



Analysis of combinatorial CRISPR screens with the Orthrus scoring pipeline

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The continued improvement of combinatorial CRISPR screening platforms necessitates the development of new computational pipelines for scoring combinatorial screening data. Unlike for single-guide RNA (sgRNA) pooled screening platforms, combinatorial scoring for multiplexed systems is confounded by guide design parameters such as the number of gRNAs per construct, the position of gRNAs along constructs, and additional features that may impact gRNA expression, processing or capture. In this protocol we describe Orthrus, an R package for processing, scoring and analyzing combinatorial CRISPR screening data that addresses these challenges. This protocol walks through the application of Orthrus to previously published combinatorial screening data from the CHyMERa experimental system, a platform we recently developed that pairs Cas9 with Cas12a gRNAs and enables programmed targeting of multiple genomic sites. We demonstrate Orthrus' features for screen quality assessment and two distinct scoring modes for dual guide RNAs (dgRNAs) that target the same gene twice or dgRNAs that target two different genes. Running Orthrus requires basic R programming experience, ~5–10 min of computational time and 15–60 min total.

Introduction

Genetic tools that systematically map genetic interactions (GIs) are powerful hypothesis-generating technologies for both basic research as well as drug discovery. GIs are defined as the phenomenon by which combinatorial mutations in multiple genes result in phenotypic effects that are greater or less than expected, given the phenotypes of the individual mutants¹. Pioneering work in yeast resulted in a nearly complete map of yeast pairwise GIs, which illuminated the functions of uncharacterized genes, revealed functional connections between pathways², and enabled the systematic characterization of compound mode-of-action^{3,4}.

The study of human GIs at scale has recently been made possible in human cell culture systems with the development of clustered regularly interspaced short palindromic repeats (CRISPR)/Cas-enabled genetic screens^{5–9}. These experiments induce mutations across any number of human genes in pooled cell culture and typically work by targeting each gene with a small number of Cas9 guides that are individually expressed in cells. Pooled CRISPR screens can be performed in various genetic backgrounds to identify different effects, such as in cancer cell lines to uncover cancer-specific genetic dependencies¹⁰ or in isogenic cell lines to map GIs¹¹. Combinatorial CRISPR screening platforms were also developed to directly identify GIs by knocking multiple genes out through the expression of multiple guides within the same cell. These platforms function by pairing individual Cas9 guide RNAs (gRNAs) with each other, combining orthologous Cas9 gRNAs, or by multiplexing multiple Cas12a gRNAs^{11–19}. We recently developed a novel combinatorial screening platform named Cas Hybrid for Multiplexed Editing and Screening Applications (CHyMERa) that instead targets genes with hybrid guide RNAs (hgRNAs) composed of Cas9 guides fused with one or more additional Cas12a guides, which can subsequently be processed into individual guides by Cas12a's RNA-processing activity¹².

Although accurate scoring of GI data requires precise quantitative measurements^{2,11}, there is a lack of computational tools that enable quantitative GI scoring for combinatorial experimental designs.

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To date, most combinatorial CRISPR screening studies have scored data by taking simple or weighted averages of log fold-change (LFC) values between start and end read counts for dual guide RNAs (dgRNAs) targeting specific gene pairs with additional corrections based on control guides or the phenotypes of similar dgRNAs^{13–16}. While this approach may be suitable for noncombinatorial screens, different Cas enzymes¹², as well as the position of gRNAs on combinatorial guide constructs¹⁶, may strongly affect guide efficiency. This necessitates the development of combinatorial scoring methods that take these effects into account.

To address issues arising from the combinatorial nature of data output from CHyMERa and other platforms, we developed a novel scoring method named Orthrus for combinatorial CRISPR screening data. The key feature of Orthrus is that it takes orientation—whether gene A is targeted by a gRNA in position 1 and gene B is targeted by a gRNA in position 2, or vice versa—into account during scoring, which is necessary to consider for Cas9 and Cas12a guides that can cause different fitness effects even when targeting the same gene. This scoring method is bundled in a well-documented R package with a variety of features to simplify combinatorial data processing, quality control (QC) and analysis²⁰, and is downloadable at <https://github.com/csbio/Orthrus>. Here we present the recommended Orthrus workflow, demonstrate its key features on previously published data from two separate combinatorial screening experiments, discuss important considerations for performing QC analyses of screening data, and detail expected results from the application of Orthrus to combinatorial screening data.

Development of the protocol

The Orthrus package implements the GI scoring schema presented in Gonatopoulos-Pournatzis et al.¹², although with several key improvements that include more sensitive scoring, a variety of QC plots and metrics, and a new user interface. Most broadly, Orthrus presents a consistent user interface for scoring any kind of combinatorial screening data stored in a delimited text file. The code that was used to score data in Gonatopoulos-Pournatzis et al.¹² does not generalize to other experiments or combinatorial screening platforms. This new user interface also provides push-button functions that automatically generate a variety of QC plots and metrics useful for assessing the quality of most types of screening data. In addition, the Orthrus package implements several features absent from the previously published scoring code. These features improve the sensitivity of scoring and user confidence in resulting hits and include guide filtering based on plasmid pool or early timepoint read counts, loess normalization of residual effects, and moderated *t*-testing in addition to the original Wilcoxon rank-sum testing. Lastly, given properly formatted input files, the Orthrus package provides several different scoring interfaces that enable partial or complete automation of the scoring process.

Comparison with other methods

We are aware of five existing scoring methods for combinatorial CRISPR screens: Δ LFC, GIMap, π -score, Norm-GI and GEMINI^{13–15,17,21}. All scoring methods compare the null model of multiplicative single-gene effects, typically derived from intragenic guides paired with intergenic controls (exonic–intergenic guides), to the observed effects of exonic–exonic guides. The simplest proposed method is Δ LFC¹³, which directly performs this comparison while deriving empirical FDRs from permuted data. GIMap, Norm-GI and π -score all perform the same comparison with some additional corrections. GIMap normalizes exonic–intergenic guides to a quadratic fit and residual effects to negative control guide effects¹⁷. Norm-GI normalizes residual effects to control guide phenotypes, as well as to the phenotypes of similar guides using a moving average across bins¹⁵. The π -score weights exonic–exonic guides, giving higher preference to guides with stronger phenotypes¹⁴. GEMINI is a Bayesian approach that explicitly models sample-independent and sample-dependent effects and uses coordinate ascent variational inference to update the posterior distributions of these effects²¹. Of these five scoring methods, only GEMINI exists in a generalized, runnable form as an R package.

Orthrus primarily differs from GEMINI in how it accounts for orientation, whether it computes effects relative to control genes, the types of guides it is designed to score, and the number of auxiliary functions it offers. GEMINI does not account for guide orientation, which is an important consideration for CHyMERa screening data. Furthermore, GEMINI computes *P*-values and false discovery rates (FDRs) relative to a specified set of negative control genes, which may be appropriate for whole-genome screens but is not necessarily appropriate for screens performed with specialized libraries. GEMINI similarly computes effect size based on a set of positive control genes, which may or may not be available depending on the library design and interrogated phenotype. Orthrus, on the

other hand, does not rely on negative or positive control genes during scoring, and can identify hits even for moderate phenotypes owing to the use of moderated *t*-testing. The combination of these choices allows Orthrus to score multiple different types of guides, as it can directly score combinatorial guides against single-targeting controls, as well as chemogenetic screens against nontreated controls or single-targeting guides. Lastly, unlike GEMINI, Orthrus presents the user with a host of data processing, QC and plotting functions to assess screen quality.

Applications of the method

The Orthrus package is generally applicable to a variety of combinatorial CRISPR screening settings. While Orthrus' orientation-based filtering is designed to minimize false positives or discrepancies due to guide orientation, this scoring approach requires double the amount of hypothesis testing per gene pair and may be overly conservative in some settings. For applications such as CHyMERa screens, this conservative approach should be adopted (see Step 15 of Procedure 2). For multiplexing screens with single nucleases (either Cas9 or Cas12a), the user should instead reduce the amount of hypothesis tests performed per gene pair by ignoring orientation-specific effects (see Step 4 of Procedure 3). In addition to flexible scoring functions, Orthrus provides an extensive selection of data processing and QC functions that are applicable to any combinatorial screening dataset and can be applied irrespective of downstream scoring methods.

Limitations

The Orthrus package currently supports the analysis of data from negative selection CRISPR screens with combinatorial guide libraries targeting (A) a single gene of interest twice, (B) two different genes and (C) a gene paired with a control region. It has been tested on CHyMERa data for all of these guide types as well as for combinatorial-targeting guides from a different multiplexed Cas12a platform, as described in the Procedures below. Genes of interest could be targeted in either exonic or intronic regions depending on the screens' experimental design, although Orthrus has only been tested to score the effects of gRNAs targeting exonic regions. While Orthrus is currently the only scoring package that accounts for orientation-specific effects, this feature can be disabled to increase statistical power for screens where guide orientation is less relevant. Thus, Orthrus is flexible enough to score any combinatorial data for all guide types listed above. While Orthrus is not designed to score combinatorial positive selection or drug rescue experiments, a combination of stringent guide filtering and scoring with Wilcoxon rank-sum hypothesis testing may be appropriate for these types of screens. However, further testing on genome-scale combinatorial positive selection screens is necessary to assess Orthrus' ability to score this type of data.

Expertise needed to implement the protocol

Basic experience with R programming is required. Experience analyzing CRISPR screen data is recommended, but not required.

Input format

Orthrus requires one mandatory and two optional (but encouraged) tab-separated input files: a mandatory reads file, and the two optional sample and batch files. The *reads* file contains read count information for all screens, and is required for Orthrus to function. Its construction from raw sequencing data is detailed in Procedure 1. The *sample* file maps replicate columns to their matching screens, but unlike similar files required by other packages, it also maps screens to other screens they must be normalized against (e.g., reference time point screens, plasmid pools). The *batch* file maps screens to other screens they must be scored against, such as for drug treatment screens against control screens. While Orthrus provides manual options detailed in Procedure 2 that offer precise input methods for specifying the information contained in sample and batch files, the sample and batch files can drastically reduce the complexity of data processing and scoring—as demonstrated in Procedures 2 and 3—and are thus highly encouraged. In addition to the detailed descriptions below, please refer to this video tutorial for a conceptual overview of these three input files, which is available to view as Supplementary Video 1 or at <https://youtu.be/w8mGWnQ-9Wo>.

Reads file

Like other packages that score CRISPR screening data, Orthrus requires input data formatted as a delimited text file where rows correspond to guides and columns correspond to metadata and raw

Table 1 | Example of a properly formatted reads file for mock data of one gene pair

gene1	gene2	Cas9 guide	Cas12a guide	T0 reads	T18 reads
ARID1A	ARID1B	AATG	TTGC	45	0
ARID1B	ARID1A	CGAC	TATT	54	1
ARID1A	NegControl	AATG	CGCT	70	60
NegControl	ARID1A	GGTA	TATT	82	75
ARID1B	NegControl	CGAC	CGCT	61	87
NegControl	ARID1B	GGTA	TTGC	76	92
ARID1A	None	AATG	TATT	91	42
ARID1B	None	CGAC	TTGC	63	53

Lines 1 and 2 represent combinatorial-targeting guides knocking out both ARID1A and ARID1B simultaneously, lines 3–6 represent single-targeting guides knocking out either ARID1A (lines 3–4) or ARID1B (lines 5–6) paired with a negative control guide in both orientations, and lines 7–8 represent dual-targeting guides cutting either ARID1A (line 7) or ARID1B (line 8) twice. Guide sequences and read counts are mock data for illustrative purposes.

read counts for all screens. However, unlike alternative scoring methods, Orthrus requires two gene label columns whose position reflects the orientation of each guide. In detail, Orthrus requires the following assumptions about the input file's format to be met.

- 1 All guides in the dataset (every row) must have non-empty labels for the genomic regions they target contained in two separate columns.
- 2 These columns must be labeled `gene1` and `gene2`. Although these will typically contain gene symbol annotations, they may contain any identifier desired by the user.
- 3 The `gene1` column must contain gene labels for the first guide in the dataset, and the `gene2` column must contain gene labels for the second guide in the dataset. The definition of 'first' and 'second' is left to the user. For example, the dataset analyzed in Procedure 2 contains genes targeted by Cas9 guide sequences in the `gene1` column and Cas12a guide sequences in the `gene2` column, whereas the dataset analyzed in Procedure 3 contains genes targeted by Cas12a in both columns.
- 4 To enable Orthrus' default scoring mode, each gene label column must map to a respective guide ID column. For example, the `gene1` and `gene2` columns for the dataset analyzed in Procedure 2 map to the columns `Cas9.Guide` and `Cpf1.Guide` (Cas12a was previously named `Cpf1`), respectively. Orthrus does not assume a standardized name for guide ID columns, and instead, the user passes in the name of guide ID columns during the processing step. The first guide ID column name passed in maps to `gene1` and the second maps to `gene2`. For libraries where single-targeting guides do not share guide IDs with combinatorial-targeting guides, the less-sensitive Wilcoxon rank-sum scoring approach implemented in Orthrus does not require this information.
- 5 For 'dual-targeting' guides that target the same gene twice, the `gene1` column must contain the name of the targeted gene and the `gene2` column must contain the string `None`.
- 6 For single-targeting guides paired with a standardized negative control, such as an intergenic region for the dataset analyzed in Procedure 2 or a nonessential gene for the dataset analyzed in Procedure 3, the control must be named `NegControl` in the corresponding gene label column.

While these assumptions may require users to manually preprocess their reads file, this format is a concise way to represent both orientation-specific information as well as guide type information for any combinatorial screen. Although alternative scoring methods also require tab-delimited files with identifying gene labels and guide ID columns, they do not take either orientation or guide type into account (to which assumptions 3–6 above relate). An example format for a reads file containing mock data is presented in Table 1. This table contains the gene name columns `gene1` and `gene2`, the guide ID columns `Cas9 guide` and `Cas12a guide`, and reads for two technical replicates in the columns `T0 reads` and `T18 reads`.

