

1 A framework for evaluating edited cell 2 libraries created by massively parallel 3 genome engineering

4 **Author list:** Simon Cawley¹, Eric Abbate¹, Christopher G. Abraham¹, Steven Alvarez¹, Mathew
5 Barber¹, Scott Bolte¹, Jocelyne Bruand¹, Deanna M. Church¹, Clint Davis¹, Matthew Estes¹,
6 Stephen Federowicz^{1,2}, Richard Fox^{1,3}, Miles W. Gander^{1,4}, Andrew D. Garst^{1,3}, Gozde Gencer¹,
7 Andrea L. Halweg-Edwards¹, Paul Hardenbol¹, Thomas Hraha¹, Surbhi Jain¹, Charlie Johnson¹,
8 Kara Juneau¹, Nandini Krishnamurthy¹, Shea Lambert¹, Bryan Leland¹, Francesca Pearson¹, J.
9 Christian J. Ray¹, Chad D. Sanada¹, Timothy M. Shaver¹, Tyson R. Shepherd¹, Joshua
10 Shorenstein¹, Eileen C. Spindler¹, Craig A. Struble¹, Maciej H. Swat¹, Stephen Tanner¹, Tian
11 Tian¹, Ken Wishart¹, Michael S. Graige¹

12 ¹Inscripta, Inc., Boulder, CO 80301

13 ²IDEAYA Biosciences, South San Francisco, CA 94080

14 ³Infinome Biosciences, Boulder, CO, 80301

15 ⁴Absci Corp, Vancouver, WA 98683

16 **Abstract**

17 Genome engineering methodologies are transforming biological research and discovery.
18 Approaches based on CRISPR technology have been broadly adopted and there is growing
19 interest in the generation of massively parallel edited cell libraries. Comparing the libraries
20 generated by these varying approaches is challenging and researchers lack a common
21 framework for defining and assessing the characteristics of these libraries. Here we describe a
22 framework for evaluating massively parallel libraries of edited genomes based on established
23 methods for sampling complex populations. We define specific attributes and metrics that are
24 informative for describing a complex cell library and provide examples for estimating these
25 values. We also connect this analysis to generic phenotyping approaches, using either pooled
26 (typically via a selection assay) or isolate (often referred to as screening) phenotyping
27 approaches. We approach this from the context of creating massively parallel, precisely edited
28 libraries with one edit per cell, though the approach holds for other types of modifications,
29 including libraries containing multiple edits per cell (combinatorial editing). This framework is a
30 critical component for evaluating and comparing new technologies as well as understanding
31 how a massively parallel edited cell library will perform in a given phenotyping approach.

32 **Introduction**

33 Genome engineering methodologies are transforming biological research and discovery.
34 Approaches based on CRISPR technology have been broadly adopted due to the relative ease
35 of targeting defined genomic regions using specific guide RNAs (gRNAs) (Jinek et al. 2012).
36 While there has been a large focus on modifying one or a small number of sites for translational
37 research and therapeutics, there is growing interest in the generation of massively parallel
38 edited cell libraries (Ding et al. 2014; Frangoul et al. 2020; Wilkinson et al. 2021). These libraries
39 can accelerate the pace of genome discovery or cell engineering by allowing for the
40 simultaneous interrogation of hundreds to thousands of loci in a single experiment. Current
41 genome-wide approaches typically either leverage knock-out libraries – largely relying on
42 error-prone repair processes for sequence disruptions – or rely on transcriptional modulation
43 by tethering a nuclease-deficient Cas9 with a transcriptional repressor or activator to modulate
44 gene expression (Mali et al. 2013; Cong et al. 2013; Gilbert et al. 2014). Recently, the generation
45 of genome-wide libraries of precise edits has been described in microbes and human (Garst et
46 al. 2017; Sadhu et al. 2018; Bao et al. 2018; Sharon et al. 2018; Hanna et al. 2021). This ability to
47 make more refined changes will provide greater precision and information around genotype-
48 phenotype relationships. Comparing the libraries generated by these varying approaches is
49 challenging and groups typically take different approaches and measures in reporting their
50 work. What is currently lacking is a common framework for defining and assessing the
51 characteristics of these libraries.

52 The evaluation of these complex libraries can be challenging. The library represents a mixed
53 population, with some cells containing the desired edit and the remaining cells constituting a
54 Burden Population (Table 1) of cells containing incomplete, unintended or no edits. The
55 population of cells containing the designed edits will also be a mosaic, with individual edit
56 representations being driven by the representation of the design in the reagent pool, the
57 functionality of the guide, the edit rate at different loci and any fitness effects an edit may have
58 on an individual cell. Frequently the efficiency of massively parallel editing experiments is
59 extrapolated based on experiments where editing has been performed in isolates rather than
60 in a pooled manner (Sadhu et al. 2018; Sharon et al. 2018). Although this methodology is more
61 experimentally tractable, it is not necessarily predictive of performance in a pooled setting.
62 Additional biological factors can strongly affect outcomes, such as differential growth rates of
63 cells that have undergone the editing process, the introduction of edits that impair cell viability
64 to varying degrees, cells in which no double-stranded break (DSB) is created and which thus
65 grow faster, and cells in which a DSB is created with failure to repair leading to their depletion.
66 All of these factors impact the final library composition. In general, it is preferable for a library
67 to contain a high fraction of edited cells, with an even representation of edits. Understanding
68 the library composition is critical for assessing if a cell library is fit for a given phenotyping
69 regime, though in practice obtaining this information can be technically challenging or cost
70 prohibitive.
71 Here we describe a framework for evaluating massively parallel libraries of edited genomes
72 based on established methods for sampling complex populations. We define specific attributes
73 and metrics that are informative for describing a complex cell library and provide examples for
74 estimating these values. Obtaining all of these measures may be challenging or expensive, so

75 we also provide a theoretical framework to allow assessment of a given library in the absence
76 of some desired data points. We also connect this analysis to generic phenotyping approaches,
77 using either pooled (typically via a selection assay) or isolate (often referred to as screening)
78 phenotyping approaches. We approach this from the context of creating massively parallel,
79 precisely edited libraries with one edit per cell, though the approach holds for other types of
80 modifications, including libraries containing multiple edits per cell (combinatorial editing). This
81 framework is a critical component for evaluating and comparing new technologies as well as
82 understanding how a massively parallel edited cell library will perform in a given phenotyping
83 approach.

