
Beyond Forward Prediction: Reverse Perturbation Retrieval as a Bridge from Transcriptomic Phenotypes to Drug Targets

Anrui Wang

2023533015

wangar2023@shanghaitech.edu.cn

Jiawen Dai

2023533132

daijw2023@shanghaitech.edu.cn

Yiting Qi

2023533043

qiyt2023@shanghaitech.edu.cn

Abstract

Phenotypic drug discovery faces a fundamental "inverse problem": while therapeutic effects are observable, identifying the underlying molecular drivers remains a significant bottleneck. Although "Virtual Cell" models aim to bridge this gap through transcriptomic simulation, current benchmarks reveal that optimizing forward prediction accuracy does not inherently yield actionable mechanistic insights. To address this, we reframe the modeling objective from forward simulation to *in silico* reverse perturbation retrieval, transforming the virtual cell into a prescriptive engine that ranks potential causal targets based on observed disease phenotypes. Our evaluation uncovers a critical "geometry versus semantics" trade-off: while geometric proximity suffices for retrieving targets with strong phenotypic signals, foundation models uniquely capture the semantic context required to identify complex, low-signal targets that lie beyond simple similarity. This establishes a new paradigm where the utility of virtual cell models is defined not by their ability to simulate known outcomes, but by their capacity to close the causal loop and generate testable hypotheses for target deconvolution.

1 Introduction

1.1 Phenotypic screening creates an inverse problem

Phenotypic drug discovery has historically been a cornerstone of therapeutic development, uniquely capable of identifying compounds that rescue disease states in a physiologically relevant context without requiring prior knowledge of a specific molecular target. However, this "target-agnostic" success comes with a fundamental **inverse problem**: while the therapeutic effect—the phenotype—is observed, the underlying molecular mechanism of action (MoA) or target often remains unknown, necessitating a complex and laborious target deconvolution process. Unlike target-based approaches that proceed from cause to effect, phenotypic screening presents us with the effect, requiring us to solve for the cause.

High-throughput transcriptomic profiling has emerged as a critical bridge in this gap, offering a high-dimensional, information-rich "fingerprint" of cellular states that reflects the downstream consequences of upstream perturbations. By defining the biological phenotype as a computable vector, it becomes possible to systematically map the causal relationship between a perturbation (whether genetic or chemical) and its resulting cellular state. This capability transforms the biological

question into a data-driven inference task: if we can quantify the phenotypic shift, can we identify the perturbation that caused it?

Recent advances in artificial intelligence have begun to address this by proposing the construction of “Virtual Cells”—high-fidelity simulators learned directly from large-scale biological data. As highlighted by Roohani et al. [1], these systems are expected to learn the fundamental relationship between cell state and function, with the primary intent of predicting the consequences of perturbations—such as gene knockdowns or drug treatments—across diverse cell contexts. While much of the current effort focuses on this **forward prediction**, the ultimate utility of AI Virtual Cells (AIVCs) extends beyond mere simulation to “closing the loop” for discovery. Bunne et al. [2] articulate a vision where AIVCs serve as engines for **in silico experimentation**, capable of simulating the effects of varying interventions to propose potential causal factors behind observed phenotypes. Although computation alone may not fully resolve all causal links, AIVCs offer the critical potential to drastically “**reduce the space of possible hypotheses**”, thereby accelerating the identification of underlying mechanisms and new drug targets. This positions the Virtual Cell not just as a simulator, but as a computational partner for target deconvolution in phenotypic drug discovery.

1.2 The community’s asymmetry: optimizing forward metrics vs answering actionable questions

The field currently faces a critical asymmetry: while immense effort is poured into optimizing forward prediction metrics, empirical gains remain marginal. A systematic benchmark across diverse datasets reveals that complex deep learning models do not consistently outperform simpler alternatives, with performance often heavily confounded by perturbation effect sizes rather than architectural superiority [3]. More strikingly, recent rigorous evaluations serve as a wake-up call, demonstrating that state-of-the-art models for perturbation prediction do not yet consistently outperform simple linear baselines or even trivial mean predictions [4]. This suggests that the singular pursuit of minimizing reconstruction error (MSE) may not inherently lead to the actionable causal discovery required for drug development.

1.3 Reframing: invert the causal chain to make models actionable

To bridge the gap between model performance and biological discovery, we propose a fundamental reframing of the Virtual Cell’s utility: shifting the focus from **forward simulation** (predicting the expression profile of a known perturbation) to **reverse retrieval** (inferring the causal perturbation of an observed profile). This is not an arbitrary task invention but a rigorous operationalization of the “*in silico reverse perturbation prediction*” paradigm already emerging in foundational models. For instance, **scGPT** [5] explicitly formalizes this as a top-K retrieval task, demonstrating that foundation models can learn to rank potential genetic drivers based on their alignment with a target cell state. By adopting this retrieval-based formulation, we align the model’s output directly with the decision-making logic of drug discovery: generating a ranked list of probable targets for experimental validation.

This inversion transforms the Virtual Cell from a descriptive simulator into a prescriptive engine for **therapeutic target identification**. This utility has been notably demonstrated by **Geneformer** [6], which leveraged transfer learning on large-scale data to move beyond mere expression forecasting, successfully prioritizing candidate therapeutic targets and key network regulators for cardiomyopathy. However, reliable reverse inference requires models to possess true mechanistic reasoning rather than simple pattern matching, particularly when dealing with the combinatorial complexity of biological systems. As evidenced by **GEARS** [7], incorporating biological inductive biases—such as gene regulatory graphs—is essential for generalizing to unseen perturbations and combinatorial sets that lack direct experimental training data. Similarly, achieving robust target discovery requires models (like the transformer-based STATE [8]) to model distributions across **sets of cells**, enabling mechanistic generalization to novel cellular contexts where training data may be sparse or absent. By closing this causal loop, we move from optimizing abstract error metrics to answering the actionable question: *what perturbation caused this phenotype?*

1.4 Contributions

In this work, we close the loop between forward simulation and reverse inference, presenting a unified framework that repurposes the Virtual Cell from a predictive simulator into a prescriptive engine for target discovery. We ground this framework by establishing a strict evaluation protocol for forward perturbation prediction using the **Virtual Cell Challenge** (VCC) dataset [1]. By enforcing **perturbation-level splits** and prioritizing biologically grounded metrics (DES, PDS) over simple reconstruction error, we ensure that state-of-the-art models (including scGPT and STATE) capture true causal perturbation-response maps rather than merely fitting statistical noise. Transitioning from simulation to inference, we formally operationalize **in silico reverse perturbation prediction** as a retrieval and ranking problem. This paradigm shift—treating transcriptomic phenotypes as queries to identify causal agents—demonstrates that deep perturbation models can effectively bridge the gap between phenotypic screening readouts and genetic ground truths. Crucially, our investigation uncovers a fundamental “**Geometry vs. Semantics**” trade-off in phenotypic retrieval. While heuristic baselines like PCA-KNN excel at identifying strong, “geometrically reachable” phenotypes, pre-trained foundation models such as scGPT demonstrate superior robustness in retrieving the “long tail” of subtle signals, suggesting they uniquely capture the semantic context required for deciphering complex biological mechanisms.

Author Contributions

- **Anrui Wang:** Implemented scGPT experiments to fulfill the Virtual Cell Challenge (VCC) requirements, evaluated model performance, adapted scGPT to leverage the improved VCC evaluation metrics, and created scGPT-based models specifically for target gene identification, enabling a more biologically meaningful assessment of predictions.
- **Jiawen Dai:** Reimplemented a more biologically informed set of evaluation metrics for the Virtual Cell Challenge (VCC), conducted baseline experiments for VCC models, and extracted relevant data from the Tahoe-100M dataset.
- **Yiting Qi:** Implemented STATE experiments to fulfill the VCC requirements and evaluate performance, conducted studies on the Tahoe-100M dataset, and developed baseline models for target gene prediction on the Tahoe-100M dataset.

2 Forward validation as a credibility scaffold

2.1 Forward prediction on VCC under perturbation-level splits

We anchored our investigation by benchmarking state-of-the-art models on the **Virtual Cell Challenge** (VCC) dataset, establishing a rigorous baseline for virtual cell capabilities. Diverging from standard evaluations that rely on random cell splits—a practice that risks reducing the task to mere cellular interpolation—we enforced a strict **perturbation-level split**. This rigorous setting tasked models with predicting the transcriptional effects of genetic perturbations entirely unseen during training, effectively isolating true mechanistic generalization from simple pattern memorization.

