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Multi-task learning in perturbation modeling

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Περιεχόμενα

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1 Abstract

With the recent advancements in single-cell technology and the large scale perturbation datasets, the field of perturbation modeling has created an opportunity for a wide variety of computational methods to be leveraged to harness its potential. Multi-task learning is one of the methods that has been left unexplored in this field. In this study we aim to bridge this gap unraveling the potential of multi-task learning in single-cell perturbation modeling.

2 Introduction

The complexity of biological systems have imposed a challenge to capture the underlying mechanisms of cellular heterogeneity. Deciphering the effect of external stimuli (perturbation) at the cellular level, a field referred to as perturbation modeling [5], plays a crucial role in biomedicine and drug discovery. With the recent surge of data generation, machine learning methods aim to understand the effect of perturbations and to extrapolate on unseen events.

An overview of the models on perturbation modeling can be found on this study [3]. One of the main objectives is the out-of-distribution detection, which is the focal point of our study. The task is about predicting the perturbation response of the omics signature of cells with a specific cell type, while having observed the perturbation response of other cell types.

UnitedNet [10] is a multi-task framework that has shown its potential in multi-omics tasks such as cross modal prediction and cell type classification. We aim to extend this approach to perturbation modeling.

3 Current single-cell perturbation modeling methods

In the literature body, there are several approaches for predicting single-cell perturbation responses. To compare our multi-task method, we have chosen the models of scGen [7], scPreGAN [11], scButterfly [1], and scVIDR [6].

scGen projects the gene expression profile to a probabilistic latent space with a VAE. Then, the perturbation effect is modeled with a vector that represents the difference between the control and perturbed gene expression projections in the latent space.

scVIDR builds upon scGen by enhancing the architecture with cell type-specific knowledge. It is capable of predicting cellular responses to multiple chemical perturbations in a dose-dependent manner. As a multi-task model, scVIDR leverages data from various perturbations to improve prediction accuracy across conditions.

scPreGAN is based on a GAN and autoencoder setup. The study aims to decouple the perturbation effect from the latent space, and to apply it to the decoder stage.

scButterfly was not originally designed for perturbation modeling. However, its cross-modal architecture, which includes dual aligned VAEs, can be repurposed for perturbation tasks. Rather than predicting one omic modality from another, the model can treat perturbed and control gene expression profiles as distinct modalities.

The performance of all of these models will serve as a baseline of our multi-task learning architecture.

In a **fully-connected** network,
FiLM applies a different affine
transformation to each feature.

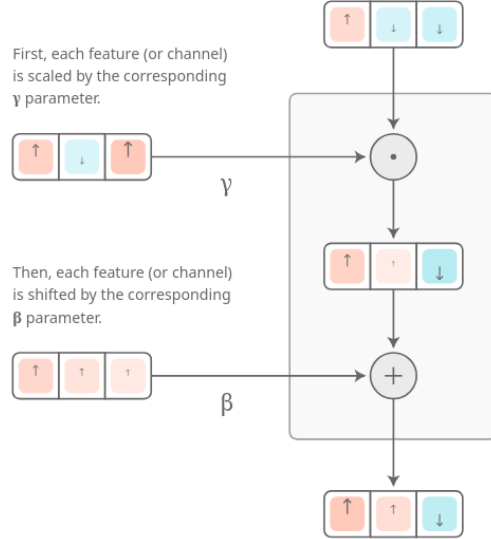


Figure 1: Illustration of the feature-wise transformation [2]

4 Method

Multi-task learning is a machine learning paradigm and its core idea is that training a model to solve multiple tasks can be more effective than training separate models for each specific task [12]. A joint architecture that shares knowledge between the tasks can lead to better generalization. The relationship of the tasks determines the positive or negative transfer to each other and the overall effectiveness of the paradigm.

Defining as a task the prediction of the gene expression given a perturbation, we will explore designing a model that can predict gene expressions after a perturbation for a set of perturbations.

One of the key problems of deep learning methods is the data demand. Another benefit of multi-task learning is the combination of data from multiple sources of informations, especially in perturbation modeling where the data is limited for a specific number of perturbations.

To integrate the tasks, we have explored the application of feature-wise transformations [2]. For this kind of transformation, we have:

$$\text{FiLM}(x) = \gamma(z) \odot x + \beta(z)$$

, where γ , and β are learnable parameters generated by a network that represent a condition z (e.g. a vector that indicates the task), and x is the input.

This particular technique is referred to as conditional affine transformation (a combination of multiplicative and additive conditioning) that shifts and scales the input element-wise. It is efficient in terms of scaling and parameters compared to multi-head architectures, where each task has its dedicated network to generate the output of the task.

