MIXTURE-OF-EXPERTS VARIATIONAL AUTOEN-CODER FOR CLUSTERING AND GENERATING FROM SIMILARITY-BASED REPRESENTATIONS

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Abstract

Clustering high-dimensional data, such as images or biological measurements, is a long-standing problem and has been studied extensively. Recently, Deep Clustering gained popularity due to the non-linearity of neural networks, which allows for flexibility in fitting the specific peculiarities of complex data. Here we introduce the Mixture-of-Experts Similarity Variational Autoencoder (MoE-Sim-VAE), a novel generative clustering model. The model can learn multi-modal distributions of high-dimensional data and use these to generate realistic data with high efficacy and efficiency. MoE-Sim-VAE is based on a Variational Autoencoder (VAE), where the decoder consists of a Mixture-of-Experts (MoE) architecture. This specific architecture allows for various modes of the data to be automatically learned by means of the experts. Additionally, we encourage the lower dimensional latent representation of our model to follow a Gaussian mixture distribution and to accurately represent the similarities between the data points. We assess the performance of our model on synthetic data, the MNIST benchmark data set, and a challenging real-world task of defining cell subpopulations from mass cytometry (CyTOF) measurements on hundreds of different datasets. MoE-Sim-VAE exhibits superior clustering performance on all these tasks in comparison to the baselines and we show that the MoE architecture in the decoder reduces the computational cost of sampling specific data modes with high fidelity.

1 INTRODUCTION

Clustering has been studied extensively (Aljalbout et al., 2018; Min et al., 2018) in machine learning. Recently, many Deep Clustering approaches were proposed, which modified (Variational) Autoencoder ((V)AE) architectures (Min et al., 2018; Zhang et al., 2017) or with varying regularization of the latent representation (Dizaji et al., 2017; Jiang et al., 2017; Yang et al., 2017; Fortuin et al., 2019).

Reconstruction error usually drives the definition of the latent representation learned from an AE or VAE. The representation for AE models is unconstrained and typically places data objects close to each other according to an implicit similarity measure that also yields favorable reconstruction error. In contrast, VAE models regularize the latent representation such that the represented inputs follow a certain variational distribution. This construction enables sampling from the latent representation and data generation via the decoder of a VAE. Typically, the variational distribution is assumed standard Gaussian, but for example, Jiang et al. (2017) introduced a mixture of Gaussian variational distribution for clustering purposes.

A key component of clustering approaches is the choice of similarity metric for the considered data objects which we try to group (Irani et al., 2016). Such similarity metrics are either defined *a priori* or learned from the data to specifically solve classification tasks via a Siamese network architecture (Chopra et al., 2005). Dimensionality reduction approaches, such as UMAP (McInnes et al., 2018) or t-SNE (van der Maaten & Hinton, 2008), allow to specify a similarity metric for projection and thereby define the data separation in the inferred latent representation.

In this work, we introduce the Mixture-of-Experts Similarity Variational Autoencoder (MoE-Sim-VAE), a new deep architecture that performs similarity-based representation learning, clustering of the data and generation of data from each specific data mode. Due to a combined loss function,



Figure 1: Overview of the proposed model MoE-Sim-VAE. Data (in panel A) gets encoded via a encoder network (B) into a latent representation (C) which is trained to be a mixture of standard Gaussians. Via a clustering network (G), which is trained to reconstruct a user-defined similarity matrix (F), the encoded samples get assigned to the data mode-specific decoder subnetworks (which we call experts) in the MoE Decoder (D). The experts reconstruct the original input data and can be used for data generation when sampling from the variational distribution (E).

it can be jointly optimized. We assess the scope of the model on synthetic data and we present superior clustering performance on MNIST. Moreover, in an ablation study, we show the efficiency and precision of MoE-Sim-VAE for data generation purposes in comparison to the most related state-of-the-art method (Jiang et al., 2017). Finally, we show an application of MoE-Sim-VAE on a real-world clustering problem in biology on multiple datasets.

Our main contributions are to

- Develop a novel autoencoder architecture for
 - similarity-based representation learning
 - unsupervised clustering
 - accurate and efficient data generation
- Embed the Mixture-of-Expert architecture into a Variational Autoencoder setup to train a separate generator for each data mode
- Show superior clustering performance of the model on benchmark dataset and real-world biological data

2 MIXTURE-OF-EXPERTS SIMILARITY VARIATIONAL AUTOENCODER

Here we introduce the Mixture-of-Experts Similarity Variational Autoencoder (MoE-Sim-VAE, Figure 1). The model is based on the Variational Autoencoder (Kingma & Welling, 2014). While the encoder network is shared across all data points, the decoder of the MoE-Sim-VAE consists of a number of K different subnetworks, forming a Mixture-of-Experts architecture (Shazeer et al., 2017). Each subnetworks constitutes a generator for a specific data mode and is learned from the data.

The variational distribution over the latent representation is defined to be a mixture of multivariate Gaussians, first introduced by Jiang et al. (2017). In our model, we aim to learn the mixture components in the latent representation to be standard Gaussians

$$\boldsymbol{z} \sim \sum_{k=0}^{K} \omega_k \mathcal{N}(\boldsymbol{\mu}_k, \boldsymbol{I})$$
 (1)

where ω_k are mixture coefficients, μ_k are the means for each mixture component, I is the identity matrix and K is the number of mixture components. The dimension of the latent representation z needs to be defined to suit the demands of Gaussian mixtures which have limitations in higher dimensions (Bishop, 1995). Similar to optimizing an Evidence Lower Bound (ELBO), we penalize the latent representation via the reconstruction loss of the data $\mathcal{L}_{reconst}$ and by using the Kullback-Leibler (KL) divergence for multivariate Gaussians (Jiang et al., 2017) on the latent representation

$$\mathcal{L}_{KL} = D_{KL}(\mathcal{N}_0, \mathcal{N}_1) = \frac{1}{2} \{ tr(\mathbf{\Sigma}_1^{-1} \mathbf{\Sigma}_0) + (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_0)^T \mathbf{\Sigma}_1^{-1} (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_0) - k + ln \frac{|\mathbf{\Sigma}_1|}{|\mathbf{\Sigma}_0|} \}$$
(2)

where k is a constant, $\mathcal{N}_0 \sim \mathcal{N}(\boldsymbol{\mu}_0, \boldsymbol{\Sigma}_0 = \boldsymbol{I})$, and \boldsymbol{I} is the identity matrix. Further, $\mathcal{N}_1 \sim \mathcal{N}(\boldsymbol{\mu}_1, \boldsymbol{\Sigma}_1 = diag(\sigma_j))$, where σ_j for j = 1, ..., D, for a number of dimensions D, is estimated from the samples of the latent representation. Finally, we assume $\boldsymbol{\mu}_0 = \boldsymbol{\mu}_1$ resulting in the following simplified objective

$$\mathcal{L}_{KL} = D_{KL}(\mathcal{N}_0, \mathcal{N}_1) = \frac{1}{2} \{ tr(\boldsymbol{\Sigma}_1^{-1} \boldsymbol{\Sigma}_0) - k + ln \frac{|\boldsymbol{\Sigma}_1|}{|\boldsymbol{\Sigma}_0|} \} ,$$
(3)

to penalize exclusively the covariance of each cluster. It remains to define the reconstruction loss $\mathcal{L}_{reconst}$, where we choose a Binary Cross-Entropy

$$\mathcal{L}_{reconst} = \sum_{i}^{N} \sum_{d}^{D} x_{i,d} \log(x_{i,d}^{reconst})$$
(4)

between the original data x (scaled between 0 and 1) and the reconstructed data $x^{reconst}$, where i iterates the batch size N and d the dimensions of the data D. Finally the loss for the VAE part is defined by

$$\mathcal{L}_{VAE} = \mathcal{L}_{reconst} + \pi_1 \mathcal{L}_{KL} \tag{5}$$

with a weighting coefficient π_1 which can be optimized as a hyperparameter.

