

## GENOMICS

# Exploring genetic interaction manifolds constructed from rich single-cell phenotypes

Thomas M. Norman<sup>1,2,3\*</sup>, Max A. Horlbeck<sup>1,2,3\*</sup>, Joseph M. Replogle<sup>1,2,3</sup>, Alex Y. Ge<sup>4,5</sup>, Albert Xu<sup>1,2,3</sup>, Marco Jost<sup>1,2,3</sup>, Luke A. Gilbert<sup>4,5†</sup>, Jonathan S. Weissman<sup>1,2,3†</sup>

How cellular and organismal complexity emerges from combinatorial expression of genes is a central question in biology. High-content phenotyping approaches such as Perturb-seq (single-cell RNA-sequencing pooled CRISPR screens) present an opportunity for exploring such genetic interactions (GIs) at scale. Here, we present an analytical framework for interpreting high-dimensional landscapes of cell states (manifolds) constructed from transcriptional phenotypes. We applied this approach to Perturb-seq profiling of strong GIs mined from a growth-based, gain-of-function GI map. Exploration of this manifold enabled ordering of regulatory pathways, principled classification of GIs (e.g., identifying suppressors), and mechanistic elucidation of synergistic interactions, including an unexpected synergy between *CBL* and *CNN1* driving erythroid differentiation. Finally, we applied recommender system machine learning to predict interactions, facilitating exploration of vastly larger GI manifolds.

The complexity of cell types in multicellular organisms is driven not by a large increase in gene number but instead by the combinatorial expression of a surprisingly small number of components. Specific combinations of genes exhibit emergent properties when expressed together, enabling the generation of many diverse cell types and behaviors. Searching for such emergent properties is enabled by the quantitative study of genetic interactions (GIs), which compare the phenotypic consequences of perturbing a pair of genes alone or in combination, typically by measuring growth [although other phenotypic readouts such as reporter gene expression and transcriptional responses have been explored (1, 2)]. GIs can reveal synthetic lethal vulnerabilities in tumors, identify suppressors of inherited and acquired disorders, and guide the design of cocktails of genes to drive differentiation between cell types (3, 4). Pioneering efforts in yeast to construct systematic GI maps between all gene pairs have enabled systematic determination of gene function, identification of protein complexes, and definition of gene-regulatory networks in a principled and unbiased manner (1, 2).

Recent studies have extended such approaches to mammalian and other metazoan systems, but these efforts face two major challenges: scale and information content (1, 2). For example, mapping pairwise interactions among the ~10,000 tran-

scribed genes in a human cell would require measuring ~50 million double mutants. This scale has necessitated the use of highly parallelizable readouts of phenotype, such as growth rate, that obscure the mechanistic or molecular basis for any particular interaction. Put simply, there are many ways for cells to appear equally “unfit”: the reprogramming of a pluripotent cell to a terminally differentiated neuron may affect growth as much as induction of apoptosis or cell cycle arrest. Furthermore, many metazoan cell types are quiescent or postmitotic and as such are not amenable to growth-based screens. Finally, bulk measures of their properties may obscure important cell-to-cell variability.

Emerging high-throughput approaches for monitoring rich phenotypes of individual cells (e.g., imaging or droplet single-cell transcriptomics) present a potential solution to these problems. For example, Perturb-seq pairs CRISPR-based screens with single-cell RNA sequencing (5–8). Each individual cell is in effect an independent experiment connecting a genetic perturbation to its transcriptional consequences, allowing hundreds of thousands of parallel measurements (9, 10). It has been suggested that the rich phenotypes enabled by Perturb-seq can be used to better interpret the impact of genetic interactions (7).

In this study, we exploit the scalability and rich transcriptome readout of Perturb-seq to implement a principled approach for systematically studying mammalian GIs. Each transcriptional measurement, and effectively each genetic perturbation (single or combinatorial), can be viewed as defining a point in high-dimensional space. By measuring many GIs, we therefore effectively map out a surface—a manifold in mathematical terminology—that describes the transcriptional states that a cell can occupy upon perturbation (Fig. 1A). We argue that this GI manifold carries much more information about interactions than

a traditional GI map and is intrinsically more interpretable in several ways, including the ability to resolve the distinct outcomes underlying GIs and to model the different ways genetic perturbations combine to yield new phenotypes.

## An overexpression strategy identifies strong genetic interactions

Most previous studies of GIs have focused on loss-of-function perturbations, but many important cellular processes such as differentiation are also associated with activation of gene expression (e.g., *MYOD1* in muscle cells). Genes that exhibit phenotypes when expressed alone have a higher rate of genetic interactions with other genes (1, 2). To identify mechanistically diverse GIs arising from gene activation, we therefore selected 112 “hit” genes whose activation enhances or retards growth of K562 cells (Fig. 1A and table S1) (11), including cell cycle regulators, transcription factors, kinases, phosphatases, and genes of unknown function.

To systematically measure gain-of-function (overexpression) GIs, we adapted a technology developed for constructing fitness GI maps in human cells using CRISPR interference (CRISPRI) (12) (Fig. 1, A and B, and fig. S1). Each candidate interaction was probed by constructing a library of vectors containing pairs of single-guide RNAs (sgRNAs) (table S2). As we included two distinct sgRNAs targeting each gene, a total of 28,680 unique sgRNA pairs were tested. K562 cells stably expressing the SunTag CRISPR activation (CRISPRa) system (11) were transduced with the CRISPRa GI library, and sgRNA pair abundance was compared at the start of the screen and after 10 days of growth to measure fitness phenotypes. GI scores were assigned by measuring deviation between the observed fitness of overexpressing both genes from the expected fitness based on the average impact of each single gene [(12); see materials and methods, fig. S2, and table S3]. Independent replicate experiments showed high levels of concordance for sgRNA-level GI scores and GI profile correlations, and independent sgRNAs targeting the same gene were much more similar than the background of all sgRNA GI correlations (median  $R = 0.50$  compared with 0.04; Fig. 1, C and D; fig. S3A; and table S4). The gene-level GI scores obtained by averaging sgRNAs targeting the same gene were also well correlated between replicates (gene-level GI  $R = 0.80$ ,  $p < 10^{-300}$ ; Fig. 1E, fig. S3A, and table S5) and followed a bell-shaped distribution but with GIs ranging well beyond the expectation from negative control sgRNAs (Fig. 1F and fig. S3B).

We then clustered genes according to the similarity of their GI profiles to produce a GI map (Fig. 1B; a larger version with gene labels is provided in fig. S4). Highly correlated genes were enriched for genes with the same DAVID term annotations (13), allowing for unbiased annotation of clusters (Fig. 1B; fig. S5, A and B; and table S6). In addition, the map contained fewer clusters than would be expected in a random map (fig. S5, C and D), consistent with the typical low-rank structure of GI maps (i.e., groups of genes interact

<sup>1</sup>Department of Cellular and Molecular Pharmacology, University of California, San Francisco, CA 94158, USA. <sup>2</sup>Howard Hughes Medical Institute, University of California, San Francisco, CA 94158, USA. <sup>3</sup>California Institute for Quantitative Biomedical Research, University of California, San Francisco, CA 94158, USA. <sup>4</sup>Department of Urology, University of California, San Francisco, CA 94158, USA. <sup>5</sup>Helen Diller Family Comprehensive Cancer Center, San Francisco, CA 94158, USA.