In our approach, we aim to decouple the perturbation effect by constructing a perturbation-free latent space, while explicitly modeling the perturbation response through a conditioning

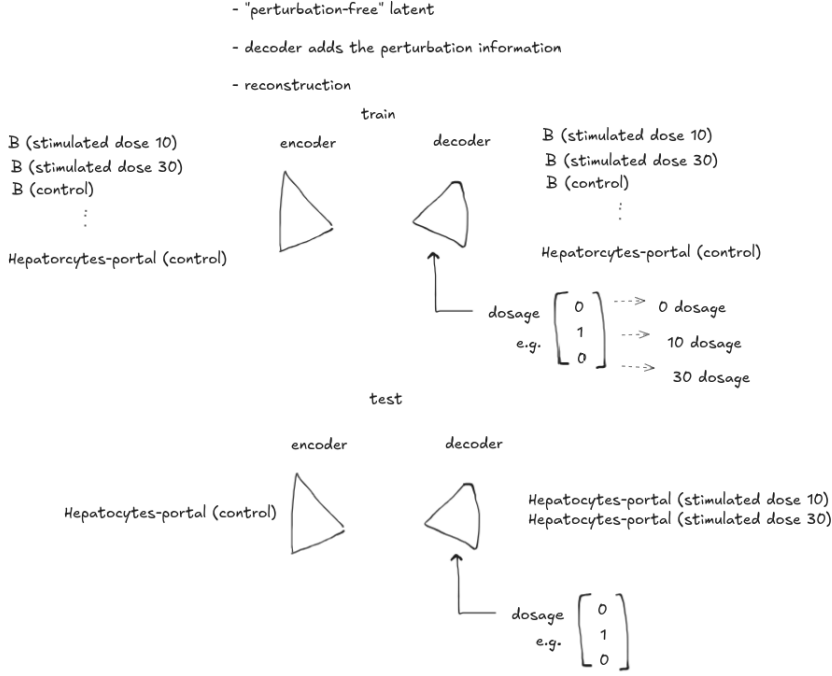


Figure 2: Illustration of the multi-task architecture. The encoder is shared across all tasks, while the decoder is conditioned by the task-specific FiLM layers.

vector. Our architecture is built around an autoencoder, where task-specific conditioning — in our case, the type of perturbation — is integrated via FiLM layers fused into the decoder (MTAe). The modulation parameters γ and β are learned independently for each fusion point.

The loss is the reconstruction loss of the autoencoder, which is the mean squared error between the input and the output of the decoder:

$$\mathcal{L}_{\text{recon}} = \frac{1}{N} \sum_{i=1}^N \|x_i - \hat{x}_i\|^2$$

, where x_i is the input gene expression profile, \hat{x}_i is the reconstructed gene expression profile, and N is the number of samples.

Regarding data splitting, we hold of the stimulated samples of the cell type of interest as a test set. The controlled ones, along with the rest of the cell types in both conditions of control and stimulated, are used for training. Thus, the autoencoder during training attempts to reconstruct the gene expressions while the condition vector is set accordingly to the type of perturbation. The condition vector is one-hot encoded, and given a dataset with N perturbations, its length is $N+1$, including the control condition.

We have explored several variations of this approach, all of which maintain the decoder architecture with the inclusion of FiLM-based conditioning. These variations can be split to three main groups, a) adversarial autoencoders, b) optimal transport, c) Variational Autoencoders (VAEs).

Regarding the first ones, we are aiming to enforce a condition in the latent space via an adversarial loss. The architecture consists of the aforementioned autoencoder scheme with the FiLM layers with the addition of the discriminator. The discriminator aims to differentiate between samples of the latent space and a target distribution, while the encoder aims to fool the discriminator via an adversarial loss to enforce the target distribution in the latent space.

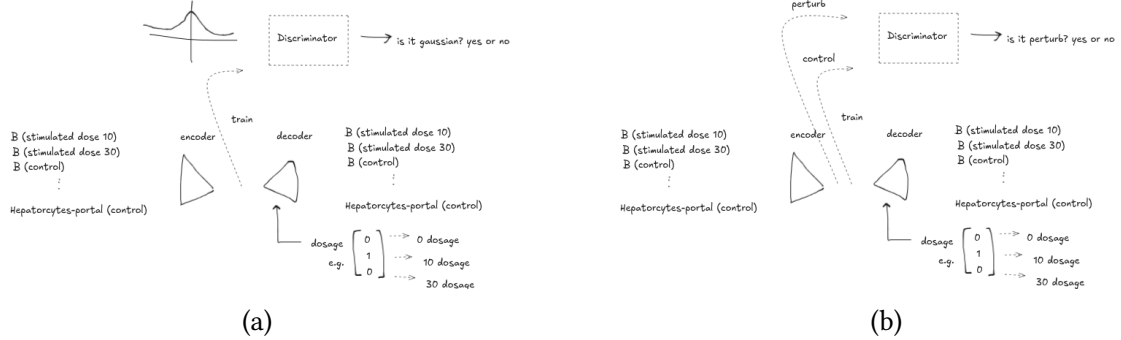


Figure 3: Adversarial autoencoders

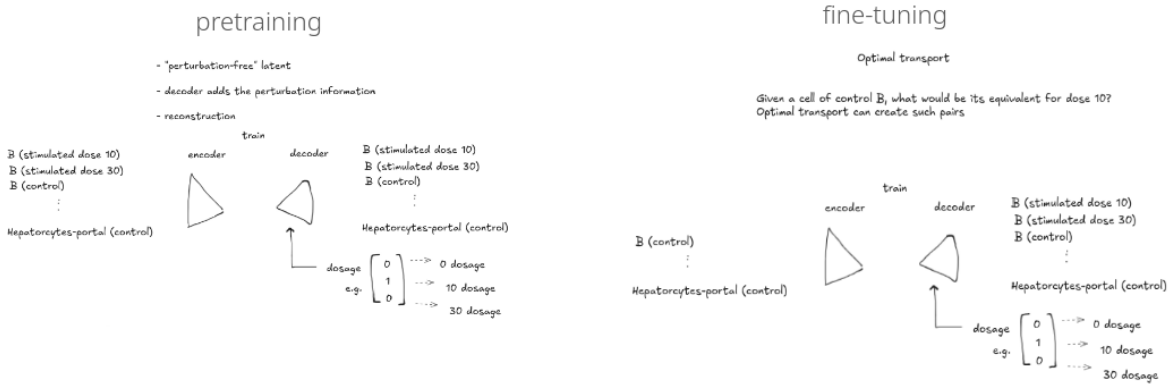


Figure 4: Using optimal transport to fine-tune the MTAe architecture (MTAePlusOT)