SIMILARITY CLUSTERING AND GATING OF LATENT REPRESENTATION

Training of a data mode-specific generator expert requires samples from the same data mode. This necessitates to solve a clustering problem, that is, mapping the data via the latent representation into K clusters, each corresponding to one of the K generator experts. We solve this clustering problem via a clustering network, also referred to as gating network for MoE models. It takes as input the latent representation z_i of sample i and outputs probabilities $p_{ik} \in [0, 1]$ for clustering sample i into cluster k. According to this cluster assignment, sample i is then gated to expert $k = \operatorname{argmax}_k p_{ik}$ for each sample i. We further define the cluster centers μ_k for $k \in \{1, \ldots, K\}$ similar as in the Expectation Maximization (EM) algorithm for Gaussian Mixture models (Bishop, 2006) as

$$\boldsymbol{\mu}_{k} = \frac{1}{N_{k}} \sum_{i=1}^{N} p_{ik} \boldsymbol{z}_{i} , \qquad (6)$$

where N_k is the absolute number of data points assigned to cluster k based on highest probability p_{ik} for each sample i = 1, ..., N. The Gaussian mixture distributed latent representation (via KL loss in Equation 3) is motivation for the empirical computation of the cluster means and further, similar as in the EM algorithm, it allows iterative optimization of the means of the Gaussians. We train the clustering network to reconstruct a data-driven similarity matrix S, using the Binary Cross-Entropy

$$\mathcal{L}_{Similarity} = \sum_{i}^{N} \sum_{j}^{N} \boldsymbol{S}_{i,j} \log((\boldsymbol{P}\boldsymbol{P}^{T})_{i,j})$$
(7)

to minimize the error in $PP^T \approx S$, with $P := \{p_{ik}\}_{i \in \{1,...,N\},k \in \{1,...,K\}}$ where N is the number of samples (e.g., batch size). Intuitively, PP^T approximates the similarity matrix S since values in PP^T are only close to 1 when similar data objects are assigned to the same cluster, similar to the entries in the adjacency similarity matrix S. This similarity matrix is derived in an unsupervised way in our experiments (e.g. UMAP projection of the data and k-nearest-neighbors or distance thresholding to define the adjacency matrix for the batch), but can also be used to include weaklysupervised information (e.g., knowledge about diseased vs. non-diseased patients). If labels are available, the model could even be used to derive a latent representation with supervision. The similarity feature in MoE-Sim-VAE thus allows to include prior knowledge about the best similarity measure on the data.

Moreover, we apply the DEPICT loss from Dizaji et al. (2017), to improve the robustness of the clustering. For the DEPICT loss, we additionally propagate a noisy probability \hat{p}_{ik} through the clustering network using dropout after each layer. The goal is to predict the same cluster for both, the noisy \hat{p}_{ik} and the clean probability p_{ik} (without applying dropout). Dizaji et al. (2017) derived as objective function a standard cross-entropy loss

$$\mathcal{L}_{DEPICT} = -\frac{1}{N} \sum_{i=0}^{N} \sum_{k=0}^{K} q_{ik} \log(\hat{p}_{ik})$$
(8)

whereby q_{ik} is computed via the auxiliary function

$$q_{ik} = \frac{p_{ik} / (\sum_{i'} p_{i'k})^{\frac{1}{2}}}{\sum_{k'} p_{ik'} / (\sum_{i'} p_{i'k'})^{\frac{1}{2}}}$$
(9)

where we refer to Dizaji et al. (2017) for exact derivation. The DEPICT loss encourages the model to learn invariant features from the latent representation for clustering with respect to noise (Dizaji et al., 2017). Looking at it from a different perspective, the loss helps to define a latent representation which has those invariant features to be able to reconstruct the similarity and therefore the clustering correctly. The complete clustering loss function $\mathcal{L}_{Clustering}$ is then defined by

$$\mathcal{L}_{Clustering} = \mathcal{L}_{Similarity} + \pi_2 \mathcal{L}_{DEPICT} \tag{10}$$

with a mixture coefficient π_2 which can be optimized as a hyperparameter.

MOE-SIM-VAE LOSS FUNCTION

Finally, the MoE-Sim-VAE model loss is defined by

$$\mathcal{L}_{MoE-Sim-VAE} = \underbrace{\mathcal{L}_{VAE}}_{\mathcal{L}_{reconst} + \pi_1 \mathcal{L}_{KL}} + \underbrace{\mathcal{L}_{Clustering}}_{\mathcal{L}_{Similarity} + \pi_2 \mathcal{L}_{DEPICT}}$$
(11)

which consists of the two main loss functions \mathcal{L}_{VAE} , acting as a regularization for the latent representation, and $\mathcal{L}_{Clustering}$, which helps to learn the mixture components based on an *a priori* defined data similarity. The model objective function $\mathcal{L}_{MoE-Sim-VAE}$ can then be optimized end-to-end to train all parts of the model.

3 Related Work

(V)AEs have been extensively used for clustering (Xie et al., 2016; Dizaji et al., 2017; Li et al., 2017; Yang et al., 2017; Saito & Tan, 2017; Chen et al., 2017; Aljalbout et al., 2018; Fortuin et al., 2019). The most related approaches to MoE-Sim-VAE are Jiang et al. (2017) and Zhang et al. (2017).

Jiang et al. (2017) introduced the VaDE model, comprising a mixture of Gaussians as underlying distribution in the latent representation of a Variational Autoencoder. Optimizing the Evidence Lower Bound (ELBO) of the log-likelihood of the data can be rewritten to optimize the reconstruction loss of the data and KL divergence between the variational posterior and the mixture of Gaussians prior. Jiang et al. (2017) motivate the use of to two separate networks for reconstruction and the generation process of the model. Further, to effectively generate images from a specific data mode and to increase image quality, the sampled points have to surpass a certain posterior threshold and are otherwise rejected. This leads to an increased computational effort. The MoE Decoder of our model, which is used for both reconstruction and generation, does not need such a threshold, as we discuss in more detail in Section 4.2.1.