Additionally, because the CHyMERa reads file downloadable at <https://crispr.cabr.utoronto.ca/chymera/index.html> contains several formatting errors, the script used to reformat the CHyMERa data into the version bundled in the Orthrus package download is uploaded to the Zenodo repository (<https://zenodo.org/record/4527616>) and is demonstrated in Step 6 of Procedure 1.

Sample file

The sample file must be a tab-separated file containing exactly three columns named `Screen`, `Replicates` and `NormalizeTo`. The `Screen` column contains unique user-specified labels for

Table 2 | Sample table used in Procedure 2, Step 4B

Screen	Replicates	NormalizeTo
HAP1_T0	HAP1.T0	NA
RPE1_T0	RPE1.T0	NA
HAP1_T12	HAP1.T12A;HAP1.T12B;HAP1.T12C	HAP1_T0
HAP1_T18	HAP1.T18A;HAP1.T18B;HAP1.T18C	HAP1_T0
Torin_T12	HAP1.Torin.T12A;HAP1.Torin.T12B;HAP1.Torin.T12C	HAP1_T0
Torin_T18	HAP1.Torin.T18A;HAP1.Torin.T18B;HAP1.Torin.T18C	HAP1_T0
RPE1_T18	RPE1.T18A;RPE1.T18B;RPE1.T18C	RPE1_T0
RPE1_T24	RPE1.T24A;RPE1.T24B;RPE1.T24C	RPE1_T0

Each row corresponds to a single screen, with the 'Replicates' column containing the names of its technical replicates separated by semicolons and the 'NormalizeTo' column containing the name of a screen to normalize against (e.g., a T0 screen).

each screen. The `Replicates` column contains a list of read column names for all technical replicates corresponding to that screen in the reads file, separated by semicolons. The `NormalizeTo` column contains the name of a different screen in the `Screen` column to normalize the current screen against (e.g., a T0 screen or plasmid pool). The sample file for Procedure 2 is presented in Table 2.

Batch file

The batch file must be a tab-separated file containing exactly two columns named `Screen` and `Control`. The `Screen` column contains labels for screens listed in the `Screen` column of the sample file that the user wants to score against screens listed in the `Control` column. To score combinatorial-targeting guides against null models derived from single-targeting guides, instead specify `combn` in the `Control` column. The batch file for Procedure 2 is presented in Table 3.

Experimental design

Workflow

The recommended workflow for combinatorial CRISPR screen analysis with Orthrus involves the following key stages, as shown in Fig. 1. First, the computational workspace is set up with appropriate variables that describe screen information in a read count matrix. Second, read counts are processed and normalized to sequencing depth as well as to screens from earlier timepoints, if provided. During this stage, a variety of QC plots are output, which include QC plots for raw read counts as well as for normalized LFCs. Based on the results of QC plots and metrics, this stage may be repeated several times with updated filtering and normalization options. Third, LFCs are scored and analyzed in separate ways for guides that target one gene multiple times (dual-targeted scoring) or guides that target different genes with the same construct (combinatorial scoring).

Scoring modes

Orthrus offers two primary scoring modes and one additional scoring mode for different types of guides.

- 1 The *dual-targeting* mode scores guides that target one gene multiple times across different conditions. This mode is suitable for drug screening applications. For example, in previous CHyMERa screens¹², dual-targeting guides cut each gene in two separate exonic regions, and these guides are scored for differences between drug treated (Torin1) and untreated cells.
- 2 As an extension to the dual-targeting scoring mode, the *single-targeting* mode scores single-targeting guides that cut both a single gene and a control region across different conditions. The gene of interest could be cut in an exonic or intronic region, while the control gRNA typically targets an intergenic region or nonessential gene. This mode uses the same interface as the dual-targeting scoring mode with different parameters.
- 3 The *combinatorial-targeting* mode scores guides that target multiple genes. This mode scores GIs. In previous CHyMERa screens, combinatorial-targeting guides cut two different genes in a single exonic region per gene. The effect of double-knockouts induced by these guides is scored against the estimated effect of double-knockouts derived from single-targeting guides that target each gene separately (while paired with guides targeting control regions—see below). For experiments that

Table 3 | Batch table used in Procedure 2, Step 19

Screen	Control
Torin_T12	HAP1_T12
Torin_T18	HAP1_T18
HAP1_T12	combn
HAP1_T18	combn
RPE1_T18	combn
RPE1_T24	combn
Torin_T12	combn
Torin_T18	combn

Each row corresponds to a single screen, with the 'Screen' column containing the names of screens defined in the sample table and the 'Control' column containing either the names of screens to score against for the dual-targeted scoring mode or 'combn' to score them with the combinatorial-targeting mode.

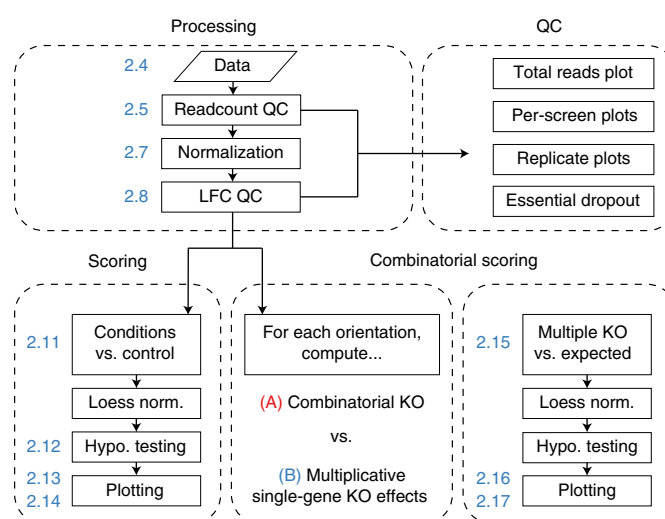


Fig. 1 | The Orthrus scoring workflow. First, combinatorial screening data is processed and depth-normalized before LFCs between late and early timepoints are computed. During this step, QC plots for raw read count data as well as LFC data are output. Second, data are scored, either for condition screens against control screens (e.g., for drug-treated screens against untreated screens), or for the effects of combinatorial knockouts against expected effects derived from single knockouts. Numeric labels, in blue, indicate the corresponding step of Procedure 2.

investigate intronic function, this mode could instead score the effect of dgRNAs that target intronic regions of multiple genes. This mode can also be configured to ignore orientation-specific effects, which increases statistical power for experiments with few observable orientation-specific signatures.

For a given gene pair's set of guides, all of Orthrus' scoring modes compare LFC values for an effect of interest against LFC values for control effects with either moderated *t*-testing or Wilcoxon rank-sum testing. Users may also loess-normalize their residual effects to account for non-normality in their data. For most purposes, we recommend that users run moderated *t*-testing with loess normalization enabled. However, for screens where guide-level residuals cannot be computed because single-targeting controls do not share guide IDs with combinatorial-targeting dgRNAs (see the 'Input format' subsection below), Wilcoxon rank-sum testing without loess normalization must be applied instead. A description of important parameters and the algorithms applied by Orthrus during the scoring process, as well as their typical use cases, is provided in Table 4.

Scoring interfaces

Orthrus provides three different interfaces to the stages listed above in the 'Workflow' subsection: a manual interface that allows fine-grained control over each stage, a batch scoring interface for users to

Table 4 | Select parameters of Orthrus' normalization and scoring functions with their associated algorithm (when applicable) and a description of their typical use case

Parameter	Algorithm	Description	Typical use case
scaling_factor	scaling factor for LFC computation	Scales raw read counts to a default value of 1e6 that forces each screen to the chosen read depth, ensuring comparability across technical replicates. The specific choice of scaling_factor is largely irrelevant	All screens
pseudocount	pseudocount for LFC computation	Adds a pseudocount to each raw read count, by default 1, as required to take log2-normalized read counts. Smaller pseudocounts, e.g., between 1 and 5, are advised to avoid deprioritizing moderate effects	All screens
test	'moderated-t' - moderated t-testing	Computes <i>P</i> -values via empirical Bayes estimate across all residuals fit with separate linear models for each gene pair. Calls limma's eBayes function on its lmFit function applied to residuals with default parameters for both ²⁶	Most screens
test	'rank-sum' - Wilcoxon rank-sum testing	Computes <i>P</i> -values via Wilcoxon rank-sum testing between effect and control LFCs	Combinatorial screens with unpaired controls
loess	'TRUE' - loess normalization with MA transformation	Normalizes by fitting a loess curve with degree 2 and a span of 0.4 to MA-transformed residuals. The MA transformation was originally developed for the analysis of microarray data ²⁷ . Here loess fits a trend for the measured residual value ([double mutant - null model] or [condition - control]) versus the sum of the two values used in computing this residual (e.g., [double mutant + null model] or [condition + control])	Most screens
fdr_method	"BY" - Benjamini-Yekutieli FDR correction	Adjusts <i>P</i> -values with Benjamini-Yekutieli FDR correction	Most screens
fdr_method	'BH' - Benjamini-Hochberg FDR correction	Adjusts <i>P</i> -values with Benjamini-Hochberg FDR correction	Low-signal screens
fdr_method	'bonferroni' - Bonferroni FDR correction	Adjusts <i>P</i> -values with Bonferroni multiple hypothesis correction	High-signal screens
filter_genes	N/A	Genes to filter out from scoring process	Remove technical controls or flagged genes
ignore_orientation	N/A	If TRUE, groups guides from both orientations for each gene pair together to reduce the amount of hypothesis testing by half	Cas12a-Cas12a or low-signal screens

process data manually and score it automatically, and a wrapper interface that runs the entire Orthrus pipeline in a single function call. After construction of the reads file from raw sequencing data in Procedure 1, the first interface is demonstrated in Procedure 2, and the latter two interfaces are demonstrated in Procedure 3. All three interfaces to Orthrus are described below, and their corresponding function calls are summarized in Fig. 2.

- 1 The *manual* interface is the most verbose and allows users to score specific screens in different ways. For instance, with this interface, users may choose to score screens with different FDR thresholds, or may alter the parameters of plots generated for different screens (e.g., to relabel hits in figure legends). This interface requires a minimum of 13 Orthrus function calls for an experiment with both dual-targeting and combinatorial-targeting guides and may require significantly more for complex experimental designs.
- 2 The *batch scoring* interface is more succinct than the manual interface and affords users a similar level of control as the manual interface. Scoring many screens with this interface requires only one line of code, for a minimum (and typically, a maximum) of seven Orthrus function calls across the entire workflow. However, this interface requires that users score all guides of a specific type with the same parameters across all screens. For large-scale experiments where scores should be computed in a standardized way, this behavior is desired.
- 3 The *wrapper* interface allows users to score their data with a single Orthrus function call. While convenient, this interface is only recommended for users with a deep understanding of both their own data as well as how specific parameters applied to different steps of the scoring process affect their results.

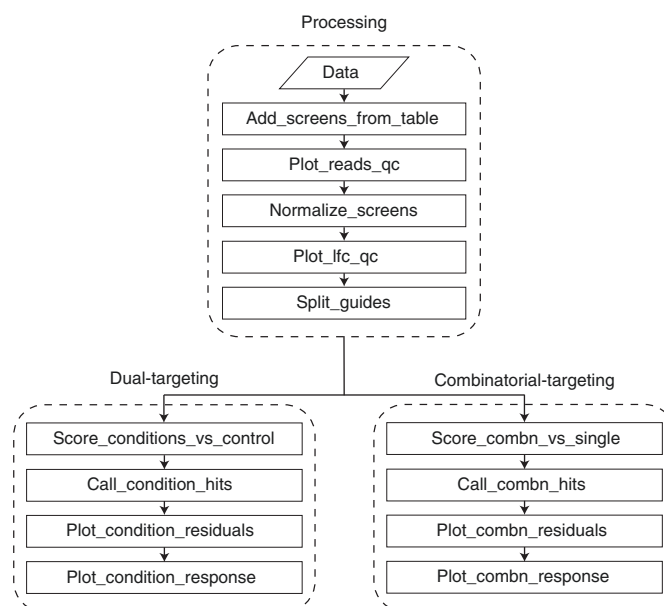


Fig. 2 | Key Orthrus functions. Specific Orthrus functions to call in order for data processing as well as dual-targeting and combinatorial-targeting scoring.

For typical experimental designs, the batch scoring interface is recommended. Orthrus separates the processing and scoring steps in order to encourage users to manually examine their data at key breakpoints during an analysis session. Ideally, after running Orthrus' processing steps, users will refer to the variety of automatically generated QC plots and metrics that Orthrus outputs to inform their parameter choices. Users will then choose to either proceed to the scoring steps with their current parameter choices or choose to reprocess their data with changed parameters (or other additions, such as manual filters for problematic guides). While the manual interface also encourages this behavior, the batch scoring interface is far more succinct and additionally forces the user to choose common parameters for scoring different screens, which facilitates the generation of results that are more comparable across screens.

Preparation stage

During the preparation stage, users first format their reads file, their sample table and their batch table as described in the 'Input format' subsection. They load these files into their workspace and proceed to the processing stage.

Processing stage

Users run Orthrus' processing and QC functions as shown in Steps 5–10 of Procedure 2 or Step 3 of Procedure 3. After running these functions and manually examining QC plots and metrics for their data, users then decide whether or not to change certain parameters (e.g., to filter low-readcount guides) and either rerun this step or proceed to the scoring stage.

Scoring stage

Finally, users call Orthrus' scoring functions either manually or with the batch scoring interface. The manual interface requires users to dual-targeting, combinatorial-targeting and single-targeting guides separately, but the recommended batch scoring interface allows users to score all three types of guides at the same time.

After scoring their data, users are encouraged to examine their final results and decide to either keep their current parameters or rerun the processing and scoring stages with different parameters. Some of the most consequential parameters include the choice of hypothesis testing (we recommend moderated *t*-testing as opposed to Wilcoxon rank-sum testing for most screens), whether or not to loess-normalize residual effects, and the choice of FDR and effect size thresholds for hit-calling. Table 4 provides more information on these and other parameters, as well as their recommended use cases.