For the MTAeAdv architecture, we have attempted to explicitly model a perturbation-free latent space, by using a discriminator to differentiate between the control and perturbed gene expression profiles. In that case, the samples were drawn from the latent space. Similarly, the MTAeAdv architecture aims to enforce a Gaussian distribution in the latent space, by using a prior Gaussian distribution for the discriminator to sample from.

Another set of variations is the inclusion of optimal transport. In single-cell RNA sequencing, we can't sequence the same cell before and after a perturbation, thus we compare distributions since we lack the pair-wise information. To mitigate this, optimal transport can be used to create these pairs, by sampling from the perturbed distribution and matching it with the sample from the control distribution. Using that technique, instead of reconstructing the input, the goal was, given a sample from the controlled distribution, to predict its pair from the perturbed distribution. The loss is the mean squared error between the input and the output of the decoder, defined as:

$$\mathcal{L}_{\text{OT}} = \frac{1}{N} \sum_{i=1}^N \|x_i - \hat{x}_i\|^2$$

, where x_i is the input gene expression profile, \hat{x}_i is the pair from the perturbed gene expression profile, and N is the number of samples. Compared to the previous architectures, the perturbed gene expression profiles are not fed in the network and used only to calculate the loss. This approach is named as MTAeOT. Additionally, we have attempted to pretrain the model with the MTAe architecture and then fine-tune it with the MTAeOT architecture. This approach is named as MTAePlusOT.

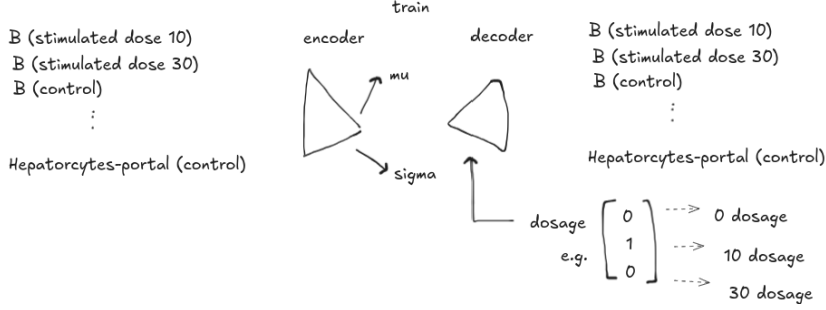


Figure 5: VAE

The last set of variations involves the inclusion of Variational Autoencoders (VAEs). The architecture builds upon the previously described autoencoder framework augmented with FiLM layers, while additionally incorporating a VAE loss to regularize the latent space. The VAE loss is defined as the sum of the reconstruction loss and the Kullback–Leibler (KL) divergence between the learned latent distribution and a standard normal prior:

$$\mathcal{L}_{\text{VAE}} = \mathbb{E}_{q_{\phi}(z|x)}[\log p_{\theta}(x|z)] - D_{\text{KL}}(q_{\phi}(z|x) \parallel p(z))$$

Here, $q_{\phi}(z|x)$ is the encoder’s approximation of the posterior over latent variables, $p_{\theta}(x|z)$ is the decoder’s likelihood of reconstructing the input, and $p(z) \sim \mathcal{N}(0, I)$ is the prior over latent variables. This model is named as MTVae, and as we have described above with the optimal transport use case, we have the MTVaeOT and MTVaePlusOT architectures.

5 Evaluation

We have tested the models on two datasets, one where human peripheral blood mononuclear cells have been stimulated by IFN- β interferon, and a multi-perturbation dataset, where liver cells have been stimulated by multiple doses of tetrachlorodibenzo-p-dioxin (TCDD) *in vivo*.

The models are evaluated on the unseen cell type, given as input the control gene expression fig. 6. Regarding the single perturbation response models, the scGen, scButterfly, scPreGAN and scVIDR’s single-task version, for the multi-perturbation dataset of ten dosages Nault et.al [8, 9], we have trained a dedicated model for each dosage. In these cases, the dataset is consisted of only two conditions the control and the perturbed one for a particular dosage. The performance is measured by comparing the predicted gene expression with the actual one.

For this comparison, we have used the count of differentially expressed genes (DEGs), the R^2 of all the highly variable genes (HVGs), and the top 100 most variable ones. To complement the evaluation, we have calculated a set of five distance metrics (euclidean, edistance, wasserstein, mean pairwise, mmd) to capture the differences between the expected and predicted perturbed gene expressions in a point-wise and distributional manner using pertpy [4].

To address the randomness of the models, we have performed the experiments three times, with three different seeds 1, 2, 19193, and the metrics have been averaged across experiments.