Zhang et al. (2017) have introduced a mixture of autoencoders (MIXAE) model. The latent representation of the MIXAE is defined as the concatenation of the latent representation vectors of each single autoencoder in the model. Based on this concatenated latent representation, a Mixture

Table 1: Performance comparison of our method MoE-Sim-VAE with several published methods.
The Table is mainly extracted from Aljalbout et al. (2018) and complemented with results of interest.
(" - ": metric not reported)

METHOD	NMI	ACC
JULE, Yang et al. (2016b)	0.915	- 10
CCNN, Hsu & Lin (2017)	0.876	-
DEC, Xie et al. (2016)	0.8	0.843
DBC, Li et al. (2017)	0.917	0.964
DEPICT, Dizaji et al. (2017)	0.916	0.965
DCN, Yang et al. (2017)	0.81	0.83
Neural Clustering, Saito & Tan (2017)	-	0.966
UMMC, Chen et al. (2017)	0.864	-
VaDE, Jiang et al. (2017)	-	0.945
TAGnet, Wang et al. (2016)	0.651	0.692
IMSAT, Hu et al. (2017)	-	0.984
Aljalbout et al. (2018)	0.923	0.961
MIXAE, Zhang et al. (2017)	-	0.945
Spectral clustering, Shaham et al. (2018)	0.754	0.717
SpectralNet (input space, Euclidean dist.), Shaham et al. (2018)	$0.687 {\pm} 0.004$	$0.622 {\pm} 0.008$
SpectralNet (input space, Siamese dist.), Shaham et al. (2018)	$0.884{\pm}0.02$	$0.826 {\pm} 0.03$
SpectralNet (code space, Euclidean dist.), Shaham et al. (2018)	$0.814{\pm}0.008$	$0.800 {\pm} 0.003$
SpectralNet (code space, Siamese dist.), Shaham et al. (2018)	$0.924{\pm}0.001$	$0.971 {\pm} 0.001$
MoE-Sim-VAE (proposed)	0.935	0.975

Assignment Network predicts probabilities which are used in the Mixture Aggregation to form the output of the generator network. Each AE model learns the manifold of a specific cluster, similarly to our MoE Decoder. However, MIXAE does not optimize a variational distribution, such that generation of data from a distribution over the latent representation is not possible, in contrast to the MoE-Sim-VAE (Figure 2).

4 EXPERIMENTS

We evaluate the MoE-Sim-VAE using synthetic data and the MNIST data set of handwritten digits (LeCun et al., 1998) for clustering and data generation. Furthermore, we performed an ablation study to demonstrate the importance of the MoE Decoder. Finally, we present experiments on a real-world application of defining cellular subpopulations from mass cytometry measurements (Bandura et al., 2009) of multiple publicly available datasets (Weber & Robinson, 2016; Bodenmiller et al., 2012). Model implementation details are reported in the appendix in section A.1

We found that our model achieves superior clustering performance compared to other models on synthetic, MNIST and real-world datasets. Moreover, we show that MoE-Sim-VAE can more effectively and efficiently generate data from specific modes in comparison to other methods.

4.1 EVALUATION OF MOE-SIM-VAE ON SYNTHETIC DATA

We evaluated our model using data sampled from a 100-dimensional multivariate Gaussian with equal mixture weights for each component. We tested two aspects of our model: Firstly, we evaluated up to how many clusters our model can fit well. Therefore, we sampled data from distributions with up to a hundred mixture components. For this experiment, we assume knowledge of the true number of clusters in the data for both methods, MoE-Sim-VAE and GMMs. Secondly, we tested if our model is able to identify the true number of clusters in the data. The similarity matrix S was defined as an adjacency matrix over the data items. Adjacency indicators were based on projecting the data via dimensionality reduction with UMAP (McInnes et al., 2018) and selecting neighbors according to a distance threshold. Details on model parameters can be found in Section A.1.1.

MoE-Sim-VAE performs better or comparable to the baseline for the number of clusters of up to 40 (Figure A1a). The model predicts with a close to perfect F-measure until reaching a true number



Figure 2: Generation of MNIST digit images. Data points from the latent representation were sampled from the variational distribution (A) which is learned to be a mixture of standard Gaussians and then clustered and gated (B) to the data-mode-specific experts of the MoE Decoder (C). (D) All samples from the variational distribution were correctly classified and therefore also correctly gated.

of clusters of 30. Within the range of true number of clusters from 30 to 40, the model performs comparable to GMMs. Further, MoE-Sim-VAE learns the true number of clusters on its own (Figure A1b). For up to 23 components in the data, MoE-Sim-VAE learns the true number of clusters even when defining a model with K = 40 experts in the MoE Decoder. This suggests that the model is robust to misspecification regarding the number of experts.

4.2 UNSUPERVISED CLUSTERING, EMBEDDING AND DATA GENERATION OF MNIST

We trained a MoE-Sim-VAE model on images from MNIST. We compared our model against multiple models which were recently reviewed in Aljalbout et al. (2018), and specifically against VaDE (Jiang et al., 2017) which shares similar properties with MoE-Sim-VAE (see Sec 3).

We compare the models with the Normalized Mutual Information (NMI) criterion but also classification accuracy (ACC) (Table 1). The MoE-Sim-VAE outperforms the other methods w.r.t. clustering performance when comparing NMI and achieves the second-best result when comparing ACC. Note that we used the number of experts k = 10 in our model to fit the existing number of digits in MNIST. Regarding the similarity measure, we decided to use as similarity a UMAP projection (McInnes et al., 2018) of MNIST and then apply k-nearest-neighbors of each sample in a batch. More details on the model are reported in Section A.1.2. In an ablation study we show the importance of the similarity matrix to create a clear separation of the different digits in the latent representation (Figure A4)

In addition to the clustering network, we can make use of the latent representation for image generation purposes. The latent representation is trained as a mixture of standard Gaussians. The means of these Gaussians are the centers of the clusters trained via the clustering network. Therefore, the variational distribution can be sampled from and gated to the cluster-specific expert in the MoE-decoder. The expert then generates new data points for the specific data mode. Results and the schematic are displayed in Figure 2 and in more detail and with greater sample size in the Appendix in Figure A2.

4.2.1 Why does a MoE Decoder actually matter?

In an ablation study, we compare the two models MoE-Sim-VAE and VaDE (Jiang et al., 2017) on generating MNIST images with the request for a specific digit. The goal is to show that a MoE decoder, as proposed in our model, is beneficial. We focus our comparison to VaDE since this model, as the MoE-Sim-VAE, resorts to a mixture of Gaussian latent representation but differs in generating images by means of a single decoder network instead of a Mixture-of-Expert decoder network. The rationale for our design choice is to ensure that smaller sub-networks learn to reproduce and generate specific modes of the data, in this case of specific MNIST digits.

To show that both models' latent representations are separating the different clusters well, we computed the Maximum Mean Discrepancy (MMD), defined in Section A.1.2. The MMD can be interpreted as a distance between distributions computed based on samples drawn from these distributions. The heatmaps of the MMDs for VaDE and MoE-Sim-VAE as well as an UMAP projection of the latent representation colored with the mixture component confirm visually the separation of the clusters in the latent representations of both models (Fig. A3). As a result, we can conclude that both latent representations can separate the clusters of respective digits well, such that the decoder gets well-defined samples to generate the requested digit. Therefore, the main difference of generating specific digits arises in the decoder/generator networks.

We evaluated the importance of the MoE-Decoder to (1) accurately generate requested digits and (2) be efficient in generating requested digits. Specifically, we sampled 10,000 points from each mixture component in the latent representation, generated images, and used the model's internal clustering to assign a probability to which digits were generated. To generate correct and high-quality images with VaDE, the posterior of the latent representation needs to be evaluated for each sample. This was done for the different thresholds $\phi \in [0.0, 0.1, 0.2, \dots, 0.9, 0.999]$. The default threshold Jiang et al. (2017) used was $\phi = 0.999$. Instead of thresholding the latent representation, we ran the generation process for MoE-Sim-VAE for each threshold with the same settings. To generate images from VaDE we used the Python implementation¹ and model weights publicly available from Jiang et al. (2017).