Chosen data for procedures

The procedures below demonstrate the application of the Orthrus package for analyzing two combinatorial CRISPR screening example datasets. The first dataset analyzed in Procedures 1 and 2 consists of both raw sequencing data as well as a preprocessed reads file from CHyMERa screens described in Gonatopoulos-Pournatzis et al.¹². The second dataset analyzed in Procedure 3 contains screens from a separate multiplexed Cas12a system described in Dede et al.¹⁸. All data for Procedures 1 and 3 are downloadable from the Zenodo repository (<https://zenodo.org/record/4527616>), whereas the processed reads file and required tables for Procedure 2 are bundled with Orthrus' download.

Materials

Equipment

Software

- Procedure 1
 - Tested with Bowtie 0.12.9 (<https://sourceforge.net/projects/bowtie-bio/files/bowtie/0.12.9/>), although any Bowtie version 1 is applicable
 - Perl 5 (<https://www.perl.org/>)
 - Bash (<https://www.gnu.org/software/bash/>)
 - R version 3.6 or greater (<https://www.r-project.org/>)
- Procedures 2 and 3
 - R version 3.6 or greater with the packages listed below and in Step 1 of Procedures 2 and 3
 - devtools
 - Orthrus
 - ggplot2
 - ggthemes
 - pheatmap
 - PRROC
 - RColorBrewer
 - BiocManager
 - limma
 - stringr

Hardware

- Procedure 1
 - CPUs: tested on a machine with a single 2.6 GHz Intel Core i7 processor
 - Memory: tested on a machine with 16 GB of random-access memory (RAM), but should run on most machines with 4+ GB of RAM
 - Operating system: tested on macOS 10.13.6 High Sierra, but should run on most Unix operating systems
 - An internet connection is required to download the required software, scripts and data
- Procedures 2 and 3
 - Memory: at least 4 GB of RAM
 - Operating system: any operating system capable of running R and installing the packages listed in the Software section above
 - An internet connection is required to download the Orthrus package

Data

The CHyMERa dataset analyzed in Procedures 1 and 2 comprises combinatorial CRISPR screens performed with the CHyMERa experimental platform in two different human cell lines, HAP1 and RPE1, across a control and a Torin1-treated condition, with read counts taken at two different timepoints for each cell line (T12 and T18 for HAP1, T18 and T24 for RPE1). While the CHyMERa dataset contains single-targeting, dual-targeting and combinatorial-targeting guides, the multiplexed Cas12a dataset analyzed in Procedure 3 only contains combinatorial-targeting guides that target 400 paralog pairs and control guides that target paralogous genes paired with nonessential genes¹⁸. These control guides are treated as single-targeting guides for scoring purposes, as they are analogous to single-targeting guides in Procedure 1 and 2's CHyMERa dataset that target paralogous genes paired with intergenic regions.

- Procedure 1
 - The subset sequencing data processed in Procedure 1 only contains reads from the WT HAP1 T18 screen
 - Downloadable in the 'Procedure1' folder of the Zenodo repository (<https://zenodo.org/record/4527616>)
- Procedure 2
 - The reads file analyzed in this Procedure contains processed reads for all screens and cell lines described above
 - Each guide in the reads file, represented by a single row, has associated read counts for the knockout of regions specified by a Cas9 and a Cas12a guide sequence
 - The first set of columns in the dataset contain metadata for guide pairs, and the remaining numeric columns contain raw read counts for the guide pairs across every screen
 - This guide library contains several different types of guides
 - *Dual-targeting* guides that target the same gene twice
 - *Combinatorial-targeting* guides that target each gene of a paralogous gene pair
 - *Single-targeting* guides that target a gene's exonic region in addition to a relatively distant intergenic region. These are required to score combinatorial-targeting guides for GIs
 - Bundled with Orthrus' download
- Procedure 3
 - The reads file analyzed in this Procedure contains processed reads for screens performed in A549, HT29 and OVCAR8 cell lines, and is formatted similarly to the reads file in Procedure 2
 - The guide library contains two different types of guides
 - *Combinatorial-targeting* guides that target pairs of paralogous genes
 - *Single-targeting* guides that target paralogous genes paired with nonessential control genes
 - Downloadable in the Zenodo repository (<https://zenodo.org/record/4527616>)

Equipment setup

Software

If R is not already installed, download R version 3.6 or greater from <https://cran.rstudio.com/>. To use R, we recommend downloading Rstudio IDE from <https://rstudio.com/products/rstudio/download/>.

Procedure 1: processing CHyMERa data

Setup ● Timing 7–8 h

- 1 *Download required data.* Download the subset sequencing data, library guide sequences, processing scripts and Bowtie. CHyMERa data was processed with Bowtie version 0.12.9, whose download link is below. While other versions of Bowtie 1 are similarly appropriate for processing short sequencing data such as for CHyMERa screens, Bowtie 2 is not recommended.

Download everything in the Zenodo repository located here and unzip the files: <https://zenodo.org/record/4527616>

Download bowtie version 0.12.9 here:

<https://sourceforge.net/projects/bowtie-bio/files/bowtie/0.12.9/>

Unzip the downloaded bowtie files into a subdirectory of the "Procedure1" folder of the Zenodo repository. Ensure that the "Procedure1" folder is the current working directory and that it contains all files in the "Procedure1" folder of the Zenodo repository. Similarly, ensure that the "Library" folder is a subdirectory of the current working directory, and that it contains all the files in the "Library" folder of the original Zenodo repository linked above.

- 2 *Preprocess sequencing data.* To identify hgRNA barcodes present in a given sample, CHyMERa screening libraries are subjected to paired-end Illumina sequencing to capture both Cas12a and Cas9 guide sequences²². To identify guide sequences within sequencing reads for two technical replicates of a single screen (HAP1 T0) using U6 promoter and Cas9 tracrRNA sequences as 'anchor' sequences, run the preprocessReadsPE.pl script as follows. This takes two FASTQ

sequencing files, representing a single technical replicate of a single screen, as input. It outputs four files, two per FASTQ file. The first, [FILENAME]_preprocessed.fastq, contains reads trimmed down to the guide sequences, and the second, [FILENAME]_failed.fastq, contains full reads where anchor sequences were not found. These anchor sequences are hard-coded U6 and tracr sequences in the R1_stem and R2_stem variables, which should be appropriate for all CHyMERa screens, but may be replaced if analyzing data from a different experimental platform. This step takes ~1 Mb of memory and 5 min to preprocess 34.4 M paired-end reads.

```
./preprocessReadsPE.pl Moffat_HH-79_S1_R1_001.fastq.gz
Moffat_HH-79_S1_R2_001.fastq.gz
```

- 3 *Align reads with Bowtie.* Next, the reads must be aligned to the screening library, which may be custom or provided by a vendor. The library used for previous CHyMERa screens is contained in the file Human_HybridGuide_Library_v3.txt, and for details on how this library was constructed, please consult Gonatopoulos et al. (2020)¹². Perform this alignment with bowtie 0.12.9, using either option A to run on a single core, or option B to run on a cluster if available. The output of this step consists of six files: two contain unmapped reads, two contain mapped reads, one is a .sam file and the last is a log file. The .sam file named 'HH-79_aligned.sam' is the only file required by the next step.

(A) Aligning reads on a single core

- (i) Change -p [N_CORES] as follows to take advantage of bowtie's parallel processing to use any number of available CPU cores. On a single core, it takes ~7–8 h to align 31.5 M preprocessed sequence reads.

```
export BOWTIE_INDEXES=./Library/
bowtie-0.12.9/bowtie -p 1 -v 3 -l 18 --chunkmbs 256 -t
paralog_library_V3 --un HH-79_unmapped.fastq --al
HH-79_mapped.fastq -1
Moffat_HH-79_S1_R1_001_preprocessed.fastq -2
Moffat_HH-79_S1_R2_001_preprocessed.fastq HH-79_aligned.sam
2> HH-79.log
```

(B) Aligning reads on a cluster

- (i) To run this on a cluster, run bowtie with the submitjob command as follows in a Bash for loop. This is useful for processing more than one screen concurrently.

```
export BOWTIE_INDEXES=./Library/
for f1 in *_R1_001_preprocessed.fastq
do
d=$(echo $f1 | sed -E
's/_S[0-9]+_R1_001_preprocessed.fastq/g')
submitjob -c 6 bowtie-0.12.9/bowtie -p 6 -v 3 -l 18 --chunkmbs
256 -t paralog_library_V3 --un $d_unmapped.fastq --al
$d_mapped.fastq -1 $f1 -2 ${f1/_R1/_R2_} $d_aligned.sam
2\> $d.log"
done
```

- 4 *Parse alignments to guide counts.* Run parseBowtieOutput.pl as follows to get guide-level read counts for the HAP1 T0 screen. This takes ~2 min running on a single core and outputs the file 'HH-79_counts.txt' required by Step 5.

```
cat HH-79_aligned.sam | perl parseBowtieOutput.pl >
HH-79_counts.txt
```

- 5 *Merge count files.* Merge all count files in the current working directory into a reads file with mergeAndAnalyzeParalogResults.R as follows. This generates a raw reads file named 'rawCounts.txt'.

```
Rscript mergeChymeraResults.R
```

- 6 *Reformat counts for Orthrus (optional).* As described in the 'Input format' section, Orthrus requires reads files to describe orientation in gene symbol columns. Both the initial library file and the file 'rawCounts.txt' with appended read counts do not represent orientation in this way. Run the following script to fix this issue for the CHyMERa paralog data. Because the reads output in Step 5 are only for the HAP1 T0 screen, the script 'prepChymeraData.R' instead runs on the included file 'paralogLibrary_rawCounts.txt'.

```
Rscript prepChymeraData.R
```

Procedure 2: scoring CHyMERa data manually

Setup ● Timing 1–5 min

- 1 *Install required R packages.* Before installing Orthrus, install all packages it requires from both the CRAN repository and Bioconductor as follows:

```
install.packages("ggplot2")
install.packages("ggthemes")
install.packages("pheatmap")
install.packages("PRROC")
install.packages("RColorBrewer")
if (!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")
BiocManager::install("limma")
```

You may have to restart R once or more during this installation process. To install the Orthrus package, install it directly from its Github repository using the `install_github` command from the `devtools` package as follows:

```
install.packages("devtools")
library(devtools)
install_github("csbio/Orthrus")
```

- 2 *Load packages.* After installing the dependencies above, load Orthrus and ggplot into the R environment as follows and rename the example data described above. For a detailed description of the dataset, please consult Gonatopoulos-Pournatzis et al¹².

```
library(orthrus)
df <- chymera_paralog
```

- 3 *Set parameters.* While Orthrus does not require any global parameters, it is helpful to create output folders for various plots, text files and spreadsheets ahead of time, as follows. Many downstream functions will take these as parameters.

```
output_folder <- file.path("orthrus_protocol")
qc_folder <- file.path(output_folder, "qc")
plot_folder <- file.path(output_folder, "scored")
lfc_folder <- file.path(qc_folder, "lfc_plots")
if (!dir.exists(output_folder)) { dir.create(output_folder,
  recursive = TRUE) }
if (!dir.exists(plot_folder)) { dir.create(plot_folder) }
if (!dir.exists(qc_folder)) { dir.create(qc_folder) }
if (!dir.exists(lfc_folder)) { dir.create(lfc_folder) }
```

- 4 *Name screens.* Lastly, Orthrus requires the user to associate technical replicate read counts with screen names, so that downstream functions operate on the screen level rather than the replicate level. To do this, build up a list of screen objects either manually with the `add_screen` function,

described in option A, or automatically from a sample file mapping screen names to replicate columns with the `add_screens_from_table` function, described in option B.

(A) Build up a list of screen objects manually

- (i) The `add_screen` function requires the user to give each screen a name, such as `HAP1_T12`, and a list of column names corresponding to technical replicates for that screen. To apply this function, list names for T0 screens, which have no technical replicates, and call parameters by their name. For subsequent calls to `add_screen`, pass previous results in as the first argument to build up the list of screens, as follows:

```
screens <- add_screen(name = "HAP1_T0", replicates = "HAP1.T0")
screens <- add_screen(screens, "RPE1_T0", "RPE1.T0")
```

- (ii) To normalize the rest of the screens to their respective T0 screens to get LFCs, add the name of the screen to normalize against in the final parameter of the `add_screen` function (the `normalize_name` parameter) as follows. All screens from later timepoints have three technical replicates, A, B and C, which are separately normalized to T0s and are automatically averaged farther downstream in the pipeline.

```
screens <- add_screen(screens, "HAP1_T12", c("HAP1.T12A",
      "HAP1.T12B", "HAP1.T12C"), "HAP1_T0")
screens <- add_screen(screens, "HAP1_T18", c("HAP1.T18A",
      "HAP1.T18B", "HAP1.T18C"), "HAP1_T0")
screens <- add_screen(screens, "Torin_T12", c("HAP1.Torin.T12A",
      "HAP1.Torin.T12B", "HAP1.Torin.T12C"), "HAP1_T0")
screens <- add_screen(screens, "Torin_T18", c("HAP1.Torin.T18A",
      "HAP1.Torin.T18B", "HAP1.Torin.T18C"), "HAP1_T0")
screens <- add_screen(screens, "RPE1_T18", c("RPE1.T18A",
      "RPE1.T18B", "RPE1.T18C"), "RPE1_T0")
screens <- add_screen(screens, "RPE1_T24", c("RPE1.T24A",
      "RPE1.T24B", "RPE1.T24C"), "RPE1_T0")
```

(B) Build up a list of screen objects automatically

- (i) Use the `add_screens_from_table` function as follows. This function requires the user to specify either a dataframe or the path to a tab-separated file mapping screen names to technical replicate names, in addition to another screen to which the given screen should be normalized during LFC computation in Step 7. This sample table is bundled with Orthrus' download and is described in Table 2, which must include the column names `Screen`, `Replicates` and `NormalizeTo`. To disable LFC computation for specific screens, such as T0 screens, specify NA for those screens in the `NormalizeTo` column.

```
sample_table <- chymera_sample_table
screens <- add_screens_from_table(sample_table)
```

Processing ● Timing 5-30 min

- 5 *Make read count QC plots.* After associating technical replicate read counts to screen names, make QC plots for pre-normalization read count data. This allows users to investigate potential issues with their screening data, such as low sequencing depth for certain screens or unexpected skew in read count distributions. One function makes all of these QC plots for all screens, which are automatically saved to the QC subfolder created earlier as either png or pdf files, as follows:

```
plot_reads_qc(df, screens, qc_folder, display_numbers = FALSE,
  plot_type = "pdf")
```

A summary of all plots made by the `plot_reads_qc` function is contained in Table 5.