To rank the models, since there could be conflicting cases between metrics, where one model could be better than the other, per model’s metric we averaged them across all the experiments. Then we scaled them to the range of 0-1 with the following formula:

$$\frac{\text{current} - \text{best}}{\text{worst} - \text{best}}$$

, that can track how a metric deviates from the best one. Then we summed all the metrics, giving a score (penalty) to each model. The model with the lowest score is considered the best one.

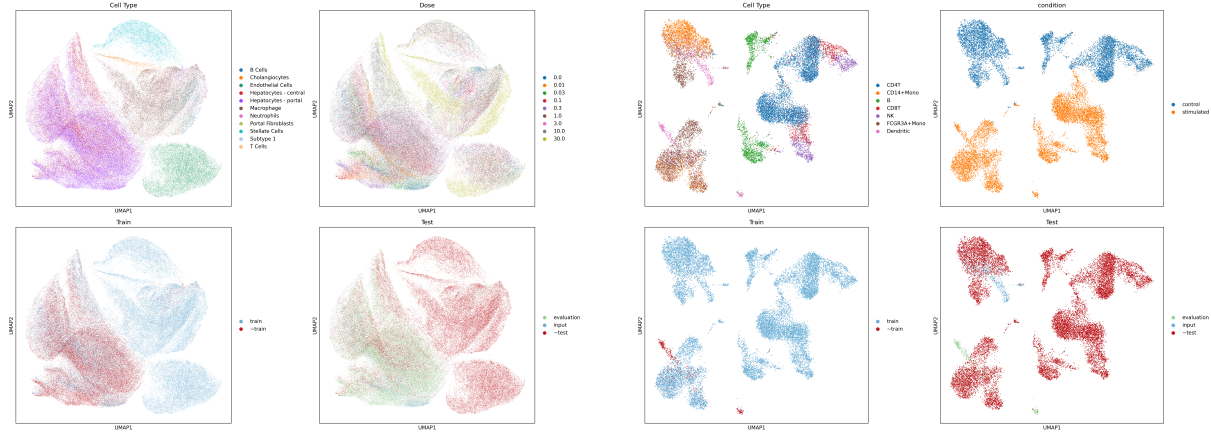


Figure 6: UMAP representations of data split

6 Results

model	DEGs	R_{HVG}^2	R_{HVG20}^2	R_{HVG100}^2	Euc	Was	E-dist	MPD	MMD
MTAe	0.000 (20.341)	0.167 (0.862)	0.267 (0.792)	0.189 (0.833)	0.056 (1.386)	0.603 (1.217)	0.116 (1.116)	0.94 5 (1.050)	0.056 (1.386)
MTAeAdv	0.368 (13.716)	0.388 (0.792)	0.518 (0.725)	0.458 (0.743)	0.025 (1.128)	0.246 (1.091)	0.045 (1.011)	0.426 (1.017)	0.025 (1.128)
MTAeAdvG	0.113 (18.307)	0.339 (0.808)	0.477 (0.736)	0.396 (0.764)	0.029 (1.164)	0.293 (1.107)	0.058 (1.030)	0.607 (1.029)	0.029 (1.164)
MTAeOT	0.650 (8.652)	0.969 (0.608)	0.829 (0.642)	0.916 (0.590)	0.001 (0.925)	0.008 (1.006)	0.005 (0.951)	0.122 (0.998)	0.001 (0.925)
MTAePlusOT	0.657 (8.519)	0.956 (0.613)	0.822 (0.644)	0.897 (0.596)	0.000 (0.917)	0.000 (1.004)	0.002 (0.948)	0.099 (0.996)	0.000 (0.917)
MTVae	0.076 (18.981)	0.339 (0.808)	0.523 (0.724)	0.428 (0.753)	0.025 (1.124)	0.273 (1.100)	0.041 (1.005)	0.413 (1.016)	0.025 (1.124)
MTVaeOT	0.677 (8.163)	0.953 (0.614)	0.830 (0.642)	0.906 (0.593)	0.001 (0.929)	0.016 (1.009)	0.006 (0.952)	0.132 (0.998)	0.001 (0.929)
MTVaePlusOT	0.647 (8.701)	0.948 (0.615)	0.816 (0.645)	0.894 (0.597)	0.000 (0.919)	0.008 (1.006)	0.003 (0.948)	0.112 (0.997)	0.000 (0.919)
scButterfly	0.196 (16.818)	0.553 (0.740)	0.633 (0.694)	0.600 (0.696)	0.008 (0.984)	0.029 (1.014)	0.000 (0.944)	0.000 (0.990)	0.008 (0.984)
scGen	0.781 (6.288)	0.000 (0.915)	0.000 (0.863)	0.000 (0.897)	0.178 (2.408)	0.637 (1.229)	0.299 (1.387)	0.805 (1.041)	0.178 (2.408)
scPreGAN	0.324 (14.511)	1.000 (0.599)	1.000 (0.596)	1.000 (0.562)	0.007 (0.972)	0.042 (1.019)	0.017 (0.969)	0.163 (1.000)	0.007 (0.972)
vidrMult	1.000 (2.352)	0.143 (0.870)	0.099 (0.837)	0.133 (0.852)	1.000 (9.295)	1.000 (1.358)	1.000 (2.425)	1.000 (1.054)	1.000 (9.295)
vidrSingle	0.920 (3.795)	0.191 (0.855)	0.247 (0.797)	0.216 (0.824)	0.061 (1.431)	0.480 (1.174)	0.117 (1.118)	0.544 (1.025)	0.061 (1.431)