As a result of this analysis we report a confusion matrix for MoE-Sim-VAE in Figure A6, the confusion matrices for each threshold for VaDE in Figure A7, the accuracy of generating a requested digit and the number of runs required in Figure A5. In summary, one can see that the MoE-Sim-VAE generates digits more accurately with fewer resources required. This can especially be seen when comparing the number of iterations required to fulfill the default posterior threshold of 0.999. VaDE needs nearly 2 million iterations to find samples that fulfill the aforementioned threshold criterion whereas the MoE-Sim-VAE only requires 10,000 for a comparable sample accuracy. In comparison the mean accuracy over all thresholds for MoE-Sim-VAE is 0.970, whereas VaDE reaches on average 0.944. VaDE reaches a maximum accuracy of 0.995, which costs the aforementioned 2 million iterations for generating 100,000 images, whereas MoE-Sim-VAE reaches a maximum accuracy of 0.971 with 100,000 runs, without accounting for a systematic generating/clustering error (confusing 5 and 8) of MoE-Sim-VAE which can be seen in the confusion matrix in Figure A6.

4.3 LEARNING CELL TYPE COMPOSITION IN PERIPHERAL BLOOD MONONUCLEAR CELLS USING CYTOF MEASUREMENTS

In the following, we want to show representation learning performance on a real-world problem in biology. Specifically, we focus on cell type definition from single-cell measurements. Cytometry by time-of-flight mass spectrometry (CyTOF) (Bandura et al., 2009) is a state-of-the-art technique allowing measurement of up to 1,000 cells per second and in parallel over 40 protein markers of the cells (Kay et al., 2013). Defining biologically relevant cell subpopulations by clustering this data is a common learning task (Aghaeepour et al., 2013; Weber & Robinson, 2016).

Many methods have been developed to tackle the problem introduced above and were compared on four publicly available datasets in Weber & Robinson (2016). The best out of 18 methods were FlowSOM (Gassen et al., 2015), PhenoGraph (Levine et al., 2015) and X-shift (Samusik et al., 2016). These are based on k-nearest-neighbors heuristics, either defined from a spanning graph or from estimating the data density. In contrast to these methods, MoE-Sim-VAE can map new cells into the latent representation, assign probabilities for cell types and infer an interpretable latent representation allowing intuitive downstream analysis by domain experts.

We applied MoE-Sim-VAE to the same datasets as in Weber & Robinson (2016) and achieve superior results in classification using the F-measure (Equation 12) in three out of four datasets. Similarly as in Weber & Robinson (2016) we trained MoE-Sim-VAE 30 times and report in Table 2 (adopted from Weber & Robinson (2016)) the means across all runs. The reproducibility of our model for each dataset can be seen in Figure A8.

Further, we trained a MoE-Sim-VAE model on 268 datasets from Bodenmiller et al. (2012) (more details on the data in A.1.3), and achieve superior classification results of cell subpopulations in the data when comparing to state-of-the-art methods in this field (PhenoGraph, X-Shift, FlowSOM).

¹https://github.com/slim1017/VaDE

Table 2: Comparison of MoE-Sim-VAE performance to competitor methods in defining cell type composition in CyTOF measurements. The results in the table are extracted from the review paper of Weber & Robinson (2016), where 18 methods are compared on four different datasets. Our model outperforms the baselines on four out of five data sets.

Method	Levine_32dim	Levine_13dim	Samusik_01	Samusik_all
ACCENSE	0.494	0.358	0.517	0.502
ClusterX	0.682	0.474	0.571	0.603
DensVM	0.66	0.448	0.239	0.496
FLOCK	0.727	0.379	0.608	0.631
flowClust	NA	0.416	0.612	0.61
flowMeans	0.769	0.518	0.625	0.653
flowMerge	NA	0.247	0.452	0.341
flowPeaks	0.237	0.215	0.058	0.323
FlowSOM	0.78	0.495	0.707	0.702
FlowSOM_pre	0.502	0.422	0.583	0.528
immunoClust	0.413	0.308	0.552	0.523
kmeans	0.42	0.435	0.65	0.59
PhenoGraph	0.563	0.468	0.671	0.653
Rclusterpp	0.605	0.465	0.637	0.613
SamSPECTRAL	0.512	0.253	0.263	0.138
SPADE	NA	0.127	0.169	0.13
SWIFT	0.177	0.179	0.202	0.208
Xshift	0.691	0.47	0.679	0.657
MoE-Sim-VAE (proposed)	0.70	0.68	0.76	0.74

Exact results can be seen in Table A1 or visualized in Figure 3. More details on the MoE-Sim-VAE setting used for all results on CyTOF data are reported in the appendix (Section A.1.3).

5 CONCLUSION

Our MoE-Sim-VAE model can infer similarity-based representations, perform clustering tasks, and efficiently as well as accurately generate high-dimensional data. The training of the model is performed by optimizing a joint objective function consisting of data reconstruction, clustering, and KL loss, where the latter regularizes the latent representation. On synthetic data, we have shown the strengths and limitations of the model. On the benchmark dataset of MNIST, we presented superior clustering performance and the efficiency and accuracy of MoE-Sim-VAE in generating high-dimensional data. On the biological real-world task of defining cell subpopulations in complex single-cell data, we show superior clustering performances compared to state-of-the-art methods on over 270 datasets and therefore demonstrate MoE-Sim-VAE's real-world usefulness.

Future work might include to add adversarial training to the MoE decoder, which could improve image generation to create even more realistic images. Also, specific applications might benefit from replacing the Gaussian with a different mixture model. So far the MoE-Sim-VAE's similarity measure has to be defined by the user. Relaxing this requirement and allowing for learning a useful similarity measure automatically for inferring latent representations will be an interesting extension to explore. This could be useful in a weakly-supervised setting, which often occurs for example in clinical data consisting of healthy and diseased patients. Minor details between a healthy and diseased patient might make a huge difference and could be learned from the data using neural networks.

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Figure 3: Comparison of MoE-Sim-VAE to the most popular competitor methods on defining cell types in peripheral blood mononuclear cell data via CyTOF measurements. On the x-axis different inhibitor treatments are listed whereas the y-axis reports the respective F-measure, defined in Equation 12, as performance measure of the methods. Each violin plot represents a run on a different inhibitor with multiple wells, whereas the line connects the means of the performance on the specific inhibitor.

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A APPENDIX

A.1 EXPERIMENTAL DETAILS

In the following sections we provide more details on model implementations, metrics used and additional result figures for the experiments described in the main text.

A.1.1 EVALUATION OF MOE-SIM-VAE ON SYNTHETIC DATA

Model and training details:

- number of experts: $\{2, \ldots, 40\}$
- batch size: 512
- code size: 10
- Number of iterations: 5000
- activation function; elu
- loss coefficient data reconstruction: 0.487
- loss coefficient clustering : 0.487
- loss coefficient mixture of Gaussian: 0.024
- learning rate: 0.001
- batch normalization
- dropout rate: 0.5
- distance threshold (perplexity parameter): 2
- depth clustering network: 5
- internal size clustering network: 100
- trainable parameters: depending on number of experts

We compare results based on F-measure (Aghaeepour et al., 2013), which is defined as follows

$$F(C,K) = \sum_{c_i \in C} \frac{|c_i|}{N} \max_{k_j \in K} \{F(c_i, k_j)\}$$
(12)

where N is the number of samples $C\{c_1, c_2, \ldots, c_n\}$ and $K\{k_1, k_2, \ldots, k_m\}$ are the cluster result and the reference cluster, respectively. Further $F(c_i, k_j)$ is the harmonic mean of precision and recall according to

$$F(c_i, k_j) = \frac{2Pr(c_i, k_j)Re(c_i, k_j)}{Pr(c_i, k_j) + Re(c_i, k_j)}$$
(13)

whereby $Pr(c_i, k_j)$ is the precision and $Re(c_i, k_j)$ is the recall. Results are shown in Table 2.