\*These authors contributed equally to this work.

†Corresponding author. Email: thomas.norman@ucsf.edu (T.M.N.); luke.gilbert@ucsf.edu (L.A.G.); jonathan.weissman@ucsf.edu (J.S.W.)

similarly so that there are fewer overall degrees of freedom than total genes). Thus, the structure of the CRISPRa GI map, like past efforts based on loss-of-function alleles, can assign function to individual genes by the similarity of GI profiles (1, 2). However, although the GI map robustly identified many strong GIs, the origins of specific interactions were difficult to deduce as each GI was characterized only by a single scalar value (i.e., deviation from expected growth rate).

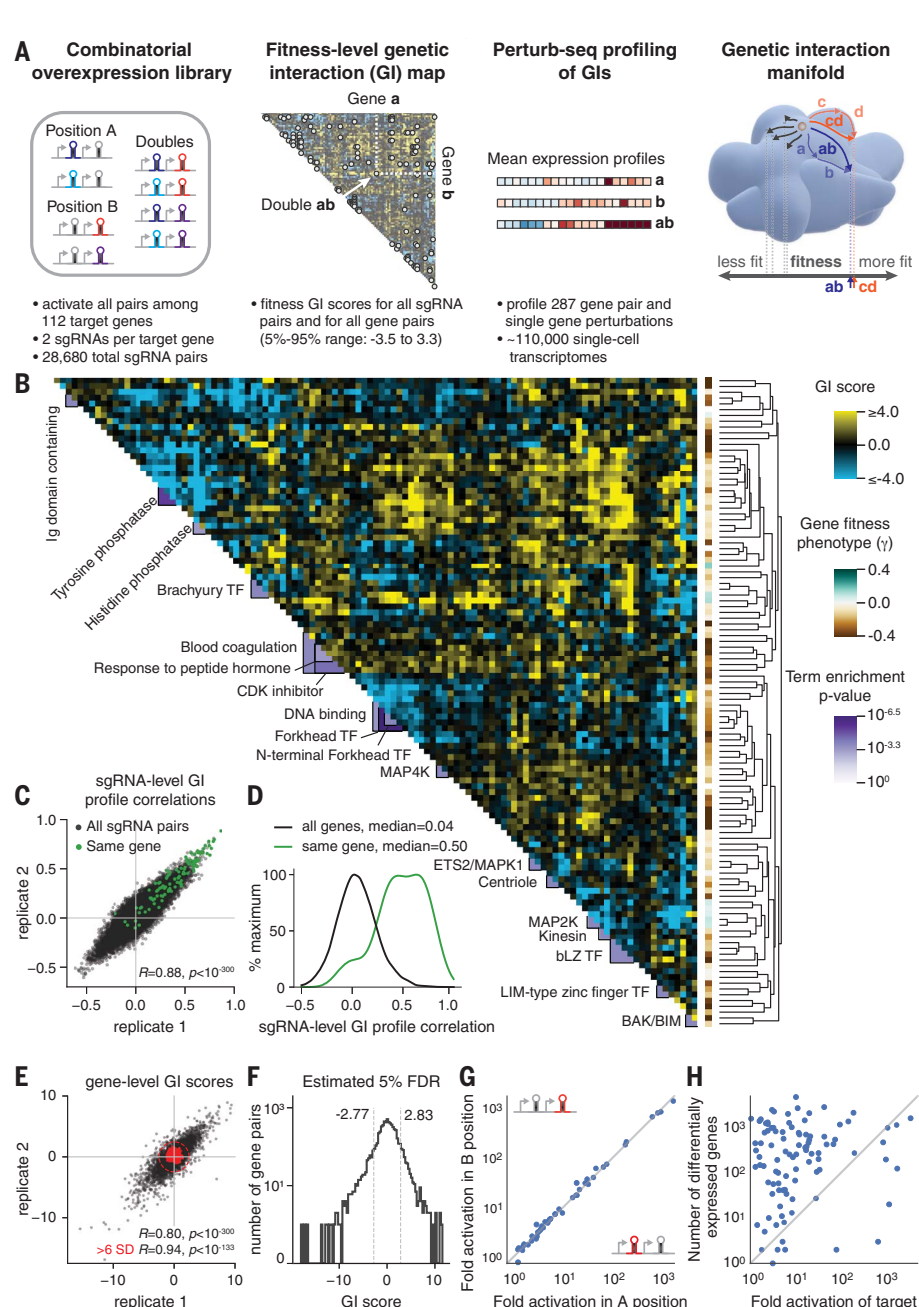
### A Perturb-seq platform for measuring GIs

We reasoned that Perturb-seq, which allows single-cell, pooled transcriptional profiling of CRISPR-mediated perturbations (5–8), would enable us to better resolve the mechanisms underlying GIs. We picked 132 gene pairs from the GI map, chosen both within and between blocks of genes with similar interaction profiles, and targeted each with CRISPRa sgRNA pairs (Fig. 1A, fig. S6A, table S2, and materials and methods). Given the low-rank structure of the fitness GI map, we reasoned that we could broadly sample the biological processes represented without measuring all gene pairs, as many GIs that fell into the “blocks” in the GI map were likely explained by similar mechanisms (materials and methods). We also profiled all single-gene perturbations to enable direct comparison of individual and combined perturbations (i.e., single gene A, single gene B, and pair AB). In total, we obtained transcriptional readouts for 287 perturbations measured across ~110,000 single cells (median 273 cells per condition; materials and methods, fig S1, and table S7) in one pooled experiment.

The Perturb-seq profiles also allowed us to directly assess the performance of our CRISPRa reagents (table S7). Levels of target gene activation spanned a broad range (Fig. 1G and fig. S6, B and C), with a general trend that poorly expressed targets were more highly induced. The A and B positions of the sgRNA cassette performed similarly (Fig. 1G and fig. S6, D and E), and expression of genes neighboring the target was generally unperturbed except when transcripts shared promoter regions (materials and methods; fig. S7, A and B; and table S8). Finally, there was minimal correlation between fold activation and the number of differentially expressed genes, implying that even small increases in the mRNA abundance of some genes can strongly alter a cell’s state ( $R = 0.07$ ; Fig. 1H). The degree of fitness defect was related to the number of differentially expressed genes (fig. S7C).

