# Exponential family measurement error models for single-cell CRISPR screens

#### Abstract

CRISPR genome engineering and single-cell RNA sequencing have transformed biological discovery. Single-cell CRISPR screens unite these two technologies, linking genetic perturbations in individual cells to changes in gene expression and illuminating regulatory networks underlying diseases. Despite their promise, single-cell CRISPR screens present substantial statistical challenges. We demonstrate through theoretical and real data analyses that a standard method for estimation and inference in single-cell CRISPR screens - "thresholded regression" - exhibits attenuation bias and a bias-variance tradeoff as a function of an intrinsic, challenging-to-select tuning parameter. To overcome these difficulties, we introduce GLM-EIV ("GLMbased errors-in-variables"), a new method for single-cell CRISPR screen analysis. GLM-EIV extends the classical errors-in-variables model to responses and noisy predictors that are exponential family-distributed and potentially impacted by the same set of confounding variables. We develop a computational infrastructure to deploy GLM-EIV across tens or hundreds of nodes on clouds (e.g., Microsoft Azure) and high-performance clusters. Leveraging this infrastructure, we apply GLM-EIV to analyze two recent, large-scale, single-cell CRISPR screen datasets, demonstrating improved performance in challenging problem settings.

Keywords: genome engineering, GWAS, GLM, latent variable model, cloud computing

## 1 Introduction

CRISPR is a genome engineering tool that has enabled scientists to precisely edit human and nonhuman genomes, opening the door to new medical therapies (Rothgangl et al. 2021; Musunuru et al. 2021) and transforming biological discovery (Przybyla and Gilbert 2021). Recently, scientists have paired CRISPR genome engineering with single-cell RNA sequencing (Dixit et al. 2016; Datlinger et al. 2017). The resulting assays, known as a "single-cell CRISPR screens," link genetic perturbations in individual cells to changes in gene expression, enabling scientists to causally map genome wide association study (GWAS) variants to their target genes at genome-wide scale (Morris et al. 2021).

Despite their promise, single-cell CRISPR screens present substantial statistical challenges. A major difficulty is that the "treatment" – i.e., the presence or absence of a CRISPR perturbation – is assigned randomly to cells and is not directly observable. As a consequence, one cannot know with certainty which cells were perturbed. Instead, one must leverage an indirect, quantitative proxy of perturbation presence or absence to "guess" which cells received a perturbation. This indirect proxy takes the form of a so-called guide RNA count, with higher counts indicating that a cell is more likely to have been perturbed. The standard approach to single-cell CRISPR screen analysis is to impute perturbation assignments onto the cells by simply thresholding the guide RNA counts; using these imputations, one can attempt to estimate the effect of the perturbation on gene expression. We call this standard approach "thresholded regression" or the "thresholding method."

We study estimation and inference in single-cell CRISPR screens from a statistical perspective, formulating the data generating mechanism using a new class of measurement error models. We assume that the response variable y is a GLM of an underlying predictor variable  $x^*$  and vector of confounders z. We do not observe  $x^*$  directly; rather, we observe a noisy version x of  $x^*$  that itself is a GLM of  $x^*$  and the same set of confounders z. The goal of the analysis is to estimate the effect of  $x^*$  on y using the observed data (x, y, z) only. In the context of the biological application,  $x^*$ , x, y, and z are CRISPR perturbations, guide RNA counts, gene expressions, and technical confounders, respectively.

Our work makes two main contributions. First, we conduct a detailed study of the thresholding method. Notably, we demonstrate on real data that the thresholding method exhibits attenuation bias and a bias-variance tradeoff as a function of the selected threshold, and we recover these phenomena in precise mathematical terms in a simplified Gaussian setting. Second, we introduce a new method, GLM-EIV ("GLM-based errors-in-variables"),

for single-cell CRISPR screen analysis. GLM-EIV extends the classical errors-in-variables model (Carroll et al. 2006) to responses and noisy predictors that are exponential family-distributed and potentially impacted by the same set of confounding variables. GLM-EIV thereby implicitly estimates the probability that each cell was perturbed, obviating the need to explicitly impute perturbation assignments via thresholding. We implement several statistical accelerations (that likely are of independent utility) to bring the cost of GLM-EIV down to within about an order of magnitude of the thresholding method.

Finally, we develop a Docker-containerized application to deploy GLM-EIV at-scale across tens or hundreds of nodes on clouds (e.g., Microsoft Azure) and high-performance clusters. Leveraging this application, we apply GLM-EIV to analyze two recent, large-scale, single-cell CRISPR screen datasets. We find that in some settings, GLM-EIV outperforms thresholded regression by a considerable margin; in other settings the two methods work best in conjunction, with GLM-EIV providing a statistically principled and empirically effective procedure for selecting the threshold.

# 2 Assay background

There are several broad classes of single-cell CRISPR screen assays, each suited to answer a different set of biological questions (Gasperini et al. 2019; Datlinger et al. 2021; Mimitou et al. 2019). In this work we focus on high-multiplicity of infection (MOI) single-cell CRISPR screens, which we motivate and describe here. The human genome consists of genes, enhancers (segments of DNA that regulate the expression of one or more genes), and other genomic elements (that are not of relevance to the current work). GWAS have revealed that the majority (> 90%) of variants associated with diseases lie outside genes and inside enhancers (Gallagher and Chen-Plotkin 2018). These noncoding variants are

thought to contribute to disease by modulating the expression of one or more diseaserelevant genes. Scientists do not know the gene (or genes) through which most noncoding variants exert their effect, limiting the interpretability of GWAS results. A central open challenge in genetics, therefore, is to link enhancers that harbor GWAS variants to the genes that they target at genome-wide scale (Gasperini et al. 2020; Morris et al. 2021).

High MOI single-cell CRISPR screens are the most promising biotechnology for solving this challenge. High MOI single-cell CRISPR screens combine CRISPR interference (CRISPRi) – a version of CRISPR that represses a targeted region of the genome – with single-cell sequencing. The experimental protocol is as follows. First, the scientist develops a library of several hundred to several thousand CRISPRi perturbations, each designed to target a candidate enhancer for repression. The scientist then cultures tens or hundreds of thousands of cells and delivers the CRISPRi perturbations to these cells. The perturbations assort into the cells randomly, with each cell receiving on average 10-40 distinct perturbations. Conversely, a given perturbation enters about 0.1-2% of cells (this work).

After waiting several days for CRISPRi to take effect, the scientist profiles each cell's transcriptome (i.e., its gene expressions) and the set of perturbations that it received. Finally, the scientist conducts perturbation-to-gene association analyses. Figure 1a depicts this process schematically, with colored bars (blue, red, and purple) representing distinct perturbations. For a given perturbation (e.g., the perturbation represented in blue), the scientist partitions the cells into two groups: those that received the perturbation (top) and those that did not (bottom). Next, for a given gene, the scientist runs a differential expression analysis across the two groups of cells, producing an estimate for the magnitude of the gene expression change in response to the perturbation. If the estimated change in expression is large, the scientist can conclude that the enhancer targeted by the perturbation

exerts a strong regulatory effect on the gene. This procedure is repeated for a large set of preselected perturbation-gene pairs. The enhancer-by-enhancer approach is valid because the perturbations assort into cells approximately independently of one another.

The genomics literature has produced a few applied methods for single-cell CRISPR screen analysis (Gasperini et al. 2019; Xie et al. 2019; Barry et al. 2021). Gasperini et al. applied negative binomial GLMs (as implemented in the Monocle software; Trapnell et al. 2014) to carry out the differential expression analysis described above. Xie et al., by contrast, applied chi-squared-like tests of independence for this purpose. Both of these approaches have limitations: the former is not robust to misspecification of the gene expression model, and the latter is unable to correct for the presence of technical confounders. Recently, Barry et al. introduced SCEPTRE, a custom implementation of the conditional randomization test (Candès et al. 2018; Liu et al. 2021) tailored to single-cell CRISPR screen data. SCEPTRE simultaneously adjusts for confounder presence and ensures robustness to expression model misspecification, overcoming limitations of the prior methods and demonstrating state-of-the-art sensitivity and specificity on single-cell CRISPR screen data. In this work we tackle a set of analysis challenges that are complimentary to those addressed by SCEPTRE. Most importantly, we seek to account for the fact that the perturbation is measured with noise, an issue that all available methods (including SCEPTRE) assume away via thresholding. Additionally, we seek to estimate (with confidence) the effect size of a perturbation on gene expression change, an objective that is challenging to attain within the nonparametric hypothesis testing framework of SCEPTRE.

