Mapping DNA damage-dependent genetic interactions in yeast via party mating and barcode fusion genetics.

J. Javier Díaz-Mejía1-4, Albi Celaj1-4, Joseph C. Mellor1-6, Atina Coté1-3, Attila Balint1,7,8, Brandon Ho1,7, Pritpal Bansal1-3, Fatemeh Shaeri1-3, Marinella Gebbia1,2, Jochen Weile1-3, Marta Verby1,3, Anna Karkhanina1-3, YiFan Zhang1-3, Cassandra Wong2, Justin Rich1-3, D’Arcy Prendergast1-3, Gaurav Gupta1-3, Sedide Öztürk5,11, Daniel Durocher2,3, Grant W. Brown1,7, Frederick P. Roth1,5,9,10*

1 Donnelly Centre, University of Toronto, Toronto, ON M5S 3E1, Canada

2 Department of Molecular Genetics, University of Toronto, Toronto, ON M5S 3E1, Canada

3 Lunenfeld-Tanenbaum Research Institute, Mt. Sinai Hospital, Toronto, ON M5G 1X5, Canada

4 Department of Computer Science, University of Toronto, Toronto, ON M5S 3E1, Canada

5 Department of Biological Chemistry and Molecular Pharmacology, Harvard Medical School, Boston, MA 02115, USA

6 Present address: seqWell, Inc. Beverly, MA 01915, USA

7 Department of Biochemistry, University of Toronto, Toronto, ON M5S 3E1, Canada

8 Present address: Center for Chromosome Stability, Department of Cellular and Molecular Medicine, University of Copenhagen, Copenhagen, Denmark
Center for Cancer Systems Biology (CCSB) and Department of Cancer Biology, Dana-
Farber Cancer Institute, Boston, MA 02215, USA

Canadian Institute for Advanced Research, Toronto, ON M5G 1Z8, Canada

Roche Sequencing Solutions, Pleasanton, CA 94588, USA

*Corresponding author.

Running title: Barcode Fusion Genetic Interactions
Abstract

Condition-dependent genetic interactions can reveal functional relationships between genes that are not evident under standard culture conditions. State-of-the-art yeast genetic interaction mapping, which relies on robotic manipulation of arrays of double mutant strains, does not scale readily to multi-condition studies. Here we describe Barcode Fusion Genetics to map Genetic Interactions (BFG-GI), by which double mutant strains generated via en masse ‘party’ mating can also be monitored en masse for growth and genetic interactions. By using site-specific recombination to fuse two DNA barcodes, each representing a specific gene deletion, BFG-GI enables multiplexed quantitative tracking of double mutants via next-generation sequencing. We applied BFG-GI to a matrix of DNA repair genes under nine different conditions, including methyl methanesulfonate (MMS), 4-nitroquinoline 1-oxide (4NQO), bleomycin, zeocin, and three other DNA-damaging environments. BFG-GI recapitulated known genetic interactions and yielded new condition-dependent genetic interactions. We validated and further explored a subnetwork of condition-dependent genetic interactions involving MAG1, SLX4, and genes encoding the Shu complex, and inferred that loss of the Shu complex leads to a decrease in the activation or activity of the checkpoint protein kinase Rad53.

Key words: Genetic interaction/ DNA barcode/ Sequencing/ Condition-dependent/ En masse
Introduction

The importance of condition-dependent genetic interactions

Genetic interactions, defined by a surprising phenotype that is observed when mutations in two genes are combined (Mani et al, 2008), are powerful tools to infer gene and pathway functions (Baryshnikova et al, 2010; Ideker & Krogan, 2012). Of the genetic interactions currently known in any species, the vast majority were found using Synthetic Genetic Array (SGA) technology in *Saccharomyces cerevisiae* (Bandyopadhyay et al, 2010; Costanzo et al, 2010, 2016; van Leeuwen et al, 2016) and these studies have yielded a rich landscape of genetic interactions. The sign of genetic interaction (defined to be negative when mutants are synergistically deleterious, and positive when the combination is less severe than would be expected from independent effects) provides clues about whether the genes act in parallel or in a concerted or serial fashion. Measuring similarity between genetic interaction profiles, both at the level of single genes and of clusters of genes, has revealed a hierarchical map of eukaryotic gene function (Costanzo et al, 2010, 2016). However, the vast majority of genetic interaction mapping has been conducted under a single standard culture condition.

The importance and qualitative nature of gene function can change with environmental fluctuation, so that a complete understanding of genetic interaction will require mapping under multiple conditions. For example, pairs of DNA repair genes had 2-4 times more genetic interactions between DNA repair genes under MMS treatment compared with rich media alone (Bandyopadhyay et al, 2010; St Onge et al, 2007; Ideker & Krogan, 2012), so that a plethora of condition-dependent
genetic interactions remain to be uncovered via gene × gene × environment studies.

**Current genetic interaction discovery technologies**

Essentially every large-scale genetic interaction mapping strategy in yeast uses a genetic marker system developed for the SGA technique, which works by mating a single-gene deletion query strain with an array of different single-gene deletion strains from the Yeast Knockout Collection (YKO) (Giaever et al., 2002). The SGA system provided genetic markers by which mated diploids can be subjected to a series of selections to ultimately yield haploid double mutants. In ‘standard’ SGA mapping, the fitness of the resulting double mutants is determined by statistical analysis of the images from each plate, yielding cell growth estimates for each separately-arrayed strain (Tong & Boone, 2005). SGA has also been used to study genetic interactions within functionally-enriched gene groups (Collins et al., 2006) and has been applied to detect environment-dependent interactions (Bandyopadhyay et al., 2010; St Onge et al., 2007). For example, St Onge et al. (2007) used the SGA markers to generate all pairwise double mutants between 26 DNA repair genes in yeast. The authors cultured each double mutant individually in microplates and monitored cell density over time to infer the fitness of double mutants and thereby identify genetic interactions in the presence and absence of MMS.

Others have measured genetic interactions via competition-based fitness measurements in liquid cultures, adding fluorescent markers for tracking cell viability, and using robotic manipulation to inoculate and measure cell growth (DeLuna et al., 2008; Garay et al., 2014). A recent technique called iSeq incorporated
barcodes into single-mutant strains, such that pairs of barcodes identifying corresponding pairs of deleted genes could be fused by Cre-mediated recombination (Jaffe et al., 2017). They demonstrated the method, showing that a pool corresponding to 9 gene pairs could be sequenced to monitor competitive growth of double-mutants en masse in different environments (Jaffe et al., 2017). Cre-mediated approaches have been used similarly to map protein-protein interactions (Yachie et al., 2016; Hastie and Pruitt, 2007; Schlecht et al., 2017).

For each of the above genetic interaction methods, double mutants were generated by individual mating of two specific yeast strains, requiring at least one distinct location for each double-mutant strain on an agar or microwell plate and necessitating robotic strain manipulation to achieve large scale. By contrast, other methods to map genetic interactions generated double mutants in a one-vs-many fashion. For example, diploid-based synthetic lethality analysis on microarrays (dSLAM) (Pan et al., 2004) disrupted a single ‘query’ gene by homologous recombination via transformation of a marker into a pool of diploid heterozygous deletion strains bearing the SGA marker. After selecting for double-mutant haploids from such ‘one-by-many’ haploid double-mutant pool, barcodes were PCR amplified from extracted double mutant DNA and hybridized to microarrays to infer the relative abundance and fitness of each double mutant. Another method, Genetic Interaction Mapping (GIM) (Decourty et al., 2008), generated a one-by-many pool of barcoded double mutants by en masse mating a single query strain to a pool of haploid gene deletion strains. Like dSLAM, GIM inferred strain abundance and fitness via barcode hybridization to microarrays. Despite the efficiency of generating one-by-many double-mutant pools, a matrix involving thousands of query strains...
would require thousands of such pools to be generated.

Each of the above methods has advantages and disadvantages. For example, measuring a growth time-course for each double-mutant strain provides high resolution fitness measurements (Garay et al., 2014; St Onge et al., 2007), but scalability is low. Standard SGA is high-throughput, but requires specialized equipment for robotic manipulation, and these manipulations must be repeated to test genetic interactions in new environments. The iSeq method shares the scaling challenge of SGA in strain construction, in that it requires many pairwise mating operations; however, once a double-mutant pool has been generated, it represents a promising strategy for measurement of competitive pools in different environments. The dSLAM and GIM methods allow generation of one-by-many pools, which reduces the number of mating operations, but both methods require customized microarrays as well as pool generation and microarray hybridization steps for every query mutation in the matrix.

Barcode Fusion Genetics to map Genetic Interactions (BFG-GI)

Here we describe BFG-GI, which borrows elements from several previous approaches. Like iSeq, BFG-GI requires generation of barcoded single-mutant strains, with only minimal use of robotics. To generate double-mutant pools, BFG-GI uses the SGA marker system and, like the GIM strategy, BFG-GI employs *en masse* mating. Unlike GIM and all other previous genetic interaction mapping strategies, BFG-GI employs many-by-many ‘party mating’ to generate all double mutants for a matrix of genes in a single mating step. All successive steps — including barcode fusion, sporulation, selection of haploid double mutants and measurement of
relative strain abundance — are also conducted *en masse*. We show that double mutants can be generated and monitored in competitive pools using BFG-GI. Like iSeq, BFG-GI infers double mutant fitness in competitively grown strain pools using next-generation sequencing of fused barcodes, and BFG-GI double mutant pools can be aliquoted and stored. Aliquots can be thawed later and challenged under specific environments (e.g. drugs) to detect condition-dependent genetic interactions without having to regenerate the double mutant strains.

We assessed BFG-GI by mapping genetic interactions of DNA repair-related genes under multiple DNA-damaging conditions, revealing many condition-dependent interactions and a discovery that perturbation of the Shu complex leads to decreased activation or activity of the Rad53 checkpoint protein kinase.
Results

**BFG-GI experimental design overview**

The first step in the BFG-GI process is generating uniquely barcoded donor and recipient strains from complementary mating types. Each donor and recipient strain contains a unique barcode locus. In the donor strain, this barcode is flanked by two distinct site-specific recombination sites (\textit{loxP}/2272 sites), while in the recipient strain, both recombination sites lie on the same side of the unique recipient barcode. After the mating step, these sites mediate barcode fusion via the Cre/Lox system, yielding chimeric barcode sites that uniquely identify specific deletion combinations. We created donors by crossing individual gene deletion strains from the YKO collection with proDonor strains that contained newly constructed pDonor plasmids (Fig 1A, Fig EV1 and Materials and Methods). We generated recipient strains by crossing individual gene deletion strains from the SGA query collection with proRecipient strains (Fig 1B, Fig EV2 and Materials and Methods). Haploid selection of double mutants followed mating of donor and recipient strains, sporulation, and \textit{in vivo} fusion of barcodes using Cre/Lox recombination (Fig 1C).