### Constructing a GI manifold reveals biological processes driving GIs

Whereas GI maps assign a scalar score to each GI, our Perturb-seq approach instead associates a transcriptional phenotype. We viewed this ensemble of measurements as defining a high-dimensional analog of a GI map, here termed a GI manifold. In our manifold analogy, each possible cellular transcriptional state defines a point on a high-dimensional surface. By applying a diverse set of perturbations and measuring the

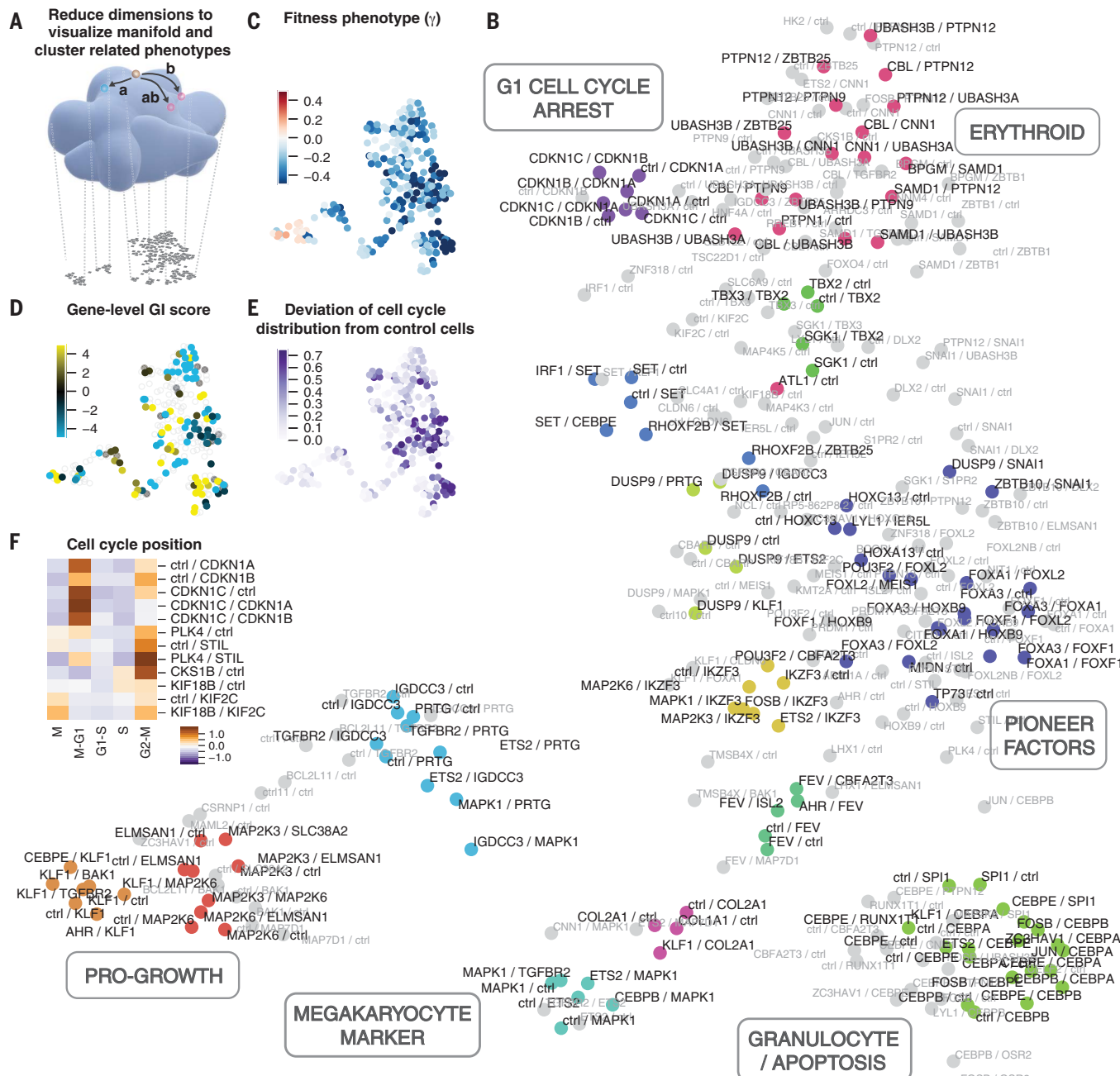


**Fig. 1. CRISPRa fitness-level GI map.** (A) Experimental strategy. Pairs of genes were systematically coactivated using dual sgRNA CRISPRa libraries and a GI map was generated from the fitness measurements. A subset of GIs was then profiled transcriptionally using Perturb-seq. These high-dimensional measurements define a surface called a GI manifold. Distinct GIs that lie in markedly different parts of the GI manifold may result in similar outcomes when viewed only at the level of fitness. (B) CRISPRa fitness-level GI map. Gene-level GI profiles were clustered by average linkage hierarchical clustering based on Pearson correlation. Clusters were annotated by assigning DAVID annotations if a DAVID term was significantly enriched in that cluster [hypergeometric  $\ln(p) \leq -7.5$ ; see materials and methods]. (C and D) GI profile correlation between pairs of sgRNAs targeting any genes (black) or the same gene (green). Data are displayed as scatter plot of replicates (C) and histogram of replicate-averaged GIs (D). (E and F) Gene-level GI scores generated by averaging all sgRNA-level GIs for each gene pair. (E) Scatter plot of replicates. Red points indicate nontargeting control sgRNA pairs, and dashed line indicates a radius of 6 SDs from nontargeting controls. (F) Histogram of gene-level GI scores with estimated empirical 5% false discovery rate threshold. (G) Comparison of fold activation of target gene measured by Perturb-seq when the targeting sgRNA is in the A or B position in the dual sgRNA expression cassette. (H) Fold activation of the target gene compared with the total number of differentially expressed genes.

resulting states (as we have), it becomes possible to infer the shape of this surface (Fig. 1A). Moreover, as each GI is characterized by a rich phenotype, we envisioned that this perspective would allow us to organize GIs by common features and globally examine their mechanistic underpinnings.

To visualize this GI manifold, we used UMAP [uniform manifold approximation and projection (14)] to project the mean expression profiles for our 287 perturbations into a two-dimensional plane (Fig. 2, A and B; a single-cell version is shown in fig. S8). This algorithm approximates

the shape of a high-dimensional surface in two dimensions by trying to conserve nearest neighbor relationships. Perturbations that induce similar transcriptional changes then naturally cluster close to each other; we defined stable clusters using the HDBSCAN algorithm [(15);



**Fig. 2. Visualization of the GI manifold.** (A) Using diverse genetic perturbations, the structure of the GI manifold can be inferred and then visualized by dimensionality reduction to a plane. (B) UMAP projection of all single-gene and gene-pair Perturb-seq profiles. Each dot represents a genetic perturbation characterized by its mean expression profile. Clusters of transcriptionally similar perturbations are colored identically, whereas gray dots are perturbations that do not fall within stable clusters.

(C) Fitness measurements from the GI map expressed as gene-pair growth phenotypes ( $\gamma$ ). (D) GI scores from the fitness-level GI map. Single-gene perturbations are not included. (E) Cell cycle deviation scores. Stronger scores indicate alteration from the distribution of cell cycle positions observed in unperturbed cells. (F) Relative enrichment or depletion of cell cycle phases relative to unperturbed cells induced by selected genetic perturbations.

materials and methods]. Both fitness and GI scores were distributed throughout the GI manifold (Fig. 2, C and D), in accordance with the idea that scalar fitness measurements collapse the much larger landscape of transcriptional states (Fig. 1A). By contrast, overlaying markers derived from the underlying transcriptional data allowed us to gain insight into the mechanisms of individual GIs by looking at common features of perturbations within each cluster. For example, one cluster of mean expression profiles (“G<sub>1</sub> cell cycle arrest”) contained canonical regulators of the cell cycle. In this case, the underlying single-cell data confirmed that these perturbations induced cell cycle arrest in the expected cell cycle stages (Fig. 2, E and F), explaining the growth defect.