We evaluated two representative model architectures: **STATE**, which leverages cell-set modeling to capture distributional shifts across cellular contexts, and **scGPT**, a foundational model that utilizes large-scale pre-training to learn generalizable gene-gene interactions. By systematically comparing these models alongside linear baselines, we defined the current performance boundary for forward prediction (Figure 1). This “forward validation” step serves as a necessary credibility scaffold; it confirms that the underlying embeddings capture sufficient biological signal to support the more complex inference tasks required for target discovery, without assuming that forward simulation is the endpoint of the modeling effort. All implementation details, including data splits and hyperparameter configurations, are detailed in Supplementary Note 1.

2.2 Why we evaluate biology, not just numbers

In evaluating these models, we depart from a reliance on raw reconstruction errors, which can be misleading in transcriptomic data. Conventional metrics like Mean Absolute Error (MAE) are frequently dominated by the high baseline expression of invariant genes, potentially awarding high

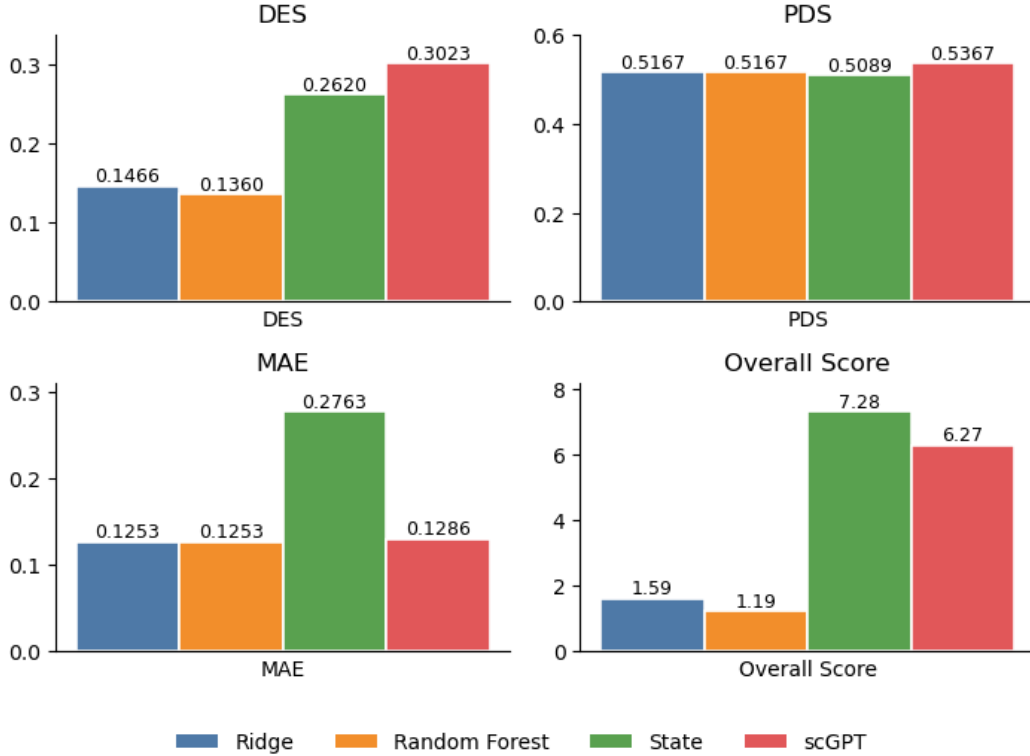


Figure 1: Evaluation Metrics and Scoring Framework. The Virtual Cell Challenge utilizes a composite scoring system derived from three complementary metrics to assess forward prediction performance. (A) Differential Expression Score (DES): Measures the model’s fidelity in recovering biologically interpretable signals by calculating the overlap (precision and recall) between predicted and ground-truth differentially expressed genes (DEGs). (B) Perturbation Discrimination Score (PDS): A ranking-based metric that evaluates whether a predicted pseudo-bulk profile is more similar to its corresponding ground truth than to other perturbations. To mitigate biases toward vectors with small norms, PDS utilizes cosine similarity to emphasize directional alignment over magnitude. (C) Mean Absolute Error (Top-2k MAE): Captures the global reconstruction accuracy by computing the element-wise error across the top 2,000 highly variable genes. (D) Overall Score: A weighted aggregation of the individual metrics used to rank model performance, enforcing a balance between global fit (MAE), directional correctness (PDS), and biological specificity (DES).

scores to trivial "mean-prediction" baselines while failing to distinguish true biological replicates from unrelated perturbations. Instead, as illustrated in Figure 1, we prioritize metrics that assess the **perturbation delta** (δ)—the specific causal shift in gene expression induced by the intervention relative to controls. We focus on two biologically interpretable metrics to serve as our primary credibility scaffold. First, the **Differential Expression Score (DES)** evaluates the model’s ability to recover the specific set of differentially expressed genes (DEGs). By measuring the overlap (precision and recall) between predicted and ground-truth DEGs, DES verifies whether the model faithfully recapitulates the biologically interpretable effects of a perturbation, rather than merely fitting global statistical distributions. Second, the **Perturbation Discrimination Score (PDS)**, adapted here to utilize cosine similarity, assesses the directional fidelity of the predicted shift. This ensures that the model correctly ranks the true perturbation as the most similar candidate among all possibilities. Unlike L1-based rankings, this approach emphasizes the alignment of up- or down-regulation vectors rather than signal magnitude, mitigating the penalty on "attenuated" predictions often seen in single-cell modeling. This evaluation philosophy aligns with the "delta-based" validation advocated in recent foundational studies, ensuring that high performance reflects the capture of causal regulatory signals rather than the preservation of static cell-type identity.

Metric Analysis and Failure Modes The selection of these metrics addresses specific failure modes in predictive modeling (Figure 1D). While MAE provides a necessary baseline for global statistical fit, it remains largely insensitive to subtle, biologically meaningful shifts. Consequently, DES serves as the strictest test of biological utility, ensuring the model identifies the precise gene sets driving the phenotype. Furthermore, the adaptation of PDS addresses the "attenuation" phenomenon, where models correctly predict the direction of regulation but with damped magnitude compared to ground truth. By prioritizing vector direction (cosine similarity) over Euclidean distance, we reward models that capture the correct regulatory mechanism even if the signal strength is conservative.

To further investigate these failure modes, we selected **MED13** from the test set for detailed analysis, as it exhibited relatively high performance scores (DES, PDS, MAE) compared to other genes. However, visual inspection reveals a critical limitation (Figure 2). While the ground truth demonstrates a significant downregulation of MED13 expression following its knockout, the STATE model predicts a distribution centered around zero change ($\Delta \approx 0$), resembling a normal distribution. We hypothesize that STATE tends to predict a conservative normal distribution around zero for unseen perturbations. This behavior likely stems from the model being trained solely on H1 cell line data with a limited set of gene perturbations. Lacking exposure to diverse cell types or combinatorial perturbations, the model adopts a conservative strategy on unknown perturbations, predicting near-control states. Furthermore, the STATE architecture does not explicitly enforce a zero-expression constraint for the target gene, further contributing to this "attenuation" artifact.

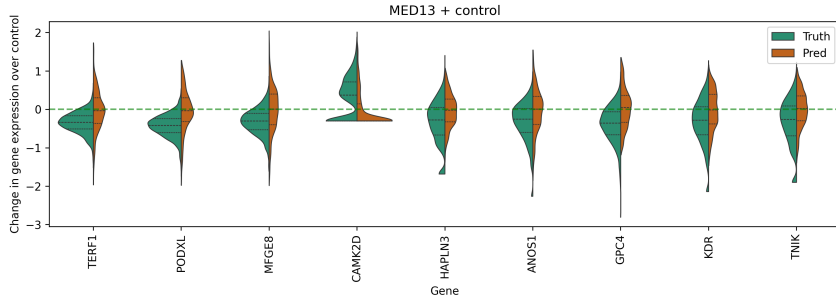


Figure 2: **Analysis of Prediction Failure on MED13.** Despite achieving high global metrics, the STATE model fails to predict the specific downregulation of the target gene MED13. The predicted expression change follows a normal distribution centered at zero (conservative prediction), contrasting with the true downregulation observed in the ground truth. This highlights the model’s tendency to revert to a mean-zero shift when encountering unseen perturbations, a phenomenon not fully captured by aggregate metrics like MAE.