? TROUBLESHOOTING

- 6 *Examine read count QC plots.* Closely examine all output QC plots to reveal screening issues that need to be manually addressed by the user. For example, certain screens may be heavily skewed

Table 5 | Files output by QC functions and their descriptions in Procedure 2, Steps 5 and 8 and Procedure 3, Step 4

QC function	Output file names	Description
plot_reads_qc	[SCREEN]_raw_reads_histogram	Histograms of log-scaled read counts
plot_reads_qc	total_reads	Total read counts for all screens with hypothetical coverage appended
plot_reads_qc	reads_heatmap	Pearson correlation between log-scaled readcounts
plot_lfc_qc	[REPLICATE1]_vs_[REPLICATE2] _replicate_comparison	Scatterplot of LFCs between all replicates
plot_lfc_qc	replicate_pcc	Tab-delimited file of correlations for all replicates

towards guides with unexpectedly high or low read counts. Additionally, for typical experiments, the user should expect T0 read counts to cluster separately from later timepoint replicates in the log-normalized read count heatmap of Pearson correlations between screens. T0 replicates that do not cluster separately could implicate overarching screen quality issues.

■ PAUSE POINT A manual examination of all read count-based QC plots and metrics should be performed before proceeding.

- 7 *Normalize read counts and compute LFCs.* Normalize read counts based on sequencing depth and compute guide LFCs with the `normalize_screens` function, as follows. All screens passed into the function will be log2-scaled and depth-normalized, including T0 screens. LFC values will additionally be computed for all screens with associated `normalized_name` parameters. In this procedure, LFCs for the RPE1_T18 and RPE_T24 screens will be computed relative to the RPE_T0 screen, and similarly for HAP1_T12, HAP1_T18, Torin_T12 and Torin_T18 to the HAP1_T0 screen.

While the `normalize_screens` function normalizes against early-timepoint screens with multiple replicates by computing the mean log2-scaled reads across early-timepoint replicates before LFC computation, because neither dataset analyzed in this protocol includes replicates for early-timepoint screens, this behavior is not demonstrated. Additionally, while it is recommended to normalize chemogenetic screens against early-timepoint screens to compute LFCs before comparing drug treatment screens with control screens, Orthrus' downstream scoring functions are applicable to log2-normalized reads as well as LFCs.

In addition to guide normalization and LFC computation, the `normalize_screens` function also automatically removes guides that are over- or underexpressed at earlier timepoints. For this procedure, to remove guides with <30 read counts in any T0 screen, pass a list of both T0 screens into the `filter_names` parameter and set the `min_reads` parameter to 30. Note that this value or more conservative values such as 40–60 are recommended for most screens, regardless of library size. Both coverage and the standard deviation of gRNA abundance for early-timepoint or plasmid pool data, however, can be taken into account when setting this parameter. Because this relationship can be complex, please refer to Imkeller et al. (2020)²³. No guides in this library meet the high-abundance threshold (defined by the default value of 10,000 reads for the `max_reads` parameter), so although high-abundance guides are also automatically filtered out, this will not affect this particular dataset.

```
df <- normalize_screens(df, screens, filter_names = c("HAP1_T0",
  "RPE1_T0"), min_reads = 30, max_reads = 10000)
```

? TROUBLESHOOTING

- 8 *Make LFC QC plots.* While read count QC plots allow for bird's-eye views of screening data, LFC-based QC reveals specific quality information on individual guides and also outputs important quantitative QC metrics. To generate LFC-based QC plots and metrics for all screens, call the `plot_lfc_qc` function, as follows:

```
plot_lfc_qc(df, screens, qc_folder, display_numbers = FALSE,
  plot_type = "png", negative_controls = c("NT"))
```

A summary of all plots made by the `plot_lfc_qc` function is contained in Table 5.

- 9 *Examine LFC QC plots.* Like for read count QC plots, closely examine all output QC plots to reveal important screen quality issues. Replicate comparison plots can reveal specific guides that are unexpectedly over- or underrepresented, or which appear to be outliers for a specific technical replicate but not others. Points are colored by whether or not both gRNAs for each dgRNA target either a nonessential gene as defined by Hart et al. (2014) or a gene in the list specified by the user in the `negative_controls` parameter²⁴. More generally, they also show technical replicates that do not appear to correlate with each other or are skewed in problematic ways. Quantitative information on replicate Pearson and Spearman correlations is also contained in the output file `'replicate_cor.tsv'`.

Because most libraries are designed with sets of positive and negative control essential genes, Orthrus reports the area under receiver operating characteristic (ROC) curves (AUCs) for essential-gene dropout for all technical replicates. Specifically, essential genes are defined by the CEG2 core essential gene set, and nonessential genes are defined by Hart et al. 2014 (ref. ²⁴). Gene effects for essential genes are computed based on the dropout of guides that target essential genes twice, two different essential genes, or an essential gene and an intergenic region, and similarly for nonessential genes. The AUC for essential genes compared with both nonessential genes as well as all other genes is reported in the output file `'essential_PR_QC.tsv'`. While for most negative selection whole-genome screens one would expect AUC values > 0.9, for specialized guide libraries in which one expects many other strong negative fitness effects outside of essential gene pairs this AUC value may be substantially lower. Moreover, because AUC values for comparisons against reference nonessential genes are typically higher than AUC values for comparisons against all genes not in the essential set, the latter AUC value tends to be more predictive of screen quality issues.

■ **PAUSE POINT** A manual examination of all read LFC-based QC plots and metrics should be performed before proceeding.

? TROUBLESHOOTING

- 10 *Parse gene pairs by type.* To support Orthrus' two different scoring modes, split the guide dataframe into different types of guides based on orientations defined by gene symbol columns named `gene1` and `gene2` with the `split_guides` function, as follows. The relationship between gene symbols and orientation, as well as their required formatting, is explained in the 'Input format' section of the Introduction. If guides are mapped to unique identifiers such as guide sequences, additionally pass in column names for those identifiers such that the column name passed in first corresponds to the `gene1` column and the column name passed in second corresponds to the `gene2` column. This enables loess correction with moderated *t*-testing, the default scoring modes supported by Orthrus, in Steps 11 and 15. If unspecified, the user must default to Wilcoxon rank-sum testing, which is not recommended for most experimental designs.

```
guides <- split_guides(df, screens, "Cas9.Guide", "Cpf1.Guide")
dual <- guides[["dual"]]
single <- guides[["single"]]
paralogs <- guides[["combn"]]
```

The output of this process is three separate lists, where each element contains all guides targeting a single gene pair. Dual-targeting guides are contained in the `dual` list, single-targeting guides are contained in the `single` list, and combinatorial-targeting guides are contained in the `paralogs` list.

Dual-targeted scoring ● Timing 5–10 min

- 11 *Score guides targeting the same gene twice.* Currently, the Orthrus package supports scoring dual-targeting guides by comparing one or more condition screens against a single control screen. This is performed via moderated *t*-testing for each gene pair, condition and orientation against corresponding guides in the control screen. As in Aregger et al.¹¹, guide-level residuals are corrected with loess normalization before performing hypothesis testing to account for skewed and non-normal distributions. Scoring guides in this way results in both an effect size measure, based on the mean of loess-normalized residuals between condition and control LFCs, as well as a measure of statistical significance (a *P*-value from moderated *t*-testing on these residuals).

One function, `score_conditions_vs_control`, performs comparisons for all gene pairs using the `dual` list of guides defined above. Score the provided dataset for Torin1-specific effects in HAP1 cells by comparing Torin1 effects at each timepoint against untreated HAP1 effects using the

Table 6 | Columns contained in scored data output from the dual-targeting scoring mode in Procedure 2, Steps 11 and 12 and Procedure 3, Step 4

Scored data column	Description
gene1	Gene symbol targeted by the first guide
gene2	Gene symbol targeted by the second guide
n_[SCREEN]	Number of guides post-filtering for the screen
mean_[SCREEN]	Mean LFC across all guides
variance_[SCREEN]	Variance for all guides
differential_[CONDITION]_vs_[CONTROL]	Loess-adjusted (if specified) differential between mean condition and control LFCs
pval_[CONDITION]_vs_[CONTROL]	<i>P</i> -value between loess-adjusted residuals for condition and control
fdr_[CONDITION]_vs_[CONTROL]	FDR-adjusted <i>P</i> -value
significant_[CONDITION]_vs_[CONTROL]	Significance calls returned from <code>call_significant_response</code>
effect_type_[CONDITION]_vs_[CONTROL]	Effect type calls returned from <code>call_significant_response</code>

SCREEN placeholders represent the names of all scored condition and control screens, whereas CONDITION and CONTROL placeholders represent the names of condition and control screens, respectively.

following code, specifying moderated *t*-testing (instead of Wilcoxon rank-sum testing, which is supported but not recommended due to its reduced statistical power) and loess normalization. Additionally, pass in 'NT' to the `filter_genes` parameter to ignore nontargeting control genes during scoring. This parameter may contain any number of genes stored in a character vector, but for the CHyMERa library only 'NT' needs to be specified.

```
temp <- score_conditions_vs_control(dual, screens, "HAP1_T12",
  "Torin_T12", test = "moderated-t", loess = TRUE,
  min_guides = 3, filter_genes = c("NT"))
dual_scores1 <- temp[["scored_data"]]
residuals1 <- temp[["residuals"]]
temp <- score_conditions_vs_control(dual, screens, "HAP1_T18",
  "Torin_T18", test = "moderated-t", loess = TRUE,
  min_guides = 3, filter_genes = c("NT"))
dual_scores2 <- temp[["scored_data"]]
residuals2 <- temp[["residuals"]]
```

The above code returns a list of two dataframes. The first dataframe in the list, named `scored_data`, contains effect size and FDR values for all gene pairs comparing Torin effects to WT Hap1 effects at either T12 or T18, as well as many additional columns described in Table 6. It is important to note that NA values in this dataframe represent genes that have too few guides remaining post-filtering based on T0 read counts and specified filter genes, for a default threshold of three guides per gene pair. The second dataframe in the list, `residuals`, contains guide-level residual values for detailed examination of a given gene pair's results, and is used for residual plotting functions in Step 13.

? TROUBLESHOOTING

- 12 *Call significant effects for dual-targeting guides.* After scoring data, for a typical experiment, the user would like to reduce a large list of significant hits down to a ranked list of high-priority hits. Prioritize hits using the function `call_condition_hits`. Call this function with a given FDR threshold and differential effect threshold to call significant positive or negative hits as follows.

Hits are called based on two criteria. Gene pairs with (a) an FDR less than the given FDR threshold and (b) an absolute value of their loess-normalized residuals that is greater than the given differential effect threshold will be called as significant positive or negative hits. When calling this function, the user may also choose to rename positive and negative hits.

```
dual_scores1 <- call_condition_hits(dual_scores1, "HAP1_T12",
  "Torin_T12", neg_type = "Sensitizer", pos_type =
  "Suppressor", fdr_threshold = 0.1, differential_threshold =
  0.5)
```

```
dual_scores2 <- call_condition_hits(dual_scores2, "HAP1_T18",
  "Torin_T18", neg_type = "Sensitizer", pos_type =
  "Suppressor", fdr_threshold = 0.1, differential_threshold =
  0.5)
write.table(dual_scores1, file.path(output_folder,
  "dual_targeting_gene_calls_t12.tsv"), sep = "\t",
  row.names = FALSE, col.names = TRUE, quote = FALSE)
write.table(dual_scores2, file.path(output_folder,
  "dual_targeting_gene_calls_t18.tsv"), sep = "\t",
  row.names = FALSE, col.names = TRUE, quote = FALSE)
```

With scoring complete, write the data to file as shown above.

- 13 *Plot residual effects (optional)*. After scoring data, users might like to see all guide-level LFC values for significant hits to visually confirm that certain genes possess consistent effects across most guides. Use the function `plot_condition_residuals` as follows to automatically generate these plots for all hits called with `call_condition_hits`, and output them in a sorted order to the given folder. Make LFC plots for Torin-specific significant hits for T12 and T18.

```
plot_condition_residuals(dual_scores1, residuals1, "HAP1_T12",
  "Torin_T12", file.path(lfc_folder, "dual_lfc_t12"), neg_type
  = "Sensitizer", pos_type = "Suppressor", plot_type = "png")
plot_condition_residuals(dual_scores2, residuals2, "HAP1_T18",
  "Torin_T18", file.path(lfc_folder, "dual_lfc_t18"), neg_type
  = "Sensitizer", pos_type = "Suppressor", plot_type = "png")
```

For these data, the plot 'neg_1_HECTD1_None.png' refers to the top negative hit in terms of differential effect (marked by 'neg_1' in the filename) for Torin T18 against WT Hap1. Similarly, the file 'pos_2_EED_None.png' refers to the second-highest ranked positive hit in terms of differential effect.

- 14 *Plot condition response*. Finally, generate plots for each condition against the control screen with `plot_condition_response`, as follows. This outputs two plots, a scatterplot and a volcano plot, to a given folder. The volcano plot displays either $-\log_{10}(P\text{-value})$ or $-\log_2(\text{FDR})$ on the y -axis based on whether 'pval' or 'FDR' is passed to the parameter `volcano_type`, respectively. These volcano plots may be helpful to determine effect size and FDR thresholds for specific datasets. As for generating residual plots, ensure that the chosen names for negative and positive effects are passed into the function.

```
plot_condition_response(dual_scores1, "HAP1_T12", "Torin_T12",
  plot_folder, neg_type = "Sensitizer", pos_type =
  "Suppressor", volcano_type = "FDR", plot_type = "pdf")
plot_condition_response(dual_scores2, "HAP1_T18", "Torin_T18",
  plot_folder, neg_type = "Sensitizer", pos_type =
  "Suppressor", volcano_type = "FDR", plot_type = "pdf")
```

■ PAUSE POINT A manual examination of all dual-targeting scoring output should be performed before proceeding.

