCTCCCGGCCTTTTATATGGGGAACTTC 1624 1625 TCCATTCCCAGTGGGATCCCAGACTTGCCCCAGGACACCTTTATCCAGTTTCCTGG ATGGACCAAGGGCCAGGTCTGGATTAATGGCTTTAACCTTGGCCGCTATTGGCCA 1626 GCCCGGGGCCCTCAGTTGACCTTGTTTGTGCCCCAGCACATCCTGATGACCTCGG 1627 CCCCAAACACCATCACCGTGCTGGAACTGGAGTGGGCACCCTGCAGCAGTGATGA 1628 TCCAGAACTATGTGCTGTGACGTTCGTGGACAGGCCAGTTATTGGCTCATCTGTGA 1629 CCTACGATCATCCCTCCAAACCTGTTGAAAAAAGACTCATGCCCCCACCCCGCAA 1630 AAAAACAAAGATTCATGGCTGGACCATGTA 1631 1632

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1634 GLB1Dup84

1635 ATGCCGGGGTTCCTGGTTCGCATCCTCCCTCTGTTGCTGGTTCTGCTGCTTCTGG GCCCTACGCGCGGCTTGCGCAATGCCACCCAGAGGATGTTTGAAATTGACTATAG 1636 1637 CCGGGACTCCTTCCTCAAGGATGGCCAGCCATTTCGCTACATCTCAGGAAGCATTC 1638 ACTACTCCCGTGTGCCCCGCTTCTACTGGAAGGACCGGCTGCTGAAGATGAAGAT 1639 GGCTGGGCTGAACGCCATCCAGACGTATGTGCCCTGGAACTTTCATGAGCCCTGG CCAGGACAGTACCAGTTTTCTGAGGACCATGATGTGGAATATTTTCTTCGGCTGGC 1640 TCATGAGCTGGGACTGCTGGTTATCCTGAGGCCCGGGCCCTACATCTGTGCAGAG 1641 TGGGAAATGGGAGGATTACCTGCTTGGCTGCTAGAGAAAGAGTCTATTCTTCTCCG 1642 1643 CTCCTCCGACCCAGATTACCTGGCAGCTGTGGACAAGTGGTTGGGAGTCCTTCTG CCCAAGATGAAGCCTCTCCTCTATCAGAATGGAGGGCCAGTTATAACAGTGCAGG 1644 TTGAAAATGAATATGGCAGCTACTTTGCCTGTGATTTTGACTACCTGCGCTTCCTGC 1645 1646 AGAAGCGCTTTCGCCACCATCTGGGGGGATGATGTGGTTCTGTTTACCACTGATGGA 1647 GCACATAAAACATTCCTGAAATGTGGGGCCCTGCAGGGCCTCTACACCACGGTGG ACTTTGGAACAGGCAGCAACATCACAGATGCTTTCCTAAGCCAGAGGAAGTGTGA 1648 1649 1650 CTTGCCCGTGGGGCGAGTGTGAACTTGTACATGTTTATAGGTGGGACCAATTTTGC 1651 CTATTGGAATGGGGCCAACTCACCCTATGCAGCACAGCCCACCAGCTACGACTAT 1652 GATGCCCCACTGAGTGAGGCTGGGGGACCTCACTGAGAAGTATTTTGCTCTGCGAA 1653 ACATCATCCAGAAGTTTGAAAAAGTACCAGAAGGTCCTATCCCTCCATCTACACCA 1654 AAGTTTGCATATGGAAAGGTCACTTTGGAAAAGTTAAAGACAGTGGGAGCAGCTCT 1655 1656 GGACATTCTGTGTCCCTCTGGGCCCATCAAAAGCCTTTATCCCTTGACATTTATCCA GGTGAAACAGCATTATGGGTTTGTGCTGTACCGGACAACACTTCCTCAAGATTGCA 1657 GCAACCCAGCACCTCTCTCTCCACCCCTCAATGGAGTCCACGATCGAGCATATGTT 1658 GCTGTGGATGGGATCCCCCAGGGAGTCCTTGAGCGAAACAATGTGATCACTCTGA 1659 1660 ACATAACAGGGAAAGCTGGAGCCACTCTGGACCTTCTGGTAGAGAACATGGGACG TGTGAACTATGGTGCATATATGGTGCATATATCAACGATTTTAAGGGTTTGGTTTCT 1661 AACCTGACTCTCAGTTCCAATATCCTCACGGACTGGACGATCTTTCCACTGGACAC 1662 TGAGGATGCAGTGTGCAGCCACCTGGGGGGGCTGGGGACACCGTGACAGTGGCCA 1663 1664 CCATGATGAAGCCTGGGCCCACAACTCATCCAACTACACGCTCCCGGCCTTTTATA TGGGGAACTTCTCCATTCCCAGTGGGATCCCAGACTTGCCCCAGGACACCTTTATC 1665 CAGTTTCCTGGATGGACCAAGGGCCAGGTCTGGATTAATGGCTTTAACCTTGGCC 1666 1667 GATGACCTCGGCCCCAAACACCATCACCGTGCTGGAACTGGAGTGGGCACCCTG 1668 1669 CAGCAGTGATGATCCAGAACTATGTGCTGTGACGTTCGTGGACAGGCCAGTTATT GGCTCATCTGTGACCTACGATCATCCCTCCAAACCTGTTGAAAAAAGACTCATGCC 1670 CCCACCCCGCAAAAAAAAAAAGATTCATGGCTGGACCATGTA 1671 1672