Table 1: Nault et al. [8, 9]

model	DEGs	R_{HVG}^2	R_{HVG20}^2	R_{HVG100}^2	Euc	Was	E-dist	MPD	MMD
MTAe	0.000 (75.714)	0.077 (0.946)	0.330 (0.871)	0.140 (0.917)	0.479 (0.488)	0.815 (0.892)	0.506 (0.651)	0.898 (0.949)	0.479 (0.488)
MTAeAdv	0.066 (72.381)	0.026 (0.961)	0.053 (0.955)	0.043 (0.948)	0.032 (0.202)	0.006 (0.604)	0.044 (0.429)	0.110 (0.800)	0.032 (0.202)
MTAeAdvG	0.195 (65.905)	0.168 (0.917)	0.305 (0.878)	0.192 (0.901)	0.503 (0.504)	0.636 (0.828)	0.570 (0.681)	0.689 (0.909)	0.503 (0.504)
MTAeOT	0.688 (41.190)	1.000 (0.657)	1.000 (0.668)	1.000 (0.648)	0.984 (0.811)	0.969 (0.947)	0.990 (0.883)	0.976 (0.963)	0.984 (0.811)
MTAePlusOT	0.768 (37.190)	0.960 (0.670)	0.983 (0.674)	0.970 (0.657)	0.982 (0.810)	0.982 (0.951)	0.983 (0.880)	0.988 (0.966)	0.982 (0.810)
MTVae	0.132 (69.095)	0.088 (0.942)	0.056 (0.954)	0.105 (0.928)	0.125 (0.261)	0.056 (0.621)	0.189 (0.499)	0.114 (0.800)	0.125 (0.261)
MTVaeOT	0.720 (39.571)	0.961 (0.669)	0.969 (0.678)	0.953 (0.663)	0.988 (0.813)	0.993 (0.955)	0.990 (0.883)	0.992 (0.966)	0.988 (0.813)
MTVaePlusOT	0.899 (30.619)	0.987 (0.661)	0.994 (0.670)	0.976 (0.655)	1.000 (0.821)	1.000 (0.958)	1.000 (0.888)	1.000 (0.968)	1.000 (0.821)
scButterfly	0.299 (60.727)	0.251 (0.891)	0.187 (0.914)	0.232 (0.889)	0.140 (0.271)	0.000 (0.601)	0.128 (0.469)	0.000 (0.779)	0.140 (0.271)
scGen	0.868 (32.143)	0.191 (0.910)	0.326 (0.872)	0.290 (0.870)	0.697 (0.627)	0.863 (0.909)	0.744 (0.765)	0.885 (0.946)	0.697 (0.627)
scPreGAN	0.796 (35.750)	0.634 (0.771)	0.376 (0.857)	0.518 (0.799)	0.496 (0.499)	0.248 (0.690)	0.572 (0.682)	0.381 (0.851)	0.496 (0.499)
vidrSingle	1.000 (25.536)	0.000 (0.970)	0.000 (0.971)	0.000 (0.961)	0.000 (0.182)	0.014 (0.606)	0.000 (0.408)	0.096 (0.797)	0.000 (0.182)

Table 2: Kang et al. [6]

model	score	baseline score	distance score
MTAeAdv	0.414099	0.188957	0.225142
MTVae	0.989313	0.381226	0.608087
vidrSingle	1.109933	1.000000	0.109933
scButterfly	1.375249	0.968395	0.406855
MTAe	3.723979	0.546259	3.177721
MTAeAdvG	3.762873	0.860886	2.901987
scPreGAN	4.519561	2.325642	2.193919
scGen	5.559246	1.674543	3.884703
MTVaeOT	8.552051	3.602547	4.949504
MTAeOT	8.590746	3.688019	4.902727
MTAePlusOT	8.598116	3.680466	4.917650
MTVaePlusOT	8.855812	3.855812	5.000000

Table 3: Kang et al. [6]

model	score	baseline score	distance score
scButterfly	2.026767	1.981569	0.045198
MTVae	2.142595	1.365570	0.777025
MTAeAdvG	2.341641	1.324716	1.016925
MTAe	2.398554	0.622043	1.776511
MTAeAdv	2.498785	1.731406	0.767379
vidrSingle	2.837696	1.573823	1.263873
scGen	2.878990	0.781217	2.097772
MTVaePlusOT	3.428329	3.305106	0.123224
MTAePlusOT	3.433685	3.332175	0.101511
MTAeOT	3.500992	3.363746	0.137247
MTVaeOT	3.522593	3.365641	0.156953
scPreGAN	3.559583	3.324068	0.235515
vidrMult	6.375357	1.375357	5.000000

Table 4: Nault et al. [8, 9]