We observed clusters of perturbations that caused cells to induce erythroid, granulocyte, or megakaryocyte markers, which is consistent with the known multilineage potential of the K562 model (Figs. 2B and 3, A and B, and fig. S9) (16, 17). These results suggested that cell differentiation or priming and a concomitant decrease in proliferation explained some of the structure of the GI manifold. For example, many interactions surrounding *CBL*, its regulators *UBASH3A/B*, and several multisubstrate tyrosine phosphatases (e.g., *PTPN9/12*) induced erythroid markers, suggesting a common mechanism in the regulation of receptor tyrosine kinase signaling (18). By contrast, the granulocyte cluster mostly contained perturbations of canonical regulators such as C/EBP- $\alpha$ , - $\beta$ , and - $\epsilon$  (*CEBPA/B/E*) and PU.1 (*SPI1*). Finally, a cluster of perturbations induced expression of the canonical megakaryocyte marker CD41, but these cells did not adopt the charac-

teristic morphological features of megakaryocytes, as seen by microscopy (fig. S9G), suggesting that they are at best primed toward megakaryocytic differentiation (16, 17).

To test our ability to better resolve specific interactions using Perturb-seq, we examined a strong synergistic interaction identified by our fitness GI map between *CBL* and *CNN1* (calponin) that belonged to the erythroid cluster. *CNN1* is a poorly characterized gene that is annotated as a smooth muscle–specific protein, although it is expressed in many cell types (19, 20). Overexpressing either gene induced similar transcriptional changes, and single-cell analysis revealed an apparent progression of phenotypes from unperturbed through singly perturbed to doubly perturbed *CBL/CNN1* cells (Fig. 3, C and D). Consistent with an erythroid transcriptional program, overexpression of *CBL* and *CNN1* caused strong induction of canonical markers: hemoglobin genes (sixfold to 39-fold), an iron importer involved in heme biosynthesis (*SLC2A5A37*, 13-fold), and the blood group antigen CD235a (*GYP4A*, twofold) (Fig. 3, C and E, and fig. S10, A and B) (17). Furthermore, overexpression of *CBL* and *CNN1* transgenes in a human erythroid progenitor model (HUDEP2 cells) individually and in combination also induced markers of erythroid differentiation (Fig. 3, F and G, and fig. S10, C and D) (21).

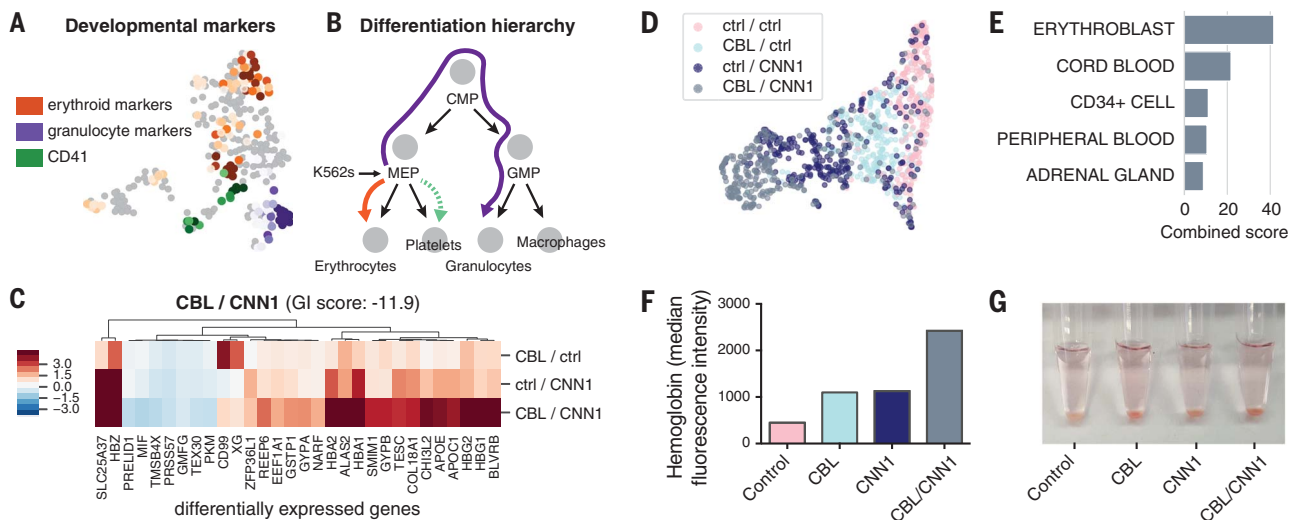
This example highlighted how Perturb-seq analysis can directly lead to a hypothesis about the biology underlying a GI even when one of the components is poorly understood. More generally, because our approach was sensitive to single-cell phenotypes and incomplete differentiation and could simultaneously detect signa-

tures of multiple differentiation states, it could facilitate higher-order combinatorial perturbation screens aimed at improving protocols for driving cells into distinct differentiation states.