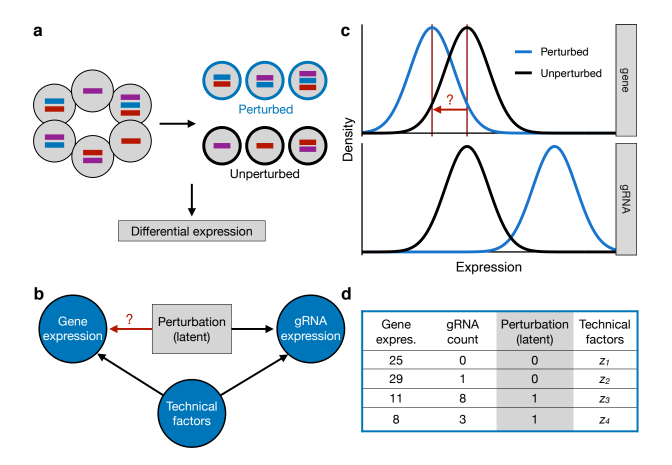


Figure 1: Experimental design and analysis challenges: a, Experimental design. For a given perturbation (e.g., the perturbation indicated in blue), we partition the cells into two groups: perturbed and unperturbed. Next, for a given gene, we conduct a differential expression analysis across the two groups, yielding an estimate of the impact of the given perturbation on the given gene. b, DAG representing all variables in the system. The perturbation (latent) impacts both gene expression and gRNA expression; technical factors act as confounders, also impacting gene and gRNA expression. The target of estimation is the effect of the perturbation on gene expression. c, Schematic illustrating the "background read" phenomenon. Due to errors in the sequencing and alignment processes, unperturbed cells exhibit a nonzero gRNA count distribution (bottom). The target of estimation is the change in mean gene expression in response to the perturbation (top). d, Example data on four cells for a given perturbation-gene pair. Note that (i) the perturbation is unobserved, and (ii) the gene and gRNA data are discrete counts.

## 3 Analysis challenges and proposed statistical model

High MOI single-cell CRISPR screens present several statistical challenges, four of which we highlight here. Throughout, we consider a single perturbation-gene pair. First, the "treatment" variable – i.e., the presence or absence of a perturbation – cannot be directly

observed. Instead, perturbed cells transcribe molecules called quide RNAs (or qRNAs) that serve as indirect proxies of perturbation presence. We must leverage these gRNAs to impute (explicitly or implicitly) perturbation assignments onto the cells (Figure 1b). Second, "technical factors" – sources of variation that are experimental rather than biological in origin – impact the measurement of both gene and gRNA expressions and therefore act as confounders (Figure 1b). Third, the gene and gRNA data are sparse, discrete counts. Consequently, classical statistical approaches that assume Gaussianity or homoscedasticity are inapplicable. Finally, sequenced gRNAs sometimes map to cells that have not received a perturbation. This phenomenon, which we "background contamination," results from errors in the sequencing and alignment processes (Replogle et al. 2020). The marginal distribution of the gRNA counts is best conceptualized as a mixture model (Figure 1c; Gaussian distributions used for illustration purposes only). Unperturbed and perturbed cells both exhibit nonzero gRNA count distributions, but this distribution is shifted upward for perturbed cells. Figure 1d shows example data on four (of possibly tens or hundreds of thousands of) cells. The analysis objective is to leverage the gene expressions and gRNA counts to estimate the effect of the (latent) perturbation on gene expression, accounting for the technical factors.

We propose to model the single-cell CRISPR screen data-generating process using a pair of GLMs. Let  $n \in \mathbb{N}$  be the number of cells assayed in the experiment. Consider a single perturbation and a single gene. For cell  $i \in \{1, ..., n\}$ , let  $p_i \in \{0, 1\}$  indicate perturbation presence or absence; let  $m_i \in \mathbb{N}$  be the number of gene transcripts sequenced; let  $g_i \in \mathbb{N}$  be the number of gRNA transcripts sequenced; let  $d_i^m \in \mathbb{N}$  be the number of gene transcripts sequenced across all genes (i.e., the library size or sequencing depth); let  $d_i^g$  be the gRNA library size; and finally, let  $z_i \in \mathbb{R}^{d-1}$  be the cell-specific technical factors

(e.g., sequencing batch, percent mitochondrial reads, etc.) The letters "m," "g", and "d" stand for "mRNA," "gRNA," and "depth," respectively.

Building on the work of several previous authors (Townes et al. 2019; Svensson 2020; Hafemeister and Satija 2019), Sarkar and Stephens (2021) proposed a simple strategy for modeling single-cell gene expression data, which, in the framework of negative binomial GLMs, is equivalent to using the log-transformed library size as an offset term. Sarkar and Stephens' framework enjoys strong theoretical and empirical support; therefore, we generalize their approach to model *both* gene and gRNA modalities in single-cell CRISPR screen experiments. To this end, we assume that the gene expression counts are given by

$$m_i|(p_i, z_i, d_i^m) \sim NB_{\theta^m}(\mu_i^m); \quad \log(\mu_i^m) = \beta_0^m + \beta_1^m p_i + \gamma_m^T z_i + \log(d_i^m),$$
 (1)

where (i)  $NB_{\theta^m}(\mu_i^m)$  is a negative binomial distribution with mean  $\mu_i^m$  and known size parameter  $\theta^m$ ; (ii)  $\beta_0^m \in \mathbb{R}, \beta_1^m \in \mathbb{R}$ , and  $\gamma_m \in \mathbb{R}^{d-1}$  are unknown parameters; and (iii)  $\log(d_i^m)$  is an offset term. Similarly, we model the gRNA counts by

$$g_i|(p_i, z_i, d_i^g) \sim NB_{\theta^g}(\mu_i^g); \quad \log(\mu_i^g) = \beta_0^g + \beta_1^g p_i + \gamma_q^T z_i + \log(d_i^g),$$
 (2)

where  $\mu_i^g$ ,  $\theta^g$ ,  $\beta_0^g$ ,  $\beta_1^g$ ,  $\gamma_g$ , and  $d_i^g$  are analogous. We use a negative binomial GLM to model the gRNA counts as well as the gene expressions because the gRNA transcripts are generated via the same biological mechanism as the gene transcripts (Datlinger et al. 2017; Hill et al. 2018). Finally, we model the marginal perturbation probability as

$$p_i \sim \text{Bern}(\pi); \quad \pi \in (0, 1/2],$$
 (3)

where  $p_i$  is unobserved. Together, (1 - 3) define the negative binomial GLM-EIV model.

The log-transformed sequencing depth  $\log(d_i^m)$  is included as an offset term in (1) so that  $\beta_0^m + \beta_1^m p_i + \gamma_m^T z_i$  can be interpreted as a relative expression. Exponentiating both sides of (1) reveals that the mean gene expression  $\mu_i^m$  of the *i*th cell is  $\exp\left(\beta_0^m + \beta_1^m p_i + \gamma_m^T z_i\right) d_i^m$ . Because  $d_i^m$  is the sequencing depth,  $\exp\left(\beta_0^m + \beta_1^m p_i + \gamma_m^T z_i\right)$  is the fraction of all transcripts sequenced in the cell produced by the gene under consideration. The target of inference  $\beta_1^m$  is the log fold change in expression in response to the perturbation, controlling for the technical factors. Fold change in this context is the ratio of the mean gene expression in perturbed cells to the mean gene expression in unperturbed cells. Hence,  $\exp(\beta_1^m) = 1$  (i.e.,  $\beta_1^m = 0$ ) indicates no change in expression, whereas  $\exp(\beta_1^m) > 1$  (i.e.,  $\beta_1^m > 0$ ) and  $\exp(\beta_1^m) < 1$  (i.e.,  $\beta_1^m < 0$ ) indicate an increase and decrease in expression, respectively.