We confirmed that barcode fusion was successful using two neutral-insertion strains as controls. Specifically, we crossed a \textit{MATalpha Donor} ho\Delta::kanMX to a \textit{MATa Recipient} ylr179c\Delta::natMX and induced Cre/Lox recombination to fuse their barcodes. After sporulation and selection of the \textit{MATalpha} haploid double mutant progeny (Materials and Methods), we extracted genomic DNA, amplified barcode fusions by PCR and confirmed their integrity by Sanger sequencing (Fig 1C).
To scale up the BFG-GI process, we optimized mating and sporulation steps to generate double mutants with unique barcodes that had been fused *en masse* (Materials and Methods). We selected hundreds of double mutants using a series of marker selection steps in a many-by-many fashion. Intermediate selection steps allowed us to fuse barcodes representing each donor and recipient parental pair within each double mutant cell (Fig 1D and Materials and Methods).

Once we generated the pool of fused-barcode double mutants, aliquots were stored at -80°C for future experiments. Amplification and next-generation sequencing of fused barcodes in the pool allowed us to infer the relative abundance of each double mutant in each condition of interest (Fig 1D and Materials and Methods). In addition to haploid double mutant pools, we sequenced fused barcodes from the heterozygous diploid double mutant pools and used those as reference ('time zero') controls for fitness and genetic interaction calculations (Materials and Methods).

**BFG-GI measures strain abundances within a heterogeneous population**

We first evaluated the ability of BFG-GI to accurately detect the abundance of pooled double mutant strains. To generate reference data for this evaluation, we used the array-based SGA strategy to generate 2,800 double mutants by individual mating of barcoded BFG-GI strains, subsequently inducing barcode fusion via the Cre/Lox system. The purpose of this experiment was to assess the extent to which quantifying growth via fused-barcode-sequencing of pooled strains could recapitulate the measurements of growth in individual cell patches (as in conventional SGA). We recorded patch sizes, scraped plates to pool all double
mutant cells, extracted genomic DNA, and sequenced the fused barcodes (Materials and Methods). The resulting numbers of sequencing reads for each strain was strongly correlated with the corresponding colony sizes ($r = 0.92$, Fig 2A).

Importantly, colonies that were very small or absent often corresponded to double mutants with very few or no sequencing reads. These results show that BFG-GI detects the abundance of specific double mutants in pools of cells, with results comparable to an array-based method.

**Generating a DNA repair-focused double-mutant strain pool**

To test whether BFG-GI can accurately map genetic interactions, we generated a double mutant pool focused on DNA repair genes and compared BFG-GI results to those of other validated genetic interaction assays. We began by generating donor and recipient strains by crossing 35 YKO ($yfg1\Delta::kanMX, MATa$) single gene deletion strains to 65 BFG-GI proDonor strains, and 38 SGA query ($yfg2\Delta::natMX, MATalpha$) single gene deletion strains to 71 BFG-GI proRecipient strains. The set of deleted genes to which these strains correspond include 26 DNA repair genes from a previous condition-dependent genetic interaction study (St Onge et al, 2007), as well as 14 likely-neutral loci (i.e. the already-disrupted $HO$ locus, pseudogenes, and other loci for which single- and double-mutant phenotypes have not been previously observed). Inclusion of neutral loci allowed us to infer single mutant fitness from pools of double mutants (Materials and Methods).

To generate haploid double mutants, donor and recipient cells were scraped from plates and all subsequent steps in the BFG-GI pipeline were conducted en masse. First, the pools were combined for party mating. Seven selection steps
followed mating, including four that correspond to those in the standard SGA procedure: heterozygous diploid selection, sporulation, $MAT\text{a}$ progeny selection, and haploid double mutant selection. Additionally, before sporulation, we completed three selection steps to fuse barcodes and subsequently remove Cre to limit additional recombination events (Fig 1C and Fig EV3). This generated a pool of 4,288 haploid double mutants, which was aliquoted and stored as frozen glycerol stock. Thawed samples were used to inoculate solid media appropriate for selecting haploid double mutant cells. The media was used alone, supplemented with dimethyl sulfoxide (DMSO) as a solvent control, or supplemented with one of eight drugs targeting DNA repair pathways (Table EV1). We extracted genomic DNA, amplified and sequenced fused barcodes to infer the relative abundance of each double mutant in each condition.

To evaluate assay reproducibility, we ran all BFG-GI procedures in duplicate, starting from the mating step (technical replicates) and also barcoded multiple strains representing the same gene (biological replicates). Biological replicate strains had either the same or different parental strain origin (the parental strain for a given gene deletion might be from either the YKO or SGA query strain collection). Relative strain abundance was highly correlated between technical replicates ($r > 0.95$). Next, we used a multiplicative model to infer a genetic interaction score ($GIS$) from relative strain abundances, analogous to other methods based on strain growth (Materials and Methods). As the relative strain abundance, $GIS$ correlation between technical replicates was also high ($r = 0.96$).

Correlation of $GIS$ profiles between biological replicates representing the same gene were generally high, with 85% of replicates showing $GIS$ $r > 0.5$. We
computationally excluded from analysis 21 biological replicates (six donors and 15 recipients) showing GIS correlations with $r < 0.5$. For the remaining strains, biological replicate profiles clearly showed higher correlation than did profiles from pairs of randomly-chosen replicates carrying deletions in different genes (Fig 2B). To understand factors contributing to poorly-correlated replicate pairs we sequenced the genomes of 20 strain pairs. Ten of those pairs corresponded to strains with $GIS\ r < 0.5$ and other 10 with $GIS\ r > 0.5$. We found that all 10 strain pairs with $GIS\ r < 0.5$ had chromosome V duplicated in one of the two strains, in agreement with the report of iSeq strains showing low strain profile reproducibility, owing to this same chromosome V duplication (Jaffe et al, 2017). Chromosome V contains the CAN1 locus, the locus at which both BFI-GI recipients and iSeq strain constructs are inserted. By contrast, only 3 out of 10 strain pairs with $r > 0.5$ showed aneuploidies in just one strain in the pair (for these strains, the aneuploidies were also in chromosome V). All BFG-GI strains showing aneuploidies were recipients. This suggests that future versions of BFG-GI recipients for which selection markers are carried by plasmids may increase reproducibility, as we found for our Donor strains. Furthermore, we removed strains with poor representation in the heterozygous diploid pool, because GIS profiles from these strains yielded neutral scores even for controls ('same-gene' pairs described below) that should behave like strong negative interactions, presumably due to poor statistical power to detect fitness effects (Fig. EV4B). Our final dataset consisted of 3,232 double mutants, with 59 Donors and 56 Recipients, representing 39 genes (25 DNA repair genes and 14 neutral genes; Figure EV4A and Table EV2). Replicates representing swc5Δ showed very low relative abundance in the sequencing results and were removed from
subsequent analyses. Finally, GIS measurements for technical and biological replicates (Table EV3) were combined into a single score for each gene pair (Table EV4; Methods).

We next assessed the ability of BFG-GI to infer the fitness for three classes of double mutant strains. First, we measured the abundance of strains carrying two differently-barcoded mutations corresponding to the same gene. Compound heterozygous diploids bearing a mutation at both loci for a given gene (e.g. $MMS4/mms4\Delta::kanMX mms4\Delta::natMX/MMS4$) can survive in media supplemented with selective antibiotics; however, haploid cells derived from this parental diploid should not survive because they should only carry one locus for each gene and therefore only one of the two antibiotic resistance markers required to survive the selection. Thus, haploid strains for ‘same-gene pairs’ are expected to exhibit reduced fitness, behaving like synthetic lethal combinations, and be depleted from the pools. The calculated GIS agreed with this expectation (Fig 2C). Second, we assessed the abundance of double mutants representing pairs of linked genes (<75 kbp apart; Fig EV4C). Independent segregation is reduced between linked genes, and as expected our GIS indicated these double mutants were also depleted from the pools (Fig 2C). Third, we analyzed double mutants representing unlinked genes and we found that their GIS distribution is clearly distinguishable from same-gene and linked gene pairs (Fig 2C).

Finally, we sought to compare BFG-GI results against a dataset of genetic interactions (St Onge et al, 2007), both to obtain an overall evaluation of our method and as a way to calibrate our GIS score thresholds for calling genetic interactions. We first compared BFG-GI GIS scores with the Epsilon scores reported by St Onge et
al (2007) under both no-drug and MMS conditions, for pairs of DNA repair genes that had been tested in both studies. We found that GIS and Epsilon scores correlated well with each other in both no-drug ($r = 0.8$) and MMS ($r = 0.85$) conditions (Fig 2D and Fig 2E). Taking both conditions together, and using GIS score thresholds with an estimated 5% false positive rate, BFG-GI captured 56% of the positive genetic interactions reported by St. Onge et al. and 66% of the negative genetic interactions (Fig 2F), while reporting an additional 23 positive and 20 negative interactions not reported by St. Onge et al (2007).

Taken together, these results provide evidence that BFG-GI offers a powerful means of generating double mutants by en masse mating and monitoring strain abundance in a multiplexed fashion to infer condition-dependent genetic interactions.

**BFG-GI reveals condition-dependent genetic interactions**

Having determined that BFG-GI can accurately detect genetic interactions, we analyzed the same double mutant pool under seven additional culture conditions to more broadly explore condition-dependent genetic interactions (see Fig 3C legend for condition names and Table EV1 for details). To call positive and negative interactions, we first standardized GIS by the estimated error ($Z_{GIS}$; Materials and Methods), and used the distribution of $Z_{GIS}$ amongst unlinked barcode pairs containing a neutral gene (‘neutral pairs’, Fig EV4A) to calculate a false discovery rate (FDR) at a given $Z_{GIS}$ cutoff (Fig EV4 D-E). To call interactions, we used both a $Z_{GIS}$ cutoff corresponding to FDR = 0.01 in each condition, and an additional effect-size cutoff ($\lvert GIS \rvert > 0.075$) to filter out interactions of high confidence but low
magnitude. At these cutoffs, 91% of the called negative interactions and 77% of the
called positive interactions were also observed in a previous study (St Onge et al,
2007), while 64% of the previously-reported negative and 44% of the previously-
reported positive interactions were reproduced by BFG-GI (Fig EV4F; Table EV4).