Combinatorial scoring ● Timing 5–10 min

- 15 *Score guides targeting multiple genes*. Orthrus also supports scoring guides that target multiple genes, such as paralog pairs, by comparing the effect of double-knockouts against an expected model derived from single-knockout effects. In detail, all guides that target gene A with Cas9 and gene B with Cas12a comprise the observed model for one orientation. All multiplicative combinations (additive in log-space) of single-targeting guides that target gene A with Cas9, target gene B with Cas12a, and match guide sequences with the observed model comprise the expected model for the same orientation (Fig. 3). Like for dual-targeted scoring, the residuals between the observed and expected models are computed for each orientation and loess-normalized before performing moderated t -testing for each gene pair and condition.

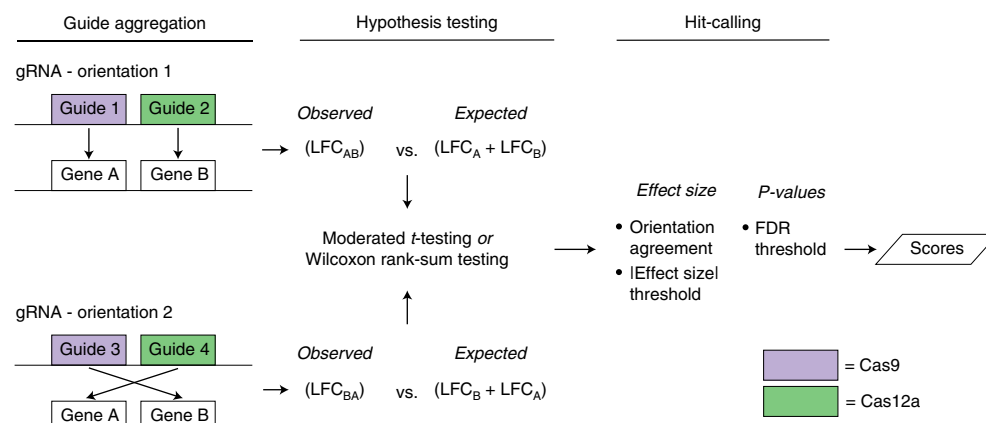


Fig. 3 | Schematic demonstrating how Orthrus accounts for guide orientation during combinatorial scoring by scoring guides from different orientations separately. Orientation is represented as guides that occupy different positions along a guide construct that targets both gene A and gene B. For both orientations, combinatorial knockouts are compared with expected effects derived from the sum of matching single knockout LFCs. After hypothesis testing, filters for absolute value of effect size, FDR and whether or not both orientations' effects have the same effect sign are applied to call significant hits.

One function, `score_combn_vs_single`, scores combinatorial-targeting guides for any number of conditions by passing in the 'dual' list of guides defined above. Call this function and score the provided dataset for combinatorial-targeting GIs for the HAP1 T12 and the RPE1 T24 screens as follows. As for dual-targeted scoring, additionally pass in 'NT' to the `filter_genes` parameter to ignore nontargeting control genes during scoring. Write scores to file afterwards.

```
screens_to_score <- c("HAP1_T12", "HAP1_T18", "RPE1_T18",
  "RPE1_T24", "Torin_T12", "Torin_T18")
temp <- score_combn_vs_single(paralogs, single, screens,
  screens_to_score, test = "moderated-t",
  return_residuals = TRUE, filter_genes = c("NT"))
paralog_scores <- temp[["scored_data"]]
paralog_residuals <- temp[["residuals"]]
paralog_scores <- call_combn_hits(paralog_scores,
  screens_to_score, neg_type = "Negative GI", pos_type =
  "Positive GI", fdr_threshold = 0.2, differential_threshold =
  0.5)
write.table(paralog_scores, file.path(output_folder,
  "paralog_gene_calls.tsv"), sep = "\t", row.names = FALSE,
  col.names = TRUE, quote = FALSE)
```

Like for `score_conditions_vs_control`, the above code returns a list of two dataframes. The first dataframe in the list, named `scored_data`, contains FDR values for all gene pairs comparing observed double-knockout effects with single-knockout effects as described in Procedure 2, Steps 11–12, for all six screens. It also contains many additional columns for each screen, listed in Table 7 and, like for `score_conditions_vs_control` NA values in this dataframe, represents genes that have too few guides remaining post-filtering based on T0 read counts. This guide threshold is also controlled by the `min_guides` parameter. The second dataframe in the list, `residuals`, contains guide-level residual values to enable the detailed examination of a given gene pair's results. For analyzing combinatorial-targeting CHyMERa data, we recommend raising the FDR threshold from the default of 0.1 to 0.2 with the `fdr_threshold` parameter as shown above. The specific choice of threshold is flexible, however, and is dependent on the expected signal-to-noise ratio in the dataset.

? TROUBLESHOOTING

- 16 *Plot combinatorial residual effects (optional).* After scoring data, some users would like to see guide-level residual LFC values for significant combinatorial hits. Like for dual-targeted scoring, the

Table 7 | Columns contained in scored data output from the combinatorial-targeting scoring mode obtained in Procedure 2, Steps 15 and 19, and Procedure 3, Step 4

Scored data column	Description
gene1	Gene symbol targeted by the first guide
gene2	Gene symbol targeted by the second guide
n_combn_[SCREEN]	Number of combinatorial-targeting guides post-filtering for the screen
n_single_[SCREEN]	Number of single-targeting guides post-filtering for the screen
mean_combn_[SCREEN]	Mean LFC across all combinatorial-targeting guides and both orientations
mean_single_[SCREEN]	Mean LFC across all combinations of single-targeting LFC sums that match combinatorial-targeting guide IDs, for both orientations
var_combn_[SCREEN]	Variance for all combinatorial-targeting guides across both orientations
var_single_[SCREEN]	Variance for all combinations of single-targeting LFC sums across both orientations
orientation_agree_[SCREEN]	True if both orientations' differential effects have the same sign, false otherwise
differential_combn_vs_single_[SCREEN]	Loess-adjusted (if specified) differential between mean combn and single LFCs
pval1_combn_vs_single_[SCREEN]	<i>P</i> -value between loess-adjusted residuals for the first orientation of combn and single LFCs
pval2_combn_vs_single_[SCREEN]	<i>P</i> -value between loess-adjusted residuals for the second orientation of combn and single LFCs
fdr1_combn_vs_single_[SCREEN]	FDR-adjusted <i>P</i> -value for the first orientation
fdr2_combn_vs_single_[SCREEN]	FDR-adjusted <i>P</i> -value for the second orientation
significant_[SCREEN]	Significance calls returned from <code>call_significant_response_combn</code>
effect_type_[SCREEN]	Effect type calls returned from <code>call_significant_response_combn</code>

SCREEN placeholders represent the names of all scored screens.

`function plot_combn_residuals` automatically generates these plots for all hits called with `call_combn_hits`, and outputs them in sorted order to the given folder as described in Step 13. Make LFC plots for HAP1_T12 and RPE1_T24 significant hits as follows.

```
residual_folder <- file.path(lfc_folder, "HAP1_T12_combn")
plot_combn_residuals(paralog_scores, paralog_residuals,
  "HAP1_T12", residual_folder, neg_type = "Negative GI",
  pos_type = "Positive GI")
residual_folder <- file.path(lfc_folder, "RPE1_T24_combn")
plot_combn_residuals(paralog_scores, paralog_residuals,
  "RPE1_T24", residual_folder, neg_type = "Negative GI",
  pos_type = "Positive GI")
```

- 17 *Plot condition response.* Finally, generate plots for each condition against the control screen as well as volcano plots for each screen with `plot_combn_response`, as follows. Ensure that the chosen names for negative and positive effects are passed into the function. For plotting effects from the Torin screen, additionally set the color of hits also significant in the WT HAP1 screens to gray by specifying the name of the respective control screen in the `filter_name` parameter.

```
plot_combn_response(paralog_scores, "HAP1_T12", loess = TRUE,
  plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI")
plot_combn_response(paralog_scores, "HAP1_T18", loess = TRUE,
  plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI")
plot_combn_response(paralog_scores, "RPE1_T18", loess = TRUE,
  plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI")
plot_combn_response(paralog_scores, "RPE1_T24", loess = TRUE,
  plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI")
plot_combn_response(paralog_scores, "Torin_T12", loess = TRUE,
```

```
plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI", filter_name = "HAP1_T12")
plot_combn_response(paralog_scores, "Torin_T18", loess = TRUE,
  plot_folder, neg_type = "Negative GI", pos_type =
  "Positive GI", filter_name = "HAP1_T18")
```

■ **PAUSE POINT** A manual examination of all combinatorial-targeting scoring output should be performed before proceeding.

Single-targeting scoring ● Timing 1-5 min

- 18 *Score guides targeting the same gene twice.* Orthrus allows users to score single-targeting guides, the same guides contained in the 'single' list used to construct the expected guide set for combinatorial scoring, by treating these as a single orientation for the dual-targeting scoring mode. To score single-targeting guides in this way, call `score_conditions_vs_control` with the `separate_orientation` argument set to `TRUE`, as follows.

This returns a list of two dataframes of scored data, one for each orientation. For this library, the first dataframe in the list contains scores for guides where Cas9 targets an exonic region, and the second dataframe contains scores where Cas12a targets an exonic region. Due to the CHyMERa library design that aimed to target each gene with three Cas9 guides and five Cas12a guides, as well as due to guides filtered from their low representation in T0 screens, the scored Cas9 single-targeting data contain too few significant hits. Accordingly, analyze only the single-targeting Cas12a data.

After scoring single-targeting guides, remove scored genes with too few guides remaining after T0 read count filters, and call significant hits with desired FDR and read count thresholds as follows:

```
single_scores <- score_conditions_vs_control(single, screens,
  "HAP1_T18", "Torin_T18", separate_orientation = TRUE)
single_scores <- single_scores[[2]][["scored_data"]]
to_keep <- !is.na(single_scores$n_HAP1_T18)
cat(paste("Removing", nrow(single_scores) - sum(to_keep), "sparse
  single-targeting genes\n"))
single_scores <- single_scores[to_keep,]
single_scores <- call_condition_hits(single_scores, "HAP1_T18",
  "Torin_T18", neg_type = "Sensitizer", pos_type =
  "Suppressor", fdr_threshold = 0.2, differential_threshold =
  0.5)
write.table(single_scores, file.path(output_folder,
  "single_targeting_gene_calls_t18.tsv"), sep = "\t",
  row.names = FALSE, col.names = TRUE, quote = FALSE)
```

■ **PAUSE POINT** A manual examination of all single-targeting scoring output should be performed before proceeding.

? TROUBLESHOOTING

Batch scoring

- 19 *Score dual and combinatorial-targeting guides in batch (optional).* Instead of scoring data manually, Orthrus provides the option to score different types of guides at the same time using its batch scoring mode. Use the following two function calls. This requires the creation of a batch file formatted as described in the 'Input format' section of the Introduction that maps screens to other screens they should be scored against, or to 'combn' to perform combinatorial scoring. This batch table is bundled with Orthrus' download, and is also presented in Table 3. The expected output of this process is largely equivalent to the output of Steps 5–17 above, with key differences including the use of standardized FDR and effect size thresholds across both dual-targeted and combinatorial-targeted scoring and that single-targeting scores are not computed automatically.

```
batch_table <- chymera_batch_table
batch_output_folder <- file.path("orthrus_protocol_batch")
if (!dir.exists(batch_output_folder))
```

```
{ dir.create(batch_output_folder) }
score_conditions_batch(dual, screens, batch_table, batch_output_
folder,
  test = "moderated-t", loess = TRUE, filter_genes = c("NT"),
  neg_type = "Sensitizer", pos_type = "Suppressor",
  fdr_threshold = 0.1, differential_threshold = 0.5)
score_combn_batch(paralogs, single, screens, batch_table,
  batch_output_folder, test = "moderated-t", loess = TRUE,
  filter_genes = c("NT"),
  neg_type = "Sensitizer", pos_type = "Suppressor",
  fdr_threshold = 0.2, differential_threshold = 0.5)
```

Procedure 3: analyzing dual Cas12a gRNA data with the batch scoring interface

Setup ● Timing 5–10 min

- 1 *Install and load required R packages.* As in Procedure 1, before installing Orthrus, install all packages it requires from both the CRAN repository and Bioconductor. Then load `devtools` to install Orthrus from the development Github repository, and finally load both Orthrus and `ggplot2` into the R environment, as follows:

```
install.packages("devtools")
install.packages("ggplot2")
install.packages("ggthemes")
install.packages("pheatmap")
install.packages("PRROC")
install.packages("RColorBrewer")
install.packages("stringr")
if (!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")
BiocManager::install("limma")
library(devtools)
install_github("csbio/Orthrus")
library(orthrus)
library(stringr)
```

- 2 *Download and load datasets.* Download and unzip the required datasets for this procedure from the Zenodo repository. Ensure that the working directory in R contains a subdirectory named ‘`dede_input`’ with input files that mirror the contents of the Zenodo repository. Load the reads file containing three combinatorial screens with two technical replicates each performed with a dual-Cas12a system¹⁸, in addition to sets of reference essential and nonessential gene standards used to process the data and to calculate QC metrics, into your R environment.