In this work we analyze two large-scale, high MOI, single-cell CRISPR screen datasets published by Gasperini et al. (2019) and Xie et al. (2019). Gasperini (resp., Xie) targeted approximately 6,000 (resp., 500) candidate enhancers in a population of approximately 200,000 (resp., 100,000) cells. Gasperini additionally designed several hundred positive control, gene-targeting perturbations and 50 non-targeting, negative control perturbations to assess method sensitivity and specificity.

# 4 Analysis of the thresholding method

We study thresholding from empirical and theoretical perspectives, highlighting several limitations of the approach. In the context of the negative bionomial GLM-EIV model introduced above (1-3), the thresholding method leverages the gRNA counts (2) to impute the latent perturbation indicator (3), thereby reducing the full data generating process to

a single, gene expression model (1). We study Gasperini et al.'s variant of the thresholding method (i.e., thresholded negative binomial regression), as this version of the thresholding method relates most closely to GLM-EIV. The method is defined as follows:

- 1. For a given threshold  $c \in \mathbb{N}$ , let the imputed perturbation assignment  $\hat{p}_i \in \{0, 1\}$  be given by  $\hat{p}_i = 0$  if  $g_i < c$  and  $\hat{p}_i = 1$  otherwise.
- 2. Assume that  $m_i$  is related to  $\hat{p}_i, d_i^m$ , and  $z_i$  through the following GLM:

$$m_i | (\hat{p}_i, z_i, d_i^m) \sim NB_{\theta^m}(\mu_i^m); \quad \log(\mu_i^m) = \beta_0^m + \beta_1^m \hat{p}_i + \gamma_m^T z_i + \log(d_i^m).$$
 (4)

The model (4) is equivalent to the model (2), but the latent perturbation indicator  $p_i$  has been replaced by the imputed perturbation indicator  $\hat{p}_i$ .

3. Fit a GLM to (4) to obtain an estimate and CI for the target of inference  $\beta_1^m$ .

## 4.1 Empirical challenges of the thresholding method

To shed light on empirical challenges of the thresholding method, we applied thresholded negative binomial regression to analyze the set of positive control perturbation-gene pairs in the Gasperini dataset. The positive control pairs consisted of perturbations that targeted gene transcription start sites (TSSs) for inhibition. Repressing the TSS of a given gene decreases its expression; therefore, the positive control pairs *a priori* are expected to exhibit a strong decrease in expression.

To investigate the sensitivity of the thresholding method to threshold choice, we deployed the method using three different choices for the threshold: 1, 5, and 20. We found that the chosen threshold substantially impacted the results (Figure 2a-b): estimates for

fold change produced by threshold = 1 were smaller in magnitude (i.e., closer to the baseline of 1) than those produced by threshold = 5. (Figure 2a.) On the other hand, estimates produced by threshold = 5 and threshold = 20 were more concordant (Figure 2b).

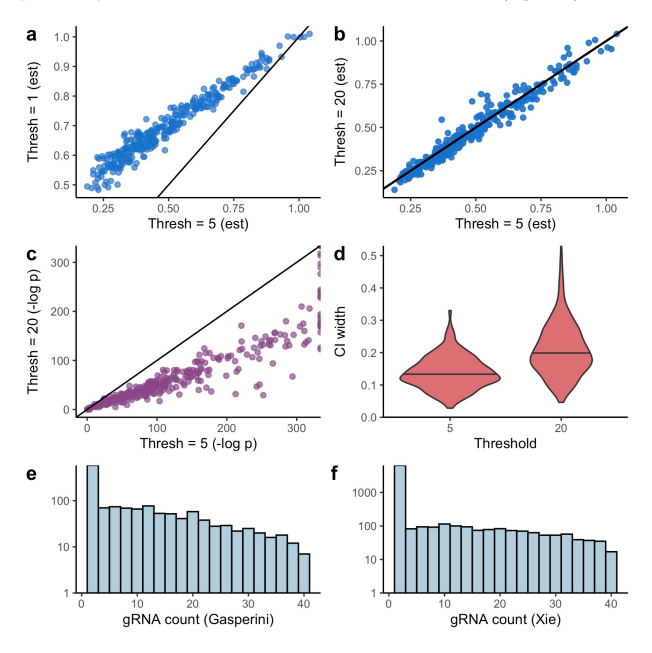


Figure 2: **Empirical challenges of thresholded regression.** a-b, Fold change estimates produced by threshold = 1 versus threshold = 5 (a) and threshold = 20 versus threshold = 5 (b). The selected threshold substantially impacts the results. c-d, p-values (c) and CI widths (d) produced by threshold = 20 versus threshold = 5. The latter threshold yields more confident estimates. e-f, Empirical distribution of randomly-selected gRNA from Gasperini (e) and Xie (f) data (0 counts not shown). The gRNA data do not appear to imply an obvious threshold selection strategy.

We reasoned that thresholded regression systematically underestimated true effect sizes on the positive control pairs, especially for threshold = 1. For a given perturbation, the majority (> 98%) of cells are unperturbed. This imbalance leads to an asymmetry: misclassifying unperturbed cells as perturbed is intuitively "worse" than misclassifying perturbed cells as unperturbed. Misclassified unperturbed cells contaminate the set of truly perturbed cells, leading to attenuation bias; by contrast, misclassified perturbed cells are swamped in number and "neutralized" by the truly unperturbed cells. Setting the threshold to a large number reduces the unperturbed-to-perturbed misclassification rate, decreasing bias.

We hypothesized, however, that the reduction in bias obtained by selecting a large threshold causes the variance of the estimator to increase. To investigate, we compared p-values and confidence intervals produced by threshold = 5 and threshold = 20 for the target of inference  $\beta_1^m$ . We found that threshold = 5 yielded smaller (i.e., more significant) p-values and narrower confidence intervals than did threshold = 20 (Figures 2c-d). We concluded that the threshold controls a bias-variance tradeoff: as the threshold increases, the bias of the estimator decreases and the variance increases.

Finally, to determine whether there is an "obvious" location at which to draw the threshold, we examined the empirical gRNA count distributions and checked for bimodality. Figures 2e and 2f display the empirical distribution of a randomly-selected gRNA from the Gasperini and Xie datasets, respectively (counts of 0 omitted). The distributions peak at 1 and then taper off gradually; there does not exist a sharp boundary that cleanly separates the perturbed from the unperturbed cells. Overall, we concluded that the thresholding method faces several challenges: (i) the threshold is a tuning parameter that significantly impacts the results; (ii) the threshold mediates an intrinsic bias-variance tradeoff; and (iii) the gRNA count distributions do not imply a clear threshold selection strategy.