2.3 Takeaway: forward is necessary but not sufficient

While forward benchmarks verify that a model can simulate *effects*, therapeutic discovery necessitates inferring *causes* from observed effects. This fundamental asymmetry motivates an explicit inversion of the perturbation–phenotype mapping, moving from verifying simulation accuracy to validating retrieval capability.

3 Inverting the chain: reverse perturbation retrieval on Norman

3.1 Task definition: from expression phenotype to a ranked list of causal factors

To bridge the gap between transcriptomic modeling and therapeutic target discovery, we reframe the modeling objective from forward prediction to **Reverse Perturbation Retrieval**. While forward models approximate the causal function $f(x, p) \rightarrow y$, drug discovery typically demands the inverse: given a disease signature y , identify the perturbation p that drives it. We operationalize this task as a retrieval problem (Box 1).

Box 1: Task Definition – Reverse Perturbation Retrieval

Query: An observed post-perturbation transcriptomic phenotype (signature), denoted as y_{obs} .

Database: A discrete search space of candidate targets \mathcal{P} (e.g., genome-wide gene knockouts).

Output: An ordered list of candidates ranked by probability, where the objective is to maximize the presence of true causal factors within the top- K predictions.

This formulation aligns with the "reverse retrieval" setting proposed in recent foundation model studies (e.g., scGPT [5]), establishing **Top-K Retrieval** as the standard for evaluation. Consequently, our primary contribution in this section is not the loss function design—which is detailed in Supplementary Note 1—but the rigorous benchmarking of model utility. We shift the evaluation focus from minimizing generation error (MSE) to maximizing actionable "Target Discovery" metrics, specifically **Hit@K**, **Recall@K**, and **Mean Reciprocal Rank (MRR)**.

3.2 Main empirical finding: geometry vs. semantics

We evaluated the retrieval performance of fine-tuned scGPT against robust linear and non-linear baselines (PCA-KNN, Random Forest, XGBoost) on the Norman et al. dataset. Our results reveal a distinct trade-off between local precision and global recall, suggesting a dichotomy in how different models access causal mechanisms.

Phenomenon: The Cross-over Effect. Figure 3 illustrates the "Geometry vs. Semantics" trade-off observed in the target identification task. The bars represent Relevant Hit@K (the probability of retrieving at least one correct driver), while the dashed lines track Recall@K (the fraction of ground-truth targets recovered). At strictly local retrieval windows ($K=1$ to $K=10$), shallow baselines such as PCA-KNN (light blue) and Random Forest (red) outperform scGPT, validating their efficiency in identifying "geometrically reachable" targets with strong phenotypic signatures. However, a decisive performance crossover occurs as the window expands: at $K=40$, scGPT (dark blue) surpasses all baselines, achieving a Recall of approximately 0.52 compared to 0.26 for PCA-KNN. This trend demonstrates that while baselines excel at high-precision "head" retrieval, scGPT possesses the "semantic reachability" necessary to retrieve the "long tail" of complex or subtle perturbation signals that simpler models miss.

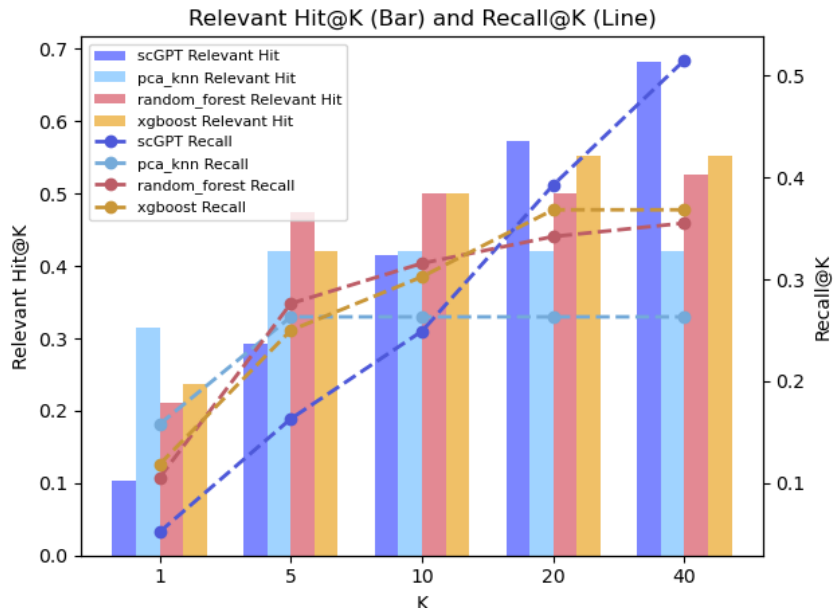


Figure 3: **The "Cross-over" Effect in Reverse Perturbation Retrieval on the Norman Dataset.** Comparison of foundation model (scGPT) performance against geometric (PCA-KNN) and ensemble (Random Forest, XGBoost) baselines across varying retrieval window sizes (K).

Explanation: Geometric vs. Semantic Reachability. To interpret this divergence, we introduce two distinct modes of perturbation retrieval. **Geometric Reachability** characterizes perturbations with strong, high-variance phenotypic signatures. In these cases, the perturbed state lies structurally close to the target definition in the raw gene expression space. Simple geometric neighbors (KNN) are sufficient to lock onto these targets. Conversely, **Semantic Reachability** characterizes perturbations with weak signals, combinatorial effects, or context-dependent regulation. Here, the raw expression distance is noisy or misleading. Retrieving these targets requires "semantic" inference—leveraging learned gene regulatory networks and context embeddings to map a subtle phenotype back to its driver, a capability unique to deep generative models.

Implication for System Design. These findings suggest that the debate should not be framed as "Deep Learning vs. Baselines," but rather as a functional decomposition of the retrieval task. An optimal target discovery system requires a tiered approach: employing geometric models for high-precision, low-cost screening of dominant effects, followed by semantic models (like scGPT) to expand recall and capture robust targets hidden within complex, non-linear regulatory landscapes.

3.3 Biological plausibility checks

To validate the biological utility of the semantic retrieval capabilities, we specifically examined performance on combinatorial perturbations. A critical challenge in perturbation biology, as highlighted by GEARS [7], is predicting and deconstructing genetic interactions.

Our analysis of **Relevant Hit@20** (the probability of retrieving at least one correct driver in a combinatorial pair) demonstrates the practical advantage of the deep learning approach. scGPT achieves a score of 0.5731, compared to 0.4211 for PCA-KNN (Supplementary Table 2). In a drug discovery context, identifying a single valid driver from a combinatorial phenotype is often sufficient to initiate a hit expansion campaign. This suggests that scGPT is more effective at generating valid mechanistic hypotheses for complex phenotypes, even if it does not always rank the complete perturbation pair perfectly at the top.

3.4 Positioning within the benchmark discourse

Our results directly address the growing controversy regarding the utility of large biological models. Recent benchmarks have correctly pointed out that for simple expression reconstruction (minimizing MAE), deep models often fail to consistently outperform linear baselines.

However, we argue that prediction error is a proxy metric that does not necessarily reflect downstream utility. By shifting the evaluation to retrieval, we align model performance with decision-making processes in R&D. Our findings demonstrate that while linear baselines are efficient for geometric interpolation, they fundamentally lack the capacity for the semantic inference required to retrieve complex targets. Thus, the value of deep models lies not in beating linear regression at curve fitting, but in providing actionable recall where geometric heuristics fail.

4 From genes to drugs: an honest boundary on Tahoe

4.1 Why drug target deconvolution is harder than CRISPR gene attribution

While the reverse retrieval of genetic perturbations (Section 3) yielded promising results, extending this logic to pharmacological perturbations reveals a fundamental escalation in complexity. Unlike CRISPR-mediated interventions, which possess a relatively direct "expression → target gene" causality, small-molecule drugs operate primarily at the protein level. The transcriptomic signature observed in scRNA-seq is merely a downstream echo of the initial drug-protein binding event, often confounded by dosage effects, temporal delays, and off-target toxicity.