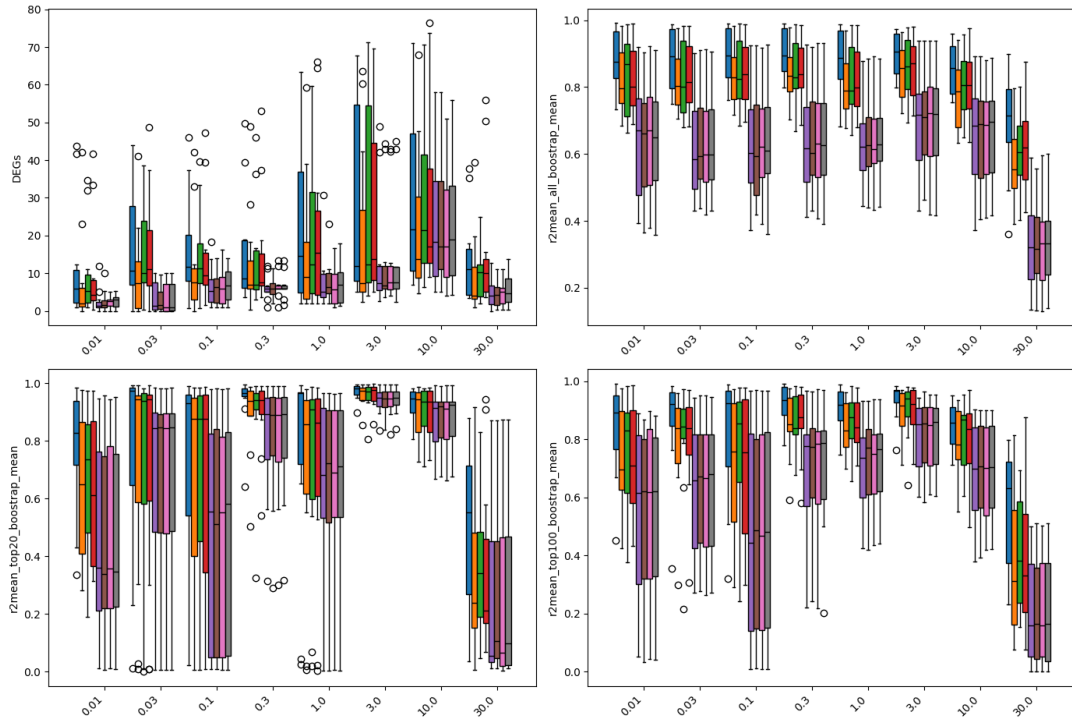


Figure 7: Baseline metrics of multi-task models for the Kang et al. [6] dataset across cell types

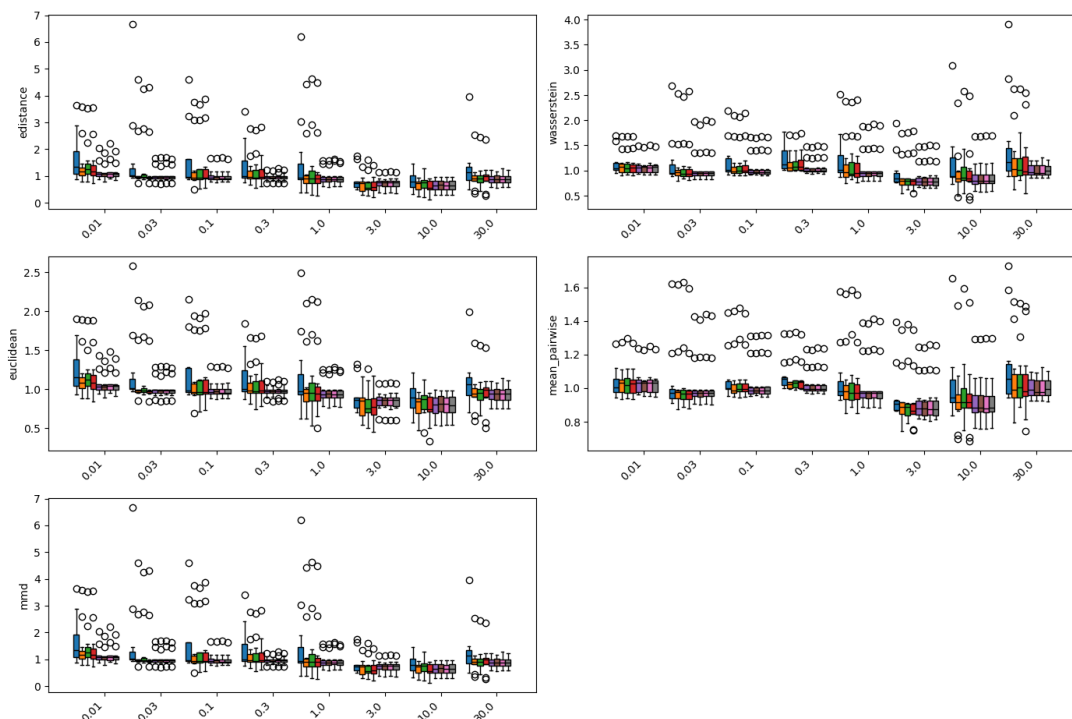


Figure 8: Distance metrics of multi-task models for the Kang et al. [6] dataset across cell types