Analyzing BFG-GI results further, we found that all DNA repair genes showed
at least one genetic interaction, and that some genes showed markedly more
interactions than others. For example, we found that the DNA helicase gene SGS1
yielded negative interactions with MMS4, MUS81 or SLX4 (all of which participate in
template switching during break-induced replication) in all nine conditions (Fig 3A,
Table EV4). Another DNA helicase gene, SRS2, interacted negatively with both SGS1
and the DNA translocase gene RAD54 in all nine conditions. By contrast, a third DNA
helicase/ubiquitin ligase gene, RAD5, showed positive genetic interactions with
SGS1 in six conditions. SGS1 and SRS2 are involved in error-free DNA damage
tolerance, while RAD5 is involved in recombinational repair of double-strand
breaks. These findings coincide with previous reports showing SGS1 and SRS2
centrality in DNA repair pathways in both unperturbed and MMS-induced stress
conditions (St Onge et al, 2007).

We next examined condition-dependent changes in genetic interactions.
First, genetic interaction differences between conditions were calculated (ΔGIS).
Then, using a similar approach to calling genetic interactions within each condition,
ΔGIS was standardized by the estimated error (ΔZGIS), and the distribution of ΔZGIS
amongst neutral pairs was used to calculate an FDR for each differential interaction
(Fig EV5A; Materials and Methods). At a ΔZGIS cutoff corresponding to FDR = 0.01
and an effect size cutoff of |GIS| > 0.1, we identified 2,932 differential interactions
amongst DNA damage genes, and further considered only the subset of 2,335
differential interactions that changed between interaction type (i.e., between the
three classes of positive, negative and neutral) for further analysis. For any given
pair of conditions, an average of 9% of all gene pairs exhibited differential
interaction. For example, we found \textit{mus81Δ/rad5Δ} displayed a negative genetic
interaction in DMSO, a positive genetic interaction in MMS, and a significant
difference between the two conditions. This change is shown as a red edge in Fig 3B,
panel \textit{i}, and agrees with a previous report (St Onge \textit{et al}, 2007). By contrast, most
changes in genetic interaction between DMSO and MMS were from neutrality in one
condition to either a positive or negative genetic interaction in the other (Fig 3B,
panels \textit{i} and \textit{iv}). Generalizing this observation to all pairwise condition comparisons,
a large majority of significant differential genetic interactions were neutral in one
condition and either positive or negative in the other (94%), and thus only 6% of
significant genetic interactions changed sign between conditions (Fig 3C and Table
EV\#5).

Genes differed both in the number of total number differential genetic
interactions in which they participated (Fig EV5B) and in the number of their
differential genetic interactions involving a change in sign (Fig EV5C). Genetic
interactions involving \textit{RAD5} were especially dynamic - \textit{RAD5} participated in 233
significant differential genetic interactions (out of 1224 comparisons; Fig EV5B),
and 55 of these involved sign reversals (Fig EV5C). Out of 55 sign-reversed
differential genetic interactions involving \textit{RAD5}, 48 involved \textit{MMS4, MUS81, RAD51,}
\textit{RAD54}, or \textit{RAD55} (Fig EV5D). \textit{MUS81} and \textit{MMS4} encode a heterodimer which
cleaves nicked intermediates in recombinational DNA repair (Schwartz \textit{et al}, 2012),
while \textit{RAD51} binds ssDNA to facilitate homologous recombination, and requires \textit{RAD54} and \textit{RAD55} for its activity (Sugawara et al, 2003). Genetic interactions of \textit{RAD5} were often positive for all five of these genes in 4NQO and MMS, and negative with all five in other tested conditions (Fig EV5D). These findings are consistent with previously-reported negative interactions of \textit{RAD5} with these genes in MMS and positive interactions when no drug stress is added (St. Onge \textit{et al}, 2007; Table EV4). The dynamic interactions of \textit{RAD5} with these two gene groups may reflect the previously-reported multifunctional nature of \textit{RAD5} and its ability to coordinate repair events and replication fork progression differently in response to different types of lesions (Choi \textit{et al}, 2015).

We assessed similarity between growth conditions as measured by similarity between patterns of \textit{GIS} profiles. As expected, the two conditions most similar to each other were no-drug and DMSO, which also yielded no significant between-condition differential interactions (Table EV#5). A hierarchical clustering of conditions by their \textit{GIS} profiles (Fig 3C) showed that pairs of drugs with similar mechanisms of action clustered together. For example, bleomycin and zeocin, which are members of the same family of glycopeptides that intercalate into DNA to induce double strand breaks (Claussen & Long, 1999) were grouped as nearest neighbours and also had the least number of differential interactions between any two drug pairs (26, compared to an average of 67 across all condition pairs).

Interestingly, MMS and 4NQO were also grouped as nearest neighbours. Although there were a large number of differential interactions between them (75), the vast majority (73) showed neutrality in one condition and negative genetic interaction in the other. MMS and 4NQO are members of different drug classes but
both are DNA alkylating agents (Svensson et al, 2012; Xiao & Chow, 1998). Both MMS and 4NQO cause checkpoint modulated fork stalling (Minca & Kowalski, 2011; Iyer & Rhind, 2017) that appears to facilitate replication of damaged templates allowing forks to quickly pass lesions (Iyer & Rhind, 2017). Furthermore, strains carrying deletion of genes involved in postreplication repair (PRR) processes, such as MMS2, RAD5 and UBC13 are significantly hypersensitive to both MMS and 4NQO (Lee et al, 2014), suggesting that PRR acts on both MMS and 4NQO lesions. DNA lesions caused by these drugs are typically corrected by either base-excision repair (MMS) or nucleotide-excision repair (4NNO), and these pathways are synergistic with each other in genetic backgrounds like mag1Δ (Xiao & Chow, 1998). We believe that these mechanistic similarities between MMS and 4NQO contributed to the similarity between their GIS profiles in in comparison to those from other drugs we tested.

The most divergent condition pairs (those yielding the highest number of differential interactions) were MMS vs. doxorubicin (104 changes) and MMS vs. bleomycin (110 changes). These results are consistent with the fact that MMS, doxorubicin, and bleomycin have different mechanisms of action and cause DNA lesions that are repaired by different pathways.

A condition-dependent subnetwork of MAG1, SLX4 and Shu complex genes

The Shu complex (a heterotetrameric protein complex consisting of Csm2, Psy3, Shu1, and Shu2) promotes Rad51 filament formation and homologous recombination during error-free lesion bypass, double strand break repair, and
meiosis (Ball et al, 2009; Bernstein et al, 2011; Godin et al, 2013; Mankouri et al, 2007; Sasanuma et al, 2013) (Fig 4A). Our BFG-GI results indicated that genes encoding all four members of the Shu complex showed negative genetic interactions with both MAG1 and SLX4 during exposure to MMS. Additionally, the Shu complex genes interacted negatively with SLX4 during treatment with 4NQO, bleomycin, and zeocin (Fig 4B). Mag1 is a 3-methyladenine DNA glycosylase that removes alkylated bases from DNA to initiate base-excision repair (BER), thereby protecting cells against alkylating agents like MMS (Berdal et al, 1990; Chen et al, 1990). Slx4 promotes the activity of three structure-specific endonucleases (Flott et al, 2007; Fricke & Brill, 2003; Gritenaite et al, 2014; Toh et al, 2010) and, upon exposure to MMS, plays a key role in down-regulating phosphorylation of the checkpoint kinase Rad53 (Jablonowski et al, 2015; Ohouo et al, 2013). We generated double mutants for each Shu complex member in combination with either MAG1 or SLX4 and tested fitness on media containing DMSO or various genotoxins using spot dilution assays (Fig 4C). Our results validated the MAG1-Shu complex interaction in MMS that we detected with BFG-GI, and are consistent with a previous study (Godin et al, 2016). The negative interactions between MAG1 and Shu complex members are explained (Godin et al, 2016) by the fact that these double mutants have simultaneously lost Mag1-mediated BER (which directly removes alkylated bases) and have a diminished capacity for error-free lesion bypass, a major pathway used during MMS-induced blocks in DNA replication (Huang et al, 2013)(Fig 4A). Our spot dilution assays also confirmed that MAG1 interacts negatively with SLX4 during MMS treatment (Fig 4C). This result is also consistent with a previous study showing that BER is unlikely to be the major function of SLX4 (Flott et al, 2007). Of
particular interest, we validated the BFG-GI interactions between Shu complex members and SLX4 during treatment with MMS, 4NQO, bleomycin, and zeocin (Fig 4C).

As the nature of the SLX4 interactions with genes encoding Shu complex proteins is unknown, we studied them in more detail. That there are negative genetic interactions between SLX4 and Shu complex members in MMS was unexpected, given that the Shu complex promotes error free lesion bypass (Ball et al, 2009; Godin et al, 2016; Mankouri et al, 2007; Xu et al, 2013) and SLX4 is epistatic to genes that regulate error-free lesion bypass during MMS treatment (Flott et al, 2007). A major role for Slx4 under MMS conditions is down-regulating phosphorylation and activation of Rad53, which occurs by Slx4 competing with Rad9 for binding to Dpb11 in order to limit the formation of Rad9-Dpb11 complexes that activate Rad53 (Cussiol et al, 2015; Jablonowski et al, 2015; Ohouo et al, 2013; Pfander & Diffley, 2011). Alternatively, levels of phosphorylated Rad53 are increased in cells deleted for PPH3, which encodes the catalytic subunit of the protein phosphatase PP4 complex that binds and dephosphorylates Rad53 during MMS treatment (O’Neill et al, 2007). Deletions of either SLX4 or PPH3 or both genes result in hyperactivation of Rad53 and hypersensitivity to MMS (Jablonowski et al, 2015). This phenotype is suppressed by expression of a hypomorphic rad53-R605A allele (Cussiol et al, 2015; Jablonowski et al, 2015; Ohouo et al, 2013). To determine whether the genetic interactions between SLX4 and Shu complex members (Fig 4C) reveal an unanticipated role for the Shu complex regulating activation of Rad53 (Fig 4D), we tested the sensitivity of pph3Δ/Shu complex double mutants to MMS using spot dilution assays. Combining pph3Δ with deletion of any of the Shu complex
genes resulted in a dramatic increase in MMS sensitivity relative to the single mutants (Fig 4E), indicating negative genetic interactions similar to those seen between \textit{SLX4} and Shu complex members (Fig 4C), or between \textit{SLX4} and \textit{PPH3} (Jablonowski \textit{et al}, 2015).