84 **Library Characterization**

85 Massively parallel genome engineering results in a library of cells, where most cells contain
86 design reagents (that is, the combination of gRNA and repair template) encoding distinct edits.
87 Each design reagent is represented in hundreds to thousands of cells. In microbial libraries,
88 these reagents are often maintained as plasmids, while in mammalian libraries, episomes or
89 genome-integrating vectors, such as lentivirus, must be used if the reagents are to be
90 maintained within the population over the course of an experiment. In many cases, the
91 reagents are attached to a barcode, or are used as a barcode themselves to track which cells
92 contain specific reagents. If selection pressure is applied to the library, these reagents may also
93 serve as a proxy for genotyping the specific edit. A percentage of the population will contain
94 the desired edits, while the remaining population constitutes a Burden Population. In order to
95 characterize such a library, we must define and measure several characteristics. Table 1
96 provides a list of terms and measures useful for characterizing libraries.

97 **Table 1:** terms and definitions useful for characterizing complex cell libraries

TERM	DEFINITION
BURDEN POPULATION	The population of cells in a library that is either unedited or contains unintended edits.
COMPLETE INTENDED EDIT	A precise edit that includes all modifications specified in the repair template (sometimes referred to as the homology arm) with no additional unintended modifications (Figure 1).
EDIT COEFFICIENT OF VARIATION (EDIT CV)	An aggregate measure across all the edits in a library, the coefficient of variation for the frequencies of the Complete Intended Edits in the edited cells of the library, defined as the standard deviation of edit frequencies normalized to their mean.
EDIT FRACTION	The fraction of cells in a library containing the Complete Intended Edit at the locus of interest (in a precise editing library) or an edit in the target region (in an imprecise editing library).

EDIT FRACTIONAL RICHNESS	The Edit Richness (see below) scaled by the library size, a value in the range [0, 1].
EDIT RICHNESS	The number of unique Complete Intended Edits present in a sample.
INTENDED EDIT	The modification of specific bases in a defined region of a genome. (https://www.nist.gov/programs-projects/nist-genome-editing-lexicon#3.4)
REAGENT COEFFICIENT OF VARIATION (REAGENT CV)	An aggregate measure across all the editing reagents in a library, the coefficient of variation for the frequencies of the editing reagents (typically plasmids, episomes or virus) in the library. Defined as for Edit CV
REAGENT FRACTIONAL RICHNESS	The Reagent Richness (see below) scaled by the library size, a value in the range [0, 1].
REAGENT RICHNESS	The number of unique reagents present in a sample.
SCREENER'S SCORE	The predicted Edit Fractional Richness for a 1x screen (number of isolates screened = number of designs in library) assuming a 30% Edit Fraction.
SELECTOR'S SCORE	The predicted Edit Fractional Richness for a selection assuming 1×10^6 cells and 30% Edit Fraction.
TRACKABILITY	The ability to identify the reagents associated with a cell in a complex cell library.

98 **Definitions Useful for Library Characterization**

99 **Defining an edit**

100 When using CRISPR-Cas based systems to generate a desired sequence variant through
101 precise editing, a guide and repair template are defined (commonly through software). In
102 many cases, auxiliary edits to the PAM site are included to prevent the nuclease from recutting
103 the edited locus. We define a 'Complete Intended Edit' as an instance where the repair
104 template sequence (the desired variant and any auxiliary edits) is faithfully and completely
105 placed into the genome (Figure 1) and no other changes occur in the genome. Cases where
106 only part of the repair template sequence is conferred to the genome are classified as
107 incomplete edits and are considered part of the burden, though there will be differences from
108 the reference sequence. Unintended events (off-target editing, reagent integration, or other
109 large-scale genome rearrangement), either occurring at the edit locus or elsewhere in the
110 genome, are also considered part of the Burden Population along with unedited cells.

111 When producing imprecise edits, such as in the case of non-homologous end joining (NHEJ)-
112 mediated knockout libraries, the concept of a Complete Intended Edit is not relevant.
113 However, in this case, the desired events would be insertion-deletion events occurring at the
114 target site. Events that do not lead to a true loss of functional protein (knockout) or that
115 happen outside of target region would fall into the Burden Population. In this framework, only

116 Complete Intended Edits (in precise editing) or target site changes leading to a knockout (in
117 imprecise editing) are considered edits. A formal definition of what is meant by an edit allows
118 us to develop a more rigorous framework by which to evaluate these complex cell libraries. In
119 the discussion that follows, the term “edit” refers to Complete Intended Edit unless indicated
120 otherwise.

Repair template

Reference Sequence	target				auxiliary				
	T	A	C	G	G	T	T	T	C
Designed Sequence	A	C	T	G	G	A	T	T	T
Complete and Intended	A	C	T	G	G	A	T	T	T
<i>Complete, but unintended</i>	A	C	A	G	G	A	T	T	T
<i>Incomplete and unintended</i>	A	C	A	G	G	T	T	T	C
Burden	<i>Incomplete</i>	T	A	C	G	G	A	T	T
		A	C	T	G	G	T	T	T
		T	A	C	G	G	T	T	T
<i>unedited</i>	overlap with/outside repair template								

121

122 **Figure 1.** Challenges of edit identification in a large pool of precisely edited cells. A complete and intended edit
123 occurs only when the complete repair template is faithfully placed in the genome; this includes the desired edit
124 and any auxiliary edits made to prevent recutting of the edited locus. Cases where only part of the repair template
125 are incorporated into the genome are considered incomplete and count as burden rather than an edit, even if they
126 include the desired variant. Any other unintended or unedited cells are also considered part of the burden.