Download and unzip the zenodo directory from the following link as follows and ensure that the folder “`dede_input`” is a subdirectory of the current working directory: <https://zenodo.org/record/4527616>

```
input_folder <- file.path("dede_input")
prepped_file <- file.path(input_folder,
  "prepped_dede_paralog.tsv")
df <- read.csv(file.path(input_folder,
  "original_dede_paralog.txt"), sep = "\t",
  header = TRUE, stringsAsFactors = FALSE)
essentials <- read.csv(file.path(input_folder,
  "control_essentials.csv"),
  header = TRUE, stringsAsFactors = FALSE)
nonessentials <- read.csv(file.path(input_folder,
  "control_nonessentials.csv"),
  header = TRUE, stringsAsFactors = FALSE)
```

- 3 *Prep readcount dataset.* Instead of using intergenic-targeting guides like the CHyMERa library, this library targets nonessential genes as negative controls. To allow Orthrus to recognize this experimental design during scoring, after splitting the dataset's single gene symbol and guide ID columns into two, rename all nonessential-targeting guides as 'NegControl' in both gene symbol columns, as follows. Because gene symbols in this dataset accurately reflect guide orientations, further alterations to the gene symbol columns are unnecessary.

```
# Preps dataset
essentials <- unlist(essentials)
nonessentials <- unlist(nonessentials)
split <- str_split_fixed(df$GENE, ":", 2)
df$gene1 <- gsub("\\.*", "", split[,1])
df$gene2 <- gsub("\\.*", "", split[,2])
df$gene1[df$gene1 %in% nonessentials] <- "NegControl"
df$gene2[df$gene2 %in% nonessentials] <- "NegControl"
# Adds guide columns
split <- str_split_fixed(df$GENE_CLONE, "_", 4)
df$Guide1 <- split[,2]
df$Guide2 <- split[,4]
# Writes to file
write.table(df, prepped_file, sep = "\t", row.names = FALSE,
  col.names = TRUE, quote = FALSE)
```

- 4 *Score combinatorial-targeting guides.* Orthrus provides two ways to automatically score data: a batch scoring mode that first requires the user to process their data with Orthrus manually, and a wrapper function that runs the entire Orthrus pipeline in a single function call. Use option A to apply the batch scoring mode to the Dede et al. dataset, and option B to apply the wrapper function to the Dede et al. dataset. Both outputs are equivalent except for the choice of whether or not to loess-normalize residuals: option A demonstrates the output of loess normalization, and option B demonstrates the linear fit computed without loess normalization. For this dataset, loess normalization is not recommended as a visual examination of loess correction on scored data displays overfitting for points with a wide spread in the bottom two quadrants of each scatterplot, implicating the presence of both false positives and false negatives (Fig. 4).

(A) Batch scoring mode

- (i) To run batch scoring, process the Dede et al. dataset as described below, which corresponds to Steps 4–10 of Procedure 2 (with a few changes as explained further below). Afterwards, call the `score_combn_batch` function to automatically score combinatorial-targeting guides in the dataset. This function requires that the user first specify either a dataframe or a batch.tsv file that maps screen names to their respective controls (for dual-targeted scoring) or to a derived null model from single-targeting effects (for combinatorial scoring). This batch table is located in the 'dede_input' folder of the Zenodo repository here (<https://zenodo.org/record/4527616#.YK9S7k2ouUk>). An additional function, `score_conditions_batch`, scores dual-targeting guides automatically and takes the same batch file as input. However, because the Dede et al. dataset contains no dual-targeting guides, call `score_conditions_batch` to demonstrate its use, but expect no output from it.

Three changes to parameters compared with values shown in Procedure 2 are suggested to accurately score combinatorial-targeting guides in this dataset. First, change the list of negative controls to include 'NegControl'. Second, because the early-timepoint readcounts to normalize against for this dataset consist of plasmid pool readcounts with relatively high sequencing depth and low dropout, the `min_reads` parameter described in Step 7 of Procedure 1 can be safely tightened to 40 reads instead of the default 30 reads (this threshold could be further tightened if the user desires). Third, set the `ignore_orientation` parameter to `TRUE` to enable combinatorial scoring that aggregates guides across both orientations before running moderated *t*-testing. This reduces the amount of *t*-testing from two tests per gene pair to one test per gene pair, and is advised for this dataset because the dual-Cas12a combinatorial system is less likely to be influenced by guide orientation than the CHyMERa experimental system. By default, hits

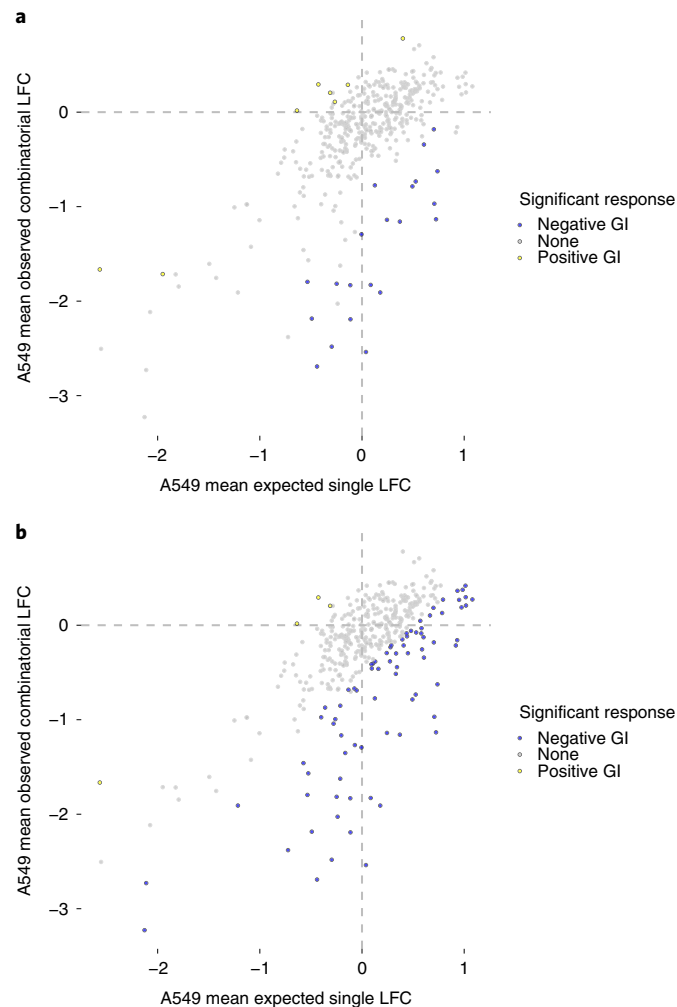


Fig. 4 | Summary plots of mean LFC. The plots are colored by significant effects, for all $n = 403$ gene pairs with combinatorial- and single-targeting guides for the Dede et al.¹⁸ A549 screen analyzed in Procedure 3. Gene-level GIs are shown as colored points that significantly deviate from the computed null model. Blue points are negative GIs with mean residual effects < -0.5 and Benjamini and Yekutieli FDRs < 0.2 , while yellow points are positive GIs with mean residual effects > 0.5 and Benjamini and Yekutieli FDRs < 0.2 . **a**, Scores with loess normalization enabled for 8 positive GIs and 21 negative GIs. **b**, Scores with loess normalization disabled for 4 positive GIs and 78 negative GIs. For this dataset, we conclude that loess normalization is not recommended due to potential false positives and negatives introduced in the bottom two quadrants of **a**.

from this scoring mode are still filtered based on whether or not the signs of their orientation-specific effects agree. The results of this step for the A549 screen are shown in Fig. 4a.

```
# Sets important paths
sample_file <- file.path(input_folder, "dede_sample_table.tsv")
batch_file <- file.path(input_folder, "dede_batch_table.tsv")
output_folder <- file.path("dede_output_batch")
qc_folder <- file.path(output_folder, "qc")
if (!dir.exists(output_folder)) { dir.create(output_folder) }
if (!dir.exists(qc_folder)) { dir.create(qc_folder) }
# Processes data
screens <- add_screens_from_table(sample_file)
plot_reads_qc(df, screens, qc_folder, display_numbers = TRUE,
  plot_type = "png")
```

```
df <- normalize_screens(df, screens, filter_names = "T0",
  min_reads = 40)
plot_lfc_qc(df, screens, qc_folder, display_numbers = TRUE,
  plot_type = "png", negative_controls = c("NegControl"))
guides <- split_guides(df, screens, "Guide1", "Guide2")
dual <- guides[["dual"]]
single <- guides[["single"]]
combn <- guides[["combn"]]
# Scores data with batch scoring functions
score_conditions_batch(dual, screens, batch_file,
  output_folder,
  test = "moderated-t", loess = FALSE)
score_combn_batch(combn, single, screens, batch_file,
  output_folder, test = "moderated-t", loess = FALSE,
  filter_genes = c("NegControl"), neg_type = "Sensitizer",
  pos_type = "Suppressor", fdr_threshold = 0.2,
  differential_threshold = 0.5)
```

(B) Using the wrapper function

- (i) To automatically run the entire Orthrus package with the wrapper function, call `orthrus_wrapper` as follows and pass in paths to the properly formatted reads file, the sample file and the batch file downloaded in Step 2. This automatically outputs all plots, metrics and scored data discussed for the data processing and combinatorial-scoring steps of Procedure 2 to a specified output folder. In addition, for this step, set the `loess` flag to `FALSE` to disable the loess correction of residuals. The results for this step are shown in Fig. 4b.

```
output_folder <- file.path("dede_output")
sample_file <- file.path(input_folder, "dede_sample_table.tsv")
batch_file <- file.path(input_folder, "dede_batch_table.tsv")
orthrus_wrapper(prepped_file, sample_file, batch_file,
  output_folder, id_col1 = "Guide1", id_col2 = "Guide2",
  filter_names = "T0", min_reads = 40,
  display_numbers = TRUE, negative_controls = nonessentials,
  test = "moderated-t", loess = FALSE, fdr_method = "BY",
  fdr_threshold = 0.2, differential_threshold = 0.5,
  plot_type = "png", ignore_orientation = TRUE)
```

? TROUBLESHOOTING

Troubleshooting

Troubleshooting advice can be found in Table 8.

Table 8 | Troubleshooting table

Procedure-Step	Problem	Possible reasons	Solutions
2-7 and 3-4	Many guides with too few reads in early-timepoint screens are filtered out	Various experimental issues early on in the screen, such as with T0 samples	Orthrus' scoring accounts for this, but users may consider applying the 'ignore_orientation' flag during scoring
2-9 and 3-4	Technical replicates correlate poorly with each other	Mislabeling in the sample file or any number of experimental issues	Fix technical replicate labels in the sample file, consider filtering early-timepoint guides more stringently, or redo the problematic screen
2-9 and 3-4	Guides appear to drop out stochastically between replicates in QC scatterplots	Dosage for a drug screen was too high or other experimental issues	Remove guides that completely drop out in any late-timepoint replicate, or redo the problematic screen

Table continued

Table 8 (continued)

Procedure-Step	Problem	Possible reasons	Solutions
2-5 and 3-4	The function <code>plot_reads_qc</code> returns the error 'Error in <code>check_screen_params</code> (df, screens): replicate [REPLICATE] not in df, remove screen [REPLICATE] with <code>remove_screens</code> '	Replicate name set in <code>add_screen</code> function does not exist in input dataframe	Remove screen containing the offending replicate from the screen list with the function <code>remove_screen</code> and re-add with correct replicate names
2-11, 2-15, 2-18 and 3-4	Many rows in scored data contain NA values	Guide filtering based on early-timepoint readcounts removed too many guides to score gene pairs with NA values (gene pairs with fewer guides than the <code>min_guides</code> parameter are not scored)	Consider relaxing guide filtering threshold, lowering the <code>min_guides</code> parameter, or investigating issues in early-timepoint screens
2-11, 2-15, 2-18 and 3-4	Scoring takes much longer than expected given the number of screens	Control gene pairs with many guides are not filtered out	Add controls such as 'NT' guides to the <code>filter_genes</code> scoring parameter in a vector

Common screen quality issues reflected by Orthrus' output as well as common issues encountered by users.

Table 9 | Time taken during different steps of the pipeline for processing increasing numbers of screens

Screens	Processing time (min)	Dual-targeted scoring time (min)	Combinatorial scoring time (min)
1	1.66	1.14	0.40
5	3.86	6.57	2.10
10	5.60	10.28	3.28
20	8.99	19.43	5.71
50	20.11	48.42	14.76

Values obtained using the script `test_at_scale.R`, which implements Procedure 2 in a loop and is available in the Zenodo repository (<https://zenodo.org/record/4527616>).

Timing

Procedure 1 demonstrates Orthrus' ability to process data for 92,746 guides across six screens. On a Windows machine running R with a single core and 16 GB of memory, the procedure run as a script took slightly under 5 min of runtime. Because Orthrus can be run on different screens sequentially, memory usage is not anticipated to be a bottleneck.

To test Orthrus' runtime for different numbers of screens, we ran Orthrus on a single screen duplicated 5, 10, 20, 50 and 100 times and timed the processing (Procedure 2, Steps 1–10), dual-targeting guide scoring (Procedure 2, Steps 11–14) and combinatorial scoring (Procedure 2, Steps 15–17) stages separately. For the processing stage, all guides were processed in the same function calls. However, because guide scoring does not take into account information between screens, both guide scoring stages were run as a loop to score each screen separately. Timing results are summarized in Table 9.

Anticipated results

All anticipated results for the protocol, as well as an R script containing the provided code, are available at <https://zenodo.org/record/4527616>.

Procedure 1

The files output by Procedure 1 are as follows. Step 2 outputs four files, two per FASTQ file, which contain reads trimmed down to the guide sequences and full reads where anchor sequences were not found. These are named based on the input.fastq.gz files with the format [FILENAME]_preprocessed.fastq and [FILENAME]_failed.fastq, respectively. Step 3 outputs six files, where two contain

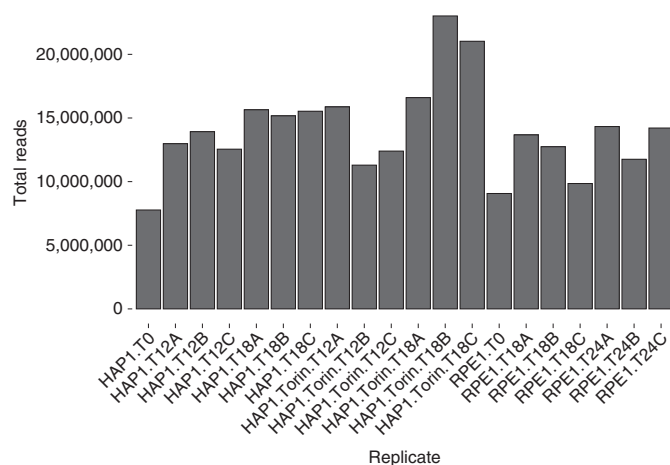


Fig. 5 | Total read counts for CHyMERa screens. Total read counts for all technical replicates in the example CHyMERa dataset ($n = 20$), output in Procedure 2, Step 5.