This distinction creates a significant "inference gap." As highlighted in recent perspectives on the Virtual Cell and AIVC, drug perturbations are characterized by polypharmacology (one drug, multiple targets) and cell-line specificity, where the same compound elicits divergent responses depending on the proteomic context. Consequently, the assumption that a drug's target can be simply retrieved as the "most altered gene" in the transcriptome is biologically naive. The challenge here is not merely computational but mechanistic: the signal required to identify the target may be attenuated or absent at the mRNA level by the time sequencing occurs.

4.2 Empirical bottleneck: annotation scarcity and label ambiguity

To empirically map this boundary, we conducted an exploratory benchmark using the Tahoe-100M dataset. Due to the lack of explicit target annotations in the original release, we integrated external drug-target databases, resulting in a curated subset of 52 drugs with verifiable targets for our experimental pilot.

Our analysis of this subset reveals a stark reality: standard expression-based baselines (Differential Expression ranking and Logistic Regression) failed to reliably identify true targets, performing close to random guessing. We posit that this performance plateau represents a data limitation rather than a model failure. The “ground truth” labels available for these public datasets are often binary and context-independent, failing to account for whether the target is actually expressed or functional in the specific cell line assayed.

Therefore, we frame the current state of Tahoe-100M not as a ready-to-use benchmark, but as a case study in annotation scarcity. The primary conclusion of this exploration is that large-scale perturbation atlases currently lack the “verifiable ground truth” density required to train or evaluate precise reverse-engineering models. Future progress demands a shift in data curation strategy: moving from aggregating massive, noisy datasets to constructing smaller, high-fidelity benchmarks where the specific drug-target interaction is biochemically validated within the assayed context.

4.3 A constructive path forward

To elevate drug target discovery from an exploratory endeavor to a rigorous benchmark, we propose three pivotal shifts in dataset curation and metric design. We advocate for High-confidence subset auditing, arguing that future benchmarks must prioritize the curation of “gold-standard” subsets—such as highly specific inhibitors with defined mechanisms—over the sheer volume of drugs, thereby ensuring that performance bottlenecks stem from modeling limitations rather than label noise. Complementing this cleaner data, evaluation frameworks should adopt Mechanism-level ground truth. Metrics must evolve from strict “Hit@Gene” precision to broader “Hit@Pathway” or “Hit@Mechanism” standards, acknowledging that a drug’s transcriptomic footprint often manifests as the modulation of a downstream functional module rather than a direct expression shift of the binding target. Ultimately, bridging the inference gap for proteome-targeted interventions necessitates the Integration of orthogonal readouts. Since unimodal scRNA-seq captures only downstream echoes of protein-level events, future benchmarks must align with the multimodal vision of foundational models like scGPT, integrating orthogonal data streams—such as proteomics or morphological profiles—to fully reconstruct the causal chain.

5 Discussion

5.1 The central message

We argue that the most actionable use of virtual-cell modeling for drug discovery is not only to simulate perturbation effects, but to invert them into testable target hypotheses. While recent advances have focused on constructing high-fidelity simulators to predict cellular responses, the ultimate utility of AI Virtual Cells (AIVCs) lies in “closing the loop” for discovery. By reframing the biological question from a forward prediction task into a data-driven inference task, we transform the Virtual Cell from a descriptive simulator into a prescriptive engine capable of systematically mapping observed disease phenotypes back to their causal drivers.

5.2 Implications for model evaluation

Our findings suggest that the community’s current reliance on forward prediction metrics creates a misalignment between model optimization and pharmaceutical utility. While we utilized forward validation (e.g., DES, PDS) as a necessary “credibility scaffold” to ensure models capture biological signals rather than statistical noise, high performance on reconstruction error (MSE) does not inherently translate to actionable causal discovery. As highlighted by recent benchmarks, complex deep learning models often fail to consistently outperform simple linear baselines in pure expression forecasting.

However, we posit that this “linear vs. deep” debate is partially a byproduct of the evaluation metric itself. By shifting the objective to Reverse Perturbation Retrieval, we align the evaluation with the decision-making logic of R&D: prioritizing candidates for experimental validation. In this retrieval context, our results demonstrate that while linear baselines are efficient for geometric interpolation, they lack the capacity for the semantic inference required to retrieve complex targets. Thus, the value of deep models lies not in beating linear regression at curve fitting, but in providing actionable recall where geometric heuristics fail.

5.3 A hybrid system hypothesis

Based on the “Geometry vs. Semantics” trade-off uncovered in our retrieval benchmarks, we propose a functional decomposition of the target discovery task. Our results reveal that heuristic baselines (e.g., PCA-kNN) excel at “head” precision for strong phenotypes via geometric similarity, whereas foundation models like scGPT demonstrate superior robustness in retrieving the “long tail” of subtle or complex signals.

This suggests the optimal path forward is not a monolithic model, but a tiered hybrid system. Such a framework would employ geometric models for the high-precision, low-cost screening of dominant effects, while leveraging deep generative models to decipher robust targets hidden within non-linear regulatory landscapes. This approach mirrors the success of Geneformer, which demonstrated that transfer learning can prioritize therapeutic candidates beyond mere expression forecasting. Furthermore, to support this semantic reasoning, future systems must incorporate biological inductive biases—such as the gene regulatory graphs utilized by GEARS or the cell-set distribution modeling of STATE—to improve generalization to unseen combinatorial perturbations.

5.4 Limitations

Validating the feasibility of in silico reverse perturbation establishes a foundational proof-of-concept, yet bridging the divide to industrial application requires navigating three distinct critical boundaries. The most formidable challenge lies in the **“Inference Gap” in pharmacological perturbations**. While genetic target retrieval proved successful on the Norman dataset, extending this logic to pharmacological contexts (Tahoe-100M) exposes a fundamental escalation in complexity. Unlike the direct mechanism of CRISPR, drugs operate at the protein level, producing transcriptomic signatures that are merely downstream echoes confounded by polypharmacology. Bridging this gap demands transcending unimodal scRNA-seq by integrating orthogonal readouts, such as proteomics, to mechanically link protein-level binding with transcriptional outcomes.

Compounding this difficulty is the **Annotation Scarcity in Large-Scale Atlases**. Our analysis of Tahoe-100M reveals that current massive perturbation datasets lack the “verifiable ground truth” density required for rigorous benchmarking; the fact that standard baselines perform near random guessing points to a data limitation rather than a model failure. Meaningful progress therefore necessitates a strategic shift: moving away from the aggregation of massive, noisy datasets and toward the curation of “gold-standard” subsets where drug-target interactions are biochemically validated within the specific assay context.

Beneath these data challenges lies a technical limitation in **Ranking Precision for Deep Models**. Although scGPT excels at recalling complex targets, it trails simple baselines in Mean Reciprocal Rank (MRR), suggesting that while it captures the correct semantic signal, it struggles to isolate it from the inherent noise of single-cell logits to achieve top-tier ranking. Future architectural iterations must prioritizing specialized ranking loss functions or denoising objectives capable of refining these broad semantic signals into precise, top-k predictions.

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Appendix A Supplementary Note 1: Methods

Appendix A.1 Transcriptome-wide Perturbation Prediction

Problem Formulation. The core task of the Transcriptome-wide Perturbation Prediction serves as the **Forward Problem**: predicting the transcriptomic consequence of a defined intervention. Formally:

$$\text{given } (x, p) \Rightarrow \hat{y} = f(x, p)$$

where:

- x denotes the *pre-perturbation cell state*, serving as the baseline. This is empirically represented by the distribution of control cells within the same experimental batch or cell type. For instance, in scGPT benchmarks on the Norman dataset, x is approximated by the mean expression vector of all control cells ($1 \times M$ genes).
- p represents the *known perturbation* (e.g., gene knockout, CRISPRi, or drug). This is typically encoded as a discrete gene identifier or a multidimensional perturbation embedding.
- \hat{y} is the *predicted post-perturbation state*. The model is optimized to minimize the divergence between \hat{y} and the experimentally observed expression profile y .