To assess MMS-induced Rad53 activation in Shu complex mutants more directly, we monitored Rad53 phosphorylation (which is a proxy for Rad53 activation) using western blot assays. Consistent with the role of \textit{SLX4} in dampening Rad53 activation (Balint \textit{et al}, 2015; Jablonowski \textit{et al}, 2015; Ohouo \textit{et al}, 2013), \textit{slx4Δ} cells challenged with MMS showed an increase in Rad53-P levels relative to wild type (Fig 4F). Interestingly, three of the Shu complex mutants (\textit{csm2Δ}, \textit{psy3Δ}, and \textit{shu1Δ}) also showed an increase in Rad53-P levels upon treatment with MMS (Fig 4F), indicating that these Shu complex mutants, like \textit{slx4Δ} and \textit{pph3Δ} cells, display hyperactivated Rad53 under exposure to MMS. We asked whether the MMS-sensitivity of Shu complex mutants could be suppressed by expression of the \textit{rad53-R605A} allele. Expression of \textit{rad53-R605A}, which is not effectively hyper-activated, suppresses the MMS sensitivity of \textit{slx4Δ} and \textit{pph3Δ} (Jablonowski \textit{et al}, 2015; Ohouo \textit{et al}, 2013). Similarly, the MMS sensitivity of \textit{csm2Δ}, \textit{psy3Δ}, \textit{shu1Δ} and \textit{shu2Δ} mutants was partially suppressed by \textit{rad53-R605A} (Fig 4G). Together, our data indicate that the Shu complex, like Slx4 and Pph3, leads to a decrease in Rad53 activation or activity in response to MMS treatment, as revealed by unique condition-dependent genetic interactions detected by BFG-GI.
Discussion

We developed a new technology, called BFG-GI, in which pools of double mutant yeast strains corresponding to a matrix of target genes are generated en masse through many × many ‘party’ mating. These pools are induced to form double-mutant-identifying chimeric barcodes by intra-cellular site-specific recombination, and assayed for growth via next-generation sequencing. Aliquots of these pools can be stored, and later cultured with different drugs to identify condition-dependent genetic interactions. To our knowledge, BFG-GI is the first method to generate haploid double-mutant strains en masse for a many × many matrix of genes without the requirement for multiple mating steps, thus enabling large-scale conditional genetic interaction mapping without extensive use of robotics.

BFG-GI showed good agreement with a previous genetic interaction mapping method (St Onge et al, 2007). Quantitatively, our GIS show a correlation of $r = 0.8 - 0.85$ with the epsilon scores obtained in St Onge et al (2007). Considering only significant interactions, 91% of the negative and 77% of the positive interactions found by BFG-GI were also observed by St Onge et al (2007), and 44 – 64% of St. Onge et al (2007) interactions were reproduced by BFG-GI. The contrast between the FDR estimated by gene pairs including neutral loci (0.01) and the validation rate by an orthogonal method suggests that the latter is a too-conservative measure of precision, and that many of the novel interactions are bona fide interactions despite not having been seen in by St. Onge et al (2007).

We detected and validated unanticipated interactions between SLX4 and Shu complex genes, which mirrored the genetic interactions observed between $PPH3$
and the Shu complex. We further found that presence of a functional Shu complex corresponded to reduced activation or activity of Rad53 during MMS treatment.

By calculating similarity between the genetic interaction profiles of different drugs, we found that those with similar mechanisms of action, like zeocin and bleomycin, are considerably more alike than comparisons between compounds with different mechanisms of action, e.g., the comparison between MMS and either zeocin or bleomycin. This suggests the potential of BFG-GI to shed light on drug mechanisms through measurement of gene-gene-environment interactions.

One advantage of BFG-GI is its cost-effectiveness. BFG-GI uses fewer reagents and less robotic assistance than other technologies to map genetic interactions. Like other pool-based technologies, BFG-GI requires less media, plates, and drugs than array-based technologies, resulting in a substantial cost advantage. For example, the amount of media used in 1,536 spot arrays on OmniTrays is reduced 50-fold by studying the same number of gene-pairs in 100 OD pooled cultures in 143 cm² Petri dishes, which is the optimal cell density we calculated for pooled double mutant selections (Materials and Methods). BFG-GI is also more cost-effective than other barcode-sequencing technologies because in BFG-GI, strains are pooled at the mating step, rather than generating double mutants using robotically manipulated strain arrays.

The reproducibility of BFG-GI indicates that it is a robust technology. Technical replicates in BFG-GI are highly reproducible, and 85% of the biological replicates correlated well with each other ($GIS r > 0.5$). The remaining 15% of biological replicates showing low correlations could be identified and removed computationally. We concur with the iSeq study (Jaffe et al, 2017) that aneuploidies
in chromosome V are the main factor contributing to the replicates with low reproducibility. Chromosome V carries both CAN1 and URA3 loci, which were replaced by selection markers in the iSeq protocol (Jaffe et al, 2017), while CAN1 was replaced by the recipient constructs in BFG-GI. Thus, de novo structural variation around these loci during strain construction could explain the low correlation between some pairs of biological replicates. This possibility is supported by our observation that almost all BFG-GI strains showing GIS r < 0.5 were recipients, whereas donors –for which constructs are carried on plasmids– showed GIS r > 0.5. In the BFG-GI protocol, once the donor and recipient barcodes are fused, the ‘relic’ donor plasmid is counter-selected with 5-FOA to reduce the chance of undesired recombination events. We concur with Jaffe et al. (Jaffe et al, 2017) who suggest that future protocols using constructs located on plasmids, such as the one we used with the proDonor strains, or at other chromosomal loci could eliminate this issue. Despite this issue, the BFG-GI method proved to be highly accurate in comparisons with previous benchmark studies.

Although this study focused on a relatively small matrix (34 × 38 genes), we elaborated on previous studies to optimize the two main bottlenecks of pooled cultures: mating (Soellick & Uhrig, 2001) and sporulation (Codon et al, 1995). We calculated that to cover a yeast genome-scale matrix of 5,500 × 5,500 genes, with 1,000 representative cells for each cross, we would need ~3 x 10^{10} cells at each step along the BFG-GI procedure. Furthermore, using the optimal conditions that we established for mating (22%) and sporulation (18%), an experiment covering all 5,500 × 5,500 crosses would need to culture pools in ~27 Bioassay 500cm² dishes for mating and ~10 L of liquid media for sporulation. Thus, in principle, BFG-GI
could be extended to genome-scale studies.

BFG-GI is a flexible technique that can be used in the future to identify genetic interactions in many different settings. Generation of BFG-GI proDonor and proRecipient strains is one of the most time consuming steps in our pipeline because it includes sequence verification of both $\textit{loxP/lox2272}$ sites and barcodes. However, once generated, these proDonor and proRecipient “toolkits” can be used many times to create donor and recipient strains representing different genes with minimal robotic manipulation. We anticipate that BFG-GI will be a valuable technology to map condition-dependent genetic interactions in yeast and, as next-generation sequencing costs continue to decrease, BFG-GI can be expanded to interrogate pools of double mutants representing bigger sets of gene pairs, including full genome combinations, across multiple conditions.
Materials and Methods

Selected DNA repair and neutral gene strains

We retrieved strains representing 26 DNA repair genes whose null mutants were sensitive to MMS (St Onge et al, 2007) from the YKO and SGA query collections. Additionally, 14 other deemed-neutral loci were selected, based on lack of evidence that their null mutations affected cell fitness (Table EV2). These 14 loci have few or no genetic interactions in genome-scale screens (Costanzo et al, 2010) and we did not find growth defects upon deletion of any of them.

BFG-GI toolkit strains

Donor toolkit construction

We constructed 60 donor strains by generating two DNA fragments with overlapping ends. These were co-transformed into yeast where they recombined to generate pDonor constructs (Fig EV1). The first fragment, called preD1, contained the hygromycin resistance gene (HygR) driven by the Schizosaccharomyces pombe TDH1 promoter and terminator, a barcode locus bearing a 20 bp unique barcode flanked by loxP/2272 sites, and flanking primer sites. First, we used Gibson assembly (Gibson, 2009) to produce plasmid pFR0032 with the $P_{spTDH1}^{-}HygR-T_{spTDH1}$ backbone. Then, we used three consecutive PCRs to add barcodes, priming sites, loxP/2272 loci, and in-yeast recombination adapters (Fig EV1A). The second fragment, preD2, contained the URA3 marker and Cre recombinase driven by $P_{tetO-CMV}$. We generated this fragment by Gibson assembly of pFR0026, followed by a PCR to add in-yeast recombination adapters (Fig EV1B). Then preD1 and preD2...
fragments were co-transformed into yeast strain RY0771 (derived from BY4742) and merged by in-yeast assembly to generate pDonor plasmids (Fig EV1C). We arrayed transformant strains to extract DNA and sequenced the preD1 loci, and proceeded with those strains containing confirmed preD1 loci. We mated selected MATα proDonors with MATα deletion strains of interest (i.e. DNA repair or neutral genes) from the YKO collection (Fig EV1D). A series of selective passages (Fig EV1D and Fig EV3) resulted in Donor strains with the relevant genotype:

\[ \text{MATα lyp1Δ::P}_{\text{STE3}}\text{-LEU2 his3Δ1 leu2Δ0 met17Δ0 ura3Δ0 yfg1Δ::kanMX} \]

\[ p\text{Donor(P}_{\text{tetO-CMV-Cre}}\text{lox2272 P}_{\text{TDH1-HygR-T}}\text{TDH1 barcode loxP P}_{\text{URA3}}\text{-URA3 CEN/ARS P}_{\text{AmpR}}\text{-AmpR ori)}. \]