127 **Estimation of the Edit Fraction**

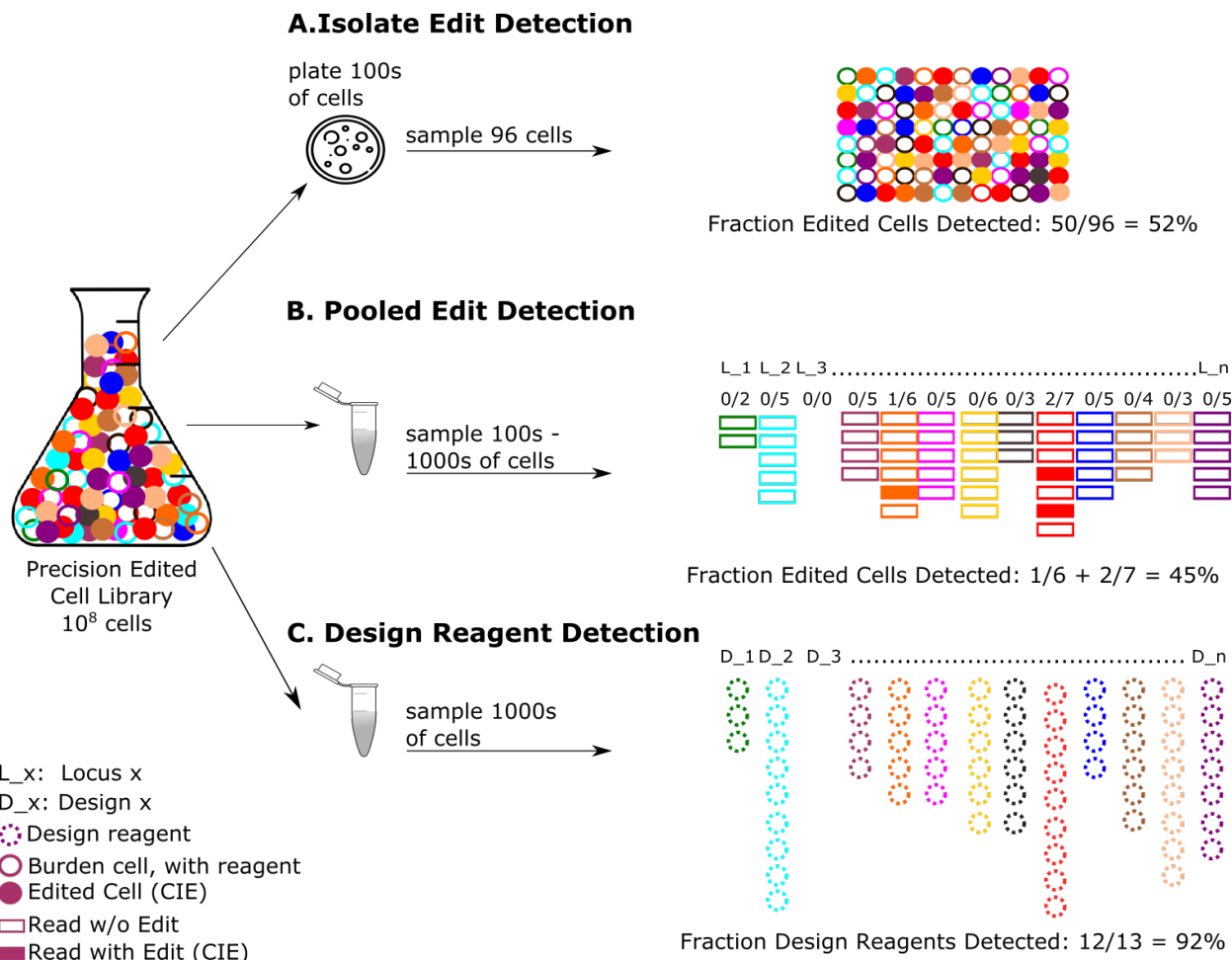
128 The Edit Fraction is a critical component of characterizing a massively parallel genome
129 engineered library. Ideally, we would like to identify all edits that occurred within a population.
130 In practice, this is challenging because of the mosaic nature of the library; at any given locus,
131 the count of reference sequence representation will far exceed the count of edit-containing
132 sequences. Fortunately, determination of the overall Edit Fraction does not require complete
133 evaluation of all members of the library. We describe two approaches for identifying the Edit
134 Fraction in a library: a shallow sampling of the library by deeply sequencing isolates or a deeper
135 sampling of the library by shallow sequencing of a pool of cells (Figure 2).

136 One way to assess the Edit Fraction is to sample isolates selected from the population (Figure
137 2A). After sufficient cell divisions, standard sequencing approaches, such as whole genome
138 shotgun (WGS) of each isolate, can be employed. This requires only collection and growth of
139 isolates (typically by low density plating and picking single colonies into a 96-well plate) and

140 library preparation. While this produces a large number of reads outside of the targeted locus
141 that do not contribute to edit detection, these reads can be assessed for off-target events.
142 Alternatively, one could take an approach to identify the design reagent in each isolate (see
143 below), and then use a targeted sequencing approach, such as hybrid capture or genomic
144 amplification, to confirm the validity of the edit. This approach has the benefit of more
145 efficiently utilizing sequencing reads but takes longer and requires two library preparations, in
146 addition to the creation of custom reagents for each edit locus. Regardless of whether whole
147 genome or targeted sequencing is performed, this isolate evaluation approach generally
148 results in very shallow sampling of a library.

149 An alternative approach to characterizing the Edit Fraction in a library employs limited WGS on
150 the entire population of cells at a shallow read depth, an approach we term pooled WGS
151 (pWGS) (Figure 2B). While the population of cells used as input for this analysis may number in
152 the millions, the cost of sequencing will typically limit the number of cells ultimately sampled,
153 often in the range of a few hundred to a few thousand. For example, if an experiment involves
154 sequencing to an average genomic coverage depth of 1000x, it will profile approximately 1000
155 cells' worth of DNA at each targeted edit locus. In contrast to isolate sampling, the pooled
156 approach limits the manual work of colony isolation and growth at the expense of greater
157 complexity in sequence analysis. If a pWGS assay is tuned to sequence roughly 1000 genomes'
158 worth of DNA per locus, then for an edit library of 1000 or more members, the assay should be
159 viewed as a sampling of mainly the right tail of the edit frequency distribution. Sampling
160 deeper would require substantially more sequencing, on the order of billions of read pairs or
161 more (Figure 3D and supplemental section 8). Even though the pWGS sampling depth is
162 typically shallow and thus incapable of providing reliable data on a per-design basis, the sum of
163 the per-design Edit Fractions produces a reliable estimate of the overall Edit Fraction in the
164 library (Figure 3A). In either the isolate or pWGS approach, many edits that are present in the
165 pool will be missed in the sequencing results due to being present at very low frequency
166 relative to the per-locus sampling depth. Despite the absence of many of the edits in the
167 sample, making the assumption that the underlying edit frequencies follow a parametric
168 distribution can allow for reliable estimation of the Edit CV (Table 1 and Figure 3D). In
169 situations where the edits are clustered in a subset of the genome, targeted sequencing
170 approaches can provide a more cost-efficient readout of the edit frequencies. Assay replicates
171 will provide differing parameter estimates due to sampling biases in the context of shallow
172 coverage; therefore, inspection of confidence intervals is helpful to guide appropriate
173 interpretation.