Appendix A.1.1 Evaluation metrics: PDS, MAE, DES

Perturbation Discrimination Score (PDS). The Perturbation Discrimination Score (PDS) measures whether the predicted gene expression changes induced by different perturbations preserve the correct global directionality across all genes. For each perturbation $k \in \{1, \dots, N\}$, we first construct pseudobulk expressions by averaging all cells under the same perturbation, yielding the ground-truth pseudobulk $y_k \in \mathbb{R}^G$, the predicted pseudobulk $\hat{y}_k \in \mathbb{R}^G$, and the control (NTC) pseudobulk $y_{ntc} \in \mathbb{R}^G$. Perturbation-induced expression changes are then defined as $\delta_k = y_k - y_{ntc}$ and $\hat{\delta}_k = \hat{y}_k - y_{ntc}$. For each predicted perturbation $\hat{\delta}_k$, we compute cosine similarities to all true perturbation deltas $\{\delta_j\}_{j=1}^N$ as

$$S_{k,j} = \frac{\hat{\delta}_k \cdot \delta_j}{\|\hat{\delta}_k\|_2 \|\delta_j\|_2}.$$

These similarities are ranked, and the rank of the correct perturbation k is denoted by R_k . The final score is reported as the mean rank $\text{PDS}_{\text{rank}} = \frac{1}{N} \sum_{k=1}^N R_k$, or its normalized version $\text{nPDS}_{\text{rank}} = \frac{1}{N^2} \sum_{k=1}^N R_k$.

MAE on Top 2000 Genes by Ground-Truth Fold Change. To focus evaluation on biologically meaningful signals, we additionally compute the mean absolute error (MAE) restricted to the genes with the largest true perturbation effects. For each perturbation k , we start from the raw (unnormalized) average counts $c_k \in \mathbb{R}^G$ and control counts $c_{ntc} \in \mathbb{R}^G$, and compute the absolute \log_2 fold change for each gene g using a pseudocount of 1:

$$\text{LFC}_{k,g} = |\log_2(c_{k,g} + 1) - \log_2(c_{ntc,g} + 1)|.$$

Genes are ranked by $\text{LFC}_{k,g}$ in descending order, and the top 2000 genes form the index set Ω_k . Using log1p-normalized pseudobulk expressions y_k and \hat{y}_k (the same representation as in PDS), the MAE is computed only over these selected genes and then averaged across perturbations:

$$\text{MAE}_{\text{top2k}} = \frac{1}{N} \sum_{k=1}^N \left(\frac{1}{2000} \sum_{g \in \Omega_k} |\hat{y}_{k,g} - y_{k,g}| \right).$$

This metric emphasizes prediction accuracy on genes that exhibit the strongest true responses to each perturbation.

Differential Expression Score (DES). The Differential Expression Score (DES) measures how accurately a model recovers perturbation-induced differential gene expression. For each perturbation $k \in \{1, \dots, N\}$, we first perform differential expression analysis between perturbed and control cells for both the predicted and ground-truth data using the Wilcoxon rank-sum test with tie correction,

and identify significantly differentially expressed (DE) genes by controlling the false discovery rate at level $\alpha = 0.05$ with the Benjamini–Hochberg procedure. This yields a predicted DE gene set $G_{k,\text{pred}}$ and a ground-truth DE gene set $G_{k,\text{true}}$, with sizes $n_{k,\text{pred}} = |G_{k,\text{pred}}|$ and $n_{k,\text{true}} = |G_{k,\text{true}}|$, respectively. If $n_{k,\text{pred}} \leq n_{k,\text{true}}$, the DES for perturbation k is defined as the size of the intersection between the predicted and true DE gene sets, normalized by the size of the true set,

$$\text{DES}_k = \frac{|G_{k,\text{pred}} \cap G_{k,\text{true}}|}{n_{k,\text{true}}}.$$

If $n_{k,\text{pred}} > n_{k,\text{true}}$, to avoid overpenalizing predictions that overestimate differential expression, we construct a reduced predicted set $\tilde{G}_{k,\text{pred}}$ by selecting the $n_{k,\text{true}}$ genes with the largest absolute log fold changes (relative to control cells) from $G_{k,\text{pred}}$, and compute

$$\text{DES}_k = \frac{|\tilde{G}_{k,\text{pred}} \cap G_{k,\text{true}}|}{n_{k,\text{true}}}.$$

The final DES is obtained by averaging DES_k across all perturbations.

Overall score We make a scale of each metric based on the cell-mean baseline model and calculate the overall score:

$$S = \frac{\text{DES}_{\text{scaled}} + \text{PDS}_{\text{scaled}} + \text{MAE}_{\text{scaled}}}{3} \times 100$$

Appendix A.1.2 cell-mean model

This baseline predicts gene expression by returning the global mean expression profile computed from all perturbed cells in the training data. It ignores the input perturbation and cell-specific information, and simply broadcasts the pre-computed mean expression vector to all samples at inference time.

Appendix A.1.3 ridge regression

We apply a linear ridge-regression model in a PCA-reduced gene expression space. Pseudo-bulk mean expression profiles are computed for each perturbation, and a regularized linear mapping is learned to predict genome-wide expression changes from the perturbed gene, with the global perturbed mean added back to obtain the final expression profile.

Appendix A.1.4 random forest regression

The preprocessing and PCA representation are identical to the ridge regression baseline. A Random Forest regressor is used in place of the linear model to predict perturbation effects in the latent space, which are then mapped back to gene space to generate the final expression profile.

Appendix A.1.5 STATE Implementation

We implement the STATE model following the official open-source implementation released by the authors. In this work, we focus on the *State Transition (ST)* module of STATE, which is responsible for predicting perturbation-induced transcriptomic responses given control cells and perturbation conditions.

Perturbation features. Perturbation identities are represented using pretrained protein embeddings. Specifically, gene perturbations are encoded using ESM2-based perturbation feature vectors provided by the official STATE repository. These embeddings allow the model to incorporate prior biological information about gene identity without directly accessing expression-level supervision.

Model architecture. The ST module takes as input control cell expression profiles, perturbation embeddings, and batch embeddings to account for technical variability (Figure 4). All inputs are projected into a shared hidden space and processed by a Transformer backbone, which models how perturbations induce shifts in cellular state distributions. The output representations are mapped back to the gene expression space to generate predicted perturbed transcriptomes.

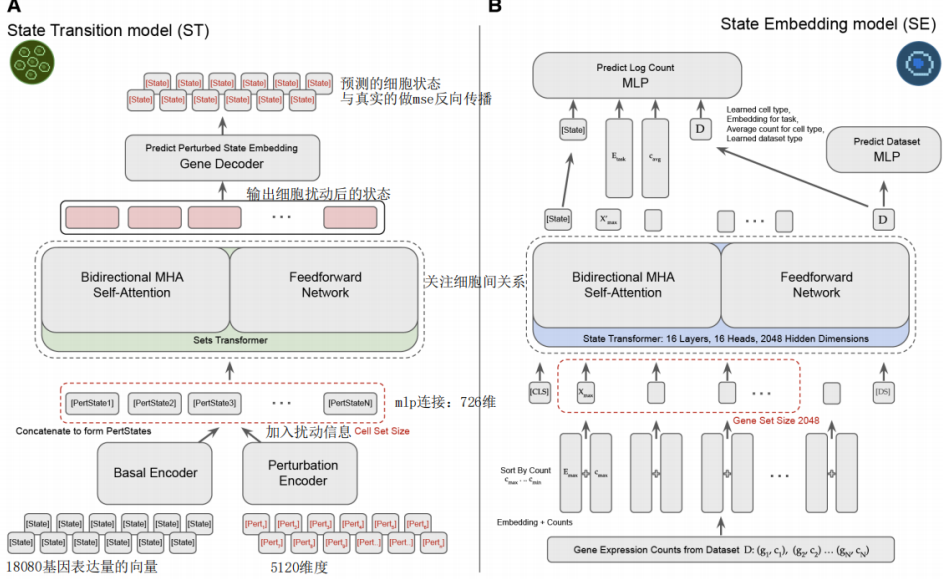


Figure 4: The architecture of the STATE model. The State Transition (ST) module integrates control expression, perturbation embeddings, and batch information to predict post-perturbation transcriptomic states.

Appendix A.1.6 scGPT Finetune

To adapt the scGPT foundation model for the specific task of predicting cellular responses to perturbations, we employ a transfer learning strategy that balances the retention of generalized biological knowledge with the flexibility to learn condition-specific dynamics.

Model Initialization and Parameter Freezing. We initialize the model using pre-trained scGPT checkpoints (Figure 5). To mitigate catastrophic forgetting and ensure computational efficiency, we adopt a partial freezing strategy. The core representation learning modules—specifically the gene token encoder, the continuous value encoder, and the central transformer encoder layers—are frozen. This preserves the model’s pre-learned understanding of gene-gene interactions and cellular states.