### 4.2 Theoretical challenges of the thresholding method

Next, we studied the thresholding method from a theoretical perspective, recovering in a simplified Gaussian setting phenomena revealed in the empirical analysis. Suppose we observe gRNA expression and gene expression data  $(g_1, m_1), \ldots, (g_n, m_n)$  on n cells from the following linear model:

$$m_i = \beta_0^m + \beta_1^m p_i + \epsilon_i; \quad g_i = \beta_0^g + \beta_1^g p_i + \tau_i; \quad p_i \sim \text{Bern}(\pi); \quad \epsilon_i, \tau_i \sim N(0, 1),$$
 (5)

where  $p_i, \tau_i$ , and  $\epsilon_i$  are independent. For a given threshold  $c \in \mathbb{R}$ , the imputed perturbation assignment  $\hat{p}_i$  is  $\hat{p}_i = \mathbb{I}(g_i \geq c)$ . The thresholding estimator  $\hat{\beta}_1^m$  is the OLS solution, i.e.  $\hat{\beta}_1^m = \left[\sum_{i=1}^n (\hat{p}_i - \overline{\hat{p}})^2\right]^{-1} \left[\sum_{i=1}^n (\hat{p}_i - \overline{\hat{p}})(m_i - \overline{m})\right]$ . We derive the almost sure limit of  $\hat{\beta}_1^m$ :

**Proposition 1.** The almost sure limit (as  $n \to \infty$ ) of  $\hat{\beta}_1^m$  is

$$\hat{\beta}_1^m \xrightarrow{a.s.} \beta_1^m \left( \frac{\pi(\omega - \mathbb{E}[\hat{p}_i])}{\mathbb{E}[\hat{p}_i](1 - \mathbb{E}[\hat{p}_i])} \right) \equiv \beta_1^m \gamma(\beta_1^g, \pi, c, \beta_0^g), \tag{6}$$

where  $\mathbb{E}[\hat{p}_i] = \zeta(1-\pi) + \omega\pi$ ,  $\omega \equiv \Phi(\beta_1^g + \beta_0^g - c)$ , and  $\zeta \equiv \Phi(\beta_0^g - c)$ .

The function  $\gamma: \mathbb{R}^4 \to \mathbb{R}$  does not depend on the gene expression parameters  $\beta_1^m$  or  $\beta_0^m$ . The asymptotic relative bias  $b: \mathbb{R}^4 \to \mathbb{R}$  of  $\hat{\beta}_1^m$  is given by

$$b(\beta_1^g, \pi, c, \beta_0^g) \equiv \frac{1}{\beta_1^m} \left( \beta_1^m - \lim_{\text{a.s.}} \hat{\beta}_1^m \right) = 1 - \gamma(\beta_1^g, \pi, c, \beta_0^g).$$

Having derived an exact expression for the asymptotic relative bias of  $\hat{\beta}_1^m$ , we can prove several results about this quantity. We fix  $\pi$  to 1/2 for simplicity. (In reality,  $\pi$  is smaller, but the relevant statistical phenomena emerge for  $\pi = 1/2$ .) First, the thresholding

estimator strictly underestimates (in absolute value) the true value of  $\beta_1^m$  over all choices of the threshold c and over all values of the regression coefficients ( $\beta_0^m, \beta_1^m$ ) and ( $\beta_0^g, \beta_1^g$ ). This phenomenon, called attenuation bias, is a common attribute of estimators that ignore measurement in errors-in-variables models (Stefanski 2000). Second, the magnitude of the bias decreases monotonically in  $\beta_1^g$ , comporting with the intuition that the problem becomes easier as the gRNA mixture distribution becomes increasingly well-separated. Third, the Bayes-optimal decision boundary  $c_{\text{bayes}} \in \mathbb{R}$  (i.e., the most accurate decision boundary for classifying cells) is a critical value of the bias function. Finally, and most subtly, there is no universally applicable rule for selecting a threshold that yields minimal bias: when  $\beta_1^g$  is small, setting the threshold to an arbitrarily large number yields smaller bias than setting the threshold to the Bayes decision boundary; when  $\beta_1^g$  is large, the reverse is true. Appendix A contains detailed proposition statements and proofs of these results.

Next, we studied the variance of the thresholding estimator, considering a slightly simpler model for this purpose. Suppose the intercepts in (5) are fixed at 0 (i.e.,  $\beta_0^m = \beta_0^g = 0$ ). For notational simplicity we write  $\beta_m = \beta_1^m$  and  $\beta_g = \beta_1^g$ . The thresholding estimator  $\hat{\beta}_m$  is the no-intercept OLS solution  $\hat{\beta}_m = \left[\sum_{i=1}^n \hat{p}_i^2\right]^{-1} \left[\sum_{i=1}^n \hat{p}_i m_i\right]$ . The following proposition derives the scaled, asymptotic distribution of  $\hat{\beta}_m$ :

**Proposition 2.** The limiting distribution of  $\hat{\beta}_m$  is

$$\sqrt{n}(\hat{\beta}_m - l) \xrightarrow{d} N\left(0, \frac{\beta_m \omega \pi (\beta_m - 2l) + \mathbb{E}[\hat{p}_i](1 + l^2)}{(\mathbb{E}[\hat{p}_i])^2}\right),$$

where

$$l \equiv \beta_m \omega \pi / [\zeta(1-\pi) + \omega \pi]; \quad \mathbb{E}[\hat{p}_i] = \pi \omega + (1-\pi)\zeta; \quad \omega \equiv \Phi(\beta_g - c); \quad \zeta \equiv \Phi(-c).$$

This proposition yields an asymptotically exact bias-variance decomposition for  $\hat{\beta}_m$ : as the threshold tends to infinity, the bias decreases and the variance increases (Appendix A). Overall, our empirical and theoretical analyses indicate that thresholded regression poses a basic difficulty: selecting a good threshold is challenging, and even if we have selected a threshold that is (in some sense) optimal, the resulting estimator still may suffer from high bias or high variance. These considerations motivate our research question: does modeling the gRNA counts directly, thereby circumventing thresholding, facilitate estimation and inference in single-cell CRISPR screen analysis?

# 5 GLM-based errors-in-variables (GLM-EIV)

We introduce the general GLM-EIV model, which generalizes the negative binomial GLM-EIV model (1-3) to arbitrary exponential family response distributions and link functions, thereby providing much greater modeling flexibility. We derive efficient methods for estimation and inference in this model and develop a pipeline to deploy the model at-scale. Appendix C develops a parallel methodology in which the gRNA counts are zero-inflated.

#### 5.1 Model

General model This section is more technical than the previous ones; a first-time reader can skip to Section 5.2 without loss of information relevant to the high-level narrative. Let  $\tilde{x}_i = [1, p_i, z_i]^T \in \mathbb{R}^d$  be the vector of covariates (including an intercept term) for the *i*th cell. (We use the tilde as a reminder that the vector is partially unobserved.) Let  $\beta_m = [\beta_0^m, \beta_1^m, \gamma_m]^T \in \mathbb{R}^d$  and  $\beta_g = [\beta_0^g, \beta_1^g, \gamma_g]^T \in \mathbb{R}^d$  be the unknown coefficient vectors corresponding to the gene and gRNA expression models, respectively. Finally, let  $o_i^m$  and  $o_i^g$  be the (possibly zero) offset terms for the gene and gRNA models; in practice, we typically

set  $o_i^m$  and  $o_i^g$  to the log-transformed library sizes (i.e.,  $\log(d_i^m)$  and  $\log(d_i^g)$ , respectively).

We use a pair of GLMs to model the gene and gRNA expressions. Considering first the gene expression model, let the *i*th linear component  $l_i^m$  of the model be  $l_i^m \equiv \langle \tilde{x}_i, \beta_m \rangle + o_i^m$ . Next, let the mean  $\mu_i^m$  of the *i*th observation be  $r_m(\mu_i^m) \equiv l_i^m$ , where  $r_m : \mathbb{R} \to \mathbb{R}$  is a strictly increasing, differentiable link function. Let  $\psi_m : \mathbb{R} \to \mathbb{R}$  be the differentiable, cumulant-generating function of the selected exponential family distribution. We can express the canonical parameter  $\eta_i^m$  in terms of  $\psi_m$  and  $r_m$  by  $\eta_i^m = ([\psi_m']^{-1} \circ r_m^{-1})(l_i^m) \equiv$  $h_m(l_i^m)$ . Finally, let  $c_m:\mathbb{R}\to\mathbb{R}$  be the carrying density of the selected exponential family distribution. The density  $f_m$  of  $m_i$  conditional on the the canonical parameter  $\eta_i$  is  $f_m(m_i; \eta_i^m) = \exp\{m_i \eta_i^m - \psi_m(\eta_i^m) + c_m(m_i)\}$ . The function  $c_m$  appears as a constant in the log likelihood of  $m_i$ ; therefore, the only functions relevant to inference are  $\psi_m$  and  $r_m$ . Let the terms  $l_i^g, o_i^g, \mu_i^g, \eta_i^g, \psi_g, r_g, h_g$  and  $c_g$  be defined in an analogous way for the gRNA model, i.e.  $l_i^g \equiv \langle \tilde{x}_i, \beta_g \rangle + o_i^g, r_g(\mu_i^g) \equiv l_i^g$ , and  $\eta_i^g = \left( [\psi_g']^{-1} \circ r_g^{-1} \right) (l_i^g) \equiv h_g(l_i^g)$ . The density  $f_g$  of  $g_i$  given the canonical parameter is  $f_g(m_i; \eta_i^g) = \exp\{g_i \eta_i^g - \psi_g(\eta_i^g) + c_g(g_i)\}$ . Finally, the unobserved variable  $p_i$  is assumed to follow a Bernoulli distribution with mean  $\pi \in (0,1/2]$ . Its marginal density  $f_p$  is given by  $f_p(p_i) = \pi^{p_i}(1-\pi)^{1-p_i}$ . The unknown parameters in the model are  $\theta = [\beta_m, \beta_g, \pi]^T \in \mathbb{R}^{2d+1}$ .