(b)

Figure A1: Testing MoE-Sim-VAE on data sampled from a Gaussian mixture model with random sampled parameters. Figure A1a: Testing with exact numbers of experts. When comparing to classification results with GMMs one can see that our model achievs better results until around 30 mixture components and is still compatitive until 40 mixture components. With more then 40 mixture components ourr MoE-Sim-VAE is not able anymore to compete with a GMM. Figure A1b: Testing for specific number of synthetic mixture components and iterating number of experts. Until a number of GMM components of 23 MoE-Sim-VAE is very precise in learning the real number of clusters even when allowing the model to have 40 experts.

A.1.2 UNSUPERVISED CLUSTERING, EMBEDDING AND DATA GENERATION OF MNIST

Model and training details:

- number of experts: 10
- batch size: 128
- code size: 68
- Number of iterations: 20000
- activation function; elu
- loss coefficient data reconstruction: 0.487

- loss coefficient clustering : 0.487
- loss coefficient mixture of Gaussian: 0.024
- learning rate: 0.0001
- batch normalization
- dropout rate: 0.5
- k from kNN (perplexity parameter): 10
- depth clustering network: 3
- internal size clustering network: 200
- trainable parameters: 1619446

One estimator of the Maximum Mean Discrepancy (MMD) (Gretton et al., 2008) is defined as

$$\widehat{MMD}^{2}(X,Y) = \frac{1}{\binom{m}{2}} \sum_{i \neq i'} k(X_{i}, X_{i'}) + \frac{1}{\binom{m}{2}} \sum_{j \neq j'} k(Y_{j}, Y_{j'}) - \frac{2}{\binom{m}{2}} \sum_{i,j} k(X_{i}, Y_{j})$$
(14)

where $X = {\hat{x}_1, \dots, \hat{x}_m} \stackrel{iid}{\sim} P, Y = {\hat{y}_1, \dots, \hat{y}_m} \stackrel{iid}{\sim} Q$ are samples from two distributions (e.g. samples from two different clusters of the latent representation, for MNIST of two different digits) and k is a kernel function, where we use the popular RBF kernel. Based on that estimator Sutherland et al. (2019) introduced the hypothesis test

$$H0: P = Q \tag{15}$$

$$H1: P \neq Q \tag{16}$$

using the statistic $\widehat{mMMD}^2(X, Y)$. The distribution for P and Q is not required to be known. Sutherland et al. (2019) used MMD and this test to train a Generative Adversarial Network (GAN) and also to evaluate the generative performance of the model. In this work we use $\widehat{MMD}^2(X,Y)$ to test if samples of different clusters of the latent representation are similar, or in other words the distance of the distributions. We used the Python implementation² from Sutherland et al. (2019).

A.1.3 LEARNING CELL TYPE COMPOSITION IN PERIPHERAL BLOOD MONONUCLEAR CELLS USING CYTOF MEASUREMENTS

Model and training details for all experiments on CyTOF data:

- number of experts: 25 (Weber & Robinson, 2016), 15 (Bodenmiller et al., 2012)
- batch size: 128
- code size: 9
- Number of iterations: 30000 (Weber & Robinson, 2016), 20000 (Bodenmiller et al., 2012)
- activation function: relu
- loss coefficient data reconstruction: 1
- loss coefficient clustering : 1
- loss coefficient mixture of Gaussian: 0
- learning rate: 0.001 (Weber & Robinson, 2016), 0.005 (Bodenmiller et al., 2012)
- batch normalization
- dropout rate: 0.5
- distance threshold (perplexity parameter): 2
- distance metric: correlation
- depth clustering network: 5
- internal size clustering network: 9

²https://github.com/dougalsutherland/opt-mmd/blob/master/two_sample/ mmd_test.py



Figure A2: More detailed overview of results and generated samples of MNIST images. The plot on the left side shows the latent representation where the red crosses are the cluster centers. Those can be used as a mean to sample from a standard Gaussian for data generation via the MoE Decoder. The boxplots on the right show the clustering and gating result of each sample from the variational distribution.

• trainable parameters: 37563 (Weber & Robinson, 2016), 22228 (Bodenmiller et al., 2012)

Results are computed setting the loss coefficient for the KL loss 3 equal to zero, since we do not intend to generate any data, but rather give the chance to the AE to pick up the correct subpouplations.









Figure A3: Comparison of two sample MMD test (Sutherland et al., 2019) on the distributions from the different mixture components in the latent representation. The heatmaps on the left side show the estimation of the MMD which can be seen as the distance between pairs of distributions. The figures on the right side show the separation of the cluster in the latent representation based on a dimensionality reduction via UMAP (McInnes et al., 2018). Figure A3a shows the results for the clusters of VaDE at a posterior threshold of 0.8 which is the first threshold which shows total separation of all clusters. Figure A3b shows the separation of the clusters in latent space learned from MoE-Sim-VAE. For both methods, all distributions belonging to clusters of different respective digits show a larger distance compared to the diagonal of matching distributions, such that we generate images from a well-separated latent representation for both methods and therefore the main difference comes from the decoders.

Also here we use the F-measure defined in Equation 12 as metric to evaluate the models. For the data compared in Weber & Robinson (2016) we ran each model 30 times and report reproducability of our results in A8. The model was trained on all data and validated on the on with labels.

For the data from Bodenmiller et al. (2012) we run each model on one time on the each of the 268 datasets. Hereby we focused on the following surface markers: CD3(110:114)Dd, CD45(In115)Dd, CD4(Nd145)Dd, CD20(Sm147)Dd, CD33(Nd148)Dd, CD123(Eu151)Dd, CD14(Gd160)Dd, IgM(Yb171)Dd, HLA-DR(Yb174)Dd, CD7(Yb176)Dd. The subpopulations were originally defined via the SPADE algorithm (Qiu et al., 2011), which is a visualization tool using Agglomerative







(b)

Figure A4: Ablation study on the similarity matrix S. Both figures show the MMD statistic and UMAP (McInnes et al., 2018) projection of reconstructed MNIST digits computed on the latent representation. Figure A4a shows the results on MoE-Sim-VAE trained with the similarity matrix. The different digits separate well which can also be seen in the heatmap showing the MMD statistics between all digits. In comparison, Figure A4b shows results of the MoE-Sim-VAE model ignoring the similarity matrix setting the loss coefficient to zero. One can observe that the MMD statistic, which can be seen as a similarity measure of two distributions, is way lower compared to the model including the similarity matrix. Further, also the UMAP projection confirms less separation in the latent representation between the different digits.

hierarchical clustering and minimum spanning trees. The gating of the cells is done manually via coloring of the tree leaves. With MoE-Sim-VAE we try to reconstruct the defined manually defined subpopulations. Bodenmiller et al. (2012) performed experiments on multiple well plates were different inhibitors and their effect was tested. We selected for each well plate row A to test our model on. We decided for all methods to discard subpopulations which are smaller then 30 cells. As a similarity measure for MoE-Sim-VAE we reduced the dimension of the data using UMAP (McInnes et al., 2018) using the Canberra distance

$$d(\mathbf{p}, \mathbf{q}) = \sum_{i=1}^{n} \frac{|p_i - q_i|}{|p_i| + |q_i|}$$
(17)