### Quantitative modeling of GIs defines mechanisms of interaction

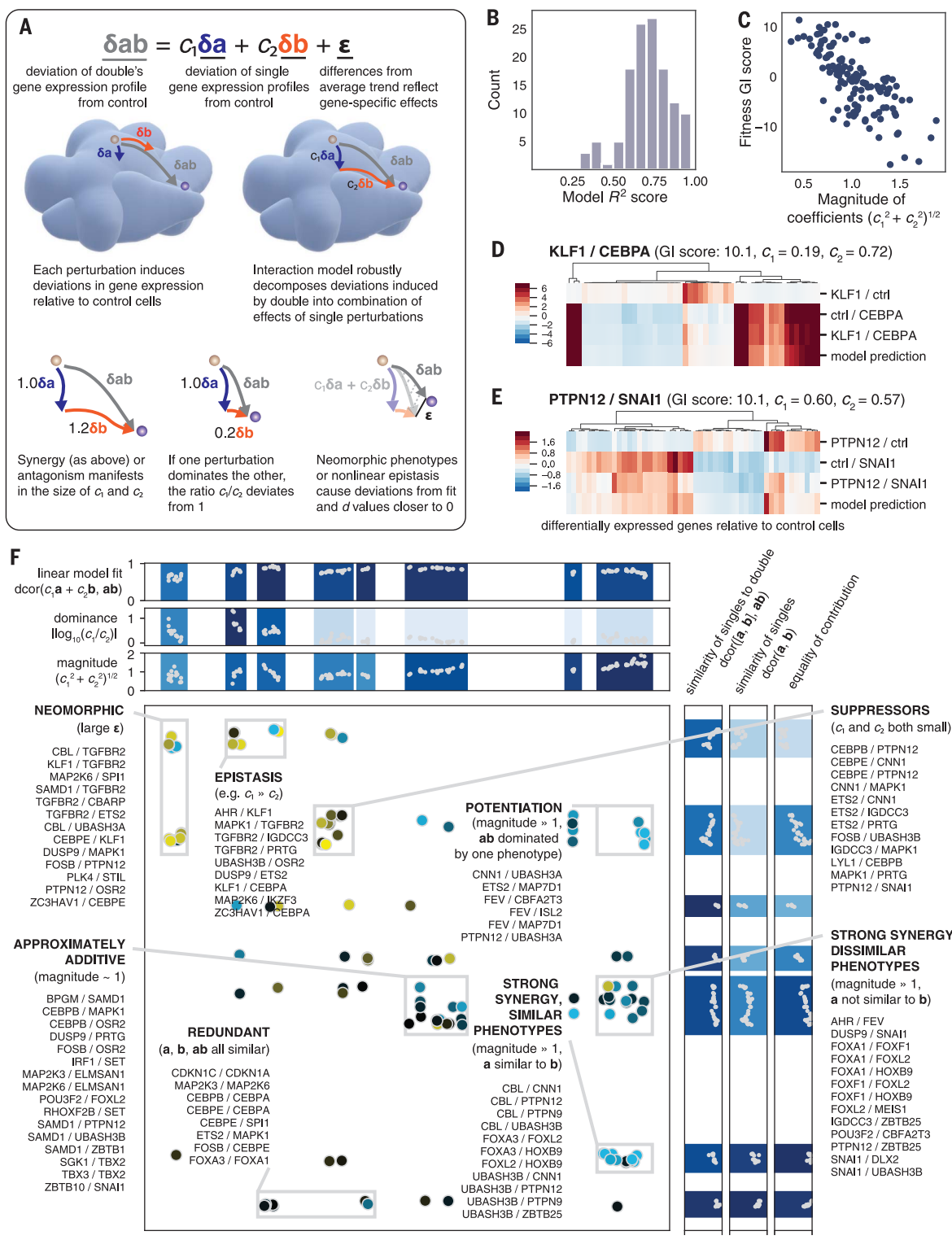
Our large collection of matched single and double overexpression transcriptional phenotypes provides us with the opportunity to quantitatively model GIs directly from transcription profiles without using the fitness GI map. We devised an approach based on fitting a regression model,  $\delta\text{ab} = c_1\delta\text{a} + c_2\delta\text{b} + \epsilon$ , which decomposes the transcriptional changes observed in doubly perturbed cells ( $\delta\text{ab}$ ) as a linear combination of the transcriptional changes induced by the two single perturbations of genes  $a$  and  $b$  ( $c_1\delta\text{a} + c_2\delta\text{b}$ ) and an error term ( $\epsilon$ ) that contains unmodeled or nonlinear effects. The coefficients  $c_1$  and  $c_2$  then effectively measure how much of the phenotype is accounted for by each single perturbation (Fig. 4A). This linear model of transcriptional GIs explained, on average, more than 70% of the variance in gene expression (Fig. 4B; mean  $R^2 = 0.71$ ).

GI maps traditionally classify interactions as either buffering (indicating antagonism, GI score positive) or synthetic sick/lethal (SSL, indicating synergy, GI score negative). We observed a robust anticorrelation ( $R = -0.72$ ) between the magnitude of the coefficients  $c_1$  and  $c_2$  and the fitness-based GI score (Fig. 4C). An intuitive explanation was that buffering interactions travel less “far” (smaller coefficients) along the GI manifold, whereas SSL interactions travel farther (bigger coefficients) (Fig. 4A).



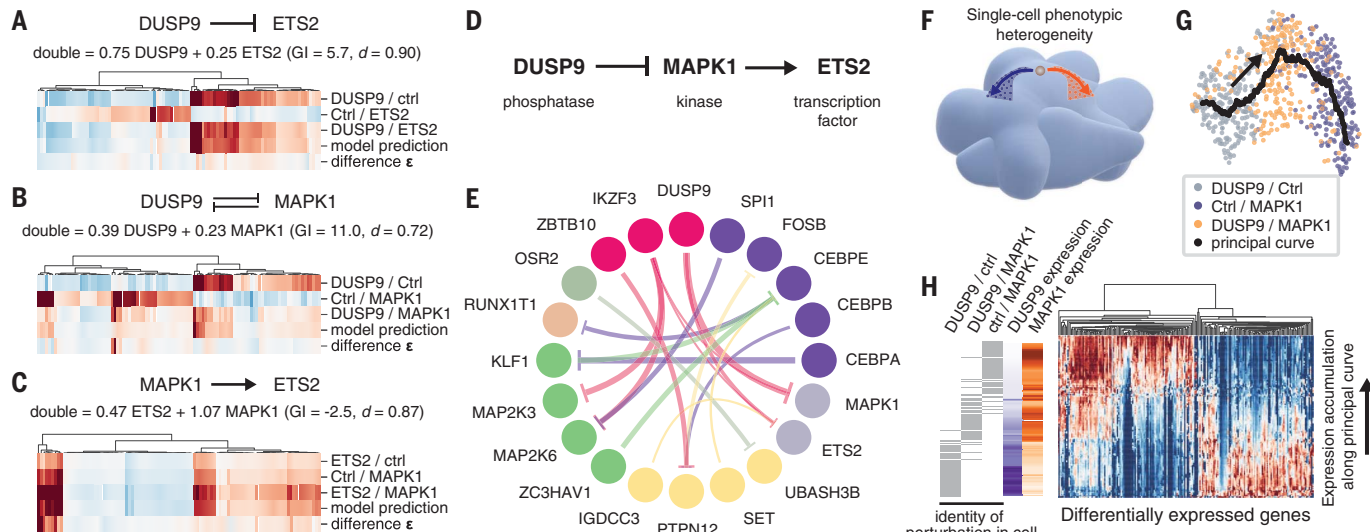
**Fig. 3. Dissecting a genetic interaction using Perturb-seq.** (A) Expression of marker genes for different hematopoietic cell types in the GI manifold UMAP projection. Color is scaled by mean expression Z-score of a marker gene panel. (B) Hematopoietic differentiation hierarchy. K562 cells are a poorly differentiated, erythroid-like cancer cell line. (C) Perturb-seq profiling of the *CBL/CNN1* GI. Average transcriptional profiles for the two constituent single perturbations are compared with the double perturbation. Heat maps show deviation in gene expression relative to unperturbed

cells. (D) UMAP projection of single-cell Perturb-seq data in the *CBL/CNN1* interaction. Each dot is a cell colored according to genetic background. (E) ARCHS4 (35) cell-type term enrichment for genes showing large expression changes in *CBL/CNN1* doubly perturbed cells. (F) Expression of hemoglobin in HUDEP2 cells upon cDNA overexpression of *CBL* or *CNN1*. Hemoglobin was labeled with anti-fetal hemoglobin antibody and measured by flow cytometry. (G) Pelleted HUDEP2 cells. Hemoglobin expression appears red.