Notation We briefly introduce notation that we will use throughout. For  $j \in \{0, 1\}$ , let  $\tilde{x}_i(j) \equiv [1, j, z_i]^T$  denote the value of  $\tilde{x}_i$  that results from setting  $p_i$  to j. Next, let  $l_i^m(j)$ ,  $\eta_i^m(j)$ , and  $\mu_i^m(j)$  be the values of  $l_i^m$ ,  $\eta_i^m$ , and  $\mu_i^m$ , respectively, that result from setting  $p_i$  to j, i.e.,  $l_i^m(j) \equiv \langle \tilde{x}_i(j), \beta_m \rangle + o_i^m$ ,  $\eta_i^m(j) \equiv h_m(l_i^m(j))$ , and  $\mu_i^m(j) \equiv r_m^{-1}(l_i^m(j))$ . Let the corresponding gRNA quantities  $l_i^g(j)$ ,  $\eta_i^g(j)$ , and  $\mu_i^g(j)$  be defined analogously. Next, let  $X \in \mathbb{R}^{n \times (d-1)}$  be the observed design matrix, and let  $\tilde{X} \in \mathbb{R}^{n \times d}$  be the augmented design

matrix that results from concatenating the column of (unobserved)  $p_i$ s to X, i.e.

$$X \equiv \begin{bmatrix} 1 & z_1 \\ \vdots & \vdots \\ 1 & z_n \end{bmatrix}; \quad \tilde{X} \equiv \begin{bmatrix} 1 & p_1 & z_1 \\ \vdots & \vdots & \vdots \\ 1 & p_n & z_n \end{bmatrix} = \begin{bmatrix} \tilde{x}_1^T \\ \vdots \\ \tilde{x}_n^T \end{bmatrix}.$$

Furthermore, for  $j \in \{0,1\}$ , let  $\tilde{X}(j) \in \mathbb{R}^{n \times d}$  be the matrix that results from setting  $p_i$  to j for all  $i \in \{1,\ldots,n\}$  in  $\tilde{X}$ , and let  $[\tilde{X}(0)^T, \tilde{X}(1)^T]^T$  denote the  $\mathbb{R}^{2n \times d}$  matrix that results from vertically concatenating  $\tilde{X}(0)$  and  $\tilde{X}(1)$ . Furthermore, define  $m := [m_1, \ldots, m_n]$ , and let  $g, p, o^m$ , and  $o^g$  be defined analogously. Finally, let  $[m, m]^T \in \mathbb{R}^{2n}$  be the vector that results from concatenating m to itself, i.e.  $[m, m]^T \equiv [m_1, \ldots, m_n, m_1, \ldots, m_n]$ , and let  $[g, g]^T$ ,  $[o^g, o^g]^T$ , and  $[o^m, o^m]^T$  be defined similarly.

Log likelihood and model properties We derive the log-likelihood of the GLM-EIV model. We conduct estimation and inference conditional on the library sizes and technical factors  $l_i^m, l_i^g$ , and  $z_i$ ; therefore, we treat these quantities as fixed constants. We assume that the gene expression  $m_i$  and gRNA expression  $g_i$  are conditionally independent given the perturbation  $p_i$ . The joint density f of  $(m_i, g_i, p_i)$  given  $\theta$  is

$$f(m_i, g_i, p_i; \theta) = f_p(p_i) f_m(m_i | p_i) f_g(g_i | p_i) = \pi^{p_i} (1 - \pi)^{1 - p_i} f_m(m_i; \eta_i^m) f_g(g_i; \eta^g).$$
 (7)

Integrating over the unobserved variable  $p_i$ , we can write the density f of  $(m_i, g_i)$  as

$$f(m_i, g_i; \theta) = (1 - \pi) f_m(m_i; \eta_i^m(0)) f_g(g_i; \eta_i^g(0)) + \pi f_m(m_i; \eta_i^m(1)) f_g(g_i; \eta_i^g(1)).$$
 (8)

Therefore, the log-likelihood is

$$\mathcal{L}(\theta; m, g) = \sum_{i=1}^{n} \log \left[ (1 - \pi) f_m(m_i; \eta_i^m(0)) f_g(g_i; \eta_i^g(0)) + \pi f_m(m_i; \eta_i^m(1)) f_g(g_i; \eta_i^g(1)) \right]. \tag{9}$$

We see from (8) that the GLM-EIV model is equivalent to a two-component mixture of *products* of GLM densities. Additionally, the GLM-EIV model is a generalization the simple errors-in-variables model (when the predictor is binary); the latter is defined as follows:

$$y_i = \beta_0 + \beta_1 x_i^* + \epsilon_i; \quad x_i = x_i^* + \tau_i,$$
 (10)

where,  $x_i^* \sim \text{Bern}(\pi)$ ,  $\epsilon_i$ ,  $\tau_i \sim N(0, 1)$ , and  $\epsilon_i$ ,  $\tau_i$ , and  $x_i^*$  are independent. GLM-EIV extends (10) in at least three directions: first, GLM-EIV allows  $y_i$  and  $x_i$  to follow exponential family (i.e., not just Gaussian) distributions; second, GLM-EIV allows  $y_i$  and  $x_i$  to be related to  $x_i^*$  through arbitrary (i.e., not just linear) link functions; and finally, GLM-EIV allows confounders  $z_i$  to impact both  $x_i$  and  $y_i$ . Therefore,  $x_i$  and  $y_i$  can be conditionally dependent given  $x_i^*$ , enabling GLM-EIV to capture more complex dependence relationships between  $x_i$  and  $y_i$  than is possible in (10) or other standard measurement error models.

#### 5.2 Estimation and inference

We derive an EM algorithm (Algorithm 1) to estimate the parameters of the GLM-EIV model. The E step entails computing the membership probability (i.e., the probability of perturbation) in each cell. The membership probability  $T_i(1)$  of cell  $i \in \{1, ..., n\}$  given the current parameter estimates  $(\beta_m^{(t)}, \beta_g^{(t)}, \pi^{(t)})$  and observed data  $(m_i, g_i)$  is

$$T_i(1) = \mathbb{P}(p_i = 1 | M_i = m_i, G_i = g_i, \beta_m^{(t)}, \beta_g^{(t)}, \pi^{(t)}).$$

We can calculate this quantity by applying (i) Bayes rule, (ii) the conditional independence property of  $M_i$  and  $G_i$ , (iii) the density of  $M_i$  and  $G_i$ , and (iv) a log-sum-exp-type trick to ensure numerical stability. Next, we produce updated estimates  $\pi^{(t+1)}$ ,  $\beta_g^{(t+1)}$ , and  $\beta_m^{(t+1)}$  of the parameters by maximizing the M step objective function. It turns out that maximizing this objective function is equivalent to setting  $\pi^{(t+1)}$  to the mean of the current membership probabilities and setting  $\beta_g^{(t+1)}$  and  $\beta_m^{(t+1)}$  to the fitted coefficients of a GLM weighted by the current membership probabilities (Algorithm 1). We iterate through the E and M steps until the log likelihood (9) converges (Appendix B). Our EM algorithm is reminiscent of (but distinct from) that of Ibrahim (1990), who also applied weighted GLM solvers to carry out an M step of an EM algorithm.