(b)

Figure A5: Comparison of data generation process between Moe-Sim-VAE and VaDE (Jiang et al., 2017). Figure A5a shows the accuracy of how certain a specific digit can be generated from the respective cluster in the latent representation whereas Figure A5b compares the number of runs until a sample from the latent representation satisfied the posterior criterion from VaDE. It needs to be mentioned that MoE-Sim-VAE does not require any thresholding such that we ran the data generation process multiple times with the same settings to compare with VaDE. In total 10000 samples are generated for each digit.

where $p = (p_1, p_2, ..., p_n)$ and $q = (q_1, q_2, ..., q_n)$. Cells were defined to be similar in MoE-Sim-VAE when the distance between the cells in the UMAP-projection was smaller then a threshold. We trained and tested MoE-Sim-VAE on a splitted dataset with rations 0.8/0.2 and evaluated the performance on the unseen test dataset. In comparison the compatitor methods were trained and



Figure A6: Confusion map for data generation using MoE-Sim-VAE. Besides the systematic error of confusing digit 5 and 8, which can also depend on the clustering network, the digit generation of our model performs very precise with a high accuracy of generating the digit asked for. In comparison to VaDE (Jiang et al., 2017) our model does not need any threshold on samples from the latent representation which reduces the computational costs by far.

tested on all the data, which is an advantage in comparison to our model, but still MoE-Sim-VAE outpreforms the compatitors.







(c)

Figure A7: Confusion maps for data generation using VaDE. Figure A7a Posterior threshold 0.0. Figure A7b Posterior threshold 0.1. Figure A7c Posterior threshold 0.2.







(f)

Figure A7: Confusion maps for data generation using VaDE. Figure A7d Posterior threshold 0.3. Figure A7e Posterior threshold 0.4. Figure A7f Posterior threshold 0.5.







(i)

Figure A7: Confusion maps for data generation using VaDE. Figure A7g Posterior threshold 0.6. Figure A7h Posterior threshold 0.7. Figure A7i Posterior threshold 0.8.



Figure A7: Confusion maps for data generation using VaDE. Figure A7j Posterior threshold 0.9. Figure A7k Posterior threshold 0.999. (default for Jiang et al. (2017))



Figure A8: The boxplots show similar as in Weber & Robinson (2016) the reproducibility of MoE-Sim-VAE on the four datasets when running MoE-Sim-VAE 30 times. The variance on defining the correct subpopulations of MoE-Sim-VAE is quite small and therefore also an improvment to many methods compared in Weber & Robinson (2016).

Table A1: Results of MoE-Sim-VAE on data published in Bodenmiller et al. (2012). CyTOF measurements from peripheral blood mononuclear cells (PBMCs) were taken and the goal is to define the different cell types present in the data. The ground truth was definied using the SPADE algorithm (Qiu et al., 2011), which can visualize the high dimensional data in such a way to be able to manual gate the cells. We compare to other fully unsupervised methods as FlowSOM, X-shift and PhenoGraph and achieve in most cases the best F-measure, which is defined as in Equation 12.

Inhibitor	Well	MoE-Sim-VAE	FlowSOM	X-shift	PhenoGraph
AKTi	A02	0.7666	0.5147	0.5704	0.6588
AKTi	A03	0.7541	0.4793	0.546	0.6026
AKTi	A04	0.6815	0.6405	0.5298	0.5974
AKTi	A05	0.7127	0.7108	0.6089	0.6104
AKTi	A06	0.6711	0.7383	0.572	0.6611
AKTi	A07	0.7233	0.7034	0.5583	0.6981
AKTi	A08	0.7901	0.7024	0.4541	0.5287
AKTi	A09	0.7604	0.4292	0.5014	0.6414
AKTi	A10	0.7275	0.4952	0.4144	0.677
AKTi	A11	0.7540	0.6456	0.6673	0.6302
BTKi	A02	0.7261	0.698	0.7136	0.7478
BTKi	A03	0.7982	0.6643	0.6012	0.7141
BTKi	A04	0.7835	0.6864	0.6983	0.7103
BTKi	A05	0.7484	0.6397	0.7454	0.7474
BTKi	A06	0.8196	0.703	0.7625	0.7949
BTKi	A07	0.7976	0.6729	0.6841	0.7102
BTKi	A08	0.8108	0.6715	0.5887	0.6884
BTKi	A09	0.7789	0.5299	0.6426	0.7236
BTKi	A10	0.7726	0.6319	0.6775	0.7148
BTKi	A11	0.7857	0.6078	0.5939	0.6786
BTKi	A12	0.6600	0.5503	0.6028	0.6308
Crassin	A01	0.6727	0.6488	0.6315	0.6237
Crassin	A02	0.8225	0.557	0.6435	0.7165
Crassin	A03	0.8346	0.5736	0.6628	0.7085
Crassin	A04	0.8446	0.5348	0.7146	0.7045
Crassin	A05	0.8462	0.7444	0.6227	0.7202
Crassin	A06	0.8569	0.7448	0.7078	0.6972
Crassin	A07	0.8170	0.5164	0.6546	0.6309
Crassin	A08	0.8431	0.8283	0.5504	0.6546
Crassin	A09	0.8412	0.5814	0.6027	0.6684
Crassin	A10	0.8527	0.7537	0.6586	0.6338
Crassin	A11	0.8453	0.7174	0.6437	0.7358
Crassin	A12	0.7320	0.6161	0.6436	0.6949
Dasatinib	A01	0.7235	0.4466	0.554	0.6725
Dasatinib	A02	0.8019	0.516	0.6238	0.701
Dasatinib	A03	0.7864	0.5108	0.5366	0.6566
Dasatinib	A04	0.6661	0.4796	0.5527	0.647
Dasatinib	A05	0.7910	0.5014	0.5804	0.6904
Dasatinib	A06	0.7979	0.5167	0.6258	0.6707
Dasatinib	A07	0.8105	0.5215	0.6016	0.6809
Dasatinib	A08	0.8047	0.6928	0.5802	0.633
Dasatinib	A09	0.7485	0.5203	0.5958	0.6861
Dasatinib	A10	0.8062	0.5158	0.5742	0.6503
Dasatinib	A11	0.7837	0.5066	0.6331	0.6813
GDC-0941	A01	0.5632	0.6434	0.5987	0.6279
GDC-0941	A02	0.8257	0.7291	0.7349	0.7507
GDC-0941	A03	0.8268	0.7321	0.6822	0.7853
GDC-0941	A04	0.8389	0.7115	0.7569	0.7421
GDC-0941	A05	0.8382	0.7946	0.7171	0.7735
GDC-0941	A06	0.8463	0.6125	0.6858	0.764
GDC-0941	A07	0.8382	0.6061	0.7776	0.7612

