

¹ gimap: An R Package for Genetic Interaction Mapping in Dual-Target CRISPR Screens

³ **Candace Savonen**  ^{1,2}, **Phoebe Parrish**  ¹, **Kate Isaac**  ¹, **Daniel Grosø** ¹,
⁴ **Marissa Fujimoto** ¹, **Siobhan O'Brien**  ¹, and **Alice Berger**  ¹

⁵ **1** Fred Hutchinson Cancer Center, United States **2** Synthesize Bio, United States

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Software

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⁶ Summary

⁷ The gimap (Genetic Interaction MAPping) R package addresses a fundamental challenge in genomic research: the difficulty of understanding combinatorial interactions among genes. ⁸ Gene redundancy makes traditional single-gene knockout methods ineffective for identifying therapeutic targets, as backup genes can mask the effects when a single gene is disabled. ⁹ gimap offers a solution by providing a comprehensive framework for analyzing dual-target CRISPR screening data, where two genes are simultaneously disabled to reveal their backup relationships. This software implements the methods used by Parrish et al. (2021). The package processes raw count data through a multi-step pipeline that includes normalization, calculation of expected and observed CRISPR scores, computation of genetic interaction scores, and statistical analysis to identify significant interactions. Unlike general tools, gimap is specifically tailored for paired guide CRISPR data with built-in quality control reporting and visualization tools. The package makes best practices the default options and is available on GitHub with comprehensive documentation to support the research community in extracting meaningful insights from complex genetic screening experiments.

²¹ Statement of Need

²² When multiple genes have the same function, a common result of evolutionary processes, it becomes challenging to isolate their true functions. This redundancy means that many possible therapeutic targets are missed by traditional methods that disable just one gene at a time (De Kegel & Ryan, 2019; Parrish et al., 2021). A more complementary approach involves disabling two genes simultaneously to reveal these backup relationships (Thompson et al., 2021).

²⁷ Recent advances in CRISPR technology now allow researchers to knock out gene pairs at once, offering a powerful solution to this problem (Gonatopoulos-Pournatzis et al., 2020). Although software solutions exist for single knockout CRISPR, such as MAGeCK, there is no standardized software solution for paired gene CRISPR studies (Li et al., 2014).

³¹ The R package, called gimap (Genetic Interaction MAPping), was developed specifically for analyzing these dual-target CRISPR experiments. It helps researchers identify important relationships between genes, such as when two genes work together or when disabling both creates a dramatic effect that wouldn't occur by disabling either one alone.

³⁵ gimap is specifically tailored to handle the unique characteristics of paired guide CRISPR data, including the distinction between single-targeting and double-targeting constructs and the need to account for differential double-strand break effects. The package seamlessly integrates with data generated using a specialized pgPEN library but can be adapted for most paired guide CRISPR screening approaches (Parrish et al., 2021).

40 Implementation

41 gimap addresses this need by providing a comprehensive analytical framework for dual-target
 42 CRISPR screening data. The package performs several critical functions: (1) normalization of
 43 read count data to account for variable sequencing depth and technical biases, (2) calculation of
 44 CRISPR scores that reflect the effect of gene knockouts on cell proliferation, (3) determination
 45 of expected CRISPR scores for gene pairs based on single-gene effects, (4) computation of
 46 genetic interaction scores that quantify deviations from expected effects, and (5) statistical
 47 analysis to identify significant interactions.

48 Overall design philosophy

49 In order to ensure usability for the research community we built gimap using the following
 50 design philosophy.

- 51 1. Making best practices as default options and including warning messages for when
 52 alternative options are chosen (e.g. if filtering has not been applied).
- 53 2. Using elements from familiar packages such as fastqc reports (our `run_qc()` function
 54 creates such a report) (*Babraham Bioinformatics - FastQC A Quality Control Tool for*
55 High Throughput Sequence Data, n.d.).
- 56 3. Trying to document and inform users of the statistics and decisions that have been made
 57 by the software clearly.