After fitting the model, we perform inference on the estimated parameters. The easiest approach, given the complexity of the log likelihood, would be to run a bootstrap. This strategy, however, is prohibitively slow, as the data are large and the EM algorithm is iterative. Therefore, we derive an analytic formula for the asymptotic observed information matrix using Louis's Theorem (Louis 1982; Appendix B). Leveraging this analytic formula, we can calculate standard errors quickly, enabling us to perform inference in practice on real, large-scale data.

## 5.3 Statistical accelerations and computational infrastructure

A downside of the EM algorithm (Algorithm 1) is that it requires fitting many GLMs. Assuming that we run the algorithm 15 times using randomly-generated pilot estimates (to improve chances of convergence to the global maximum), and assuming that the algorithm iterates through E and M steps about 10 times per run, we must fit approximately 300 GLMs. (These numbers are based on exploratory applications of the method to real and

#### **Algorithm 1** EM algorithm for GLM-EIV model.

```
Input: Pilot estimates \beta_m^{\text{curr}}, \beta_g^{\text{curr}}, and \pi^{\text{curr}}; data m, g, o^m, o^g, and X; gene expression
   distribution f_m and link function r_m; gRNA expression distribution f_q and link function
   while Not converged do
                                                                                                                                    ⊳ E step
         for i \in \{1, ..., n\} do
              T_i(1) \leftarrow \mathbb{P}\left(p_i = 1 | M_i = m_i, G_i = g_i, \beta_m^{\text{curr}}, \beta_q^{\text{curr}}, \pi^{\text{curr}}\right)
              T_i(0) \leftarrow 1 - T_i(1)
         end for
         \pi^{\mathrm{curr}} \leftarrow (1/n) \sum_{i=1}^n T_i(1)
                                                                                                                                   ⊳ M step
         w \leftarrow [T_1(0), T_2(0), \dots, T_n(0), T_1(1), T_2(1), \dots, T_n(1)]^T
         for k \in \{g, m\} do
              Fit a GLM GLM_k with responses [k, k]^T, offsets [o^k, o^k]^T, weights w, design matrix
              [\tilde{X}(0)^T, \tilde{X}(1)^T]^T, distribution f_k, and link function r_k.
              Set \beta_k^{\text{curr}} to the estimated coefficients of GLM_k.
         Compute log likelihood using \beta_m^{\text{curr}}, \beta_q^{\text{curr}}, and \pi^{\text{curr}}.
   \hat{\beta}_m \leftarrow \beta_m^{\text{curr}}; \, \hat{\beta}_g \leftarrow \beta_g^{\text{curr}}; \, \hat{\pi} \leftarrow \pi^{\text{curr}}.
   return (\hat{\beta}_m, \hat{\beta}_q, \hat{\pi})
```

simulated data.) We instead devised a strategy to produce a highly accurate pilot estimate of the true parameters, enabling us to run the algorithm once and converge upon the MLE within a few iterations. The strategy involves layering several statistical "tricks" (that could be of independent utility for accelerating other single-cell methods) on top of one another; details are deferred to Appendix D. Overall, the statistical accelerations reduce the number of GLMs that we must fit to < 10 in most cases.

Next, we developed a computational infrastructure to apply GLM-EIV to large-scale, single-cell CRISPR screen data. The infrastructure leverages Nextflow, a programming language that facilitates building data-intensive pipelines (DI Tommaso et al. 2017), and ondisc, an R/C++ package that we developed (in a separate project) to facilitate large-scale computing on single-cell data (preprint forthcoming). Nextflow and ondisc together enable the construction of highly portable single-cell pipelines: one can analyze data out-of-memory on a laptop or in a distributed fashion across tens or hundreds of nodes on

a cloud (e.g., Microsoft Azure, Google Cloud) or high-performance cluster. Leveraging these technologies, we built a Docker-containerized pipeline for deploying GLM-EIV at-scale. The pipeline aggressively recycles computation when possible, saving a considerable amount of compute; see Appendix D.3 for details. Overall, the statistical accelerations and computational infrastructure make the deployment of GLM-EIV to large-scale single-cell CRISPR screen quite feasible.

# 6 Simulation study

We conducted a simulation study to compare the empirical performance of GLM-EIV to that of the thresholding method. We generated data on n=150,000 cells from the GLM-EIV model using realistic parameter values, setting the the target of inference  $\beta_1^m$  to  $\log(0.25)$  and the probability of perturbation  $\pi$  to 0.02.  $\beta_1^m = \log(0.25)$  represents a decrease in gene expression by a factor of 4, which is a fairly large effect size on the order of what we might observe for a positive control pair. We included "sequencing batch" (modeled as a Bernoulli-distributed variable) as a covariate and sequencing depth (modeled as a Poisson-distributed variable) as an offset. We varied the log-fold change in gRNA expression,  $\beta_1^g$ , over a grid on the interval  $[\log(1), \log(4)]$ ;  $\beta_1^g$  controls problem difficulty, with higher values corresponding to easier problem settings. Finally, we generated the gene expression and gRNA count data from two response distributions: Poisson and negative binomial (size parameter fixed at  $\theta=20$  for the latter). For each parameter setting (defined by a  $\beta_1^g$ -distribution pair), we synthesized  $n_{\text{sim}}=500$  i.i.d. datasets. Section E presents additional simulation results on Gaussian response distributions.

We applied three methods to the simulated data: "vanilla" GLM-EIV, accelerated GLM-EIV, and thresholded regression. We used the Bayes-optimal decision boundary for

classification as the threshold for the thresholding method. We ran all methods on the negative binomial data twice: once treating the size parameter  $\theta$  as a known constant and once treating  $\theta$  as unknown. In the latter case we used the glm.nb function from the MASS package to estimate  $\theta$  before applying the methods (Ripley et al. 2013). We display the results of the simulation study in Figure 3. Columns correspond to distributions (i.e., Poisson, NB with known  $\theta$ , and NB with unknown  $\theta$ ), and rows correspond to performance metrics (i.e., bias, mean squared error, CI coverage rate (nominal rate 95%), CI width, and method execution time). The problem difficulty parameter  $\beta_1^g$  is plotted on the horizontal axis, and the methods are depicted in different colors (GLM-EIV masked by accelerated GLM-EIV in several panels).

First, we observed that GLM-EIV dominated thresholded regression on all statistical metrics: GLM-EIV exhibited lower bias (row 1) and mean squared error (row 2) than thresholded regression; additionally, GLM-EIV had superior confidence interval coverage (row 3) despite having produced generally narrower confidence intervals (row 4). Intuitively, GLM-EIV outperformed the thresholding method because (i) GLM-EIV leveraged information from both modalities (rather than the gRNA modality alone) to assign perturbation identities to cells, and (ii) GLM-EIV produced soft rather than hard assignments, capturing the inherent uncertainty in whether a perturbation occurred. We additionally found that accelerated GLM-EIV performed as well as vanilla GLM-EIV on all statistical metrics (rows 1-4) despite having substantially lower computational cost (bottom row). In fact, the execution time of accelerated GLM-EIV was almost within an order of magnitude of that of the thresholding method (bottom row).