Concurrently, we designate specific components as trainable to capture perturbation-specific features. A dedicated embedding layer (`pert_encoder`) is optimized to represent distinct perturbation flags. The output assembly, specifically the affine expression decoder (`ExprDecoder`), remains trainable. This module utilizes coefficient and bias heads to reconstruct gene expression values. If the model is configured for explicit zero-probability modeling, the corresponding zero-prob heads are also optimized.

Composite Loss Function. The fine-tuning process minimizes a composite loss function designed to align predicted expression profiles with ground truth data while preserving biologically relevant differential expression signals. For each training batch, the total loss \mathcal{L}_{total} is calculated as a weighted sum of four specific objectives.

The Sliced Wasserstein-1 distance (\mathcal{L}_{sw1}) is employed as the Distribution Alignment component to minimize the distributional discrepancy between the predicted and actual gene expression profiles. A ProtoInfoNCE loss (\mathcal{L}_{proto}) is applied to pseudobulk deltas for Contrastive Learning, encouraging the model to distinguish between different perturbation effects in the latent space. When a Differential Expression (DE) gene map is available, two auxiliary losses are included: \mathcal{L}_{de_rank} , which penalizes errors in the ranking of DE genes to prioritize biologically significant changes, and \mathcal{L}_{dir} , which enforces the correct directionality of regulation (up-regulation vs. down-regulation).

In this study, the default weighting coefficients are set to $\lambda_{sw1} = 0.60$, $\lambda_{proto} = 0.25$, $\lambda_{rank} = 0.10$, and $\lambda_{dir} = 0.05$.

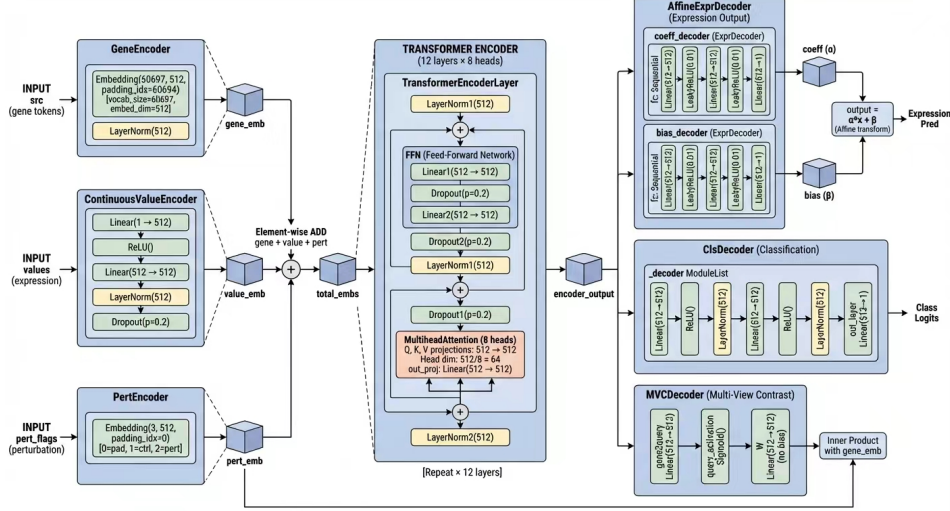


Figure 5: Overview of the scGPT model adaptation for perturbation prediction. The model is initialized with pre-trained weights, with core modules frozen to retain biological knowledge, while the perturbation encoder and decoders are fine-tuned.

Optimization Procedure. Optimization is performed using Automatic Mixed Precision (AMP) combined with gradient scaling to maximize training throughput and numerical stability. To prevent overfitting, we implement an early stopping mechanism that monitors a validation metric (`overall_score`). Training is halted if this score fails to improve within a specified patience window, and the checkpoint corresponding to the best validation performance is saved for inference.

Appendix A.2 Target Gene Identification

Problem Formulation. While forward prediction tests a model’s understanding of causality, the actionable goal in drug discovery is the inverse: inferring the mechanism that explains an observed phenotype. We formulate this through two related inverse problems:

1. Reverse Perturbation Prediction (Inverse Problem). This task asks: *given an observed cellular response, what perturbation caused it?* Mathematically, given a control state x and a perturbed state y , we seek the perturbation \hat{p} that best explains the transition:

$$\text{given } (x, y) \Rightarrow \hat{p} = g(x, y)$$

Ideally, this is the perturbation that minimizes the discrepancy between the forward model’s prediction and the observation:

$$\hat{p} = g(x, y) = \arg \min_p d(f(x, p), y)$$

2. Target Discovery (Therapeutic Goal). This extends the inverse framework to therapeutic intervention. Here, we aim to transform a disease state x_{abn} into a desired healthy state y_{desired} :

$$p^* = g(x_{\text{abn}}, y_{\text{desired}}) = \arg \min_p d(f(x_{\text{abn}}, p), y_{\text{desired}})$$

Here:

- x_{abn} : The abnormal (e.g., disease-associated) cell state.
- y_{desired} : The target state (e.g., healthy control or a specific functional profile).
- p : The candidate perturbation/target (e.g., a gene KO from a discrete set).
- $f(x_{\text{abn}}, p)$: The forward intervention effect model.
- $d(\cdot, \cdot)$: A distance metric measuring how close the intervention brings the cell to the desired state.

We explore two methodological approaches to solve these problems:

Route A: Forward Model-Based Retrieval. This approach directly utilizes the forward simulator f . For every candidate perturbation $p \in \mathcal{P}$: 1. Predict the outcome $\hat{y}_p = f(x_{\text{abn}}, p)$. 2. Compute the distance to the target $d(\hat{y}_p, y_{\text{desired}})$. 3. Rank candidates by minimizing this distance. Here, g is implicitly defined by the optimization over f .

A critical limitation of the route A framework employed by original scGPT paper is its reliance on a pre-defined, closed candidate set composed of enumerated gene combinations. While this approach is feasible for a restricted subspace—such as 20 genes yielding 210 distinct single- and double-gene perturbation classes—it suffers from prohibitive combinatorial explosion when applied to larger gene pools or when the specific number of target genes is unknown. As the search space expands to accommodate genome-wide candidates or higher-order interactions, the number of potential combinatorial targets increases exponentially, rendering the exhaustive generation and retrieval of candidate profiles computationally intractable. Consequently, due to these scalability constraints and the infeasibility of defining a discrete label space for unknown perturbation complexities, we exclude this combinatorial classification route from our methodology.

Route B: Direct Inverse Mapping. This approach learns a parameterized inverse function g_θ (e.g., a classifier or regressor) to directly predict the optimal perturbation:

$$p^* \approx g_\theta(x_{\text{abn}}, y_{\text{desired}})$$

Instead of explicitly modeling the transition, the model learns the mapping from (state, target) pairs to perturbations directly from data. This absorbs the forward dynamics and distance metric into the model weights.

Evaluation Metrics. To quantitatively assess the performance of the reverse perturbation prediction models, we employ a suite of ranking-based metrics. Let $\mathcal{C}_{\text{test}}$ denote the set of test perturbation conditions. For each condition $c \in \mathcal{C}_{\text{test}}$, let \mathcal{G}_c^* be the set of ground-truth target genes, and let \mathcal{G}_c^K represent the set of the top- K genes identified by the model based on the predicted scores.

1. Mean Reciprocal Rank (MRR). The Mean Reciprocal Rank evaluates the model’s ability to place the correct perturbation targets at the top of the ranking list. Since a perturbation may involve multiple target genes (e.g., in combinatorial perturbations), we define the reciprocal rank based on the *highest-ranked* (best) true target.

$$\text{MRR} = \frac{1}{|\mathcal{C}_{\text{test}}|} \sum_{c \in \mathcal{C}_{\text{test}}} \max_{g \in \mathcal{G}_c^*} \frac{1}{\text{rank}(g)}$$

where $\text{rank}(g)$ denotes the position of gene g in the predicted descending order list (1-indexed). A higher MRR indicates that the first correct target appears earlier in the candidate list.

2. Exact Hit@K. Exact Hit@K is a stringent metric that measures the proportion of test cases where the model successfully retrieves the *entire* set of target genes within the top- K predictions. This metric is particularly relevant for identifying complete combinatorial pairs.

$$\text{Exact Hit@K} = \frac{1}{|\mathcal{C}_{\text{test}}|} \sum_{c \in \mathcal{C}_{\text{test}}} \mathbb{I}(\mathcal{G}_c^* \subseteq \mathcal{G}_c^K)$$

where $\mathbb{I}(\cdot)$ is the indicator function, which equals 1 if the condition is true and 0 otherwise.