58 gimap data handling

59 gimap implements a multi-step analysis pipeline:

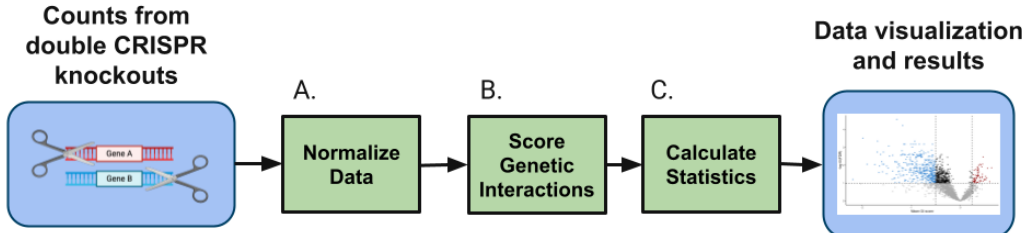


Figure 1: gimap workflow completes 3 main steps. Part A, B, and C of the figure show the major steps of the workflow which are to normalize the data through a multi step process, score genetic interactions based on the expected versus observed CRISPR scores, and finally to calculate statistics to identify statistically significant genetic interactions.

60 1. **Normalize Data:** Raw count data is transformed into log2 counts per million (CPM)
 61 and adjusted by subtracting pre-treatment values to obtain log2 fold changes. These
 62 are further normalized based on the distribution of negative (e.g. safe-targeting or non-
 63 targeting controls) and positive controls (pgRNAs targeting known essential genes). This
 64 scaling normalization is analogous to the normalization methods employed by the Cancer
 65 Dependency Map (depmap.org) (Arafeh et al., 2025; DepMap, Broad, 2025).

66 a. *Log2 Counts Per Million (CPM) Transformation:*

- 67 ▪ Let $C_{i,j}$ be the raw count for gene i in sample j
- 68 ▪ Let N_j be the total number of counts in sample j

$$L_{i,j} = \log_2 \left(\frac{C_{i,j} \times 10^6}{N_j} + 1 \right)$$

69 (The +1 is often included to avoid log(0) issues)

70 b. *Adjustment by Pre-treatment Values:*

- 71 ▪ Let $L_{i,j}^{post}$ be the log2 CPM value post-treatment
- 72 ▪ Let $L_{i,j}^{pre}$ be the log2 CPM value pre-treatment

$$LFC_{i,j} = L_{i,j}^{post} - L_{i,j}^{pre}$$

73 c. *Normalization Based on Controls:*

- 74 ▪ Let $LFC_{i,j}$ be the log2 fold change calculated above
- 75 ▪ Let μ_{neg} and σ_{neg} be the mean and standard deviation of negative controls (safe-targeting or non-targeting)
- 76 ▪ Let μ_{pos} and σ_{pos} be the mean and standard deviation of positive controls (pgRNAs targeting essential genes)

$$Z_{i,j} = \frac{LFC_{i,j} - \mu_{neg}}{\mu_{neg} - \mu_{pos}}$$

79 Or alternatively, using a more complex normalization that accounts for the distributions of
80 both control types:

$$Z_{i,j} = \frac{LFC_{i,j} - \mu_{neg}}{\sigma_{neg}} \times \frac{\sigma_{pos}}{\mu_{neg} - \mu_{pos}}$$

81 This equation represents the transformation from raw count data to normalized log2 fold
82 changes, calibrated against both negative and positive control distributions.

83 2. **Score Genetic Interactions:** The goal of this step is to quantify deviations from expected
84 additive effects when two genes are simultaneously targeted, which allows us to identify
85 true genetic interactions beyond what would be predicted from single-gene effects alone.

86 We model a control distribution for non-interacting genes by assuming that in the absence of
87 genetic interactions, the effect of simultaneously targeting two genes should be additive (i.e.,
88 the sum of their individual effects). We further assume that the observed genetic interaction
89 (GI) score is a linear transformation of the expected GI score plus a consistent error term
90 sampled from an approximately normal distribution. This error distribution is shared across all
91 genes and is homoscedastic (constant variance) across expected GI scores, allowing us to use
92 linear modeling approaches to account for systematic biases.

93 For double-targeting constructs, expected CRISPR scores are calculated as the sum of the
94 corresponding single-targeting scores. For single-targeting constructs, the expected score
95 combines the single-target effect with the mean effect of control constructs. Interaction scores
96 represent the difference between observed and expected CRISPR scores, adjusted using a linear
97 model to account for systematic biases.