**Recipient toolkit construction**

We constructed 56 recipient strains using a method based on the previously-described delitto perfetto construct (Storici & Resnick, 2006) to enhance homologous recombination of constructs as follows. First, we used consecutive PCRs to produce a fragment preR1, containing the *Kluyveromyces lactis* URA3 gene, flanked by loxP/2272 sites, 20 bp unique barcodes and a sequence complementary to the *S. cerevisiae* CAN1 locus (Fig EV2A). Second, we incorporated the *P*_{STE2-}spHis5-*T*_{STE2} into the CAN1 locus of the strain BY4741. Then the delitto perfetto construct was inserted upstream of the MATa selection reporter of the same strain (Fig EV2B) to enhance homologous recombination of preR1 fragments. This generated a pool of RY0766 proRecipient strains (Fig EV2C). We isolated and arrayed monoclonal proRecipient strains then sequenced and selected strains with intact preR1 loci. Selected MATa proRecipients were mated with MATα strains of the SGA query.
collection representing DNA repair and neutral genes (Fig EV1D). A series of selective passages (Fig EV2D and Fig EV3) resulted in recipient strains with the relevant genotype:

\[\text{MAT}^a \text{his3}^{\Delta 1} \text{leu2}^{\Delta 0} \text{met17}^{\Delta 0} \text{lyp1}^{\Delta} \text{ura3}^{\Delta 0} \text{can1}^{\Delta} : \text{barcode} \ \text{loxP klURA3 lox2272} \]

\[\text{P}_{\text{STE2}}^{-}\text{spHis5-T}_{\text{STE2}} \text{P}_{\text{CMV}}^{-}\text{rtTA I-SceI P}_{\text{GAL1}}^{-}\text{I-SceI yfg2::natMX} \]

**Generation of BFG-GI double mutants**

We took several steps to reduce the chance of undesired strains in BFG-GI from taking over pooled cultures. This included optimization of both mating and sporulation, and adapting protocols and molecular constructs that have been reported to improve the selection of the \text{MAT}^a double mutant progeny in SGA. Mating and sporulation are the two primary population bottlenecks when generating haploid double mutants by meiotic segregations. As described below, we sought to optimize cultures at these stages to maintain a pool complexity which was large enough to interrogate all desired gene-gene combinations. Optimizing these two processes is also important to reduce potential jackpot effects in pool cultures (i.e. to avoid strains with genetic anomalies to take over the entire pool growth).

**Mating optimization for en masse BFG-GI**

We focused on optimization of cell density for \textit{en masse} party mating because previous evidence shows cell density influences mating efficiency (Soellick & Uhrig, 2001). We determined the optimal cell density for \textit{en masse} party mating by inoculating mating Petri dishes with a mixture of two neutral strains (\text{MAT}^{\alpha} \text{Donor ho}\Delta::\text{kanMX}, and \text{MAT}^a \text{Recipient ylr179c}\Delta::\text{natMX}) at cell densities varying
from 30 OD to 300 OD and counting the colony forming units (CFUs). After generating mating mixtures, we took samples at 0 and 12 hours (hrs) of incubation at 23 °C, and inoculated plates with either non-selective or heterozygous diploid double mutant selective media and counted CFUs. The ratio of CFUs in non-selective vs. selective media indicated that inoculating a 58cm$^2$ Petri dish with 30 ODs of mating mixture resulted in 22% mating efficiency. In contrast, 100 ODs of mating mixture resulted in 13% mating efficiency, and 300 ODs of mating mixture resulted in 3% mating efficiency. Hence, we used 0.51 ODs of mating mixture per cm$^2$ of plate for further en masse party matings.

To generate pools of double mutants, we arrayed BFG-GI donors and recipients in their respective selective media and cultured at 30 °C for 48 hrs (Fig EV3). We made one pool for each mating type by scraping cells from plates into liquid media and normalized cell densities with sorbitol 1M to have equal number of cells per strain (50 ODs per mL) for each pool. Then, we lightly sonicated cells to disrupt clumps (Branson microtip sonicator, 10% duty cycle, output 2, 25 bursts, pause of 3 sec., and a second 25 bursts). We mixed the two pools together by stirring them in a flask for 10 min. Finally, we inoculated two Bioassay dishes (500cm$^2$) with 259 ODs each of the mating mixture, and mating cultures were incubated for 12 hrs at 23 °C (Fig EV3).

**Generation of heterozygous diploid double mutants, induction of barcode fusion and pDonor elimination**

Generation of heterozygous diploid double mutants required passaging the mating progeny every 24 hrs into fresh selective media. Passages included selection
of heterozygous diploid double mutants, induction of the Cre/Lox system with
doxycycline, counter-selection of the relic pDonor with 5-FOA, and recovery from 5-
FOA counter-selection to increase sporulation efficiency (Fig EV3).

**Sporulation optimization for en masse BFG-GI**

We used cultures recovered from 5-FOA counter selection to inoculate liquid
PRE5 pre-sporulation media (Codon *et al.*, 1995) for 2 hrs at 30 °C to induce
exponential growth, then spun down the cells and transferred them to SPO2
sporulation media (Codon *et al.*, 1995) supplemented with histidine, leucine,
methionine and uracil to mask BFG-GI strain auxotrophies at concentrations used in
the SGA sporulation protocol (Tong & Boone, 2005). We incubated sporulation
cultures at 21 °C for 12 days. This resulted in ~18% sporulation efficiency, as
evaluated by counting CFU's in non-selective and selective media and tetrad
visualization. Shorter incubation periods reduced the sporulation efficiency (~4% at
5 days, ~13% at 7 days).

**Selection of MATα haploid double mutants with fused barcodes**

We selected *MATa* haploid progeny from sporulation cultures, followed by
haploid double mutant selection (Fig EV3). Aliquots were stored in glycerol at -80
degrees for future use. We used the *STE2* and *STE3* promoters currently used for
SGA to select for haploid cells, as markers with these promoters have been reported
to perform better than earlier alternatives (e.g. *MFA1/MFA2* promoters) (Tong &
Boone, 2007). We used these constructs to first select the *MATa* progeny from
sporulation cultures and then the haploid double mutants. Using *STE2/STE3*
promoters, optimizing mating and sporulation, and using an intermediate $MAT_a$ selection step between sporulation and haploid double mutant selection together likely reduced the number of mitotic crossover survivors and jackpot mutation effects in our pools.

Exposure of pooled cultures to drugs

Before challenging haploid double mutant pools to drugs we identified the appropriate drug concentration for our experiment by exposing a neutral BFG-GI haploid double mutant ($ho\Delta::kanMX/\text{ylr179c}\Delta::\text{natMX}$) in growth assay liquid cultures to various drug concentrations. We selected drug doses corresponding to 20% of the minimal inhibitory concentration for the neutral test strain (Table EV1). To expose mutant strains to drugs we thawed frozen haploid double mutant pools, allowed the pools to recover for 2 hrs in haploid double mutant liquid media at 30 °C, and then used 100 ODs of this culture to inoculate 143cm$^2$ petri dishes containing solid media supplemented with each DNA repair drug. We cultured pools at 30 °C for 24 hrs and then collected samples to sequence fused barcodes and thus infer each double mutant abundance.

Generation of BFG-GI double mutants in an array format

Mating and selecting donor and recipient strains in an array format was similar to the pool-based en masse party mating assay described above, but in this case we used robotic assistance to pairwise mate each donor with an array of recipients. We completed all steps, including sporulation, on solid media, and imaged the final haploid double mutant selection plates. We scraped cells from the
final selection plates to sequence the fused-barcode population which allowed us to compare cell patch sizes with numbers of sequencing reads.

**Next-generation sequencing and mapping of fused barcode pairs**

The BFG-GI technology relies on the Cre/Lox system to recombine the complementary donor and recipient loxP/lox2272 sites that serve to introduce the donor barcode adjacent to the recipient barcode (Fig 1). We multiplex-sequenced the fused barcodes from pools of cells using the following steps: 1) genomic DNA extraction using glass beads and phenol/chloroform, 2) PCR amplification of the 325 bp barcode fusion product including the two 20 bp barcodes and the multiplexing sequencing adapters (one index for each condition, for each technical replicate), 3) concentration and gel purification of amplicons using 2% E-Gel EX agarose 2% (Invitrogen), DNA Clean & Concentrator Kit (Zymo Research) and MinElute Gel Extraction Kit 50 (Qiagen), 4) normalization of DNA libraries using Qubit Fluorometric Quantitation (Invitrogen), 5) mix of libraries at equal concentrations, 6) quantification of the pooled DNA library mix by qPCR, and 5) sequencing by Illumina 75 cycles NextSeq paired-end technology, including 25 cycles for each barcode and 6 cycles for the multiplex index. We mapped sequencing *.fastq files against the library of expected barcode sequences using the program Segemehl (v0.1.7, -A 85) and custom scripts; 97% of all sequencing reads mapped to expected barcodes.

**Whole-genome sequencing and detection of chromosome duplications**

Ten strain pairs with one strain with GIS < 0.5 and another with GIS > 0.5
with other replicates for the same gene were selected for genome sequencing.

Genomic DNA from 20 strains was extracted via cell wall disruption with Zymolyase 100T 10mg/ml (Amsbio) and purification using AMPure beads (Agilent). gDNA was quantified with Quant-it Picogreen dsDNA assay kit (Invitrogen) and normalized to 2ng/ul for DNA fragmentation and library normalization with a Nextera XT DNA Library Prep Kit, using a transposase (Tn5) for tagmentation. A limited-cycle PCR was used to add Illumina sequencing adapters and indices i5 and i7. PCR amplicons with size between 400 and 800 bp were gel purified using a 2% E-Gel EX agarose 2% (Invitrogen) and MiniElute Gel Extraction kit (Qiagen). Whole genome sequencing was conducted on an Illumina NextSeq 500 using a HighOutput 150 cycles v2 kit with 40x coverage. Sequencing results were mapped against the reference genome UCSC sacCer3 (SGD vR64.1.1), corrected for GC content, and chromosomal duplications detected with the HMMcopy R package (Ha et al, 2012)

Retesting double mutant construction and spot dilution assays

We generated double mutant strains for retesting in spot dilution assays by mating single mutant MATalpha SGA queries with MATa YKO collection strains, the exceptions being the MATa RAD53 (MBS1437) and rad53-R605A (MBS1440) strains with the RAD53 loci linked C-terminally to a 6xHis-3xFLAG-kanMX6 tag and resistance marker (Ohouo et al, 2013). Next, we induced sporulation of heterozygous diploid double mutants as we did for BFG-GI strains. To confirm segregation of kanMX and natMX markers, we manually dissected haploid double mutants from tetrads and verified segregation using both selective media and PCR. Sanger sequencing confirmed the proper identity of residue 605 in intact RAD53 and
rad53-R605A strains. We grew strains overnight to saturation in liquid media, diluted them 1:10, and then used 1:5 serial dilutions for the spot assays. All cultures used YPD media supplemented with indicated drug concentrations.