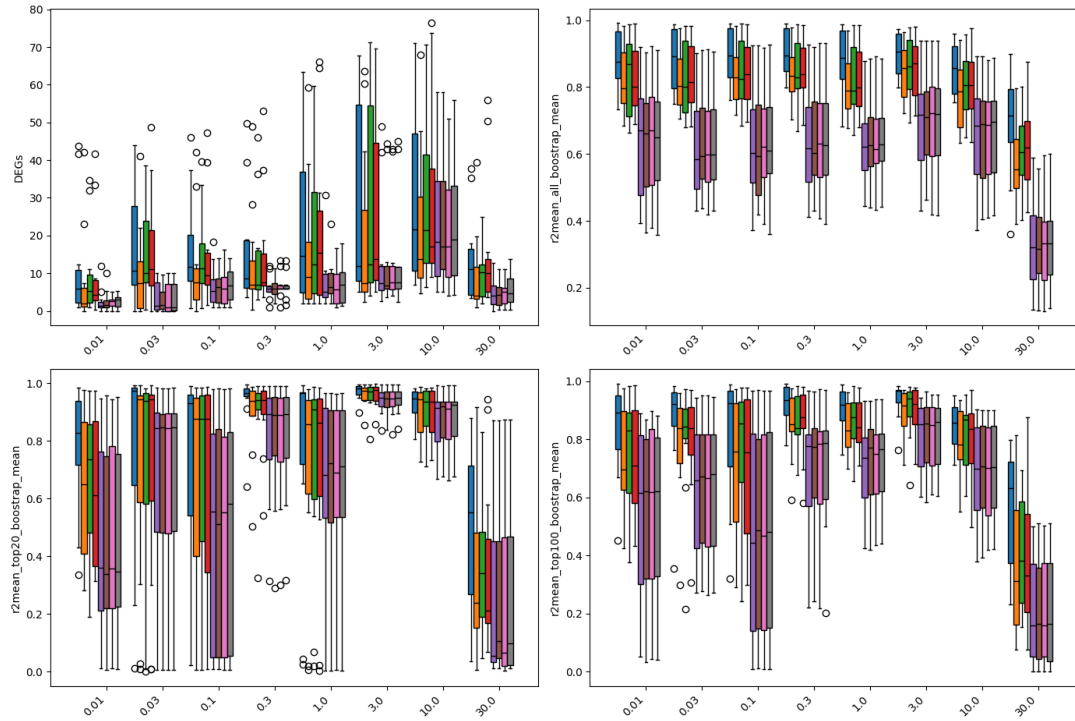


Figure 9: Baseline metrics of multi-task models for the Nault et al. [8, 9] dataset across dosages

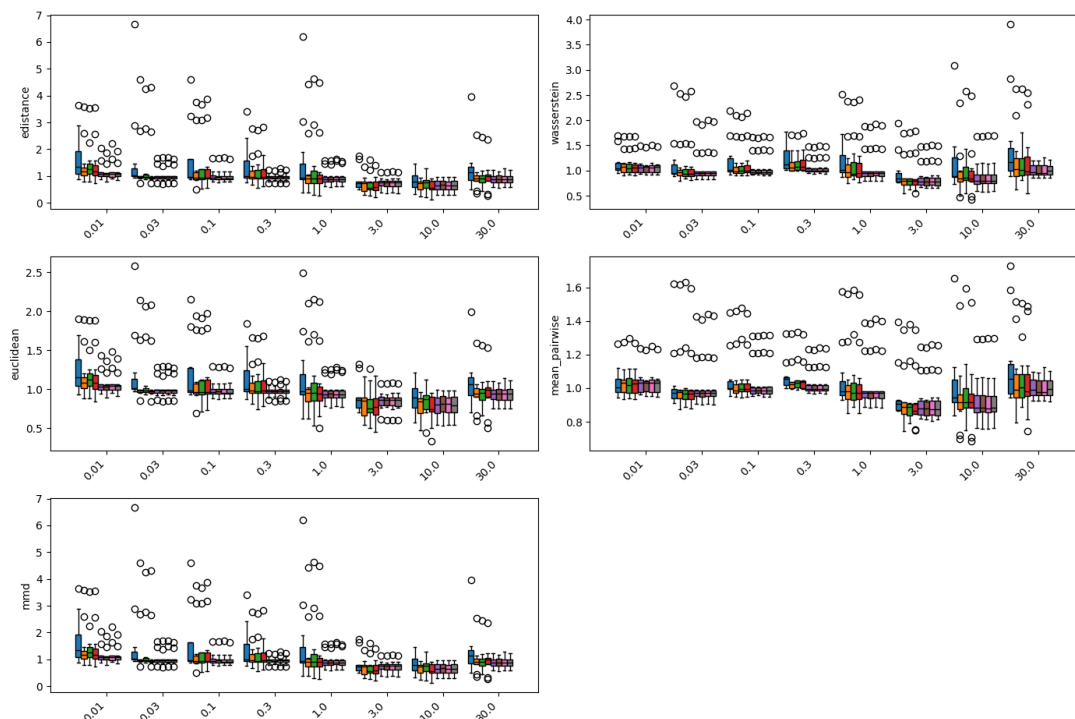
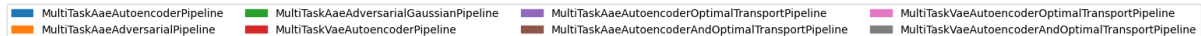


Figure 10: Distance metrics of multi-task models for the Nault et al. [8, 9] dataset across dosages