1674 **PORCNwt**

1675 ATGGCCACCTTTAGCCGCCAGGAATTTTTCCAGCAGCTACTGCAAGGCTGTCTCCT GCCTACTGCCCAGCAGGGCCTTGACCAGATCTGGCTGCTCCTTGCCATCTGCCTC 1676 GCCTGCCGCCTCCTCGGAGGCTCGGGTTGCCATCCTACCTGAAGCATGCAAGCA 1677 CCGTGGCAGGCGGGTTCTTCAGCCTCTACCACTTCTTCCAGCTGCACATGGTTTG 1678 1679 GGTCGTGCTGCTCAGCCTCCTGTGCTACCTCGTGCTGTTCCTCTGCCGACATTCCT 1680 1681 ATGCACATGGTAGACACCGTGACATGGCACAAGATGCGAGGGGGCACAGATGATTG TGGCCATGAAGGCAGTGTCTCTGGGCTTCGACCTGGACCGGGGCGAGGTGGGTA 1682 1683 CGGTGCCCTCGCCAGTGGAGTTCATGGGCTACCTCTACTTCGTGGGCACCATCGT CTTCGGGCCCTGGATATCCTTCCACAGCTACCTACAAGCTGTCCAAGGCCGCCCA 1684 CTGAGCTGCCGGTGGCTGCAGAAGGTGGCCCGGAGCCTGGCACTGGCCCTGCTG 1685 1686 TGCCTTGTGCTGTCCACTTGCGTGGGCCCCTACCTCTTCCCGTACTTCATCCCCCT 1687 CAACGGTGACCGCCTCCTTCGCAAGGGCACCATGGTAAGGTGGCTGCGAGCCTA 1688 CCACGGCCACGTTGGCGGGGGGCTGGCTTTACCGAGGAGAAGGATCACCTGGAAT 1689 GGGACCTGACGGTGTCCAAGCCACTGAATGTGGAGCTGCCTCGGTCAATGGTGG 1690 AAGTTGTCACAAGCTGGAACCTGCCCATGTCTTATTGGCTAAATAACTATGTTTTCA 1691 1692 AGAATGCTCTCCGCCTGGGGGACCTTCTCGGCTGTGCTGGTCACCTATGCAGCCAG CGCCCTCCTACATGGCTTCAGTTTCCACCTGGCTGCGGTCCTGCTGTCCCTGGCT 1693 TTTATCACTTACGTGGAGCATGTCCTCCGGAAGCGCCTGGCTCGGATCCTCAGTG 1694 CCTGTGTCTTGTCAAAGCGGTGCCCGCCAGACTGTTCGCACCAGCATCGCTTGGG 1695 1696 CCTGGGGGTGCGAGCCTTAAACTTGCTCTTTGGAGCTCTGGCCATCTTCCACCTG GCCTACCTGGGCTCCCTGTTTGATGTCGATGTGGATGACACCACAGAGGAGCAGG 1697 GCTACGGCATGGCATACACTGTCCACAAGTGGTCAGAGCTCAGCTGGGCCAGTCA 1698 CTGGGTCACTTTTGGATGCTGGATCTTCTACCGTCTCATAGGC 1699 1700 1701 PORCNDup20 ATGGCCACCTTTAGCCGCCAGGAATTTTTCCAGCAGCTACTGCAAGGCTGTCTCCT 1702 1703 GCCTACTGCCCAGCAGGGCCTTGACCAGATCTGGCTGCTCCTTGCCATCTGCCTC 1704 GCCTGCCGCCTCCTCTGGAGGCTCGGGTTGCCATCCTACCTGAAGCATGCAAGCA CCGTGGCAGGCGGGTTCTTCAGCCTCTACCACTTCTTCCAGCTGCACATGGTTTG 1705 GGTCGTGCTGCTCAGCCTCCTGTGCTACCTCGTGCTGTTCCTCTGCCGACATTCCT 1706