**Fig. 4. Quantitative model for high-dimensional GIs.** (A) Model of transcriptional genetic interactions. Different transcriptional states define points on the surface of the GI manifold, and genetic perturbations define vectors of travel. The model decomposes double perturbations as a linear combination of the two constituent single perturbations. (B) Model fit across all GIs measured with Perturb-seq. (C) Magnitude of model coefficients compared with GI score from the fitness-level GI map. (D and E) Application of the model to selected GIs. For each GI, transcriptional profiles for the two constituent single

perturbations are compared with the double perturbation and the model fit. Heat maps show deviation in gene expression relative to unperturbed cells. (F) Visualization of all measured GIs in Perturb-seq experiment. Each GI was characterized using features derived from the model (x-axis) and by measures of similarity among the transcriptional profiles (y-axis). These two viewpoints were each clustered and collapsed to a single dimension using t-SNE to define the two axes. The features defining the two axes are plotted next to them. Categories of GIs are annotated based on features shared within the clusters.

**Fig. 5. Inferring gene-regulatory logic underlying GIs. (A to C)**

Application of linear genetic interaction model to GIs among *DUSP9*, *MAPK1*, and *ETS2*. (D) Order of pathway inferred from model fits. (E) Epistatic buffering interactions oriented using the genetic interaction model. Each arrow denotes a genetic interaction originating in the gene that dominates when the two genes are simultaneously perturbed. Arrow size denotes the degree of dominance as measured by asymmetry of model coefficients. Genetic perturbations with similar transcriptional profiles are colored identically. (F) Stochastic heterogeneity can cause individual cells (dots) bearing a given genetic perturbation to explore the space on the GI manifold surrounding the

average direction of travel (arrows). (G) UMAP projection of single cells with overexpression of *DUSP9* and/or *MAPK1*. Black line represents the principal curve, which tracks the primary direction of variation in the dataset that can be used to order all cells. (H) Gene expression averaged along the principal curve. Each row denotes a cell ordered according to position along the principal curve. The left three columns indicate that cell's genetic background. At each point, cells that are close on the principal curve are averaged to produce a local estimate of median gene expression. The heat map shows normalized expression of differentially expressed genes. The *DUSP9* and *MAPK1* expression columns show the same data for the targeted genes.

To explore the ability of Perturb-seq to better resolve GIs [by analogy with past efforts in other systems (1, 22, 23)], we examined two strong buffering interactions that each had scores of +10.1 in our fitness GI map but appeared to behave differently on a transcriptional level. Analysis by Perturb-seq revealed that the GI between *KLF1* and *CEBPA* resulted from genetic epistasis (i.e., one single overexpression phenotype masking the other), resulting in smaller, asymmetrical coefficients ( $c_1 = 0.19$ ,  $c_2 = 0.72$ ; Fig. 4D and fig. S11A). By contrast, the *PTPN12/SNAI1* GI resulted from genetic suppression (i.e., when combined overexpression of two genes attenuated each other's individual phenotypes), resulting in two smaller coefficients (e.g., *PTPN12/SNAI1*,  $c_1 = 0.60$ ,  $c_2 = 0.57$ ; Fig. 4E and fig. S11B). Finally, as discussed above, synergistic or synthetic lethal interactions tended to result in two larger coefficients (e.g., *CBL/CNN1*,  $c_1 = 1.24$ ,  $c_2 = 0.8$ ; Fig. 3C and fig. S11C).

A central question when considering GIs is how often new or unexpected (neomorphic) phenotypes emerge through the combined action of genes. A relatively small number of GIs (lower mode in Fig. 4B) deviated from the expectation given by the linear model (which we quantified by distance correlation  $d$ ; materials and methods and fig. S11D). A common neomorphic behavior similar to an example observed in yeast (22) occurred when a perturbation that had little transcriptional effect on its own appeared to enhance the effects of a second perturbation (e.g., *FEV/CBFA2T3*,  $d = 0.74$ ; fig. S11E). We also observed

relatively rare instances in which double phenotypes appeared to be completely unexpected [e.g., the physical interactors *PLK4/STIL* (24)  $d = 0.53$ ; fig. S11F]. The model's parameters thus provided a simple, useful summary of how perturbations combine (table S9).

To look for structure among interactions, we used a two-dimensional visualization and clustering technique (25) (table S9 and materials and methods). One axis grouped interactions according to properties derived from the model coefficients ( $c_1$ ,  $c_2$ ,  $d$ ), whereas the other grouped interactions according to how correlated the underlying transcriptional responses were (materials and methods). The resulting figure (Fig. 4F) identified numerous distinct categories of interaction, showing that the model can serve as a generalization of the one-dimensional “buffering versus synthetic lethal” paradigm that has typically been used to categorize genetic interactions.

### Ordering genes into linear pathways using Perturb-seq

The linear GI model allowed us to make hypotheses about gene regulation because it can identify, for example, which single perturbation phenotype better explains the double perturbation phenotype. We examined the strong GIs among the genes *DUSP9*, *ETS2*, and *MAPK1*. In the *DUSP9/ETS2* interaction (Fig. 5A), the *DUSP9* phenotype dominated, suggesting that *DUSP9* overexpression antagonized *ETS2*. Similarly, *DUSP9* and *MAPK1* antagonized each other's activities (Fig. 5B). Fi-

nally, *ETS2* and *MAPK1* induced similar phenotypes, and *ETS2* transcription was activated in all backgrounds (9.3-fold in *MAPK1*, 9.2-fold in *ETS2*, and 35.8-fold in *MAPK1/ETS2* overexpression; Fig. 5C). This type of interaction, in which a perturbation (*MAPK1*) acts at least partly by upregulating its partner (*ETS2*), was uncommon in our dataset (fig. S11G). Taken together, these results suggested a linear regulatory pathway in which *DUSP9* (a phosphatase) inhibits *MAPK1* (a kinase) that activates *ETS2* (a transcription factor), which is consistent with the known biology of these gene families (Fig. 5D) (26). Following similar logic, the model allowed us to orient all the buffering interactions in which one perturbation is epistatic to another (Fig. 5E).

### Single-cell heterogeneity reveals the trajectory of GIs

The single-cell resolution afforded by Perturb-seq can reveal phenotypic heterogeneity for some GIs that we reasoned could yield further insight into their mechanism (Fig. 5F; compare fig. S8). For example, cells overexpressing both *DUSP9* and *MAPK1* showed a range of phenotypes spanning the transcriptional states observed in cells overexpressing either *DUSP9* or *MAPK1* alone (Fig. 5G). In particular, we observed cells in which *DUSP9* and *MAPK1* appeared to suppress each other's activity entirely.

We reasoned that we could therefore identify *DUSP9* or *MAPK1* regulatory targets that showed differing sensitivity to the levels of these proteins

by exploiting single-cell data. To order cells in an unbiased way by “phenotype,” we computed a principal curve measuring the path of maximum variation in the dataset [(27); Fig. 5G and materials and methods]. Examining median-filtered gene expression (materials and methods) along this curve revealed distinct classes of transcripts regulated by *DUSP9* or *MAPK1* activity (e.g., *GYP*) appeared to be more sensitive to *DUSP9* activity than did *HBZ*; Fig. 5H and fig. S12A). This variation did not appear to be the result of stable differences in the expression of *MAPK1* and *DUSP9* (fig. S12B), suggesting a possible role either for historical differences or stochastic gene expression. Perturb-seq can therefore reveal graded phenotypes resulting from antagonism between two proteins.