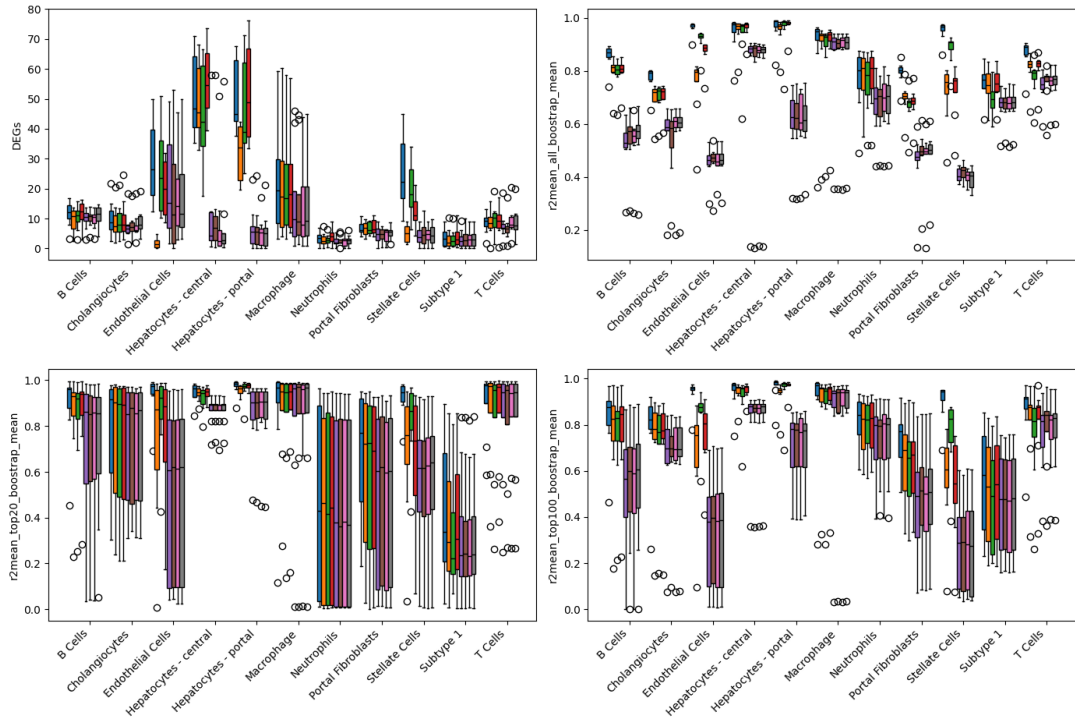


Figure 11: Baseline metrics of multi-task models for the Nault et al. [8, 9] dataset across cell types

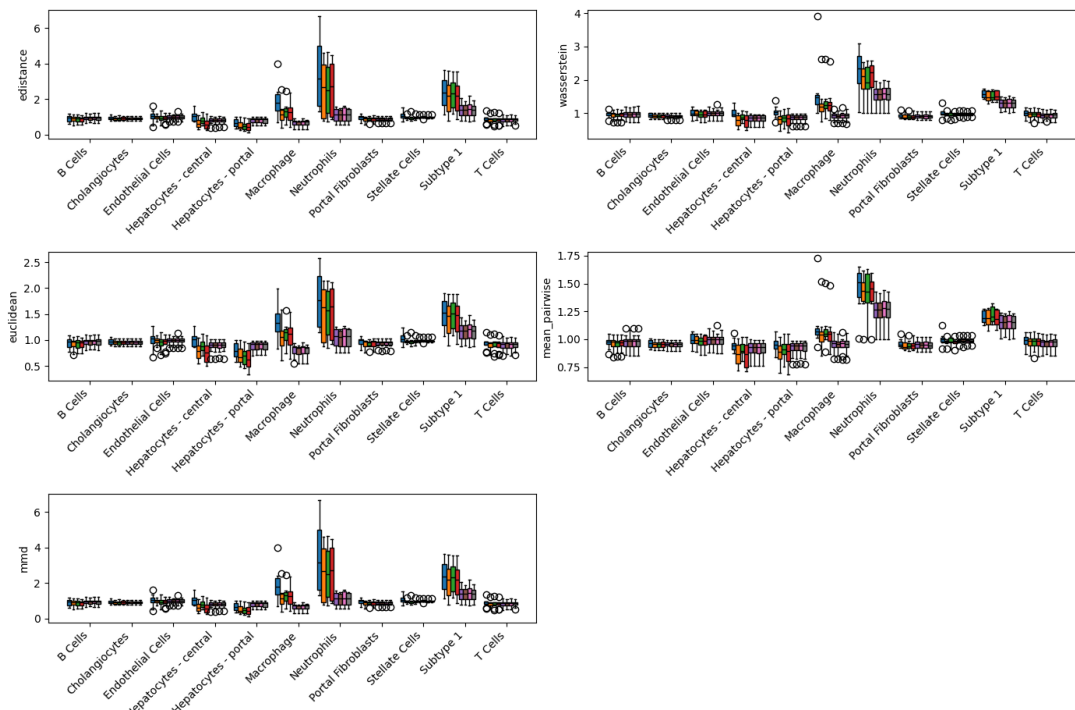
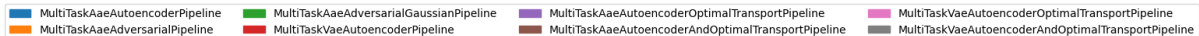


Figure 12: Distance metrics of multi-task models for the Nault et al. [8, 9] dataset across cell types

6.1 Knowledge transfer

which tasks and why they are important?

6.2 TODO

- batch effect
- interpretability
- explainability
- integration with multiple omics

7 Conclusions

8 Future work

A Ακρωνύμια και συντομογραφίες

LAN Local Area Network

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