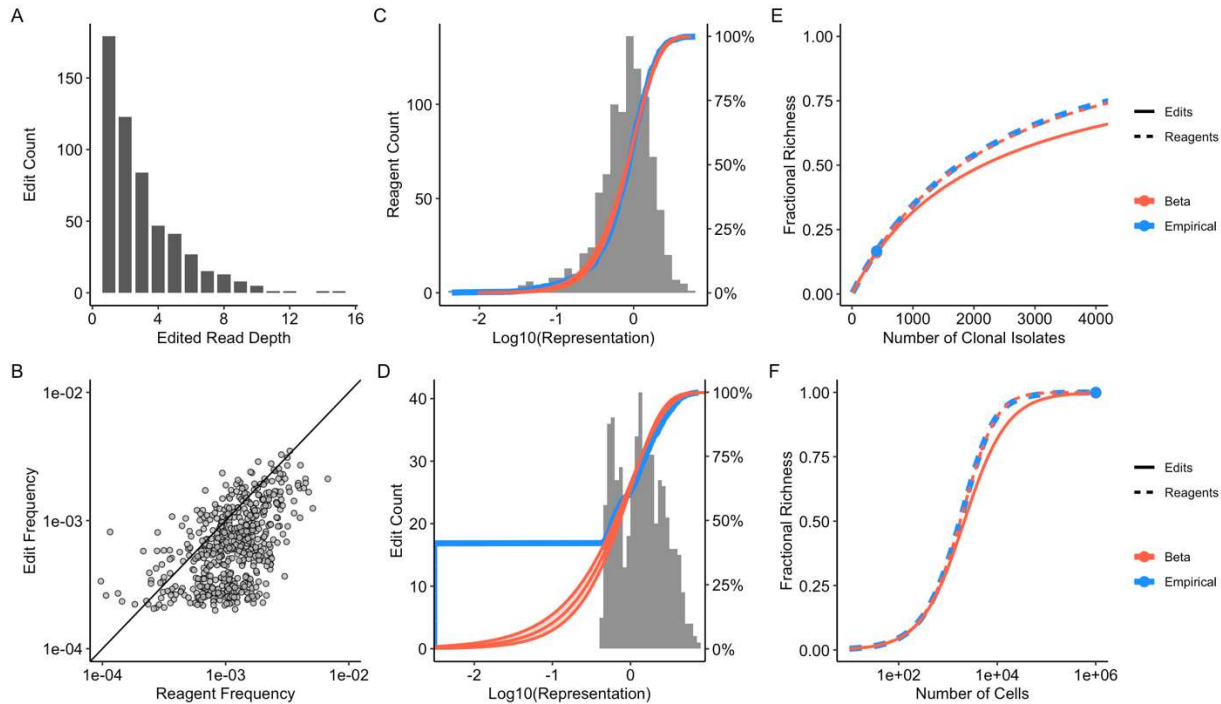
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177 **Figure 2.** Measurements of interest when evaluating a multiplex precisely edited library. This simplified example
178 is based on a contrived library targeting 13 distinct edits, with half of the cells in the pool containing a Complete
179 Intended Edit and 12 of the designs represented. Open circles represent cells of the Burden Population, most of
180 which will contain editing reagents if selection pressure is maintained or if the trackability reagent is intentionally
181 integrated into the genome. Dashed circles represent the design reagent. Rectangular boxes represent sequence
182 reads, open are wild type while filled are Complete Intended Edit-containing reads. **A.** A shallow library sampling
183 but deep sequencing approach involves edit detection by selecting isolates and performing whole genome
184 shotgun (WGS) analysis. For the isolates selected, this can provide detailed edit data, as well as information on
185 any unintended events, but the approach samples only a small number of cells in the library. It is important to use
186 sufficiently deep sequencing on each isolate to provide good power for detecting edits. **B.** An alternative
187 approach involves doing a broad library sampling but shallow sequence assessment of the library to obtain an
188 estimate of the fraction of cells containing an edit. As with the previous approach, many individual edits that are
189 present in the pool will be absent from the sample; nevertheless, an estimate of Edit Fraction f can be obtained by
190 summing the fraction of edited reads at each locus (designated by L_n). At approximately 1000x coverage and
191 with Edit Fraction f , 1000 f edited cells will be sampled. Increasing read depth will increase the number of cells
192 sampled, but very high coverage would be required to deeply assay at each edit locus. **C.** Design distribution can
193 be measured directly from the reagents, typically through a short-read sequencing (NGS) assay using
194 amplification handles. The reagents will be detected in both the edited and Burden Populations, and this assay
195 will not distinguish those populations in the absence of strong selection for edited cells.



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198 **Figure 3:** Example usage of pWGS and design reagent amplicon sequencing assays to characterize an *E. coli* edit
 199 library. After exclusion of controls, the library consists of 928 designs including insertions, deletions and
 200 substitutions spanning the genome. The resulting edits are not expected to result in any notable effects on
 201 cellular fitness. **A:** Number of sequencing reads with exact match to expected edits in a pWGS run. The pWGS run
 202 included 157M 2x150 read pairs. After exclusion of reads failing quality filters the mean coverage depth fully
 203 spanning the targeted edits is 3434. Summing the per-locus Edit Fractions produces an estimate of 0.44 for the
 204 overall Edit Fraction in the pool, thus the pWGS run profiles approximately 1501 genomes' worth of DNA overall.
 205 A total of 1615 edited reads is seen, comprising 546 unique edits (y-axis) with read depth per edit ranging from 1
 206 to 15 (x-axis). **B:** Scatterplot comparing the edit frequencies estimated from pWGS with design reagent
 207 frequencies estimated from amplicon sequencing of reagents. **C:** Histogram and cumulative distribution function
 208 (CDF) of reagent representation (defined as the product of reagent frequency and library size), measured by
 209 amplicon sequencing of the design reagents. The assay consists of 3.0M reads. Fitting the design reagent
 210 frequencies to a beta distribution via maximum likelihood estimation (MLE), the data are well described by a beta
 211 distribution with mean 1/928 and CV 0.73. **D:** Histogram and CDF as in C, but for the representation of edits as
 212 measured by pWGS. Given that the pWGS run is sampling roughly 1501 genomes' worth of DNA per locus, it
 213 should be viewed as a sampling of mainly the right tail of the edit frequency distribution. The fraction of the edit
 214 library that is observed at least once is 0.59. Fitting edit frequencies with a beta distribution via MLE, the estimate
 215 of CV is 1.01. Observation of a greater fraction of all possible edits in the library would require substantially more
 216 sequencing. For example, if the goal were to directly observe 90% of the edits in pWGS, it would require detection
 217 of edits whose frequencies among the 44% of edited cells is around the 10th percentile of the reagent frequency
 218 distribution, or 1e-4. Aiming for an expected edit read count of 10, to have a reasonable chance of observing edits
 219 at the 10th percentile, it would take a mean coverage depth of 213K. This is 62-fold larger than the actual
 220 coverage depth for the pWGS run, which would require a total sequencing throughput of 9.8B read pairs. **E:**
 221 Screener's curve, showing the predicted Reagent Fractional Richness (solid curve) and Edit Fractional Richness
 222 (dashed curve) as a function of the number of clonal isolates phenotyped in a screening experiment. The red
 223 curves are based on a beta binomial model fit. The blue curve is a prediction based on the nonparametric estimate
 224 of the distribution of reagent frequencies, a nonparametric fit to the edit frequencies is not useful given the

225 limited sampling depth of the pWGS data. The point indicated on the curve corresponds to the Screener's score,
226 which is the predicted Edit Fractional Richness when sampling depth is equal to the library size times the Edit
227 Fraction. **F**: Selector's curve, showing the same data as in E but with the x-axis changed to log scale and domain
228 extended to cover the deep sampling that is typically relevant for the large number of cells sampled in selection
229 applications. The solid point indicated on the curve corresponds to the Selector's score, which is the predicted
230 Edit Fractional Richness when sampling 1M cells.