**Defining a Genetic Interaction Score (GIS)**

In an exponential growth model, the frequency of a double mutant strain $s_{xy}$ in a given condition at a time $t$ ($f_{s_{xy},t}$) represents its total growth from an initial number $N_{s_{xy},t=0}$ as a proportion of the total growth of all other strains in the pool:

$$f_{s_{xy},t} = \frac{N_{s_{xy},t=0}2^{g_{xy}t}}{\sum N_{s_{ij},t=0}2^{g_{ij}t}}$$

Note: Before calculating frequency, we add a pseudocount of 0.5 to the count of every strain in our analysis to avoid a zero denominator in several calculations.

Here, $g_{xy}$ is inversely related to the doubling time of strain $s_{xy}$ and $g_{xy}t$ effectively represents the number of doublings of strain $s_{xy}$. Units for $t$ can be chosen arbitrarily. In this model, a frequency at $t=0$ evaluates as:

$$f_{s_{xy},0} = \frac{N_{s_{xy},t=0}}{\sum N_{s_{ij},t=0}}$$

To remove the unknown $N_{s_{ij},t=0}$ term, we define $r_{s_{ij},t}$:

$$r_{s_{xy},t} \equiv \frac{f_{s_{xy},t}}{f_{s_{xy},0}} = \frac{\frac{N_{s_{xy},t=0}}{\sum N_{s_{ij},t=0}2^{g_{ij}t}}}{\frac{N_{s_{xy},t=0}}{\sum N_{s_{ij},t=0}2^{g_{ij}t}}}$$

We note that the $\frac{\sum N_{s_{ij},t=0}}{\sum N_{s_{ij},t=0}2^{g_{ij}t}}$ term is the ratio between the initial and final number of cells in the pool, and can be calculated by the total number of generations of pool growth ($gen_{pool}$):

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Therefore, $g_{xy}t$ can be calculated as:

$$g_{xy}t = \log_2 \left( r_{s_{xy},t} \right) + g_{\text{gen pool}}$$

To calculate $g_{\text{wt}}t$, we take the mean $g_{xy}t$ of all neutral-neutral pairs:

$$g_{\text{wt}}t = \text{mean}(g_{ij}t | ij \in \text{neutral genes})$$

We then obtain the relative growth rate $w_{xy}$ of each strain compared to the wild type by dividing their number of doublings. In a constant exponential growth model, this metric is independent of time. In practice, $g$ represents the average growth rate over the measured time period.

$$w_{xy} = \frac{g_{xy}t}{g_{\text{wt}}t} = \frac{g_{xy}}{g_{\text{wt}}}$$

To estimate the single mutant fitness $w_x$ and $w_y$ for a given pair, we use the mean estimate of $x$ or $y$ combined with neutral genes.

$$w_x = \text{mean}(w_{xj} | j \in \text{neutral genes})$$

$$w_y = \text{mean}(w_{iy} | i \in \text{neutral genes})$$

We then define the genetic interaction score ($GIS$) as the difference between $w_{xy}$ and the product of $w_x$ with $w_y$:

$$GIS_{xy} \equiv w_{xy} - w_xw_y$$

Because there is uncertainty in $w$, it is possible to calculate $w < 0$ for $w_x$, $w_y$, or $w_{xy}$. Such values are assigned as 0 when performing the $GIS$ calculation.

Normalizing Genetic Interactions and Calculating P-Values

To assign a threshold for positive and negative genetic interactions, several
additional steps are performed. $GIS_{xy}$ is converted to a standard score by calculating how many standard deviations $GIS_{xy}$ is from 0 given an estimate of $GIS_{xy}$ uncertainty ($\hat{\sigma}_{GIS_{xy}}$).

$$Z_{GIS_{xy}} = \frac{GIS_{xy}}{\sigma_{GIS_{xy}}}$$

To calculate $\hat{\sigma}_{GIS_{xy}}$, we identify various sources of uncertainty. Another way to state $GIS_{xy}$ is as such:

$$GIS_{xy} = w_{xy}t - \frac{g_{xt}g_{yt}}{gw_{t}}$$

We then define an error model to calculate the standard error $\sigma$ for each term used in this calculation:

$\hat{\sigma}_{w_{xy}t}$: this is estimated globally for each condition as the median difference between $w_{xy}t$ between the R1 and R2 technical replicates for all strains. We note that this error model only captures the general expected error between two separate runs of the same biological sample.

$\hat{\sigma}_{g_{xt}}, \hat{\sigma}_{g_{yt}}, \hat{\sigma}_{gw_{t}}$: each of these $g$ values are calculated by taking the mean of multiple strains. We use the variation of growth estimates in these strains (i.e. the standard deviation) as the uncertainty.

The standard formula for approximating the propagation of measurement uncertainty is used to combine $\hat{\sigma}_{w_{xy}t}, \hat{\sigma}_{g_{xt}}, \hat{\sigma}_{g_{yt}}$ and $\hat{\sigma}_{gw_{t}}$ into $\hat{\sigma}_{GIS_{xy}}$. This formula is also used for obtaining the other error estimates reported (i.e. $\hat{\sigma}_{w_{x}}, \hat{\sigma}_{w_{y}}, \hat{\sigma}_{w_{xy}}$).

To assign a p-value for each interaction, we then analyze the distribution of $Z_{GIS}$ in all unlinked neutral-neutral and neutral-DNA damage pairs (hereafter called
‘neutral pairs’), as few or no genetic interactions are expected to take place in this space. We model $Z_{GIS_{np}}$ as a normal distribution (Fig. EV4D shows the empirical and fitted normal distribution for each condition to validate this decision) and use the $pnorm$ function in R to calculate $p_{pos} = p\left(Z_{GIS_{neutral}} \geq Z_{GIS_{xy}}\right)$ and $p_{neg} = p\left(Z_{GIS_{neutral}} \leq Z_{GIS_{xy}}\right)$ for each pair. We then combine these single-tailed tests into a two tailed value:

$$p_{neutral} = \min(p_{pos}, p_{neg}) \times 2$$

$p_{neutral}$ represents the probability that a score as extreme as $Z_{GIS_{xy}}$ or more would be found amongst neutral pairs.

**Combining Multiple Biological Replicates and Calculating a FDR**

We consolidated multiple measurements of $w_x, w_y, w_{xy}, GIS_{xy}, Z_{GIS_{xy}}$ as well as $\sigma_{w_x}, \sigma_{w_y}, \sigma_{w_{xy}}, \sigma_{GIS_{xy}}$ and $p_{neutral}$ from multiple barcode pairs into a single value for each gene pair. $GIS_{xy}$ values were weighted by the inverse of estimated squared-error ($w = \frac{1}{\sigma_{GIS_{xy}}^2}$) and averaged to obtain $GIS_{genex,geney}$. Similarly, $w_x, w_y, w_{xy}$ were averaged by the same weight ($w$) to obtain their corresponding gene-wise value.

$\hat{\sigma}_{w_{genex}}, \hat{\sigma}_{w_{geney}}, \hat{\sigma}_{w_{genex,geney}}, \hat{\sigma}_{GIS_{genex,geney}}$ were obtained using the propagation of uncertainty when calculating a weighted average:

$$\hat{\sigma}_{gene} = \sqrt{\sum_{barcode} \left(\frac{w}{\sum_{w}}\right)^2}$$

$Z_{GIS_{genex,geney}}$ was calculated using $GIS_{genex,geney}$ and $\hat{\sigma}_{GIS_{genex,geney}}$:

$$Z_{GIS_{genex,geney}} = \frac{GIS_{genex,geney}}{\hat{\sigma}_{GIS_{genex,geney}}}$$
Finally, a gene-wise $p_{\text{neutral}}$ was calculated using Stouffer’s method weighted by $w$. The gene-wise $p_{\text{neutral}}$ values were then converted to $FDR_{\text{neutral}}$ using the $qvalue$ function in the $qvalue$ R package.

**Calling Differential Genetic Interactions**

For each gene pair, we calculated $\Delta GIS$ and $\Delta Z$ for all pairwise comparisons $(a - b)$ amongst the tested conditions. $\Delta GIS_{\text{gene}_x,\text{gene}_y,a-b}$ was calculated as

$$GIS_{\text{gene}_x,\text{gene}_y,a} - GIS_{\text{gene}_x,\text{gene}_y,b},$$

and $\Delta Z_{\text{GIS}_{\text{gene}_x,\text{gene}_y,a-b}}$ was calculated as:

$$\Delta Z = \frac{GIS_{\text{gene}_x,\text{gene}_y,a} - GIS_{\text{gene}_x,\text{gene}_y,b}}{\sqrt{\sigma^2_{GIS_{\text{gene}_x,\text{gene}_y,a}} + \sigma^2_{GIS_{\text{gene}_x,\text{gene}_y,b}}}}$$

For each pair of conditions, $\Delta Z$ was calculated for all unlinked neutral-neutral and neutral-DNA damage pairs (‘neutral pairs’) to create a null distribution for $\Delta Z_{\text{neutral}}$. $p_{\Delta_{\text{neutral}}}$ was then calculated for each pair from the $\Delta Z_{\text{neutral}}$ distribution in the same manner as calculating $p_{\text{neutral}}$. $p_{\Delta_{\text{neutral}}}$ values were then converted to $FDR_{\Delta_{\text{neutral}}}$ using the $qvalue$ function in the $qvalue$ R package.

**Data Availability**

Raw and normalized sequencing measurements and GIS for each gene pair is available in Tables EV2 to EV6 and Computer Code EV1 allows to generate Tables EV3 to EV6 from Table EV2. Any modifications post-publication will have been documented at: https://github.com/a3cel2/BFG_GL_stats.
Acknowledgements

We are grateful for helpful comments from Yong Lu, Michael Principato, Ramamurthy Mani, and Meng Xiao He at the outset of this project, to Brenda Andrews and Charles Boone and members of their labs for providing reagents and insightful comments, and to many Roth lab members for support and feedback throughout this project.

Author contributions

FPR, JJDM, JCM and ACo (Atina Coté) conceived the project; ACo, AK and SO constructed pilot strains. ACo, PB, CW and JR constructed final BFG-GI strains and performed pilot mating experiments. JJDM, FS, YZ, DAP and GG optimized mating, marker selection and sporulation protocols. JJDM, FS, ACo, MG and MV performed array and pool cultures and sequencing of fused barcodes. JJDM and JCM performed computational mapping of barcodes. JJDM and ACe (Albi Celaj) performed scoring of genetic interactions. JJDM and MG performed aneuploidy experiments and JW analyzed the results. JJDM, AB and BH performed Shu complex-related experiments. DD and GWB provided advice on DNA repair pathways. JJDM, FPR, GWB, AB and ACe wrote the manuscript. FPR supervised the project.