### Predicting GIs using a recommender system

One possible strategy to address the scale of genetic interactions is to pursue a split experi-

mental and computational approach, in which only a subset of interactions is sampled [either randomly or through “compressed” experimental designs (28)] and the remainder is predicted computationally (Fig. 6A) (7, 29, 30). Perturb-seq provides a scalable means of constraining these types of searches (7), enabling exploration of the GI landscape.

There is substantial similarity between this problem and that of predicting a person’s shopping preferences based on past buying behavior, which is commonly addressed using “recommender system” algorithms. Many of these approaches can exploit low-rank structure such as that seen in GI maps and leverage additional side information obtained by other means to improve predictive power. We examined the Perturb-seq profiles of single-gene overexpression (fig. S13A) and found that there was a modest concordance between GI profile and Perturb-seq profile correlations ( $R = 0.29$ ,  $p < 10^{-103}$ ; fig. S13B), suggesting that the transcriptional data provided a complementary,

scalable means of comparing genes that might inform GI prediction.

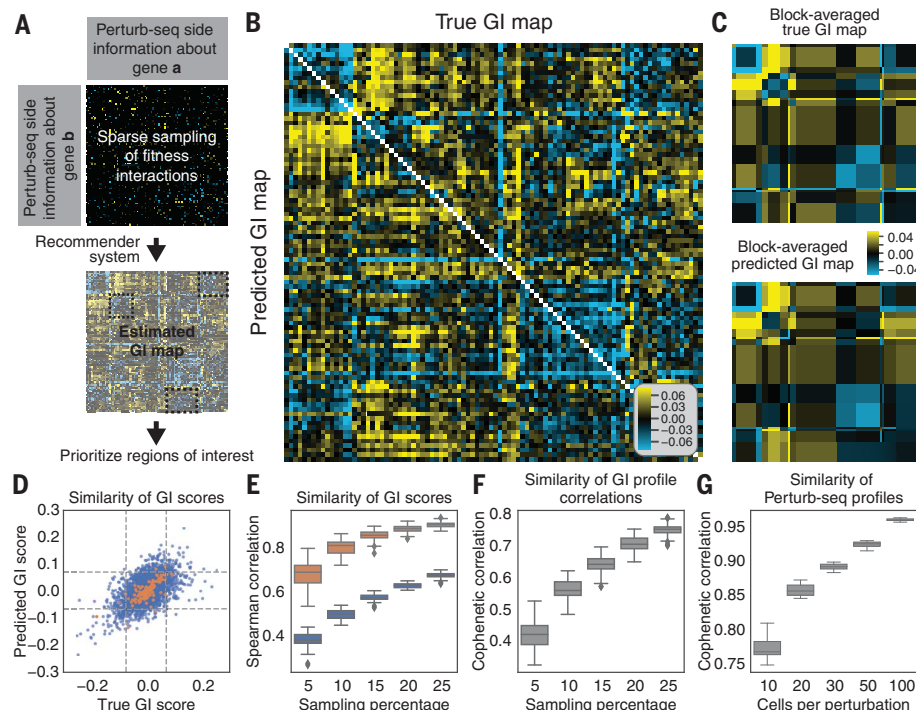
We constructed a matrix factorization model for fitness GIs and then constrained this model to encourage similar interaction profiles among genes that induced similar transcriptional changes (fig. S14A and materials and methods) (30, 31). We then predicted unobserved GIs using this model trained on different fractions of randomly subsampled interactions (Fig. 6A and materials and methods). The end result (Fig. 6, B to D) preserved much of the large-scale structure of the map as seen through block averaging of GI scores (Fig. 6, C and D; fig. S14B; and materials and methods). Our approach was also substantially better than random sampling at predicting the top 10% of interactions and reasonably preserved both the rank order of all interactions (Spearman  $\rho \approx 0.5$  at 10% sampling; Fig. 6, D and E and fig. S14C) and the pairwise similarities between GI profiles (Fig. 6F). Notably, the use of Perturb-seq-derived single perturbation profiles as side information substantially improved performance (fig. S14D). These results suggest that the hybrid approach can nominate blocks of GIs for in-depth study.

Finally, we used our data to model the minimum number of cells that would be required to perform larger experiments. By downsampling our measured perturbations and reperforming our analyses, we observed that as few as 50 cells per perturbation could be sufficient, meaning that  $\sim 10^6$  cells would be needed to collect side information for the entire set of protein-coding sequences (Fig. 6G).

### Discussion

A central goal of genetics is to understand the relationship between the set of genes that a cell expresses and its phenotype. However, this relationship is challenging to study because many phenotypes emerge only through the coordinated action of multiple genes. Here, we used Perturb-seq to manipulate a large number of gene pairs and then measure the resulting changes in cell state. This ensemble of measurements described a high-dimensional surface called a GI manifold. By interpreting and modeling the GI manifold, we can gain several insights into how complex phenotypes emerge.

First, the transcriptional profiles can distinguish distinct outcomes such as cell death, slow growth, and differentiation to a variety of cell states that would appear equivalent at the fitness level. We also identified both canonical genes (e.g., *KLF1*, *GATA1*) and unexpected genes (e.g., *CNN1*) that interacted to promote differentiation to a specific cell state (erythropoiesis). As our single-cell approach is sensitive to multiple outcomes or perturbations with incomplete penetrance, it is a natural strategy to pursue combinatorial searches for factors driving (trans)differentiation (4). Second, the shape of the GI manifold can reveal how GIs arise. We derived a simple, geometric GI model and used it to identify the different ways in which genetic perturbations combine to yield new phenotypes, allowing us to, for example,



**Fig. 6. Recommender system for exploring the GI landscape.** (A) Schematic of prediction strategy. Fitness phenotypes of a limited subset of GIs are measured. Each gene is characterized by its Perturb-seq transcriptional profile, and similarity among these profiles is used as side information to constrain a recommender system model to impute remaining fitness GI scores and highlight regions of interest. (B) True versus predicted GI map obtained by prediction from 10% of randomly sampled fitness-level GIs. (C) Block-averaged true and predicted GI maps obtained by averaging GI scores within clusters. (D) Scatter plot of true and predicted GI scores (blue dots) from (B). The dashed lines show 5 and 95% quantiles, which are used to designate strong GIs. Orange dots show equivalent scatter for block-averaged GI scores in (C). (E) Spearman correlation between true and predicted GI scores at different levels of random sampling. Fifty random subsets were measured for each sampling level. Blue and orange denote individual and block-averaged GIs, respectively. (F) Cophenetic correlation of GI profiles as a function of sampling level, measuring the similarity of correlation structure in the true and predicted GI maps. (G) To assess scaling ability, the representation of each perturbation in the Perturb-seq experiment was randomly down-sampled to different levels of representation. Plot shows cophenetic correlation between down-sampled and true transcriptional profiles used to construct the GI manifold visualization of Fig. 2.

order genes into linear pathways. We and others have established that functionally related genes interact similarly—in geometric terms, the GI manifold is therefore highly constrained, enabling imputation strategies (*1, 2, 7, 12, 29*). Our results provide a strategy for exploring large spaces of combinatorial genetic interactions by measuring only a subset of fitness-level GIs. This provides a complementary approach to efforts based on composite measurements, compressed sensing, and rich readouts to predict unmeasured GIs (*7, 28*). By intelligently measuring and exploring the GI manifold, one can start to create a global view of the nonlinear mapping between genotype and phenotype. Such approaches should enable large-scale searches for synthetic lethal interactions in cancer, the discovery of gene targets that lessen the severity of genetic disease, and, more generally, the understanding of how complex, multigenic interactions govern biological traits and disease risk.