ATGCACATGGTAGACACCGTGACATGGCACAAGATGCGAGGGGGCACAGATGATTG
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- 1710 CGGTGCCCTCGCCAGTGGAGTTCATGGGCTACCTCTACTTCGTGGGCACCATCGT
- 1711 CTTCGGGCCCTGGATATCCTTCCACAGCTACCTACAAGCTGTCCAAGGCCGCCCA
- 1712 CTGAGCTGCCGGTGGCTGCAGAAGGTGGCCCGGAGCCTGGCACTGGCCCTGCTG
- 1713 TGCCTTGTGCTGTCCACTTGCGTGGGCCCCTACCTCTTCCCGTACTTCATCCCCCT

CAACGGTGACCGCCTCCTTCGCAAGGGCACCATGGTAAGGTGGCTGCGAGCCTA 1714 1715 CCACGGCCACGTTGGCGGGGGGCTGGCTTTACCGAGGAGAAGGATCACCTGGAAT 1716 GGGACCTGACGGTGTCCAAGCCACTGAATGTGGAGCTGCCTCGGTCAATGGTGG 1717 1718 AAGTTGTCACAAGCTGGAACCTGCCCATGTCTTATTGGCTAAATAACTATGTTTTCA AGAATGCTCTCCGCCTGGGGGACCTTCTCGGCTGTGCTGGTCACCTATGCAGCCAG 1719 CGCCCTCCTACATGGCTTCAGTTTCCACCTGGCTGCGGTCCTGCTGTCCCTGGCT 1720 1721 TTTATCCCTGGCTTTTATCACTTACGTGGAGCATGTCCTCCGGAAGCGCCTGGCTC GGATCCTCAGTGCCTGTGTCTTGTCAAAGCGGTGCCCGCCAGACTGTTCGCACCA 1722 1723 GCATCGCTTGGGCCTGGGGGTGCGAGCCTTAAACTTGCTCTTTGGAGCTCTGGCC 1724 ATCTTCCACCTGGCCTACCTGGGCTCCCTGTTTGATGTCGATGTGGATGACACCAC AGAGGAGCAGGGCTACGGCATGGCATACACTGTCCACAAGTGGTCAGAGCTCAG 1725 1726 CTGGGCCAGTCACTGGGTCACTTTTGGATGCTGGATCTTCTACCGTCTCATAGGC 1727

1728 References

- 1729 1. Ceccaldi, R., Rondinelli, B. & D'Andrea, A. D. Repair Pathway Choices and Consequences at
- the Double-Strand Break. *Spec. Issue Qual. Control* **26**, 52–64 (2016).
- 1731 2. DiCarlo, J. E., Chavez, A., Dietz, S. L., Esvelt, K. M. & Church, G. M. Safeguarding
- 1732 CRISPR-Cas9 gene drives in yeast. *Nat. Biotechnol.* **33**, 1250 (2015).
- 1733 3. McVey, M. & Lee, S. E. MMEJ repair of double-strand breaks (director's cut): deleted
- sequences and alternative endings. *Trends Genet.* **24**, 529–538 (2008).
- 1735 4. Yu, A. M. & McVey, M. Synthesis-dependent microhomology-mediated end joining accounts
- 1736 for multiple types of repair junctions. *Nucleic Acids Res.* **38**, 5706–5717 (2010).
- 1737 5. Davis, A. J. & Chen, D. J. DNA double strand break repair via non-homologous end-joining.
- 1738 Transl. Cancer Res. 2, 130–143 (2013).
- 1739 6. Heidenreich, E., Novotny, R., Kneidinger, B., Holzmann, V. & Wintersberger, U. Non-
- 1740 homologous end joining as an important mutagenic process in cell cycle-arrested cells. *EMBO*
- 1741 *J.* **22,** 2274 (2003).
- 1742 7. Pfeiffer, P., Goedecke, W. & Obe, G. Mechanisms of DNA double-strand break repair and
- their potential to induce chromosomal aberrations. *Mutagenesis* **15**, 289–302 (2000).
- 1744 8. Brown, J. S. et al. Neddylation Promotes Ubiquitylation and Release of Ku from DNA-
- 1745 Damage Sites. *Cell Rep.* **11**, 704–714 (2015).
- 1746 9. Landrum, M. J. *et al.* ClinVar: Public archive of interpretations of clinically relevant variants.
- 1747 *Nucleic Acids Res.* **44**, D862–D868 (2016).
- 1748 10. Stenson, P. D. *et al.* Human Gene Mutation Database: towards a comprehensive central
- 1749 mutation database. J. Med. Genet. 45, 124 (2008).
- 1750
- 1751
- 1752