231 **Estimation of Reagent Distribution**

232 Direct detection of edits in massively parallel editing libraries is ideal for assessing library
233 diversity, but in practice it is often prohibitively expensive due to the depth of sequencing
234 required. In lieu of extensive genomic sequencing, many approaches make it relatively
235 straightforward to detect the reagents conferring edits, so profiling the reagent distribution
236 can be a useful proxy for the edit distribution. For microbes, each cell typically contains
237 multiple clonal reagent copies, and most reagents will be present in hundreds to thousands of
238 cells. For mammalian cells, the copy number of the trackable reagent is typically lower, on the
239 order of one to less than 10. Ideally, all designs would be equally represented, but in practice
240 most libraries have a distribution of representation. Every manipulation of the library (reagent
241 manufacturing, transformation, growth of the cell population) introduces an opportunity to
242 alter this distribution. Understanding the distribution of reagents is critical for interpreting
243 phenotyping results and will help define the effect size and significance of results. For example,
244 if a phenotyping approach is assessing depletion of reagents as a measure proxy for genotype
245 (a common approach in essential gene screens), designs in the extreme left tail of the
246 distribution will likely be underpowered for association with a phenotype.

247 Sequencing the reagent library throughout the experimental process provides useful insight
248 into how various manipulations can impact design reagent distribution. This approach can be
249 useful for approximating edits post-phenotyping, particularly in the case of strong selective
250 pressure. In a library containing a mixture of active and inactive gRNA-donor cassettes, the
251 number of viable edited cells is tightly coupled to gRNA activity, rate of homology directed
252 repair (HDR) and the relative survival rate of edited members of the population. DNA synthesis
253 errors that result in unintended editing events during the homology-directed repair process or
254 poor transformation efficiency can impact uniform representation of intended edits (Roy et al.
255 2018). These effects can reduce the effective diversity in an edited library, directly impacting
256 the success of phenotyping. For instance, edited variant libraries may lack the desired intended
257 diversity due to editing process failures or takeover by a sub-population of a particular
258 Complete Intended Edit, unintended edits or unedited cells. In each of these cases, the cost
259 and effectiveness of phenotypic investigations will be adversely affected.

260 Typically, short read sequencing (NGS) of the reagent is used to determine the library
261 distribution from a sample of the library (Fig 2C). Approaches that either detect a barcode
262 (Garst et al. 2017; Sadhu et al. 2018) or the reagents themselves (Bao et al. 2018; Sharon et al.
263 2018) are used. It is assumed that the read counts for a design reagent are proportional to the
264 number of cells containing that design; thus, a read count is equivalent to a design reagent
265 count. The dispersion of the distribution is measured by the Reagent CV (Table 1, Figure 3C).

266 Larger Reagent CV values indicate greater variance in the relative abundances of the designs,
267 which can lead to under- or overrepresentation of individual designs. Prior to applying selective
268 pressure, a small Reagent CV is preferable for all phenotyping approaches, though libraries
269 with larger Reagent CVs can still be useful for some experiments. It is important to note that
270 while the Reagent CV is a useful and accessible metric, what matters most for many
271 applications is the **Edit CV** (Table 1). If every design reagent has an equal probability of
272 producing an edit, the Reagent CV and Edit CV will be equal to one another. In most real-world
273 situations there are various sources of bias, including those mentioned above, which result in
274 the Edit CV being larger than the Reagent CV, to an extent that will depend on the
275 experimental context (Figure 3D).

276 We have introduced measures that can be useful for describing aspects of a massively parallel
277 edited cell library. We next introduce approaches for combining these measures to produce
278 metrics that can be utilized for evaluating these libraries.

279 Metrics for Library Evaluation

280 In this section we define several concepts that utilize the above measurements to provide a
281 fuller characterization of a library. Neither Edit Fraction nor reagent distribution alone can fully
282 characterize the utility of a library. When sampling a library with a high Edit Fraction but poor
283 representation of some or many library members, any phenotyping regime will be continually
284 sampling only a small subset of the desired variation. Alternatively, even representation of the
285 designs with a poor Edit Fraction will lead to over-sampling of the Burden Population. Different
286 phenotyping approaches will be more or less tolerant to deviations in either Edit Fraction or
287 design reagent distribution. Below, we describe metrics that combine these two measures into
288 a score that can be used to quickly assess the utility of a given library.

289 Edit Library Richness

290 When sampling cells or isolates from an engineered cell library, the quantity that is typically
291 most important is the number of unique edits represented in the sample. Borrowing from the
292 ecological literature, the term “richness” is used to refer to the number of unique edits in the
293 sample from the library (Levin et al. 2012). The expected richness μ_m of a sample of m cells or
294 isolates from a library of S edits can be predicted given f , the fraction of cells that contain an
295 edit, and the frequencies p_i of each edit among the edited cells.