- 1	GDC-0941	A08	0.8249	0.5493	0.6058	0.7796
	GDC-0941	A09	0.8606	0.7689	0.8043	0.7206
	GDC-0941		0.8412	0.7227	0.653	0.6465
		A10				
	GDC-0941	A11	0.7859	0.5703	0.7297	0.7891
	GDC-0941	A12	0.7803	0.6326	0.69	0.6727
	G069	A01	0.6520	0.6571	0.718	0.5822
	G069	A02	0.7835	0.7693	0.6075	0.7322
	G069	A03	0.7305	0.7334	0.757	0.6414
	G069	A04	0.7640	0.7456	0.8013	0.7425
	G069	A05	0.7812	0.7555	0.7294	0.7727
	G069	A06	0.7816	0.7404	0.7437	0.6443
	G069	A07	0.7407	0.8513	0.7527	0.6811
	G069	A07 A08	0.7293	0.7338	0.6984	0.6525
	G069	A09	0.8228	0.6955	0.6985	0.7317
	G069	A10	0.7560	0.7512	0.7689	0.7071
	G069	A11	0.7565	0.7373	0.7213	0.7315
	G069	A12	0.7426	0.7086	0.7846	0.6442
	H89	A01	0.6734	0.6952	0.6003	0.6105
	H89	A02	0.7288	0.5391	0.5918	0.678
	H89	A03	0.8051	0.5414	0.6856	0.6759
	H89	A04	0.8144	0.7314	0.662	0.7287
	H89	A05	0.7821	0.5468	0.6485	0.6672
	H89	A06	0.7647	0.5636	0.8281	0.7165
	H89	A07	0.7762	0.6983	0.7284	0.6442
	H89	A09	0.8131	0.5442	0.5906	0.6707
	H89	A10	0.7517	0.5549	0.6028	0.682
	H89	A10 A11	0.7417	0.3349	0.6863	0.7257
	H89	A12	0.7939	0.6934	0.5831	0.6401
	IKKi	A02	0.7945	0.6619	0.7371	0.6475
	IKKi	A03	0.6873	0.6568	0.5661	0.6895
	IKKi	A04	0.7942	0.6754	0.6386	0.7052
	IKKi	A05	0.6977	0.6569	0.6157	0.6899
	IKKi	A06	0.7442	0.6931	0.7024	0.7077
	IKKi	A07	0.7352	0.5303	0.669	0.7001
	IKKi	A08	0.7470	0.7006	0.5358	0.6869
	IKKi	A09	0.8097	0.5175	0.6299	0.6969
	IKKi	A10	0.7647	0.6308	0.657	0.7334
	IKKi	A11	0.7878	0.6365	0.6757	0.6613
	IKKi	A12	0.6673	0.5629	0.497	0.6043
ł	Imatinib	A02	0.7935	0.7571	0.6721	0.7677
	Imatinib	A03	0.7763	0.7429	0.7041	0.7499
	Imatinib	A03 A04	0.8058	0.7564	0.6921	0.7229
	Imatinib	A04 A05	0.7714	0.7559	0.6689	0.7609
	Imatinib	A06	0.7756	0.746	0.6956	0.7296
	Imatinib	A07	0.7468	0.7515	0.6974	0.7137
	Imatinib	A08	0.7631	0.7534	0.5189	0.7096
	Imatinib	A09	0.8082	0.5605	0.5819	0.7447
	Imatinib	A10	0.7964	0.5645	0.5637	0.78
	Imatinib	A11	0.7289	0.7664	0.7576	0.7395
	Imatinib	A12	0.7012	0.8451	0.6369	0.7259
ľ	Jak1i	A02	0.8210	0.5167	0.5771	0.616
	Jak1i	A03	0.7343	0.7139	0.6526	0.7133
	Jak1i	A04	0.7321	0.7066	0.6346	0.7189
	Jak1i	A05	0.7413	0.5163	0.6551	0.7089
	Jak1i	A06	0.7244	0.5525	0.6804	0.6905
	Jak1i	A07	0.7779	0.5499	0.5605	0.7099
	Jak1i	A08	0.7281	0.6995	0.6021	0.6605
	Jak1i	A08 A09	0.8043	0.5064	0.6054	0.6717
	Jak1i	A09 A10	0.7801	0.5004	0.5538	0.7015
I	JANII	A10	0.7001	0.5295	0.5550	0.7015

Jak1i	A11	0.7128	0.7307	0.7386	0.6812
Jak1i	A12	0.7204	0.6229	0.6321	0.6905
Jak2i	A01	0.6944	0.6379	0.6014	0.6207
Jak2i	A02	0.7961	0.664	0.6656	0.7083
Jak2i	A03	0.7629	0.6742	0.7138	0.7024
Jak2i	A03	0.7890	0.6716	0.6227	0.7072
Jak2i	A04 A05	0.6666	0.4689	0.5314	0.6459
Jak2i	A05 A06	0.8110	0.4089	0.6651	0.6833
Jak2i	A00 A07	0.7595	0.6818	0.0051	0.6982
Jak2i	A07 A08	0.7595	0.6601	0.7393 0.6152	0.6982
Jak2i		0.8050	0.0001	0.6132	0.6501
	A09				
Jak2i	A10	0.8030	0.6762	0.6067	0.6364
Jak2i	A11	0.8228	0.5398	0.694	0.7473
Jak2i	A12	0.6831	0.6214	0.5825	0.5687
Jak3i	A02	0.7986	0.7108	0.5666	0.6912
Jak3i	A03	0.7170	0.7116	0.6991	0.7001
Jak3i	A04	0.7983	0.5243	0.6654	0.691
Jak3i	A05	0.7087	0.6498	0.6884	0.7073
Jak3i	A06	0.7272	0.7244	0.654	0.7059
Jak3i	A07	0.7768	0.5167	0.696	0.735
Jak3i	A08	0.7196	0.6797	0.5946	0.7287
Jak3i	A09	0.7988	0.6918	0.6013	0.6826
Jak3i	A10	0.8026	0.7103	0.7104	0.7219
Jak3i	A11	0.7281	0.5107	0.6854	0.6614
Jak3i	A12	0.7511	0.6135	0.4861	0.61
Lcki	A01	0.7359	0.7582	0.6106	0.7201
Lcki	A02	0.7605	0.7453	0.6391	0.7696
Lcki	A03	0.8032	0.5608	0.6814	0.721
Lcki	A04	0.7608	0.5764	0.6788	0.7904
Lcki	A05	0.8210	0.5435	0.7204	0.7442
Lcki	A06	0.7564	0.7662	0.728	0.7556
Lcki	A07	0.8304	0.579	0.6992	0.696
Lcki	A08	0.7854	0.7457	0.5904	0.6972
Lcki	A09	0.8452	0.5859	0.6018	0.7569
Lcki	A10	0.7387	0.744	0.6598	0.6627
Lcki	A11	0.7835	0.7639	0.6836	0.7558
Lcki	A12	0.7467	0.8271	0.6888	0.6878
PP2	A02	0.7687	0.759	0.7717	0.7605
PP2	A03	0.8395	0.7644	0.7304	0.7953
PP2	A04	0.8442	0.7703	0.7116	0.7162
PP2	A05	0.8248	0.5777	0.7205	0.7547
PP2	A06	0.7866	0.7612	0.7461	0.7431
PP2	A07	0.8595	0.7616	0.724	0.7213
PP2	A08	0.8505	0.7489	0.7109	0.7195
PP2	A09	0.7902	0.5755	0.6511	0.7738
PP2	A10	0.8089	0.743	0.6635	0.7389
PP2	A11	0.7977	0.5852	0.6564	0.7846
PP2	A12	0.7667	0.6012	0.6524	0.6636
Rapamycin	A01	0.7028	0.675	0.5882	0.5677
Rapamycin	A02	0.7215	0.6831	0.6124	0.6697
Rapamycin	A03	0.7322	0.6707	0.6296	0.6861
Rapamycin	A04	0.6787	0.6696	0.6887	0.7267
Rapamycin	A05	0.7231	0.653	0.7134	0.6466
Rapamycin	A06	0.7310	0.6473	0.7009	0.6386
Rapamycin	A07	0.7595	0.6642	0.748	0.5882
Rapamycin	A08	0.7773	0.836	0.6371	0.571
Rapamycin	A09	0.7732	0.6573	0.6826	0.6615
Rapamycin	A10	0.7586	0.6702	0.7136	0.6344