3. Relevant Hit@K. Relevant Hit@K assesses the model’s capacity to discover *at least one* correct driver gene within the top- K candidates. This metric reflects the practical utility of the model in experimental screening, where identifying a subset of drivers is often sufficient for further validation.

$$\text{Relevant Hit@K} = \frac{1}{|\mathcal{C}_{\text{test}}|} \sum_{c \in \mathcal{C}_{\text{test}}} \mathbb{I}(\mathcal{G}_c^* \cap \mathcal{G}_c^K \neq \emptyset)$$

4. Recall@K. Recall@K quantifies the fraction of ground-truth target genes that are successfully retrieved within the top- K predictions. Unlike Relevant Hit@K, which is binary, Recall@K penalizes the model for missing parts of a combinatorial perturbation (e.g., finding only 1 gene of a 2-gene pair).

$$\text{Recall@K} = \frac{1}{|\mathcal{C}_{\text{test}}|} \sum_{c \in \mathcal{C}_{\text{test}}} \frac{|\mathcal{G}_c^* \cap \mathcal{G}_c^K|}{|\mathcal{G}_c^*|}$$

This metric provides a granular view of the model’s retrieval completeness, particularly as K increases (e.g., Recall@1, Recall@5, Recall@10, etc.).

Appendix A.2.1 Baseline Models

PCA + kNN Baseline. To establish a retrieval-based baseline, we first aggregate single-cell gene expression vectors into pseudobulk profiles for each perturbation condition. This yields a feature vector $\mathbf{x}_c \in \mathbb{R}^G$ over G genes, associated with a multi-hot label vector $\mathbf{y}_c \in \{0, 1\}^G$ derived from the specific perturbation targets. To prevent trivial mappings (i.e., data leakage), we optionally mask the expression values of target genes within \mathbf{x}_c . We apply Principal Component Analysis (PCA) to the training profiles, projecting them into a latent space $\mathbf{z}_c = \mathbf{W}^\top \mathbf{x}_c \in \mathbb{R}^d$. For a given query condition q , we retrieve the k nearest neighbors in the PCA space ($\mathcal{N}_k(q)$) using Cosine or Euclidean similarity. The final gene relevance scores s_q are computed via a weighted aggregation of the neighbors’ labels. Hyperparameters, including the latent dimension d and neighborhood size k , are optimized based on the Mean Reciprocal Rank (MRR) of the validation set before final evaluation on held-out test conditions.

Random Forest Baseline. Utilizing the same pseudobulk representations \mathbf{x}_c as input, we employ a multi-output Random Forest regressor to predict continuous per-gene relevance scores $\hat{\mathbf{y}}_c \in \mathbb{R}^G$ (or a restricted subset of candidate targets). Each tree in the ensemble learns non-linear splits on the input features, and the final prediction constitutes an average over T trees. Target vectors are constructed as multi-hot encodings of the condition labels, with optional input masking applied to ensure generalization. Hyperparameters such as the number of estimators T and maximum tree depth are selected via validation MRR. The predicted score vectors are subsequently ranked to compute Top-K metrics on the test split.

XGBoost Baseline. We further benchmark performance using Gradient Boosted Decision Trees (GBDT) implemented via XGBoost in a multi-output regression setting. The model predicts per-gene scores by optimizing a squared-error objective function $\mathcal{L} = \sum_c \|\mathbf{y}_c - \hat{\mathbf{y}}_c\|_2^2$ through an additive training process, where η represents the learning rate and f_m represents the m -th tree. This approach captures non-linear gene-score relationships while managing model complexity through depth constraints and subsampling. As with other baselines, inputs \mathbf{x}_c are subject to masking strategies to prevent leakage. Hyperparameters are tuned on validation MRR, and the final evaluation employs the same ranking-based metrics as the PCA+kNN and Random Forest baselines.

Appendix A.2.2 Fine-tuning scGPT for Discriminative Perturbation Prediction

This section details the methodology for the discriminative fine-tuning strategy ("Route B1"), which leverages the pre-trained scGPT backbone to identify perturbation targets at single-cell resolution. Unlike the pseudobulk baselines, this approach models the perturbation likelihood for every gene within individual cells, utilizing a control-aware embedding strategy to isolate perturbation-specific signals.

Task Formulation. The reverse perturbation prediction is formulated as a ranking problem. Given a perturbed single-cell expression profile \mathbf{x} and its associated condition context, the model learns a scoring function $f(\mathbf{x}) : \mathbb{R}^G \rightarrow \mathbb{R}^G$ that outputs a vector of logits $\hat{\mathbf{p}}$. These logits represent the likelihood of each gene being a target of the perturbation. The ground truth is represented as a multi-hot vector $\mathbf{y} \in \{0, 1\}^G$, where $y_j = 1$ if gene j is a target in the condition string (e.g., "GeneA+GeneB"), and 0 otherwise.

Architecture and Input Processing. The model architecture builds upon the scGPT transformer backbone. Single-cell expression counts are preprocessed via the standard scGPT pipeline: counts are normalized, log-transformed, and quantile-binned. Non-zero gene expressions are tokenized into gene-value pairs and prepended with a [CLS] token, yielding a sequence input for the encoder.

To mitigate confounding biological variation (e.g., cell cycle or batch effects) and emphasize the perturbation signal, we employ a control-matched delta embedding strategy:

- **Encoder Embedding:** The perturbed cell i is embedded by the scGPT encoder to obtain a latent representation \mathbf{e}_i .

- **Reference Aggregation:** We sample K matched control cells sharing the same batch and cell-type metadata. Their embeddings are averaged to form a reference baseline: $\mathbf{e}_{ref} = \frac{1}{K} \sum_{k=1}^K \mathbf{e}_{i,k}^{ctrl}$.
- **Delta Representation:** A perturbation-specific embedding \mathbf{h}_i is computed by subtracting the reference signal:

$$\mathbf{h}_i = \mathbf{e}_i - \mathbf{e}_{ref}$$

The gene-scoring head projects this delta embedding \mathbf{h}_i into the gene embedding space. Let \mathbf{g}_j denote the pre-trained embedding for gene j . The score s_{ij} for gene j in cell i is computed via a learnable projection \mathbf{W} and a dot product:

$$s_{ij} = \langle \mathbf{W}\mathbf{h}_i, \mathbf{g}_j \rangle$$

Alternatively, a multi-layer perceptron (MLP) can be applied to the concatenation $[\mathbf{h}_i; \mathbf{g}_j]$.

Optimization Objective. The training objective is a composite loss function designed to rank true perturbation targets higher than non-targets while maintaining calibration. The total loss \mathcal{L} is a weighted sum:

$$\mathcal{L} = \lambda_{\text{rank}} \mathcal{L}_{\text{rank}} + \lambda_{\text{bce}} \mathcal{L}_{\text{bce}}$$

In this study, we utilize weights $\lambda_{\text{rank}} = 0.7$ and $\lambda_{\text{bce}} = 0.1$. The components are defined as follows:

- **Pairwise Ranking Loss ($\mathcal{L}_{\text{rank}}$):** This loss enforces that the scores of ground-truth target genes ($p \in \mathcal{P}$) are higher than those of sampled negative genes ($n \in \mathcal{N}$) by a margin m :

$$\mathcal{L}_{\text{rank}} = \mathbb{E}_{p,n} [\max(0, m - (s_{i,p} - s_{i,n}))]$$

Negative samples \mathcal{N} are drawn using a mix of random sampling and "hard negative" mining (high-scoring non-targets) to facilitate robust learning.

- **Auxiliary Binary Cross-Entropy (\mathcal{L}_{bce}):** To stabilize training and provide probabilistic calibration, we apply BCE loss on the positive set plus a subset of sampled negatives. This avoids gradient dominance from the vast majority of non-target genes (class imbalance) while guiding the model to predict correct multi-hot labels.

Appendix B Supplementary Note 2: Experimental Details

Appendix B.1 Transcriptome-wide Perturbation Prediction

Appendix B.1.1 Experimental Setup

Data preprocessing. We use the ARC Institute VCC competition support dataset (competition_support_set), which provides preprocessed single-cell RNA-seq data for H1 human embryonic stem cells. Gene expression values are transformed using a log1p normalization, which stabilizes variance and reduces the dominance of highly expressed genes. Control cells corresponding to non-targeting perturbations are retained and used as a reference state.