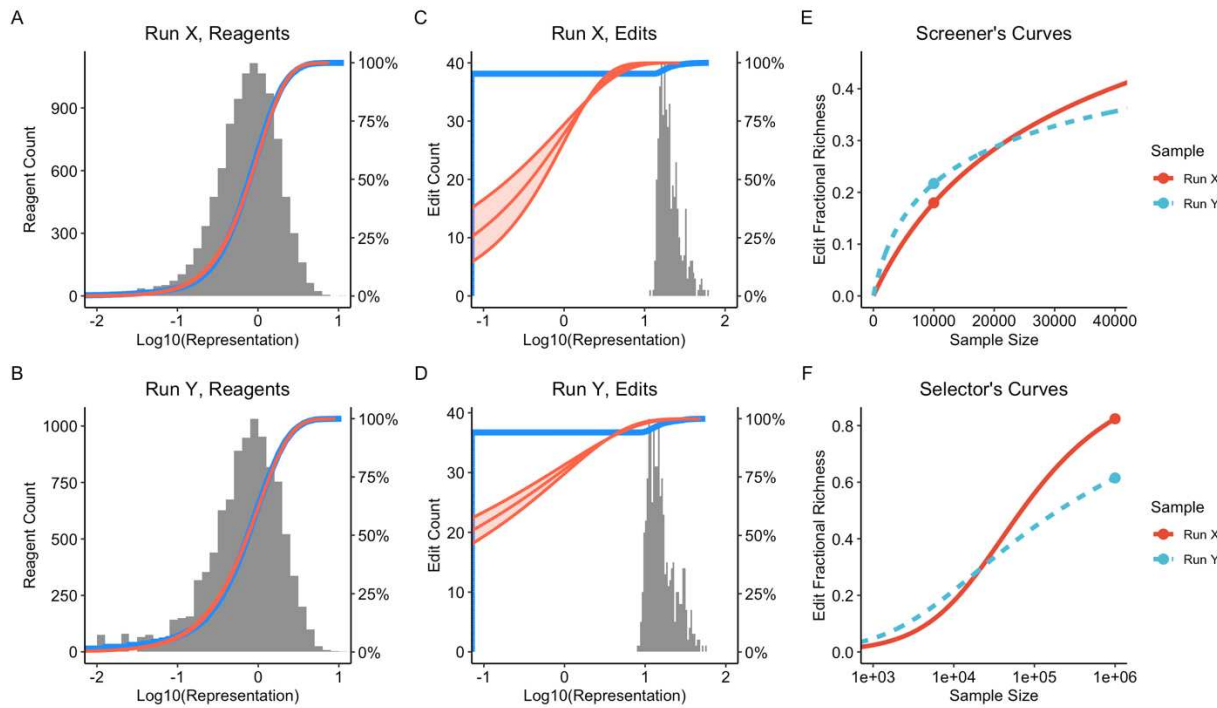
$$296 \mu_m = S - \sum_{i=1}^S (1 - fp_i)^m$$

297 As with other measures, the variance of the sample’s richness can be calculated (supplemental
298 section 1). For some approaches, a variant will need to be observed more than once to provide
299 statistical power for making the genotype-phenotype correlation. In these cases, there is a

300 tractable generalization for when richness is defined in terms of needing at least n
301 observations of each edit (supplemental section 2). This is useful in cases where the dynamic
302 range of quantification relies on a set number of observations of the edit. There is an accurate
303 approximation for the mean and variance of richness, useful both for its mathematical
304 convenience and because it reduces computational complexity from $O(n^2S^2)$ to $O(nS)$
305 (supplemental section 3).

306 Under the assumption that all designs have equal probability of conferring their edits,
307 measurements of reagent frequencies and of the Edit Fraction can be used to predict the
308 richness in a variety of circumstances. It is useful to plot the predicted richness against the
309 number of cell isolates evaluated in a screen or selection, producing a "Screener's Curve"
310 (Figure 3E, 4E and 5C) or a "Selector's Curve" (Figure 3F, 4F and 5D). These plots serve as a
311 guide to set expectations of what fraction of an edit library will be probed in a screen or
312 selection.

313 The appropriate sample size m from which to make richness predictions will depend strongly
314 on the particular situation. In some cases, the cost of phenotyping each sample is high, and the
315 sample size needs to be kept small for practical reasons. In other cases, deep sampling is
316 affordable, and many cells can be sampled. To be able to quantify a library's suitability for
317 screening and selection applications, and to be able to do so in the absence of an estimate of
318 Edit Fraction, two metrics are introduced - the Screener's Score and the Selector's Score. The
319 Screener's Score is defined as the expected Edit Fractional Richness when sampling S times (a
320 1-fold sampling of the library) and with Edit Fraction set to 0.3. The maximum possible value
321 for the Screener's Score is $1 - e^{-0.3}$ or 0.26 (supplemental section 4). The Selector's Score is
322 defined as the expected Edit Fractional Richness when sampling 10^6 times (a reasonable
323 number of input cells for a selection protocol), with the same Edit Fraction of 0.3. The
324 Selector's Score can take on any value in the range [0,1]. These scores are intended to be
325 general measures and more detailed information concerning the Edit Fraction would make this
326 estimate more accurate. Figure 4 illustrates how these concepts can be used to quantitatively
327 assess different libraries for screening and selection purposes.



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329

330 **Figure 4:** Comparative evaluation of two runs of a 10,000 member *E. coli* library, the runs are named X and Y. **A**
331 and **B**: histogram and CDF (blue) of design frequencies as determined by deep amplicon sequencing of the
332 reagents. The red curves correspond to beta distributions fit by Maximum Likelihood Estimation (MLE). The
333 estimates for Reagent CV are 0.79 and 0.90 for runs X and Y respectively. **C** and **D**: histogram and CDF (blue) of
334 genomic edit frequencies as determined by pWGS. The red curves are beta distributions fit by MLE, the shaded
335 area spans the 95% confidence interval for the edit CV estimates. The estimated edit CVs are 1.54 and 2.48 for
336 runs X and Y respectively. The pWGS assay is a shallow sampling of edits, with an estimated sampling depth of
337 488 and 724 in runs X and Y respectively, which is very small compared to the library size of 10,000. The pWGS
338 assay also enables estimation of Edit Fraction, the estimates are 0.25 and 0.57 for runs X and Y. Run X has a lower
339 Edit Fraction but also a lower edit CV compared to run Y, so determination of which run is better to use in
340 downstream applications will depend on the situation. **E**: Screener's curves plotting predicted Edit Fractional
341 Richness against sample size for the two runs. The points on the curves correspond to the Screener's Scores using
342 the estimated Edit Fractions. For a screen of 20,000 or fewer isolates (twice the library size), run Y is predicted to
343 yield greater Edit Fractional Richness, with its larger Edit Fraction making up for its larger edit CV. **F**: Selector's
344 curves, like E but with the x-axis expanded to span a range more typical for a selection application. The points on
345 the curves denote the Selector's Scores, the predicted Edit Fractional Richness when sampling 10^6 cells. The lower
346 edit CV of run X makes it a better choice for a selection application, despite it having less than half the Edit
347 Fraction of run Y.

348 When an estimate of Edit Fraction is available to complement the estimates of design reagent
349 frequencies, the Empirical Screener's Score and Empirical Selector's Score can be evaluated in
350 a similar manner, replacing the fixed assumption of 0.3 Edit Fraction with the empirically
351 determined estimate (Figure 3D). These curves aid in understanding the best phenotypic
352 approaches to take given various library characteristics and experimental goals.

353 Maximizing Library Richness

354 The four variables appearing in the expression for richness motivate different approaches for
355 maximizing the richness of a sample, though in practical applications some of the approaches
356 may be inaccessible (supplemental section 4). The first approach is the obvious one of
357 increasing the sample size – the larger the sample, the greater the richness. The second
358 approach is to increase the probability f that a design reagent confers an edit - something that
359 can be achieved, for example, by improving models for gRNA design. The third approach is to
360 increase the library size S . Lastly, the edit CV has a direct impact, with more evenly distributed
361 libraries resulting in greater richness.