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(HG004825) to JCM. FPR was also supported by a NIH/NHGRI Center of Excellence in Genomic Science (HG004233), by NIH/NHGRI grant HG001715, and by the One Brave Idea Foundation.

**Conflict of interest**

The authors declare that they have no conflict of interest.
**Figure Legends**

**Figure 1.** BFG-GI Pipeline Summary

**A** Construction of donors with unique barcodes representing each gene deletion in parental strains from the YKO collection.

**B** Construction of recipients also with unique barcodes representing genes of interest in parental strains from the SGA query collection. Pairs of recombination sites (loxp and lox2272) were located at the barcode loci of donor and recipient strains to enable *in vivo* intracellular fusion of barcode pairs at the recipient barcode locus.

**C** Donors and recipients were mated with each other to generate heterozygous diploid double mutants and barcodes were fused *in vivo* by the Cre/Lox system. The relic plasmid remaining in donors after Cre/Lox recombination was counter-selected after barcode fusion. Sporulation was induced to select for the *MATα* progeny and haploid double mutants.

**D** BFG-GI was conducted *en masse* to generate ‘many-by-many’ pools for a set of 26 DNA repair and 14 neutral genes. The resulting pool of haploid double mutants was stored as aliquots of glycerol stock. Thawed aliquots were used to inoculate media containing different chemical agents (‘drugs’). Genomic DNA was extracted and fused barcodes were amplified and sequenced to monitor double mutant abundance and to infer genetic interactions. Details of donor and recipient strain construction are shown in Fig EV1 and Fig EV2, respectively. Media details are shown in Fig EV3.
Figure 2. BFG-GI Quality Control and Benchmarking

A  Correlation between two measures of cell abundance (colony size and next-generation-sequencing-based quantification of fused barcodes) for BFG-GI double mutant strains. Histograms show distribution of abundance in the two measurements. Peaks in the histograms representing data points in the bottom-left corner of the scatter plot indicate that absent and very small colonies produced few or no sequencing reads.

B  Density plots for BFG-GI genetic interaction score (GIS) correlation between replicates of the same gene, with same or different parental origin, or pairs of different genes. Only replicates with a GIS correlation > 0.5 were retained for further analyses.

C  Histograms comparing the GIS distribution for ‘same-gene pairs’ (which are expected to behave like synthetic lethals given the SGA double-mutant selection process) with that for linked and unlinked gene pairs.

D  Comparison of BFG-GI-inferred genetic interactions in haploid double mutant media without MMS with genetic interactions identified using similar media (St Onge et al, 2007).

E  Comparison of BFG-GI-inferred genetic interactions in haploid double mutant media containing MMS with genetic interactions previously identified in similar media (St Onge et al, 2007).

F  Benchmarking of BFG-GI genetic interactions against the St. Onge et al. dataset (St Onge et al, 2007).
Figure 3. Condition Dependent Genetic Interactions Mapped by BFG-GI

A  Networks showing the number of conditions with a genetic interaction for each gene pair (using FDR < 0.01 and |GIS| > 0.075 as cutoffs). Numbers besides gene names are guides for the reader to locate nodes in networks of panels (B) and (C). Data for individual interactions are available in Tables EV3 and EV4.

B  Networks in the diagonal (subpanels ii and iii) show genetic interactions for DMSO or MMS after applying the same criteria as in (A). The network in subpanel i shows significant genetic interaction changes (FDR < 0.01, |ΔGIS| > 0.1) when comparing the DMSO and MMS treatments. Interaction types are positive (+), negative (-), or neutral (n). The barplot in subpanel iv summarizes the number of changes between interaction type in subpanel i.

C  The networks are the same as described in (B) with additional drug conditions: cisplatin (CSPL), doxorubicin (DXRB), hydroxyurea (HYDX), zeocin (ZEOC), bleomycin (BLMC) and 4NQO. The no-drug condition was omitted from this figure as it showed no significant condition-dependent genetic interactions with DMSO. GIS profiles were hierarchically clustered using maximum distance and complete linkage, with the resulting dendrogram shown on the left. Data for individual differential interactions are available in Tables EV5 and EV6.
Figure 4. Shu Complex Condition-Dependent Genetic Interactions with \textit{MAG1}, \textit{SLX4}, \textit{PPH3} and \textit{RAD53}

\textbf{A} Pleiotropic participation of the Shu complex in DNA replication and repair pathways.

\textbf{B} Network showing condition-dependent genetic interactions inferred from BFG-GI for the indicated conditions.

\textbf{C} Confirmation of interactions between the Shu complex, \textit{MAG1} and \textit{SLX4} using spot dilution assays including single and double mutants exposed to the indicated drugs for 48 hrs. Orange, blue, and red boxes indicate genetic interactions of Shu complex members with \textit{MAG1} and \textit{SLX4}, and of \textit{MAG1} with \textit{SLX4}, respectively.

\textbf{D} Schematic of potential functional connections between the Shu complex and \textit{SLX4}. As with deletion of \textit{SLX4} or \textit{PPH3}, deletion of Shu complex members may lead to hyperphosphorylation and hyperactivation of Rad53, resulting in increased sensitivity to MMS.

\textbf{E} Spot dilution assays showing genetic interactions of \textit{Shu/pph3Δ} double mutants and corresponding single mutants exposed to MMS at indicated concentration for 48 hrs.

\textbf{F} Western blot assays showing hyperphosphorylation of Rad53 in \textit{csm2Δ}, \textit{psy3Δ}, \textit{shu1Δ}, and \textit{slx4Δ} strains following treatment with 0.03% MMS. Note increased intensity of Rad53-P bands compared with the Rad53 bands.

\textbf{G} Spot dilution assays of Shu complex mutants expressing a hypomorphic \textit{rad53-R605A} allele (\textit{rad53-R605A-6xHis-3xFLAG-kanMX6}) compared with a wild type \textit{RAD53} allele (\textit{RAD53-6xHis-3xFLAG-kanMX6}). Cells were exposed to MMS at indicated concentration for 60 hrs.
**Figure EV1.** Donor Toolkit Construction

A  Two fragments were built to generate proDonor plasmids. The first, preD1, contained \(\text{loxP/lox2272}\) sites flanking two 20 bp unique barcodes and a hygromycin resistance marker. In this study only the upstream barcode was used for further steps and for simplification the downstream barcode was omitted from Fig 1.

B  The second, preD2, contained the Cre recombinase driven by the doxycycline inducible tetO-CMV, and a \(\text{URA3}\) marker.

C  The two fragments were assembled \textit{in vivo} in yeast to generate pDonors.

D  pDonors were arrayed and Sanger sequenced to confirm the integrity of the preD1 fragment. ProDonors with confirmed preD1 fragments were mated with YKO strains to generate strains carrying both a uniquely barcoded pDonor and a gene deletion of interest. Then they were sporulated and the haploid \(\text{MAT}\alpha\) progeny was selected using the mating type maker indicated in panel C. Details on selective media are shown in Figure EV3.

**Figure EV2.** Recipient Toolkit Construction

A  Two constructs were built to generate recipients. The first fragment, preR1, contained \(\text{loxP/lox2272}\) sites flanking a kl\text{URA3} marker and two 20 bp unique barcodes flanking these loci. In this study only the upstream barcode was used for further steps and for simplification the downstream barcode was omitted from Fig 1.

B  The second construct, preR2, contained the \(can1\Delta::P_{\text{STE2}}-sp\text{His}5-T_{\text{STE2}}\) mating type marker.
**C** The two fragments were assembled *in vivo* using a derivative of the *delitto perfetto* construct.

**D** Resulting proRecipients were arrayed and Sanger sequenced to confirm integrity of preR1 loci. ProRecipients with confirmed preR1 loci were mated with SGA query strains to generate strains carrying both a uniquely barcoded recipient construct and a gene-deletion of interest. Then they were sporulated and the haploid *MATa* progeny were selected using the mating type maker indicated in panel C. Details on selective media are shown in Figure EV3.

**Figure EV3.** Media Details to Generate BFG-GI Strains and Pools

Donors, recipients and double mutants used in BFG-GI were generated as shown in Figs 1, EV1 and EV2. This figure shows media details, optimal inoculum cell densities, and incubation times for pool-based cultures. All incubations were at 30°C for 24 hrs, except for mating (12 hrs at 23°C) and sporulation (12 days at 21°C). Sporulation was conducted in flasks with liquid media shaking at 200 rpm. We used the following reagent concentrations: G418 = 200µg/mL, clonNat = 100µg/mL, canavanine = 100µg/mL, thialysine = 100µg/mL, hygromycin = 200µg/mL, 5-FOA = 1mg/mL. Amino acid concentrations were as described in (Tong & Boone, 2005).

**Figure EV4.** Calling Genetic Interactions

**A** Two collections of 59 donor strains (containing 34 unique knockouts) and 56 recipient strains (containing 38 unique gene knockouts) were crossed against each other in an all-by-all pooled format. Each strain contains a knockout at either a DNA repair gene or neutral locus. Double knockout strains were divided into four spaces
based on the types of genes knocked out. Numbers in parentheses represent the
number of strains and unique gene knockouts, respectively.

B Distribution of GIS amongst strain pairs containing the same gene, split by
those which were well-measured from the heterozygous diploid stage ($C_{xy} \geq 30$)
and not well-measured from the same stage ($C_{xy} < 30$). Non-well-measured strains
(72 out of 3,305) were excluded from analysis and GIS was re-calculated after their
exclusion.

C Distribution of GIS in strains representing linked neutral pairs. Using the GIS
profiles, an empirical cutoff of 75 kbp (red dashed line) was chosen to classify
strains with knockout pairs on the same chromosome as either linked or unlinked.
GIS was then re-calculated based on this linkage criterion.

D Distribution of $Z_{GIS}$ calculated for DNA-repair pairs (space 1 in panel A, red)
and pairs involving well-measured and unlinked neutral genes (spaces 2, 3, 4 in
panel A, black). $Z_{GIS}$ for pairs involving neutral genes were used to calculate a $p$-
value.

E Distribution of $p$-values calculated by the null distribution in D. $p$-values
were combined for multiple barcode replicates of each gene-gene pair and
converted to FDR scores (see Methods). Barcode-level $p$-values are available in
Table EV3, and gene-level FDR scores are available in Table EV4.