Interestingly, thresholded regression exhibited better confidence interval coverage under estimated  $\theta$  than under known  $\theta$  (row 3). Estimating  $\theta$  leads to slight inflation bias (i.e.,

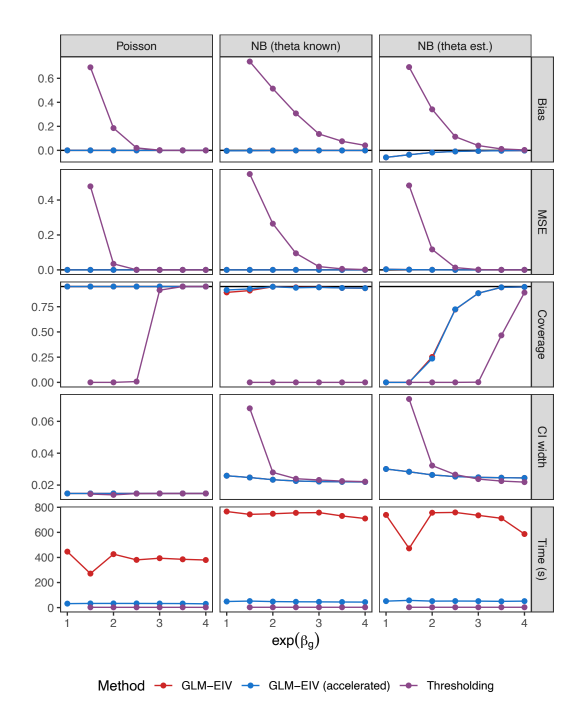


Figure 3: **Simulation study**. Columns correspond to distributions (Poisson, NB with known  $\theta$ , NB with estimated  $\theta$ ), and rows correspond to metrics (bias, MSE, coverage, CI width, and time). Methods are shown in different colors; GLM-EIV (red) is masked by accelerated GLM-EIV (blue) in several panels. GLM-EIV demonstrated superior statistical performance to the thresholding method on all metrics (rows 1-4). Accelerated GLM-EIV had substantially lower computational cost than "vanilla" GLM-EIV (bottom row), despite demonstrating identical statistical performance (rows 1-4).

overestimating the true effect size), whereas, as we showed previously, thresholding leads to attenuation bias (i.e., underestimating the true effect size). These phenomena partially canceled, yielding less biased estimates. GLM-EIV exhibited worse performance under unknown  $\theta$  than known  $\theta$ , likely due to poor  $\theta$  estimation. We note that GLM-EIV and the thresholding method in principle are compatible with  $any \theta$  estimation procedure, including those based on more sophisticated techniques, such as regularization (Hafemeister and Satija 2019). We defer rigorous investigation of the impact of different  $\theta$  estimation strategies on these methods to future work.

## 7 Data analysis

Leveraging our computational infrastructure, we applied GLM-EIV and the thresholding method to analyze the entire Gasperini and Xie datasets. We report only the most important aspects of the analysis and results in the main text; full details are available in Appendix F. We set the threshold in the thresholding method to the approximate Bayes-optimal decision boundary, as our theoretical analyses and simulation studies indicated that the Bayes-optimal decision boundary is a good choice for the threshold when the gRNA count distribution is well-separated. Operating under the assumption that the effect of the perturbation on gRNA expression is similar across pairs, we leveraged the fitted GLM-EIV models to approximate the Bayes boundary in the following way: we (i) sampled several hundred gene-perturbation pairs, (ii) extracted the fitted values  $\hat{\beta}_g$  and  $\hat{\pi}$  from the GLM-EIV models fitted to these pairs, (iii) computed the median  $\overline{\hat{\beta}_g}$  and  $\overline{\hat{\pi}}$  across the  $\hat{\beta}_g$ s and  $\hat{\pi}$ s, and (iv) used  $\overline{\hat{\beta}_g}$  and  $\overline{\hat{\pi}}$  to estimate a dataset-wide Bayes-optimal decision boundary. We repeated this procedure on both datasets, yielding a threshold of 3 for Gasperini and 7 for Xie. These thresholds were close to the thresholds used in the original publications,

which were selected in a more heuristic way.

We compared GLM-EIV to thresholded regression on the real data, focusing specifically on the negative control pairs (i.e., gene-perturbation pairs that, by design, are expected to exhibit a fold change of 1, or no association). We found that GLM-EIV and the thresholding method produced similar results (Figure 4a-b): estimates, CI coverage rates, and CI widths were concordant. CI coverage rates, which ranged from 87.7%-91.2%, were slightly below the nominal rate of 95%, likely due to mild model misspecification. The estimated effect of the perturbation on gene expression  $\exp(\hat{\beta}_1^g)$  was unexpectedly large: the 95% CI for this parameter was [4306, 5186] and [300, 316] on the Gasperini and Xie data, respectively. We reasoned that the datasets lay in an "easy" region of the parameter space, making thresholding a tenable strategy (provided the threshold is selected well). However, this was not obvious a priori and may not be the case for other datasets. We note that GLM-EIV produced outlier estimates (defined as estimated fold change < 0.75 or > 1.25) on a small (<2.5%) on Gasperini, <0.05% on Xie) number of pairs consisting of a handful of genes. likely due to non-global EM convergence. These outliers are not plotted in Figures 4a-b but were used to compute the CI coverage reported in the inset tables.

To evaluate performance of GLM-EIV versus thresholding in more challenging settings, we artificially increased the difficulty of the perturbation assignment problem by generating partially-synthetic datasets. First, for a given pair, we sampled gRNA counts directly from the fitted GLM-EIV model. Next, to simulate elevated background contamination, we sampled gRNA counts from a slightly modified version of the fitted model in which we increased the mean gRNA expression of *unperturbed* cells while holding constant the mean gRNA expression of *perturbed* cells. We defined a parameter called "excess background contamination" (normed to take values in [0,1]) to quantify the relative distance between

the unperturbed and perturbed gRNA count distributions. We held fixed the real-data gene expressions, library sizes, covariates, and fitted perturbation probabilities in all settings.

We generated partially-synthetic data in the above manner for each of the 322 positive control pairs in the Gasperini dataset, varying excess background contamination over the interval [0, 0.4]. We then applied GLM-EIV and the thresholding method to analyze the data. We present results on two example pairs (the pair containing gene *LRIF1* and the pair containing gene *NDUFA2*) in Figures 4c-d. We observed that the estimate produced by the methods on the raw data (depicted as a horizontal black line) coincided almost exactly with the estimate produced by the methods on the partially-synthetic data generated by setting excess background contamination to zero (This result replicated across nearly all pairs; average relative difference 0.003.) We additionally observed that as excess background contamination increased, the performance of thresholded regression degraded considerably while that of GLM-EIV remained stable.

We generalized the above analysis to the entire set of positive control pairs. First, for each pair we computed the "relative estimate change" (REC) as a function of excess background contamination, defined as the relative difference between the estimate at a given level of excess contamination and zero excess contamination (Figure 4d). Next, we computed the median REC across all positive control pairs (Figure 4e; upper and lower bands indicate the pointwise interquartile range of the REC). As excess background contamination increased, thresholded regression exhibited severe attenuation bias (as reflected by large median REC values); GLM-EIV, by contrast, remained mostly stable. Finally, letting  $\hat{\beta}_1^m$  denote the estimate obtained on the raw data, we computed the CI coverage of  $\hat{\beta}_1^m$  as a function of excess contamination. Under the assumption that  $\hat{\beta}_1^m$  is close to the true parameter  $\beta_1^m$ , the CI coverage of the former is similar to that of the latter. We computed the