362 For a sample of size m from a library of size S with Edit Fraction f , the maximum richness
363 possible is $S \left(1 - e^{-\frac{mf}{S}}\right)$, attained for a perfectly even library where all design reagent
364 frequencies are equal to $1/S$ (supplemental section 4).

365 Predicting Library Richness

366 The predictor of library richness introduced above requires an estimate of the frequency of
367 every member of the library. In some situations where deep sampling from the library is
368 feasible it will be possible to get good frequency estimates, but for large libraries it is often
369 desirable to be able to predict richness from shallow sampling, to help guide decisions about
370 when to proceed with deep sampling.

371 The problem of predicting future richness from an initial sampling is commonly referred to as
372 the unknown species problem in ecology, one of the earliest solutions was the Good-Toulmin
373 estimator (Good and Toulmin 1956). The Good-Toulmin estimator is a nonparametric
374 approach which works well for predicting up to twice the depth as available in the initial
375 sample but beyond that it becomes unstable. An improved nonparametric approach
376 introduced the use of rational function approximations to produce stable estimates at
377 sampling depths orders of magnitude larger than the initial sample (Daley and Smith 2013) and
378 subsequent work extended the approach to predict richness when requiring more than one
379 observation of each library member (<https://arxiv.org/pdf/1607.02804.pdf>).

380 An alternative approach is to assume a parametric model to describe the library frequencies. A
381 benefit of the parametric approach is that it can produce good estimates from shallow
382 sampling, as long as the model is a good fit for the underlying data. The beta distribution,
383 described by two parameters, is a natural model to consider and one that is often an excellent
384 fit for genome editing libraries (Figures 3, 4, S4). When using a model for design reagent
385 frequencies where the total library size is known, a constraint is needed to ensure that the
386 frequencies sum to 1, or equivalently, to ensure their mean is $1/S$; as a result, there is only one
387 free parameter. It turns out to be convenient to use the CV as the free parameter. When design
388 reagent frequencies follow a beta distribution, there is a closed-form solution available for the
389 expected Edit Fractional Richness, where Edit Fractional Richness is defined as the Edit
390 Richness scaled by the library size (supplemental section 6). For a beta model, Edit Fractional
391 Richness depends on only two parameters - the CV of the design reagent frequencies c , and

392 the sampling fraction F , defined as mf/S , which can be thought of as the effective fraction of
393 the library that is profiled in a sampling of m cells (Figure 5). The expected Edit Fractional
394 Richness $\mu_{m,n}$ where at least n observations of an edit are required, is well approximated as

$$395 \quad \frac{\mu_{m,n}}{S} = 1 - \sum_{k=0}^{n-1} \left(\frac{1}{1+Fc^2} \right)^{\frac{1}{c^2}} \left(1 - \frac{1}{1+Fc^2} \right)^k \binom{1/c^2 + k - 1}{k}$$

396 Consistent with the expression for Edit Fractional Richness, the number of observations of
397 each edit in the sample follows a negative binomial distribution with failure probability set to
398 $1/(1+Fc^2)$ and failure count set to $1/c^2$. There is also an expression for the variance of
399 richness (supplemental section 6). These expressions can be used with the delta method to
400 account for uncertainty in the estimates of CV and Edit Fraction, enabling construction of
401 confidence intervals for Screener's and Selector's curves.

402 Supplemental section 9.3 presents a comparison of parametric and nonparametric estimators
403 of richness on some empirical data.

404 Applying These Estimates and Metrics

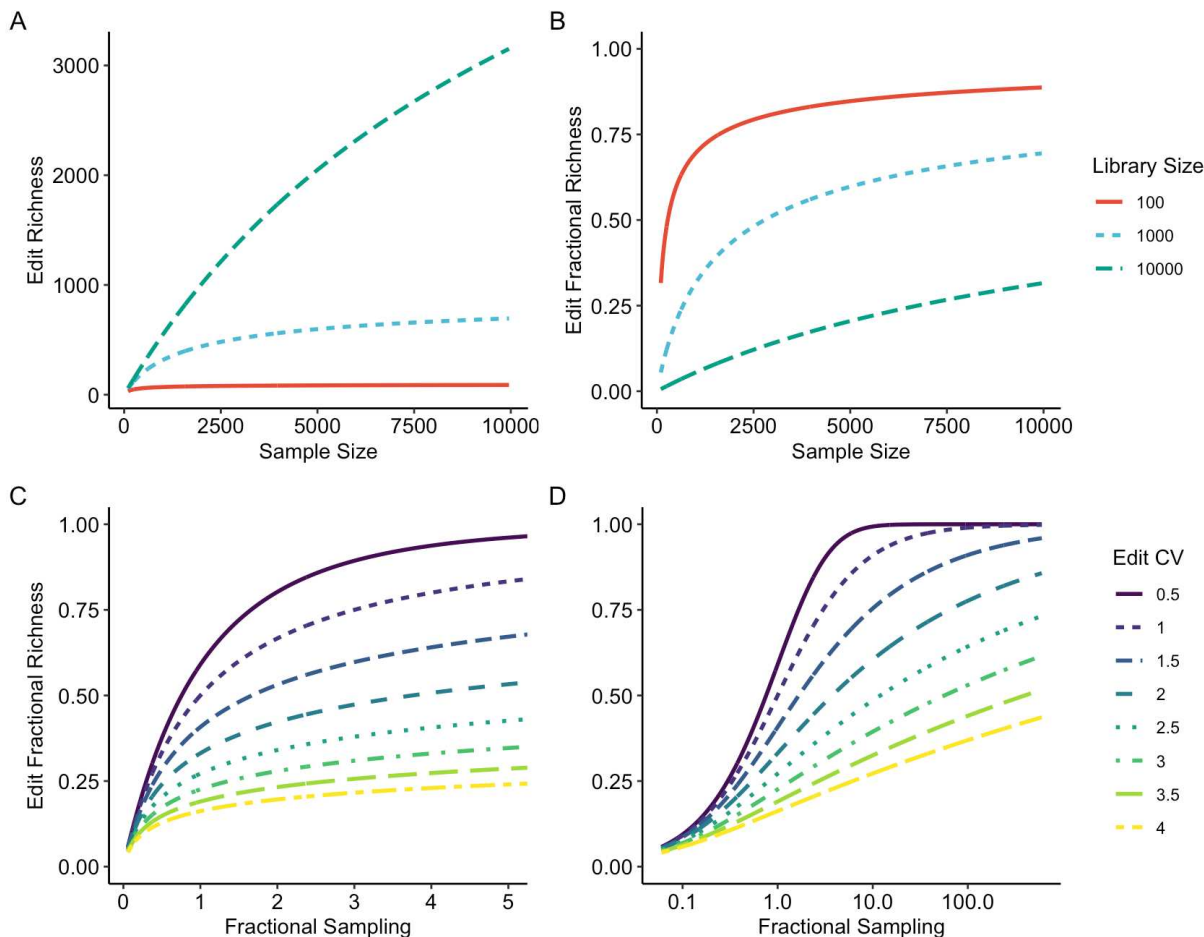
405 Massively parallel genome engineered libraries provide rich diversity for a variety of
406 applications. The framework described above can be applied to experimental design, library
407 evaluation and comparing results from different approaches. Below, we describe using this
408 framework to evaluate libraries for utility in either forward engineering or genome discovery
409 applications.