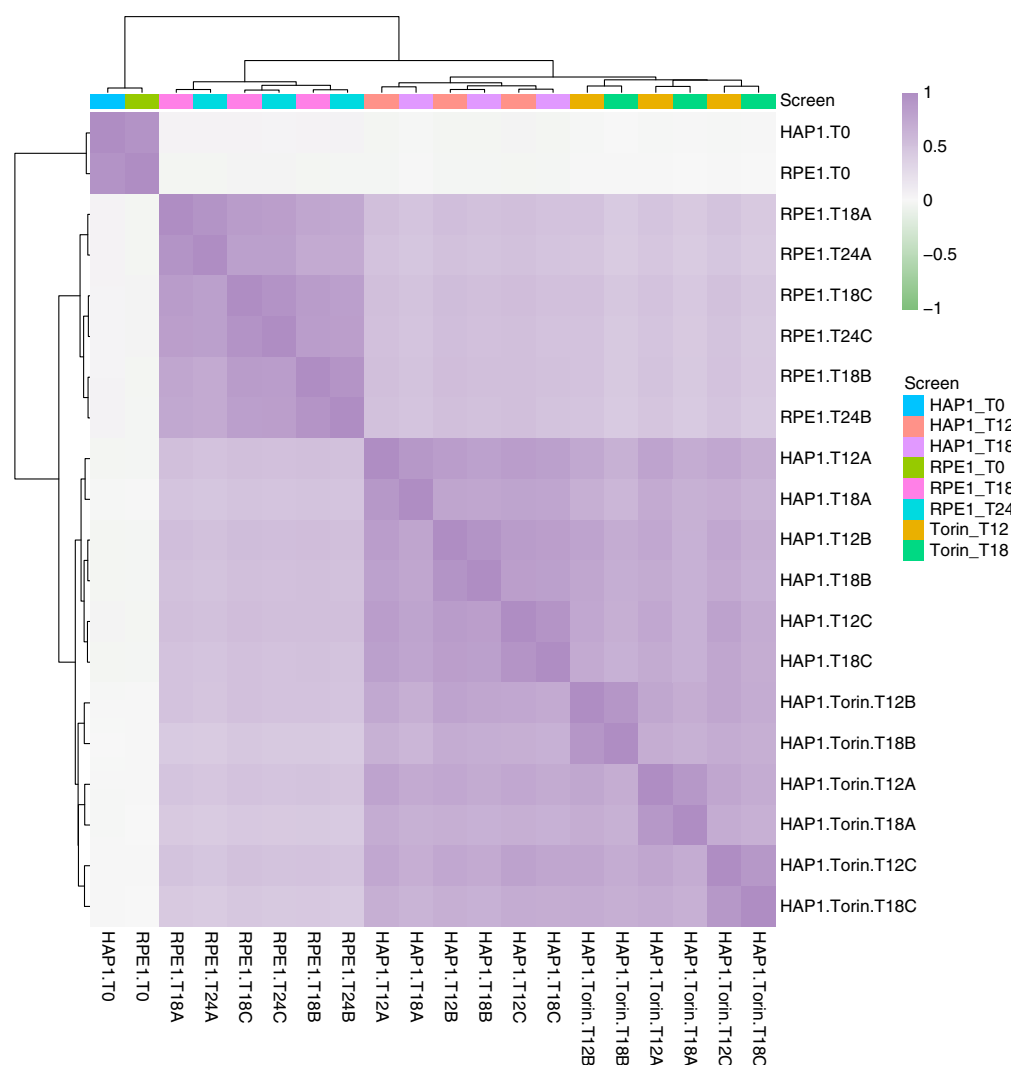


Fig. 6 | Heatmap of Pearson correlations. Shown are the correlations between LFCs for all technical replicates in the example CHyMERa dataset ($n = 20$), output in Procedure 2, Step 8. Well-correlated screens cluster together depending on the main sources of variation in the dataset. Here RPE1 screens cluster separately from HAP1 screens, which are further clustered into WT HAP1 screens and HAP1 + Torin1 screens.

Table 10 | Evaluation of recovery of essential gene LFC values for all technical replicates compared with all other genes (reported as AUC values, area under the ROC curve values) from the ChyMERa dataset analyzed in Procedure 2, Step 8

Technical replicate	Essential gene recovery AUC
HAP1 T0	0.51
RPE1 T0	0.51
HAP1 T12A	0.70
HAP1 T12B	0.70
HAP1 T12C	0.71
HAP1 T18A	0.69
HAP1 T18B	0.70
HAP1 T18C	0.70
HAP1 + Torin1 T12A	0.68
HAP1 + Torin1 T12B	0.68
HAP1 + Torin1 T12C	0.68
HAP1 + Torin1 T18A	0.67
HAP1 + Torin1 T18B	0.67
HAP1 + Torin1 T18C	0.67
RPE1 T18A	0.60
RPE1 T18B	0.60
RPE1 T18C	0.61
RPE1 T24A	0.61
RPE1 T24B	0.61
RPE1 T24C	0.61

unmapped reads, two contain mapped reads, one is a .sam file and the last is a log file. If the bowtie command run in this step fails for any reason, the log file will note the error. Otherwise, it will log how long bowtie took to run at different stages and the numbers of reads processed, reads with alignments and reads that failed to align. Out of 31.5 M reads processed, ~18 M reads should align and 13 M reads should fail to align. The unmapped and mapped reads files are intermediate files not required by subsequent steps. Step 4 outputs the file 'HH-79_counts.txt', which is processed by Step 5 into the file 'rawCounts.txt'. Lastly, Step 6 is optional because the output file is included in the Orthrus R package, but is named 'procedure1_reads.tsv'.

Procedure 2

Key results for the processing phase are QC plots, which are listed in Table 5. These include, among many other plots, a plot of total reads and a heatmap of LFCs across all replicates. The total reads for each screen should correspond to expected reads based on sequencing depth for each screen (Fig. 5). To interpret the heatmap, for typical experimental designs, the most important features to examine are whether or not T0 screens cluster separately from other screens and whether or not technical replicates cluster within their respective screens (Fig. 6). The output AUC values for recovering essential genes should hover close to 1 for whole-genome screens as aforementioned, although for this highly specialized library with few essential genes and many strong expected effects, we see values between 0.6 and 0.7 (Table 10).

Dual-targeted scoring should output scored Torin-specific GIs in HAP1 cells for two timepoints, T12 and T18. These results are plotted in the files 'torin_vs_hap1_t12.png' and 'torin_vs_hap1_t18.png', and the scored data is contained in the files 'dual_targeted_gene_calls_t12.tsv' and 'dual_targeted_gene_calls_t18.tsv'. These output plots should resemble a slightly skewed fit between data that is mostly correlated with many outliers representing GIs. The null model, indirectly shown by gray, noninteracting genes that correlate well across both screens, reflects the loess normalization of residual values performed to account for skewed data or nonlinear trends (Fig. 7).

Residual LFC values across all guides for all significant hits are also output from dual-targeted scoring (automatically with batch and wrapper scoring), and they should reflect consistent positive

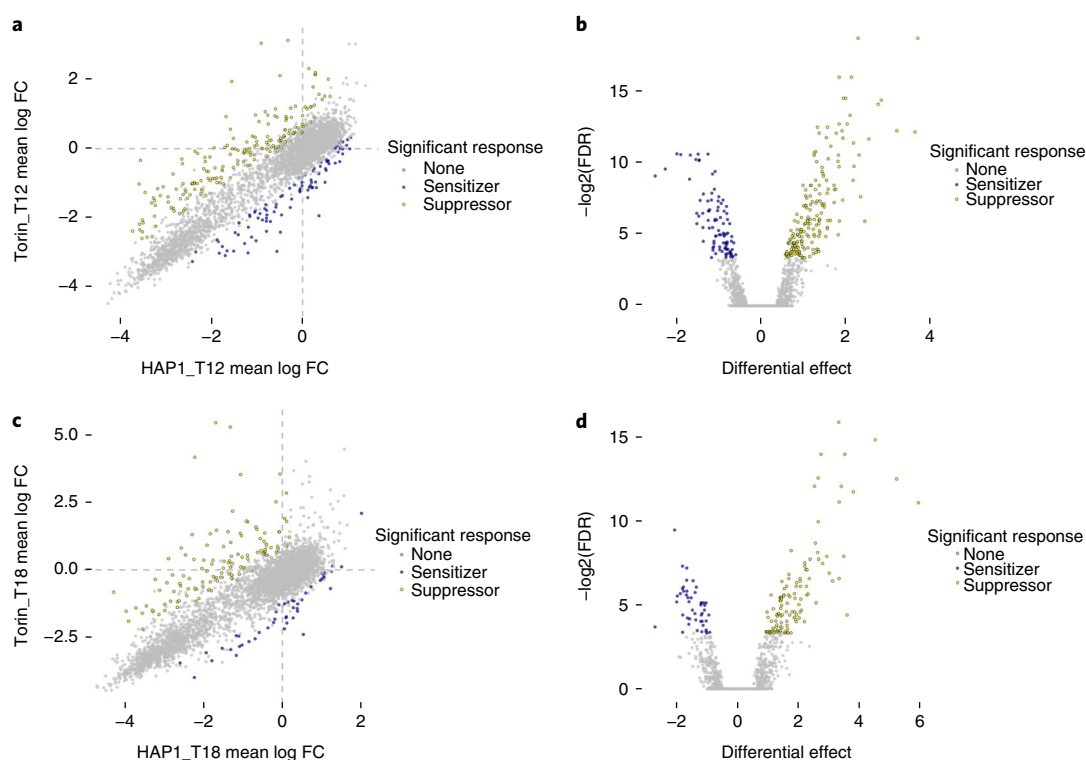


Fig. 7 | Summary plots of mean LFC for dual-targeting guides. The plots are colored by significant effects, for WT HAP1 and HAP1 + Torin1 screening data for all $n = 3,870$ gene pairs with dual-targeting guides analyzed in Procedure 2, Steps 11–14. Gene-level drug–gene interactions, where sensitizer interactions indicate that the gene’s knockout confers increased sensitivity to Torin-1 and suppressor interactions indicate that the knockout bypasses potentially deleterious effects of Torin-1 on cell fitness, are shown as colored points that significantly deviate from the computed null model. Blue points are sensitizing interactions with mean residual effects < -0.5 and Benjamini and Yekutieli FDRs < 0.1 , while yellow points are suppressor interactions with mean residual effects > 0.5 and Benjamini and Yekutieli FDRs < 0.1 . **a,b**, Scores for T12 data in a scatter plot (**a**) and a volcano plot (**b**) for 182 suppressor interactions and 93 sensitizer interactions. **c,d**, Scores for T18 data in a scatter plot (**c**) and a volcano plot (**d**) for 114 suppressor interactions and 47 sensitizer interactions. These plots allow users to contextualize effect size, effect strength and statistical significance for both WT HAP1 data at T12 and WT RPE1 data at T24.

and negative effects for well-performing guides (Fig. 7). While context affects the definition of positive and negative effects, for chemogenetic screens, negative effects typically represent sensitizers and positive effects represent suppressors, and for GI screens, negative effects typically represent synthetic sick or lethal interactions while positive effects represent buffering interactions²⁵. The top-ranked negative hit in WT HAP1 screening data at T18, HECTD1 (Fig. 8a), shows three such well-performing guides with consistent differential effects. On the other hand, the fourth-ranked negative hit, INPPL1 (Fig. 8b), shows mostly positive differential effects despite being called a negative hit. This is a red flag for the quality of this hit, and closer examination reveals that it is called as a result of how the loess-normalized null model poorly fits the handful of points with the strongest negative and positive expected effects. This hit should be ignored, and the user can consider tightening the effect size threshold to account for this during scoring. As another example, the eighth-ranked negative hit TAF5L (Fig. 8c) shows another case where three of its four guides display strong negative phenotypes, but the fourth showed little phenotype. This is not a red flag for hit quality, but could be a red flag for the guide with the smallest differential effect.

Combinatorial scoring should output scored interactions for ~700 paralog gene pairs for all six screens. Similarly, six plots summarizing scored paralog gene pairs that resemble dual-targeted scoring plots should be output, one for each screen, with the scored data available in the file ‘paralog_gene_calls.tsv’. Two of these plots, for WT HAP1 T12 and WT RPE1 T24 data, are shown in Fig. 9. Plots of residual effect values across guides for all significant hits for the HAP1 with Torin T18 and untreated HAP1 screens should also be output in the subfolders ‘HAP1_T18_combn’ and ‘Torin_T18_combn’ of the qc folder, respectively. The top negative and positive hits for WT HAP1 T12 data are shown in Fig. 10.