98 For double-targeting constructs:

- 99 ▪ Let $S_{i,j}^{obs}$ be the observed CRISPR score for a construct targeting genes i and j
- 100 ▪ Let S_i be the single-targeting score for gene i
- 101 ▪ Let S_j be the single-targeting score for gene j

102 The expected score for a double-targeting construct is:

$$S_{i,j}^{exp} = S_i + S_j$$

103 For single-targeting constructs:

- 104 ▪ Let S_i^{obs} be the observed CRISPR score for a construct targeting gene i

105 ▪ Let $\mu_{control}$ be the mean effect of control constructs

106 The expected score for a single-targeting construct is:

$$S_i^{exp} = S_i + \mu_{control}$$

107 The interaction score calculation, with adjustment for systematic biases:

- 108 ▪ Let $I_{i,j}$ be the interaction score for genes i and j
- 109 ▪ Let $S_{i,j}^{obs}$ be the observed score
- 110 ▪ Let $S_{i,j}^{exp}$ be the expected score
- 111 ▪ Let β_0 and β_1 be the intercept and slope from a linear regression of observed vs expected scores

113 The interaction score calculation:

$$I_{i,j} = S_{i,j}^{obs} - (\beta_0 + \beta_1 \cdot S_{i,j}^{exp})$$

114 Where the genetic interaction score is the difference between the observed score and the linear model prediction based on the expected score, accounting for systematic deviations between observed and expected values.

117 **3. Calculate Statistics:** T-tests compare the distribution of double-targeting genetic interaction scores for one pair against the background distribution of single-targeting scores, with false discovery rate correction for multiple hypothesis testing.

- 120 ▪ S_{double} as the set of double-targeting genetic interaction scores
- 121 ▪ S_{single} as the set of single-targeting scores (background distribution)
- 122 ▪ μ_{double} as the mean of double-targeting scores
- 123 ▪ μ_{single} as the mean of single-targeting scores
- 124 ▪ σ_{double} as the standard deviation of double-targeting scores
- 125 ▪ σ_{single} as the standard deviation of single-targeting scores
- 126 ▪ n_{double} and n_{single} as the sample size of the paired guides

127 The t-test statistic would be:

$$t = \frac{\mu_{double} - \mu_{single}}{\sqrt{\frac{\sigma_{double}^2}{n_{double}} + \frac{\sigma_{single}^2}{n_{single}}}}$$

128 For each comparison, we calculate a p-value from this t-statistic.

129 Then, to account for multiple hypothesis testing, we apply false discovery rate (FDR) correction 130 using Benjamini Hochberg procedure ([Benjamini & Hochberg, 1995](#)):

- 131 1. Order all p-values: $p_{(1)} \leq p_{(2)} \leq \dots \leq p_{(m)}$
- 132 2. For a given FDR threshold α (e.g., 0.05), find the largest k such that:

$$p_{(k)} \leq \frac{k}{m} \cdot \alpha$$

133 3. Reject the null hypothesis for all tests with p-values $\leq p_{(k)}$

134 The package also provides comprehensive visualization tools including volcano plots to highlight 135 significant genetic interactions and detailed result tables for further analysis.

136 Use Cases

137 gimap has been successfully used to identify synthetic lethal interactions among paralog genes
138 in cancer cell lines, revealing potential therapeutic targets that single-gene approaches have
139 missed. The package accommodates various experimental designs, including time-course
140 studies and treatment comparisons, offering flexibility for diverse research questions.

141 *Example applications include:*

- 142 ▪ Identification of genes that provide functional redundancy in critical cellular pathways
- 143 ▪ Discovery of context-dependent genetic interactions that emerge under specific conditions
- 144 or treatments
- 145 ▪ Systematic mapping of gene networks based on functional interactions rather than
- 146 physical associations

147 Conclusion

148 gimap provides a robust, accessible framework for analyzing paired guide CRISPR screening
149 data and identifying genetic interactions with potential biological and therapeutic significance.
150 By streamlining the computational workflow from raw counts to statistically rigorous
151 interaction scores, gimap enables researchers to efficiently extract meaningful insights
152 from complex genetic screening experiments. The package is available on GitHub
153 (<https://github.com/FredHutch/gimap>) with comprehensive documentation and tutorials to
154 facilitate adoption by the research community.

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159 Related Publications

160 This software implements methods originally described in Parrish et al. (2021). The current
161 submission focuses on the software implementation and its accessibility to the research
162 community.

163 Conflict of Interest

164 The authors declare no conflicts of interest.

165 AI Usage Disclosure

166 No generative AI tools were used in the development of this software. AI was used for minor
167 polishing in the preparation of this manuscript.

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