F Benchmarks of BFG-GI with data from St. Onge et al (2007) for strains
containing a significant genetic interaction (FDR < 0.01). Each graph shows
precision and recall compared to St. Onge et al (2007) as a function of an additional
GIS effect size cutoff (left = negative interaction performance, right = positive
interaction performance). Overlay text indicates performance at $|GIS| = 0.075$
(dashed lines), which was chosen as the effect size threshold.

**Figure EV5.** Calling Differential Genetic Interactions

A Distribution of $\Delta Z_{GIS}$ for neutral pairs compared to DNA repair pairs. The
distribution amongst neutral pairs was used to calculate a $p$-value for $\Delta Z_{GIS}$ amongst
DNA repair pairs, which was then converted to an FDR for each differential
interaction (see Methods; Table EV6). An additional effect size cutoff of $|\Delta GIS| > 0.1$
was added to call differential genetic interactions in Fig 3 and Table EV#5.

B Distribution of significant differential genetic interaction calls per gene.

C Distribution of significant differential genetic interaction calls involving a
reversal of direction (i.e. from positive to negative or vice-versa) by gene. $RAD5$ is
involved in 47 differential genetic interactions with a reversal of direction.

D Summary of significant genetic interactions of $RAD5$ with $MUS81$, $MMS4$,$RAD51$, $RAD54$, or $RAD55$ in different conditions. Edges represent genetic interaction
type, and are labeled by conditions in which significant genetic interactions were
found for the corresponding pair and direction.
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A proDonor strain

**pDonor**

\[ \text{URA3} \]

\[ \text{Cre} \]

\[ \text{barcode} \]

\[ \text{loxP} \]

\[ \text{HygR} \]

mating

**marker selections**

Fig S1

**donor strain**

**YKO strain**

\[ \text{yfg1Δ::kanMX} \]

\[ \text{MATa} \]

\[ \text{MATa} \]

\[ \times \]

\[ \text{klURA3} \]

\[ \text{klURA3loxP} \]

\[ \text{lox2272} \]

**barcode**

**rtTA**

**spHis5**

B recipient strain

**proRecipient**

\[ \text{MAT} \]

\[ \alpha \]

\[ \times \]

\[ \text{MAT} \]

\[ \alpha \]

\[ \times \]

\[ \text{pDonor} \]

\[ \text{MAT} \]

\[ \alpha \]

\[ \times \]

\[ \text{URA3} \]

\[ \text{barcode} \]

\[ \text{loxP} \]

\[ \text{lox2272} \]

\[ \text{HygR} \]

\[ \text{Cre} \]

marker selections

**mating and heterozygous double mutant selection**

**recipient strain**

**yfg2Δ::natMX**

**MATa**

\[ \times \]

**SGA query strain**

**MATa**

\[ \times \]

\[ \text{can1 locus} \]

**barcode fusion**

**Cre/Lox**

**plasmid curing**

**sporulation**

**and haploid double mutant selection**

**culture under various conditions**

**DNA extraction and PCR of fused barcodes**

**multiplexed sequencing of fused barcodes**

**for each condition, infer strain relative abundance and genetic interaction network**

**Fig S1**

**Fig S2**

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A. Barcode Fusion Genetics (read counts)

B. Genetic interaction profile similarity

C. Genetic Interaction Score

D. Epsilon (St Onge et al; no drug)

E. Epsilon (St Onge et al; MMS)

F. False Positive Rate (%)
A DNA lesion → fork stalling → double strand break

Prophase I crossover between chromosomes → error-free lesion bypass

B Genetic interactions found by BFG-GI

C

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<th>MMS 0.003%</th>
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D Increased Rad53 phosphorylation

E MMS time (min) Rad53-P Rad53 Pgk1

F MMS time (min) Rad53-P Rad53 Pgk1

G
A. Generation of fragment preD1

- pIS419 natMX

- Gibson assembly #1 to replace natMX by \( P_{spTDH1}^{hyg} - T_{spTDH1}^{spTDH1} \)

- PCR #1 to produce HygR backbone

- PCR #2 to add barcodes

- PCR #3 to add lox sites and adapters for in yeast assembly

- fragment preD1

B. Generation of fragment preD2

- pSH47

- Gibson assembly #2 to replace \( P_{GAL1} \) by \( P_{tetO-CMV} \)

- PCR #4 to produce PFR0026 backbone and add adapters for in yeast assembly

- fragment preD2

C. In-yeast assembly to generate proDonors

- strain RY0771

- Cre

- fragment preD1

D. Strain array and quality control by sequencing of preD1
A. Generation of preR1 fragment

PCR #5 to produce klURA3 backbone and add lox sites

PCR #6 to add barcodes

PCR #7 to add CAN1 upstream sequence

B. Generation of preR2 construct and integration into CAN1 locus

Insertion of PSTE2-spHis5-TSTE2 upstream of CAN1 locus of strain BY4741

Insertion of delitto perfetto construct upstream of MATα selection marker of strain BY4741

C. Insertion of preR1 and preR2 into CAN1 locus to generate proRecipients

D. Strain array and quality control by sequencing of preR1

MATα his3Δ1 leu2Δ0 met17Δ0 lyp1Δ P_{STE2}^{spHis5-T_{STE2}}

SGA query strains

details in Figure EV3
**ProDonors**

- **MATa**
- pDonor (Cre/Lox, loxP/2272, URA3)
- [pd1] SC-Ura+Hyg

**YKO MATa**

- **yfg1D:: kanMX**
- [pd2] YPD+G418

**ProRecipients**

- MATa
- **loxP/2272, klUra3**
- [pr1] SC-Ura

**SGAquery**

- **MATalpha**
- **yfg2D:: NatMX**
- [pr2] YPD+Nat

**Donors**

- [r1] SC-Ura+G418
- [SPO2] sporulation
- [d1] SC-Leu-Lys-Ura+Thi+Hyg

**Recipients**

- [r2] SC-His-Arg-Lys+Can+Thi
- [r3] SC-His-Arg-Lys-Ura+Can+Thi+Nat

**Mating all-vs-all**

- SC-Ura
- Inoculum = 0.51 ODs/cm²

**Heterozygous diploid double mutant (2 passages)**

- [dm1] SC-Ura+Hyg+G418+Nat
- Inoculum = 0.4 ODs/cm²

**Barcode Fusion (Cre/Lox induction)**

- [dm2] SC-Ura+Hyg+G418+Nat+Dox
- Inoculum = 0.4 ODs/cm²

**Relic pDonor curation**

- [dm3] SC+Hyg+G418+Nat+5-FOA+Ura+20mg/L
- Inoculum = 0.4 ODs/cm²

**Recovery**

- [dm4] SC+Hyg+G418+Nat
- Inoculum = 0.4 ODs/cm²

- [PRE5] Pre-sporulation
- [SPO2] Sporulation
- Inoculum = 1 OD/mL

**MATa progeny (2 passages)**

- [dm5] SC-His-Arg-Lys+Can+Thi
- Inoculum = 0.2 ODs/cm²

**MATa haploid double mutant (2 passages)**

- Inoculum = 0.4 ODs/cm²

**Solid media Bioassay dishes**

**Pre-sporulation and sporulation liquid media**

**Solid media Bioassay dishes**

**Thaw and recovery**

**Recovery**

- 2 hrs in liquid dm6 media

**Culture under 10 conditions in duplicate**

- 143 cm² Petri dishes

Record number of inoculated and collected OD's after culture, to be used for growth rate calculation (full list in Table EV1)

<table>
<thead>
<tr>
<th>Condition</th>
<th>Doublings</th>
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</thead>
<tbody>
<tr>
<td>NoDrug-R1</td>
<td>12.24</td>
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<tr>
<td>NoDrug-R2</td>
<td>13</td>
</tr>
<tr>
<td>DMSO-R1</td>
<td>8.2</td>
</tr>
<tr>
<td>DMSO-R2</td>
<td>8.48</td>
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<tr>
<td>MMS-R1</td>
<td>7.8</td>
</tr>
<tr>
<td>MMS-R2</td>
<td>7.88</td>
</tr>
<tr>
<td>... etc</td>
<td></td>
</tr>
</tbody>
</table>
**Neutral pairs**

(spaces 2, 3 and 4 in panel A)

**DNA repair pairs**

(space 1 in panel A)

---

**Recipients (56 strains / 38 genes)**

- DNA repair (35/25)
- Neutrals (21/13)

**Donors (59 strains / 34 genes)**

- DNA repair (36/20)
- Neutrals (23/14)

---

“Neutral pairs” - used to model null distribution for genetic interactions

---

**GIS of Same-Gene Pairs**

GIS

Density

-1.0 -0.8 -0.6 -0.4 -0.2 0.0 0.2

GIS

Distribution

- NoDrug
- DMSO
- MMS
- 4NQO
- BLM C
- ZEOC
- HYDX
- DXRB
- CSPL

---

**Negative Interactions**

- Precision: 91%
- Recall: 64%

**Positive Interactions**

- Precision: 77%
- Recall: 44%
**Figure A**

Significant Differential GIs

Number of genes

100 150 200 250

0 1 2 3 4 5

\( \Delta Z_{GIS} \)

**Figure B**

Significant Sign-Reversed GIs

Number of genes

0 10 20 30 40 50

0 1 2 3 4 5

**Figure C**

Neutral pairs

DNA repair pairs

**Figure D**

Neutral pairs

DNA repair pairs

**Table**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Positive GI</th>
<th>Negative GI</th>
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<tbody>
<tr>
<td>MMS</td>
<td>Blue</td>
<td>Orange</td>
</tr>
<tr>
<td>4-NQO</td>
<td>Blue</td>
<td>Orange</td>
</tr>
<tr>
<td>doxorubicin</td>
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<td>Orange</td>
</tr>
<tr>
<td>cisplatin</td>
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<td>Orange</td>
</tr>
<tr>
<td>MMS</td>
<td>Blue</td>
<td>Orange</td>
</tr>
<tr>
<td>4-NQO</td>
<td>Blue</td>
<td>Orange</td>
</tr>
<tr>
<td>DMSO</td>
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<td>Orange</td>
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<tr>
<td>zeocin</td>
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<td>Orange</td>
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<tr>
<td>bleomycin</td>
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</tr>
<tr>
<td>hydroxyurea</td>
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</tr>
<tr>
<td>no drug</td>
<td>Blue</td>
<td>Orange</td>
</tr>
</tbody>
</table>

**Figure Legends**

- **RAD5**: DNA repair pairs
- **MUS81**: Neutral pairs

**Figure Notes**

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