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Rapamycin	A12	0.6955	0.6361	0.6561	0.5472
SB202	A01	0.6884	0.6713	0.941	0.7101
SB202	A03	0.7869	0.7549	0.6686	0.7633
SB202	A05	0.7856	0.5564	0.7387	0.6999
SB202	A06	0.7707	0.755	0.7913	0.7869
SB202	A10	0.7559	0.7554	-	0.7749
SP6	A01	0.7033	0.6882	0.4191	0.532
SP6	A02	0.7536	0.5035	0.5104	0.657
SP6	A03	0.7387	0.6973	0.534	0.5858
SP6	A04	0.6910	0.503	0.5065	0.5975
SP6	A05	0.7210	0.5068	0.5643	0.6869
SP6	A06	0.7052	0.719	0.5063	0.6384
SP6	A07	0.7281	0.7074	0.5382	0.6501
SP6	A08	0.7301	0.6832	0.4665	0.6133
SP6	A09	0.7743	0.5001	0.4618	0.6208
SP6	A10	0.7198	0.5111	0.524	0.6773
SP6	A11	0.7494	0.493	0.5407	0.5935
SP6	A12	0.7311	0.6131	0.4488	0.6198
Sorafenib	A01	0.7185	0.7217	0.5884	0.6574
Sorafenib	A02	0.8250	0.7659	0.6658	0.7664
Sorafenib	A03	0.7689	0.7732	0.7078	0.6869
Sorafenib	A04	0.8360	0.7094	0.7114	0.7218
Sorafenib	A05	0.8304	0.5571	0.7672	0.7153
Sorafenib	A06	0.8021	0.5783	0.6991	0.7506
Sorafenib	A07	0.8461	0.7051	0.7267	0.6701
Sorafenib	A09	0.8226	0.7275	0.7522	0.7587
Sorafenib	A10	0.8103	0.7561	0.7457	0.7214
Sorafenib	A11	0.8465	0.5777	0.7192	0.7503
Sorafenib	A12	0.7715	0.6533	0.6084	0.6129
Staurosporine	A01	0.7985	0.8464	0.6057	0.5945
Staurosporine	A02	0.8347	0.8312	0.5999	0.6626
Staurosporine	A03	0.8079	0.7072	0.6704	0.6787
Staurosporine	A04	0.8418	0.8666	0.6452	0.6776
Staurosporine	A05	0.8657	0.7305	0.7071	0.7515
Staurosporine	A06	0.8694	0.516	0.6453	0.6619
Staurosporine	A07	0.8277	0.7052	0.6349	0.6657
Staurosporine	A08	0.8310	0.8316	0.6213	0.678
Staurosporine	A09	0.8319	0.5117	0.6747	0.6726
Staurosporine	A10	0.8417	0.5108	0.6211	0.7126
Staurosporine	A11	0.8246	0.8711	0.6547	0.7445
Streptonigrin	A01	0.7128	0.5689	0.6571	0.6599
Streptonigrin	A02	0.7836	0.5095	0.549	0.6155
Streptonigrin	A03	0.7776	0.547	0.6497	0.6527
Streptonigrin	A04	0.8466	0.7521	0.5762	0.7061
Streptonigrin	A05	0.8130	0.5406	0.6459	0.6928
Streptonigrin	A06	0.8031	0.7409	0.6446	0.6343
Streptonigrin	A07	0.7987	0.5353	0.5882	0.6657
Streptonigrin	A08	0.7470	0.7458	0.5864	0.6443
Streptonigrin	A09	0.7586	0.7034	0.5928	0.6196
Streptonigrin	A10	0.7159	0.6974	0.5174	0.6809
Streptonigrin	A11	0.8178	0.5649	0.593	0.6814
Streptonigrin	A12	0.7410	0.6034	0.5896	0.6286
Sunitinib	A01	0.7152	0.6622	0.5653	0.6522
Sunitinib	A02	0.8056	0.498	0.6138	0.6521
Sunitinib	A03	0.8095	0.6873	0.6889	0.6913
Sunitinib	A04	0.8142	0.6925	0.6467	0.7121
	A05	0.8157	0.6959	0.673	0.7073
Sunitinib	AU	V .() I . <i>i</i> /			
Sunitinib Sunitinib	A05 A06	0.7968	0.5061	0.6654	0.7025

Sunitinib A08 0.8186 0.6894 0.59 Sunitinib A09 0.8029 0.4886 0.66 Sunitinib A10 0.8126 0.848 0.60 Sunitinib A11 0.8241 0.8241 0.6241	99 0.6621
Sunitinib A10 0.8126 0.848 0.60	
	87 0.6713
C	
Sunitinib A11 0.8241 0.824 0.64	08 0.6811
Sunitinib A12 0.7747 0.7898 0.59	42 0.5867
Syki A02 0.7682 0.7073 0.66	36 0.685
Syki A03 0.7224 0.7042 0.64	24 0.7116
Syki A04 0.7461 0.7069 0.79	08 0.7256
Syki A05 0.7468 0.7182 0.62	63 0.6804
Syki A06 0.7381 0.7134 0.77	
Syki A07 0.7891 0.7 0.74	
Syki A08 0.7509 0.7154 0.69	
Syki A09 0.7712 0.73 0.73	
Syki A10 0.7695 0.7531 0.71	
Syki A11 0.7360 0.7311 0.75	
Syki A12 0.6717 0.6793 0.74	
U0126 A01 0.6844 0.6178 -	0.6486
U0126 A02 0.8440 0.5545 0.53	
U0126 A03 0.8340 0.5346 0.61	
U0126 A04 0.8263 0.7079 0.61	
U0126 A05 0.8535 0.5468 0.70	
U0126 A06 0.8199 0.5285 0.60	
U0126 A07 0.8079 0.5304 0.56	
U0126 A08 0.8278 0.6864 0.53	
U0126 A09 0.8331 0.5394 0.56	
U0126 A10 0.8436 0.5593 0.60	
U0126 A11 0.7654 0.5072 0.63	
U0126 A12 0.7227 0.6496 0.62	
VX680 A01 0.6930 0.4818 0.60	
VX680 A02 0.7340 0.711 0.55	
VX680 A03 0.7525 0.6976 0.56	
VX680 A04 0.8127 0.6435 0.67	
VX680 A05 0.6937 0.6742 0.73	
VX680 A06 0.7168 0.7101 0.57	
VX680 A07 0.7663 0.4944 0.53	
VX680 A08 0.7315 0.7082 0.47	
VX680 A09 0.7703 0.7054 0.58	
VX680 A10 0.7143 0.7137 0.66	
VX680 A11 0.7050 0.6773 0.72	
VX680 A12 0.7852 0.7922 0.55	83 0.6808