To examine specific hits for data scored with Orthrus, we recommend the following process. First, establish clear statistical significance and effect size thresholds when calling significant hits, which

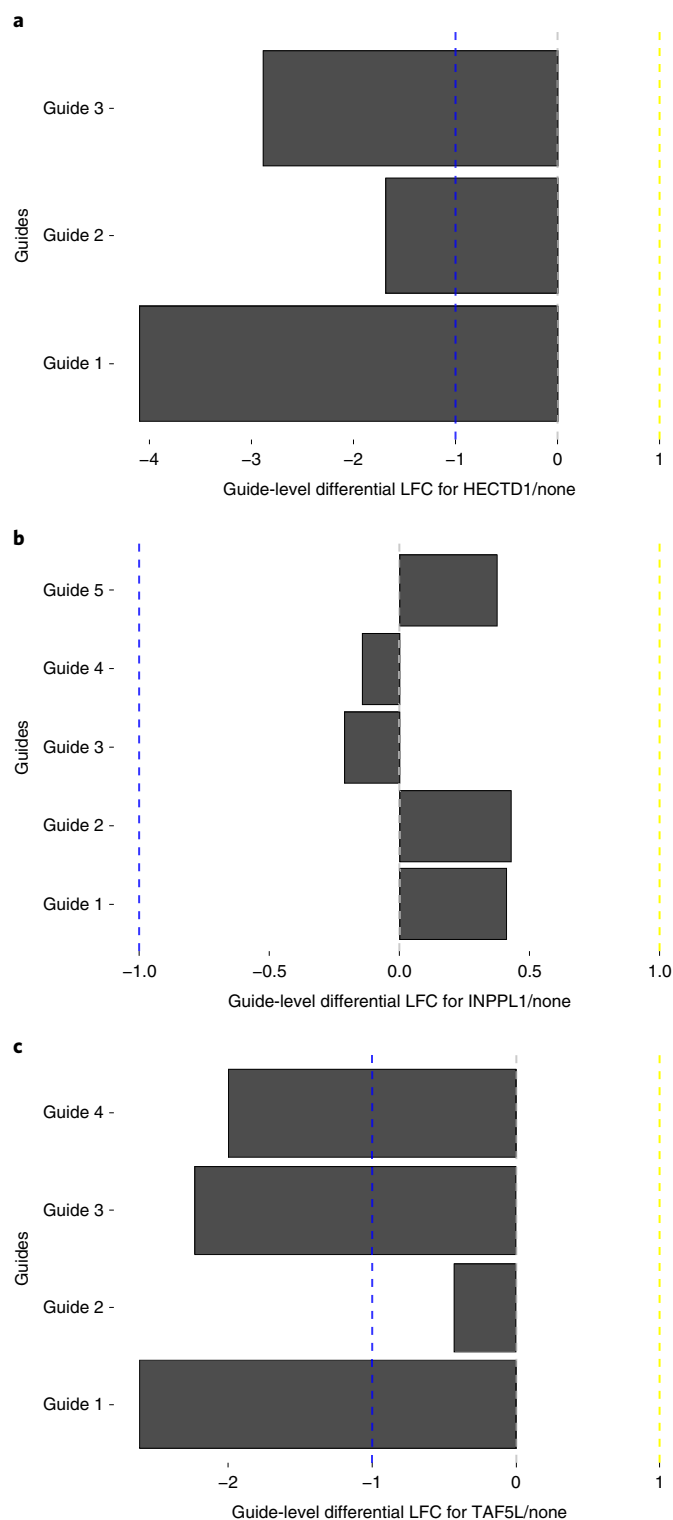


Fig. 8 | Differential LFC for WT HAP1 guides from the ChyMERa dataset analyzed in Procedure 2. Differential LFC for three significant hits of the scored dual-targeting guides at T18. Genes were scored based on deviations from a loess-corrected null model, so the agreement of individual guides of interesting hits should be examined to qualitatively confirm hit quality. Dashed lines are plotted at -1 (blue), 0 (gray) and 1 (yellow) differential LFC values for ease of interpretation across multiple plots. **a**, The top-ranked negative hit, HECTD1, with strong agreement for all three guides. **b**, The fourth-ranked negative hit, INPPL1, whose guides indicate a lower hit quality due to three out of five guides possessing positive residual LFCs. **c**, The eighth-ranked negative hit TAF5L, with strong agreement for three guides and no phenotype for the fourth.

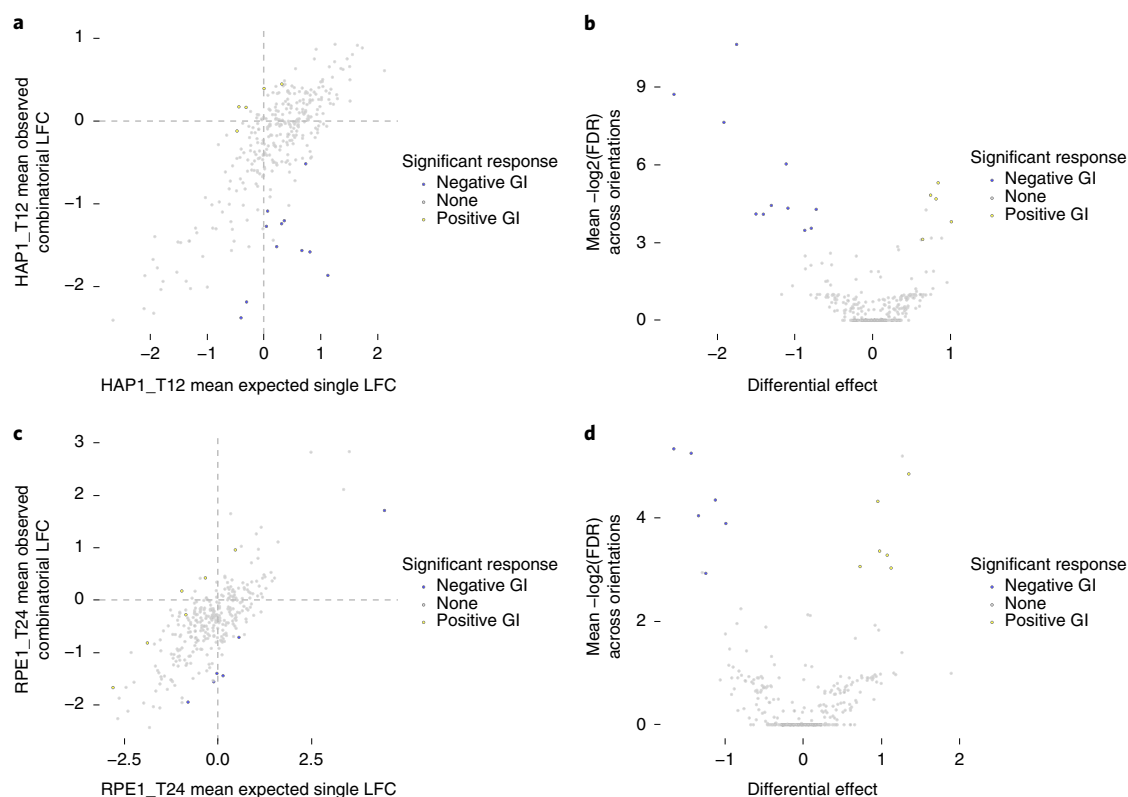


Fig. 9 | Summary plots of mean LFC for combinatorial-targeting guides. The plots are colored by significant effects for all $n = 313$ CHyMeRA gene pairs with combinatorial- and single-targeting guides analyzed in Procedure 2, Steps 15–17. Gene-level GIs are shown as colored points that significantly deviate from the computed null model. Blue points are negative GIs with mean residual effects < -0.5 and Benjamini and Yekutieli FDRs < 0.2 , while yellow points are positive GIs with mean residual effects > 0.5 and Benjamini and Yekutieli FDRs < 0.2 . **a,b**, Scores for WT HAP1 data at T12 in a scatter plot (**a**) and a volcano plot (**b**) for 5 positive GIs and 11 negative GIs. **c,d**, Scores for WT RPE1 data at T24 in a scatter plot (**c**) and a volcano plot (**d**) for six positive GIs and six negative GIs. These plots allow users to contextualize effect size, effect strength and statistical significance for both WT HAP1 data at T12 and WT RPE1 data at T24.

may be different for different guide libraries and experiments. Second, to look at either negative (e.g., synthetic lethal) or positive (e.g., buffering) GIs, subset the scored data to those that are labeled as negative or positive hits in the ‘effect_type_[CONDITION]’ column. Third, sort the data by the differential effect column, which for dual-targeting scored data is ‘differential_[CONDITION]_vs_[CONTROL]’ and for combinatorial-targeting scored data is ‘differential_combn_vs_single_[CONDITION]’.

The output for batch scoring performed in Step 19 mirrors the output for the manual scoring performed in Steps 11–17, but contains additional residual LFC plot folders, as Steps 13 and 16 only generated residual LFC plots for four out of eight screens.

Procedure 3

The anticipated results for Procedure 3 mirror the anticipated results for Procedure 2, although dual-targeting plots and scores are not output because the analyzed guide library only contains combinatorial-targeting and single-targeting guides. Like for Procedure 2, all anticipated results are provided at <https://zenodo.org/record/4527616>.

A handful of differences between the anticipated output of the two procedures exist. First, the scored data for Procedure 3 is only contained in the file ‘combn_gene_calls.tsv’. Accordingly, the ‘plots’ folder only contains three corresponding scatterplots and three volcano plots, one for each screen. Second, because the library contains fewer than ten guides that dual-target one nonessential gene or combinatorially target two nonessential genes in the Hart et al. 2014 nonessential standard²⁴, the file ‘essential_PR_QC.tsv’ reports NA values for the AUC of recovering essential genes compared with nonessential genes under ROC curves. Third, we chose not to apply loess normalization to score the dataset because of the wide spread of points at the left tail at the plot with strong negative

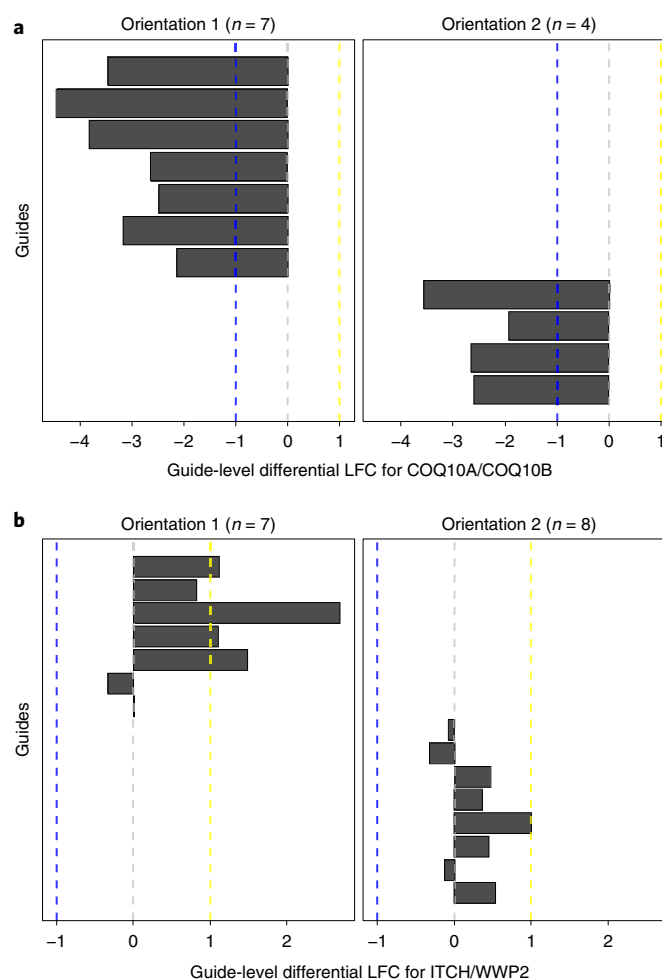


Fig. 10 | Differential LFC for WT HAP1 guides comprising two significant hits of the scored combinatorial-targeting guides at T12 from the ChyMERa dataset analyzed in Procedure 2, Steps 15–17. Genes were scored based on deviations from a loess-corrected null model, so the agreement of individual guides of interesting hits should be examined to qualitatively confirm hit quality. Dashed lines are plotted at -1 (blue), 0 (gray) and 1 (yellow) differential LFC values for ease of interpretation across multiple plots. **a**, The top-ranked negative hit, COQ10A and COQ10B, whose guides in both orientations strongly agree. **b**, The top-ranked positive hit, ITCH and WWP2, with agreement for five out of seven guides for orientation 1 and agreement for five out of eight guides for orientation 2. The remaining guides show weak phenotypes, indicating poor performance for those guides and good agreement for all other guides.

phenotypes for both the expected and observed combinatorial LFCs. Loess typically overfits points such as these, and for this dataset, a visual examination of the A549 screen shows that the linear fit to residuals compared with loess correction avoids both likely false positives and false negatives (Fig. 4). The scored data shown in the plots folder thus represent plots scored against a linear fit with Step 4B as opposed to loess-corrected residuals calculated in Step 4A.

Reporting Summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The example dataset is downloadable with the Orthrus package at <https://github.com/csbio/Orthrus>. The expected output from all procedures is provided under a CC-BY 4.0 license at <https://zenodo.org/record/4527616>.

Code availability

The Orthrus package is available at <https://github.com/csbio/Orthrus>, and the version of the code run in the protocol is available at <https://zenodo.org/record/4827171> (ref. ²⁰). All code presented from all procedures is also available in separate scripts along with their expected output, and are provided under a CC-BY 4.0 license in the Zenodo repository at <https://zenodo.org/record/4527616>. Code contained in this repository generated Figs. 4–10. The code in this protocol has been peer reviewed.

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Author contributions

H.N.W. wrote the software and performed all analyses. H.N.W. drafted the protocol, and C.L.M., M.A., T.G.-P., M.B., K.R.B. and T.K.O. provided revisions. H.N.W. and M.B. developed the scoring procedure implemented in Orthrus based on conceptual contributions from M.B. T.K.O. provided feedback to improve the software. M.A., T.G.-P., M.B., K.R.B., J.M., B.J.B. and C.L.M. developed the CHyMERa experimental platform. M.A., T.G.-P. and K.R.B. performed experiments to generate the data analyzed in Procedure 2. K.R.B. contributed data, code and text to Procedure 1. J.M., B.J.B. and C.L.M. acquired funding to support this work and provided supervision throughout the project.

Competing interests

A patent application (no. GB 1907733.8) describing the development and applications of CHyMERa, to the University of Toronto and T.G.-P., M.A., K.R.B., J.M. and B.J.B., is pending. J.M. previously performed sponsored research for Repare Therapeutics.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41596-021-00596-0>.

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Gonatopoulos-Pournatzis, T. et al. *Nat. Biotechnol.* **38**, 638–648 (2020): <https://doi.org/10.1038/s41587-020-0437-z>
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| Data analysis | The Orthrus scoring package is available at https://github.com/csbio/orthrus . All analyses presented in the protocol, which were also used to generate all panels of Figures 4-9, are available in a script format along with their expected output at https://zenodo.org/record/4527616 . |

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Sample size	Sample size for analyses was defined by the number of remaining gene pairs post-filtering for guides with low T0 read counts.
Data exclusions	The protocol focuses on analysis of only a subset of the screens present in the example CHyMERa dataset for the purposes of clarity.
Replication	Each screen consisted of three technical replicates which were extensively tested for reproducibility and expected effects for control genes as shown in the procedure.
Randomization	The analyzed screens, as detailed in Gonatopoulos-Pournatzis et al. 2020 and Dede et al. 2020, were divided into several groups based on cell type, timepoint and drug treatment. Randomization was not relevant for this experimental design.
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