## REFERENCES AND NOTES

1. M. Costanzo *et al.*, *Cell* **177**, 85–100 (2019).
2. J. Domingo, P. Baeza-Centurion, B. Lehner, *Annu. Rev. Genomics Hum. Genet.* **20**, annurev-genom-083118-014857 (2019).
3. J. L. Hartman 4th, B. Garvik, L. Hartwell, *Science* **291**, 1001–1004 (2001).
4. K. Takahashi, S. Yamanaka, *Cell* **126**, 663–676 (2006).
5. D. A. Jaitin *et al.*, *Cell* **167**, 1883–1896.e15 (2016).
6. B. Adamson *et al.*, *Cell* **167**, 1867–1882.e21 (2016).
7. A. Dixit *et al.*, *Cell* **167**, 1853–1866.e17 (2016).
8. P. Datlinger *et al.*, *Nat. Methods* **14**, 297–301 (2017).
9. M. Gasperini *et al.*, *Cell* **176**, 377–390.e9 (2019).
10. J. M. Replogle *et al.*, *bioRxiv* **503367**, (2018).
11. L. A. Gilbert *et al.*, *Cell* **159**, 647–661 (2014).
12. M. A. Horlbeck *et al.*, *Cell* **174**, 953–967.e22 (2018).
13. W. Huang, B. T. Sherman, R. A. Lempicki, *Nucleic Acids Res.* **37**, 1–13 (2009).
14. L. McInnes, J. Healy, J. Melville, UMAP: Uniform manifold approximation and projection for dimension reduction. arXiv:1802.03426v2 [stat.ML] (6 December 2018).
15. L. McInnes, J. Healy, S. Astels, *J. Open Source Softw.* **2**, 205 (2017).
16. R. Alitalo, *Leuk. Res.* **14**, 501–514 (1990).
17. S. H. Orkin, L. I. Zon, *Cell* **132**, 631–644 (2008).
18. K. Kowanetz *et al.*, *J. Biol. Chem.* **279**, 32786–32795 (2004).
19. S. J. Winder, M. P. Walsh, *Cell. Signal.* **5**, 677–686 (1993).
20. GTEx Consortium; Laboratory, Data Analysis & Coordinating Center (LDACC)—Analysis Working Group; Statistical Methods groups—Analysis Working Group; Enhancing GTEx (eGTEx) groups; NIH Common Fund; NIH/NCI; NIH/NHGRI; NIH/NIMH; NIH/NIDA; Biospecimen Collection Source Site—NDRI; Biospecimen Collection Source Site—RPCI; Biospecimen Core Resource—VARI; Brain Bank Repository—University of Miami Brain Endowment Bank; Leidos Biomedical—Project Management; ELSI Study; Genome Browser Data Integration & Visualization—EBI; Genome Browser Data Integration & Visualization—UCSC Genomics Institute, University of California Santa Cruz; Lead analysts; Laboratory, Data Analysis & Coordinating Center (LDACC); NIH program management; Biospecimen collection; Pathology; eQTL manuscript working group; A. Battle, C. D. Brown, B. E. Engelhardt, S. B. Montgomery, *Nature* **550**, 204–213 (2017).
21. R. Kurita *et al.*, *PLOS ONE* **8**, e59890 (2013).
22. S. van Wageningen *et al.*, *Cell* **143**, 991–1004 (2010).
23. N. Van Driessche *et al.*, *Nat. Genet.* **37**, 471–477 (2005).
24. M. Ohta *et al.*, *Nat. Commun.* **5**, 5267 (2014).
25. Y. Cheng, M. T. Wong, L. van der Maaten, E. W. Newell, *J. Immunol.* **196**, 924–932 (2016).
26. C. J. Caunt, S. M. Keyse, *FEBS J.* **280**, 489–504 (2013).
27. C. Trapnell *et al.*, *Nat. Biotechnol.* **32**, 381–386 (2014).
28. B. Cleary, L. Cong, A. Cheung, E. S. Lander, A. Regev, *Cell* **171**, 1424–1436.e18 (2017).
29. N. S. Madhukar, O. Elemento, G. Pandey, *Front. Bioeng. Biotechnol.* **3**, 172 (2015).
30. M. Žitnik, B. Zupan, *J. Comput. Biol.* **22**, 595–608 (2015).
31. M. Xu, R. Jin, Z.-H. Zhou in *Advances in Neural Information Processing Systems*, M. I. Jordan, Y. LeCun, S. A. Solla, Eds. (MIT Press, 2013), pp. 2301–2309.

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**contributions:** T.M.N., M.A.H., L.A.G., and J.S.W. were responsible for the conception, design, and interpretation of the experiments and wrote the manuscript. M.J. and J.M.R. provided critical feedback on the manuscript. L.A.G., M.J., A.X., and J.M.R. constructed GI vectors, cDNA vectors, and libraries. T.M.N., J.M.R., A.X., A.Y.G., M.A.H., and L.A.G. conducted the GI mapping and Perturb-seq experiments. M.A.H. performed the GI map data analysis. T.M.N. performed the Perturb-seq analysis, GI prediction analysis, and modeling of transcriptional genetic interactions. L.A.G., T.M.N., A.Y.G., and A.X. designed and conducted validation experiments. **Competing interests:** T.M.N., M.A.H., L.A.G., M.J., and J.S.W. have filed patent applications related to CRISPR/a screening, Perturb-seq, and GI mapping. J.S.W. consults for and holds equity in KSQ Therapeutics, Maze Therapeutics, and Tenaya Therapeutics. J.S.W. is a venture partner at 5AM Ventures. T.M.N., M.J., J.M.R., and M.A.H. consult for Maze Therapeutics. The remaining authors declare no competing interests. **Data and**

**materials availability:** All data are available in the manuscript, the supplementary materials, or GEO (GSE133344). Notebooks implementing analyses are available at <https://github.com/mhorlbeck> and <https://github.com/thomasmaxwellnorman>.

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Materials and Methods

Figs. S1 to S14

Tables S1 to S9

References (32–35)

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## Exploring genetic interaction manifolds constructed from rich single-cell phenotypes

Thomas M. Norman, Max A. Horlbeck, Joseph M. Replogle, Alex Y. Ge, Albert Xu, Marco Jost, Luke A. Gilbert and Jonathan S. Weissman

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### Manifold destiny

Mapping of genetic interactions (GIs) is usually based on cell fitness as the phenotypic readout, which obscures the mechanistic origin of interactions. Norman *et al.* developed a framework for mapping and understanding GIs. This approach leverages high-dimensional single-cell RNA sequencing data gathered from CRISPR-mediated, pooled perturbation screens. Diverse transcriptomic phenotypes construct a "manifold" representing all possible states of the cell. Each perturbation and GI projects the cell state to a particular position on this manifold, enabling unbiased ordering of genes in pathways and systematic classifications of GIs.

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