Data Split. Instead of using the original VCC split, we re-split 150 gene perturbations into training, validation, and test sets at the perturbation level to evaluate generalization to unseen perturbations. Specifically, 64% of perturbation genes are used for training, 16% for validation, and the remaining 20% (30 genes) form a held-out test set. Control (non-targeting) cells are included in all splits to provide a consistent reference state. Importantly, there is no overlap of perturbation genes between the training, validation, and test sets, ensuring that models are evaluated on entirely unseen perturbation targets.

Appendix B.1.2 Main Results

We select Ridge Regression and Random Forest Regression as representative baseline models, as their overall scores fall in the range of 1 to 2. Among all methods, the State Model achieves the best overall performance, with substantial improvements in both DES and PDS. The scGPT fine-tuned model also shows competitive performance in DES and overall score, but suffers from a significantly

Table 1: Performance comparison of different models.

Model	PDS	MAE	DES	Overall
Cell-mean baseline	0.5167	0.1258	0.1075	0.00
Ridge Regression	0.5167	0.1253	0.1466	1.59
Random Forest Regression	0.5167	0.1253	0.1360	1.19
State Model	0.5367	0.1286	0.3023	7.28
scGPT finetune	0.5089	0.2763	0.2620	6.27

higher MAE. Overall, more complex models tend to improve DES and PDS, but these gains do not consistently translate into lower MAE. The detailed results are reported in Table 1.

The aim of the Virtual Cell Challenge (VCC) was to predict post-perturbation single-cell gene expression in the H1 cell based on 150 available perturbations. Though the dataset itself is single-cell, the competition metrics PDS (Perturbation Discrimination Score) and MAE (Mean Absolute Error) were calculated on the pseudobulk gene expression profiles. The third competition metric, DES (Differential Expression Score), in theory, is a more challenging metric as it does consider the single-cell expression distributions and uses standard differentially expressed gene algorithms to infer differentially expressed genes.

Conceptually, DES evaluates a model’s ability to faithfully reproduce biologically interpretable perturbation effects, focusing on whether differentially expressed genes are correctly identified and whether the most influential genes are properly prioritized. As such, DES provides a direct measure of how well a model captures single-cell-level perturbation signals.

Under these metrics, our model learns meaningful perturbation patterns at the single-cell level, as evidenced by non-trivial DES scores. However, the predicted expression changes for individual cells are often small or close to zero, causing the learned effects to be attenuated when aggregated into pseudobulk representations ($\hat{\delta}_k$). This attenuation partially explains the relatively low PDS scores observed.

Furthermore, single-cell data are inherently noisy, which can interfere with effective model training. Since our model is optimized at the single-cell level rather than directly on pseudo-bulk, this mismatch likely further contributes to the lower performance on PDS.

Appendix B.2 Target Gene Identification

Appendix B.2.1 Experimental Setup

In our experiments, we instantiate these tasks (Section Appendix A.2) as follows:

- **Norman Dataset (Genetic Perturbations):** We define the task as *Reverse Genetic Perturbation Identification*. The input is the perturbed expression profile (with optional control baseline), and the output is a ranking of potential perturbation genes. Ground truth labels are derived from Perturb-seq metadata, and performance is evaluated using ranking metrics (Hit@K, MRR, NDCG).
- **Tahoe Dataset (Drug Perturbations):** We define the task as *Reverse Drug Target Identification*. The input is the drug-treated vs. DMSO control profile (or delta signature). The output is a ranking of the drug’s target gene. By focusing on the single-target subset, we ensure a clean label space for evaluation.

Dataset and Preprocessing. We evaluated our method using the Norman et al. Perturb-seq dataset, pre-processed via the GEARS framework. The dataset comprises $N = 91,205$ cells and $G = 5,045$ log-normalized Highly Variable Genes (HVGs). Gene symbols were mapped to the pre-trained scGPT vocabulary to ensure compatibility. While control cells (unperturbed) were retained to serve as references for computing perturbation-specific shifts (e.g., delta embeddings), the "control" class itself was excluded from the target label space.

Condition Filtering and Canonicalization. To ensure robust statistical modeling, we applied quality control filters to the perturbation conditions. Conditions were first canonicalized to enforce

lexicographic order (e.g., treating $A + B$ and $B + A$ as identical) and to remove redundant control strings. We subsequently filtered out conditions with insufficient cell coverage, requiring a minimum of 50 cells for single-gene perturbations and 30 cells for combinatorial (double) perturbations. This process yielded a curated set of valid conditions for splitting.

Compositional Splitting Strategy. We employed a strict condition-level splitting strategy to prevent data leakage, ensuring that no perturbation condition appeared in more than one data split (train/validation/test). To rigorously test the model's ability to generalize to novel biological contexts, we adopted a "compositional split" design:

- **Unseen Single Genes:** A fraction of single-gene perturbations (15%) was held out as "unseen."
- **Combinatorial Logic:** Combinatorial perturbations were categorized based on the training visibility of their constituent genes. "Seen2" doubles (where both genes are present in the training set as singles) were stratified by gene frequency and cell count to ensure balanced distribution across splits. Conversely, "Seen1" (one gene unseen) and "Seen0" (both genes unseen) combinations were exclusively allocated to the test set to evaluate zero-shot generalization.

Dataset Statistics. The final filtered dataset consisted of 236 unique perturbation conditions across 83,803 cells. Using a fixed random seed (42), the data was partitioned as follows:

- **Training Set:** 147 conditions (56,580 cells), comprising 81 single perturbations and 66 "Seen2" doubles.
- **Validation Set:** 23 conditions (6,560 cells).
- **Test Set:** 66 conditions (20,712 cells).

To facilitate detailed performance analysis, the test set was further stratified into four generalization tiers: 15 unseen single perturbations, 15 "Seen2" doubles, 31 "Seen1" doubles, and 5 "Seen0" doubles. This split was persisted and applied uniformly across all benchmarked models (scGPT, PCA+kNN, Random Forest, XGBoost) to ensure a standardized evaluation protocol.

Appendix B.2.2 Results

The quantitative results for the Reverse Genetic Perturbation Identification task are summarized in Table 2. Interestingly, robust baselines operating on pseudobulk profiles (PCA+kNN and Random Forest) outperform the scGPT single-cell fine-tuning approach in top-ranking metrics, such as MRR (0.3602 vs 0.1975). This indicates that for immediate precision (Hit@1, Hit@5), the aggregated signal used by pseudobulk methods is cleaner and more discriminative than noisy single-cell logits. However, as the retrieval window expands (Hit@20, Hit@40), scGPT demonstrates superior recall, significantly overtaking the baselines (Recall@40 of 0.5152 vs 0.3684). This suggests that while scGPT successfully identifies valid perturbation signals, it tends to rank them lower in the candidate list, likely due to the difficulty of optimizing rank-based objectives directly on high-variance single-cell data.

Table 2: Performance comparison of scGPT and baseline models on the Norman dataset for Reverse Perturbation Prediction. Best results are bolded.

Metric	scGPT	PCA+kNN	Random Forest	XGBoost
MRR	0.1975	0.3602	0.3232	0.3323
Exact Hit@1	0.0000	0.0000	0.0000	0.0000
Relevant Hit@1	0.1039	0.3158	0.2105	0.2368
Recall@1	0.0520	0.1579	0.1053	0.1184
Exact Hit@5	0.0526	0.1053	0.0789	0.0789
Relevant Hit@5	0.2922	0.4211	0.4737	0.4211
Recall@5	0.1631	0.2632	0.2763	0.2500
Exact Hit@10	0.0963	0.1053	0.1316	0.1053
Relevant Hit@10	0.4155	0.4211	0.5000	0.5000
Recall@10	0.2492	0.2632	0.3158	0.3026
Exact Hit@20	0.2123	0.1053	0.1842	0.1842
Relevant Hit@20	0.5731	0.4211	0.5000	0.5526
Recall@20	0.3927	0.2632	0.3421	0.3684
Exact Hit@40	0.3476	0.1053	0.1842	0.1842
Relevant Hit@40	0.6828	0.4211	0.5263	0.5526
Recall@40	0.5152	0.2632	0.3553	0.3684