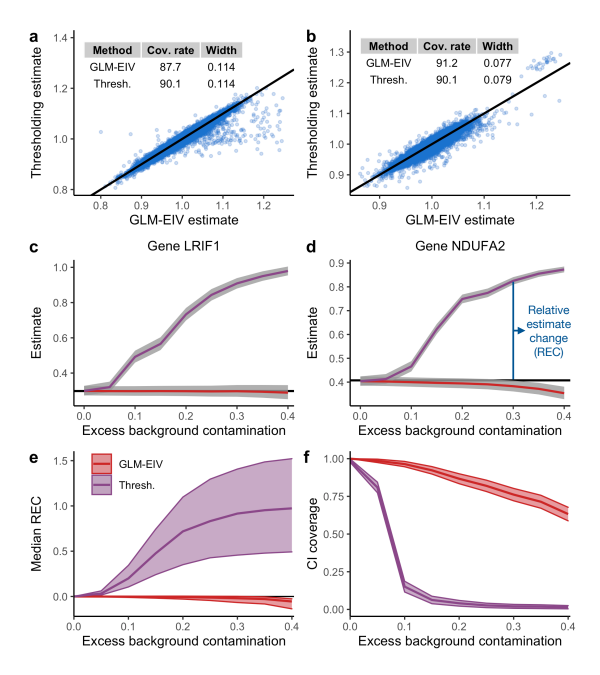


Figure 4: **Data analysis**. **a-b**, Estimates for fold change produced by GLM-EIV and thresholded regression on Gasperini (**a**) and Xie (**b**) negative control pairs. **c-d**, Estimates produced by GLM-EIV and thresholded regression on two positive control pairs – *LRIF1* (**a**) and *NDUFA2* (**b**) – plotted as a function of excess background contamination. Grey bands, 95% CIs for the target of inference outputted by the methods. **e-f**, Median relative estimate change (REC; **e**) and confidence interval coverage rate (**f**) across *all* 322 positive control pairs, plotted as a function of excess background contamination. Panels (**c-f**) together illustrate that GLM-EIV demonstrated greater stability than thresholded regression as background contamination increased.

CI coverage of  $\hat{\beta}_1^m$  by calculating each individual pair's coverage of  $\hat{\beta}_1^m$  (across the Monte Carlo replicates) and then averaging this quantity across all pairs. GLM-EIV exhibited significantly higher CI coverage than thresholded regression as the data became increasingly contaminated (Figure 4f; bands indicate 95% pointwise CIs). Coverage rates were slightly above the nominal level of 95% in some settings because we covered an *estimate* of  $\beta_1^m$  rather than  $\beta_1^m$  itself, leading to mild "overfitting." Nonetheless, this experiment was meaningful to assess the stability of both methods to elevated background contamination.

## 8 Discussion

In this work we introduced GLM-EIV ("GLM-based errors in variables"), a new model and associated method for single-cell CRISPR screen analysis. GLM-EIV extends the classical errors-in-variables model to responses and noisy predictors that are exponential familydistributed and potentially impacted by the same set of confounding variables. These extensions enable GLM-EIV to resolve novel analysis challenges posed by single-cell CRISPR screens. We demonstrated through simulation studies, real data analyses, and theory that GLM-EIV outperforms thresholded regression by a considerable margin in high background contamination settings. GLM-EIV intuitively achieves this performance gain by leveraging information from both modalities (rather than the gRNA modality alone) to assign perturbation identities to cells. On the other hand, in low background contamination settings, GLM-EIV and thresholded regression work best in conjunction, with GLM-EIV providing a statistically principled and empirically effective procedure for selecting the threshold. GLM-EIV thereby neutralizes a tuning parameter that, until this point, has been selected using heuristic procedures, with little confidence that the choice is near optimal. Figure 5 summarizes how we anticipate GLM-EIV being used in practice.

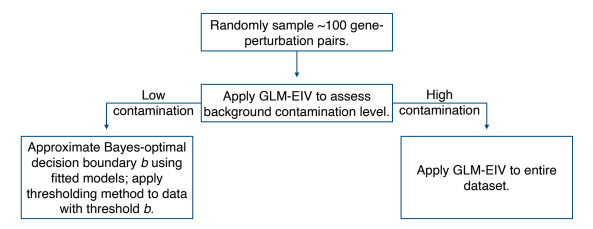


Figure 5: Use of GLM-EIV in practice. The decision tree above illustrates how we anticipate GLM-EIV could be used in practice. First, apply GLM-EIV to a set of randomly-sampled gene-perturbation pairs to assess background contamination level (positive control pairs work best for this purpose). If GLM-EIV indicates that background contamination is high (e.g.,  $\exp(\beta_1^g) \lesssim 10$ ), apply GLM-EIV to analyze the entire dataset; otherwise, approximate the Bayes-optimal decision boundary using the fitted GLM-EIV models. Next, apply a thresholding method (e.g., SCEPTRE or thresholded negative binomial regression) to analyze the data, setting the threshold to the estimated Bayes-optimal decision boundary.

To our knowledge this is the first single-cell CRISPR screen paper oriented toward a statistical audience. We hope that this work helps to introduce the broader statistics community to an emerging class of functional genomics assays that likely will exert a major impact on biological research in the coming years (Przybyla and Gilbert 2021). Additionally, this is the first work to leverage the ondisc-Nextflow-HPC/cloud technology stack, a tightly-integrated, user-friendly, and powerful set of tools for large-scale single-cell analysis. We expect this technology stack to be of interest to other single-cell researchers.

We anticipate that GLM-EIV could be applied to other types of single-cell CRISPR screen and multimodal single-cell data. For example, GLM-EIV might be extended (with some effort) to "low-multiplicity of infection" screens (Schraivogel et al. 2020) in which each cell receives one or two perturbations rather than dozens (as is the case in "high multiplicity screens," studied in this work). We also could apply GLM-EIV to analyze multimodal single-cell chromatin accessibility assays. A question of interest in such experiments is

whether chromatin state (i.e., closed or open) is associated with the expression of a gene or abundance of a protein (Mimitou et al. 2021). We do not directly observe the chromatin state of a cell; instead, we observe tagged DNA fragments that serve as count-based proxies for whether a given region of chromatin is open or closed. GLM-EIV might be applied in such experiments to aid in the selection of thresholds or to analyze whole datasets.

Several authors working on statistical methods for single-cell data recently have extended models that (implicitly or explicitly) assume Gaussianity and homoscedasticity to a broader class of exponential family distributions. For example, Lin et al. (2021) and Townes et al. (2019) (separately) developed eSVD and GLM-PCA, generalizations of SVD and PCA, respectively, to exponential family response distributions. Unlike their vanilla counterparts, eSVD and GLM-PCA can model gene expression counts directly, improving performance on dimension reduction tasks. We see our work (in part) as a continuation of this broad effort to "port" common statistical methods and models to single-cell count data. Our focus, however, is on regression rather than dimension reduction: we extend the classical errors-in-variables model in several key directions (see above), enabling its direct and natural application to multimodal single-cell data.

The closest parallels to GLM-EIV in the statistical methodology literature are Grün and Leisch (2008) and Ibrahim (1990). Grün and Leisch derived a method for estimation and inference in a k-component mixture of GLMs. While we prefer to view GLM-EIV as a generalized errors-in-variables method, the GLM-EIV model is equivalent to a two-component mixture of products of GLM densities. Ibrahim proposed a procedure for fitting GLMs in the presence of missing-at-random covariates. Our method, by contrast, involves fitting two conditionally independent GLMs in the presence of a totally latent covariate. Thus, while Ibrahim and Grün & Leisch are helpful references, our estimation and infer-

ence tasks are more complex than theirs. Next, Aigner (1973) and Savoca (2000) proposed measurement error models that consist of unobserved binary rather than continuous predictors; the latter are more commonly used in measurement error models. GLM-EIV likewise consists of a latent binary predictor, but unlike Aigner and Savoca, GLM-EIV handles a much broader class of exponential family-generated data. Finally, GLM-EIV accounts for a common source of measurement error between the predictor and response, a property not shared by classical measurement error models (Carroll et al. 2006).

GLM-EIV might be applied to areas beyond genomics, such as psychology. Many psychological constructs (e.g., presence or absence of a social media addiction) are latent and can be assessed only through an imperfect proxy (e.g., the number of times one has checked social media). Researchers might use GLM-EIV to regress an outcome variable (e.g., self-reported well-being) onto the latent construct via the imperfect proxy, potentially resolving challenges related to attenuation bias and threshold selection. Applications to psychology and other areas are a topic of future investigation.

## 9 Acknowledgments and resource availability

[This section is muted to preserve anonymity. The authors have prepared the data, results, and detailed instructions to reproduce the analyses reported in the paper.]

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