410 Forward Engineering Experiments

411 Forward Engineering of biological systems relies on effective methods to generate beneficial
412 genetic diversity to provide the fuel for evolutionary optimization (Fox and Giver 2011).
413 Screening of isolated genetic variants that drive improved phenotypes becomes an exercise in
414 maximizing richness while managing sampling depth. As noted above, increasing the library
415 size is a way of maximizing richness. Shallow screening of large libraries has proven to be an
416 efficient way to maximize the beneficial diversity rate, as most of the genotypes observed are
417 likely to be unique at lower sampling depth (Alvizo et al. 2014).

418 The effects of library size, Edit Fraction and Edit CV for screening experiments is shown in
419 Figure 5. The discovery rates for libraries with differing Edit CVs are plotted, showing the effect
420 to which libraries with higher variance in the distribution of the population forces much deeper
421 screening in order to continue to observe unique variants. For forward engineers seeking
422 simply to maximize the discovery rate of beneficial diversity, a shallow sampling from a large
423 library is a particularly effective approach. For shallow sampling, the impact of Edit CV on Edit
424 Fractional Richness is modest, as few of the sampled variants are duplicates. Conversely, with

425 deeper sampling (where researchers desire observing the highest fraction of designs) the
426 effect of a larger Edit CV becomes more limiting. As the Edit CV of the library population
427 increases, it becomes increasingly difficult to observe those designs present at the lower
428 frequencies in the population. Edit Fraction has a linear effect on screening outcomes - halving
429 the edit rate while doubling the sample size results in no net change in expected richness.



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Figure 5: Exploration of richness under the assumption that edit frequencies follow a beta distribution. **A:** Edit Richness for different library sizes, assuming an Edit CV of 1.5 and an Edit Fraction of 0.6. **B:** Edit Fractional richness for the same scenarios as used in A. **C:** Screener's curves, showing Edit Fractional Richness as a function of Fractional Sampling, with different values for edit CV. Fractional Sampling is defined as the product of sampling depth (the number of cells or isolates sampled) and Edit Fraction divided by the library size. Fractional Sampling and Edit CV are all that is required to predict Edit Fractional Richness under the beta assumption. **D:** Selector's curves, which are the same figure as C with a log-scale x-axis to enable prediction of Edit Fractional Richness with the deep sampling that is typically used for a selection experiment

441

442 **Genome Discovery**

443 While forward engineering is driven largely by the identification of desired phenotypes,
444 genome discovery is often focused on testing specific variants to determine if they drive a
445 phenotype. In this case, a researcher may be more interested in observing all, or most, variants
446 within a library several times in order to develop robust hypotheses around genotype-
447 phenotype correlations. In this case, maximizing library coverage may be the most beneficial
448 approach. When employing an isolate phenotyping approach, this will likely require minimizing
449 library size so that the edits can be sampled multiple times. When employing a selection
450 strategy, increasing library size may be appropriate if Edit CV is held low. This will be driven by
451 the number of times a researcher wants to observe edits in the left tail of the distribution. For
452 more precise genotype-phenotype correlations, assessing more libraries containing a smaller
453 number of edits will likely yield more robust results. Strategic use of the Screener's and
454 Selector's Scores in planning experiments can maximize outcomes by informing sampling
455 depth needed to robustly associate genotypic changes with phenotypes of interest.

456 **Conclusions**

457 As technology continues to improve, the ability to create larger libraries with precise edits will
458 become commonplace. To date, no common standards exist for describing and evaluating cell
459 libraries. This makes comparing libraries produced using different approaches challenging.
460 Perhaps more importantly, a lack of common standards makes planning experiments and
461 evaluating libraries as fit-for-purpose challenging, and these measures differ from lab to lab.
462 Here, we have proposed a framework for evaluating massively parallel libraries of genome
463 engineered cells. We have provided precise definitions around what constitutes an edit. While
464 previous groups have often looked at the reagents within a complex cell library, we
465 demonstrate the value of measuring the fraction of cells within the pool that actually contain
466 an edit and we introduce methodology to directly profile the distribution of edit frequencies.
467 This provides for robust characterization of library properties without needing to employ
468 expensive and labor-intensive approaches to understand editing at every target site. We
469 introduce the concept of edit library richness to more fully describe a library quantitatively, as
470 the Edit Fraction is insufficient to fully characterize a library's quality. When generating a
471 complex editing library, it is valuable to have a large percentage of the designs represented in
472 the final population, not just have a large Edit Fraction that all contain the same, or a few edits.
473 We also provide models and methods that allow predictions of library quality when some key
474 metrics, typically Edit Fraction, are not available. Development of a robust framework for
475 evaluating complex cell libraries will be necessary to inform which approaches will be useful for
476 phenotypic analysis of a library. Establishment of common methods will facilitate comparing
477 libraries created from various methods. While we have focused on libraries of precise genome
478 edits, the metrics, models and methods proposed here can be applied to any type of library
479 conforming to the general statistical assumptions introduced.

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481

482 **Supplemental Materials**

483 Mathematical derivations and deeper discussion of the metrics are available in the attached
484 Supplement. Code and data used for analyses can be accessed online at
485 https://github.com/InscriptaLabs/cell_lib_eval_paper

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489 **Author Contributions**

490 CGA, SC, CD, ME, SF, RF, MWG, ADG, MSG, ALH, PH, TH, SJ, CJ, KJ, NK, SL, BL, TMS, JS, ECS,
491 CAS, MHS, ST and TT developed the general framework for characterizing a pool of edited
492 cells and created the novel associated metrics. EA, SA, ME, GG, NK, BL, FP, CDS, TRS, and KW
493 used the Onyx™ platform to generate the pooled editing data used in this manuscript. MB,
494 DMC, SC, ME, RF, MSG, TH, BL, JCJR, TMS, CAS, and MHS wrote and/or reviewed the
495 manuscript and associated figures. JB, SC, TH, SL, TMS, CAS, MHS, and ST derived the
496 mathematical results in the main text and supplement, and implemented them in
497 bioinformatics pipelines.

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