Neural Decomposition: Functional ANOVA with Variational Autoencoders

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Abstract

Variational Autoencoders (VAEs) have become a popular approach for dimensionality reduction. However, despite their ability to identify latent low-dimensional structures embedded within high-dimensional data, these latent representations are typically hard to interpret on their own. Due to the black-box nature of VAEs, their utility for healthcare and genomics applications has been limited. In this paper, we focus on characterising the sources of variation in Conditional VAEs. Our goal is to provide a feature-level variance decomposition, i.e. to decompose variation in the data by separating out the marginal additive effects of latent variables $z$ and fixed inputs $c$ from their non-linear interactions. We propose to achieve this through what we call Neural Decomposition – an adaptation of the well-known concept of functional ANOVA variance decomposition from classical statistics to deep learning models. We show how identifiability can be achieved by training models subject to constraints on the marginal properties of the decoder networks. We demonstrate the utility of our Neural Decomposition on a series of synthetic examples as well as high-dimensional genomics data.

1 Introduction

Dimensionality reduction is often required for the analysis of complex high-dimensional data sets in order to identify low-dimensional sub-structures that may give insight into patterns of interest embedded within the data. Recently there has been particular interest in applications of Variational Autoencoders (VAEs) (Kingma and Welling, 2014) for generative modelling and dimensionality reduction. VAEs are a class of probabilistic neural-network-based latent variable models. Their neural network (NN) underpinnings offer considerable modelling flexibility while modern programming frameworks (e.g. TensorFlow and PyTorch) and computational techniques (e.g. stochastic variational inference) provide immense scalability to large data sets, including those in genomics (Lopez et al., 2018; Eraslan et al., 2019; Märtens and Yau, 2020). The Conditional VAE (CVAE) (Sohn et al., 2015) extends this model to allow incorporating side information as additional fixed inputs.

The ability of (C)VAEs to extract latent, low-dimensional representations from high-dimensional inputs is a strength for predictive tasks. For example (C)VAEs have been particularly popular as generative models for images. However, when the objective is to understand the contributions of different inputs to components of variation in the data, the black box decoder is insufficient to permit this. This is particularly important for tabular data problems where individual features might correspond to actual physical quantities (e.g. expression of a gene or a physiological measurement) and we are interested in how the variability of the multivariate output response is driven by specific changes in the inputs (where inputs may be either latent variables or observed covariates).

Functional analysis of variance (functional ANOVA, or fANOVA) are a class of statistical models that uniquely decompose a functional response according to the main effects and interactions of various factors (Sobol, 1993; Ramsay and Silvermann, 1997). For example, smoothing spline ANOVA models (Gu, 2013) express the effects as linear combinations of some underlying basis functions, and the coefficients on these basis functions are chosen to minimise a loss function balancing good-
ness of fit with a measure of smoothness of the fitted functions. More recent treatments have proposed functional ANOVA decompositions via tree-based models (Lengerich et al., 2019), and within a Bayesian nonparametric framework using Gaussian Processes (Kaufman and Sain, 2010).

In this paper, we propose the notion of neural decomposition – the integration of functional ANOVA and deep neural networks for dimensionality reduction and variance decomposition. Specifically, we develop and explore a class of (C)VAE models where the decoding network is specifically implemented to explain feature-level variability in terms of additive and interaction effects between the latent variables and any additional covariate information (Figure 1).

Thus, neural decomposition can be seen from two perspectives:

- From the classical statistics perspective, it is a scalable neural adaptation of the functional ANOVA decomposition, where additional latent variables $z$ have been introduced to capture low-dimensional structure within high-dimensional data and non-linearities implemented via deep neural nets.

- From the deep learning perspective, it is an extension of the (C)VAE framework, where the decoder has a decomposable additive structure to aid interpretability on the level of individual features.

Crucially, in order for this class of models to be useful, identifiability must be enforced to ensure the expressive power of each neural network component does not absorb variation that should be explained by others. Inference for our model relies on the imposition of functional constraints which we implement as constrained optimisation in the weight-space of neural networks.

2 Background

VAEs: Variational Autoencoders are constructed based on particular a latent variable model structure. Let $y \in \mathcal{Y}$ denote a data point in some potentially high-dimensional space and $z \in \mathcal{Z}$ be a vector of associated latent variables typically in a much lower dimensional space. We assume that the prior $p(z)$ is from a family of distributions that is easy to sample from - we will assume $z \sim \mathcal{N}(0, 1)$ throughout. Now suppose $f^\theta(z)$ is a family of deterministic functions, indexed by parameters $\theta \in \Theta$, such that $f : \mathcal{Z} \times \Theta \rightarrow \mathcal{Y}$. Given data points $y_{1:N} = \{y_1, \ldots, y_N\}$, the marginal likelihood is given by $p(y_{1:N}) = \prod_{i=1}^N \int p_\theta(y_i|z_i)p(z_i)dz_i$. We will assume a Gaussian likelihood, i.e. $y_i|z_i, \theta \sim \mathcal{N}(f^\theta(z_i), \sigma^2)$. In the VAE, these deterministic functions $f^\theta$ are given by deep neural networks (NNs). Posterior inference in a VAE is carried out via amortised variational inference, i.e. with a parametric inference model $q_\phi(z_i|y_i)$ where $\phi$ are variational parameters. Application of standard variational inference methodology, see e.g. (Blei et al., 2017), leads to a lower bound on the log marginal likelihood, i.e. the ELBO of the form:

$$
\mathcal{L}_{\phi, \theta} = \sum_{i=1}^N \mathbb{E}_{q_\phi(z_i|y_i)}[\log p_\theta(y_i|z_i)] - \text{KL}(q_\phi(z_i|y_i)||p(z_i))
$$

In the VAE, the variational approximation $q_\phi(z_i|y_i)$ is referred to as the encoder since it encodes the data $y$ into the latent variable $z$ whilst the decoder refers to the generative model $p_\theta(y|z)$ which decodes the latent variables into observations.

Training a VAE seeks to optimise both the model parameters $\theta$ and the variational parameters $\phi$ jointly using stochastic gradient ascent. This typically requires a stochastic approximation to the gradients of the variational objective which is itself intractable. Typ-
ically the approximating family is assumed to be Gaussian, i.e. \( q_\theta(z_i|y_i) = \mathcal{N}(z_i|\mu_\theta(y_i),\sigma^2_\theta(y_i)) \), so that a reparameterisation trick can be used (Kingma and Welling, 2014; Rezende et al., 2014).

**Conditional VAEs:** The Conditional VAE (Sohn et al., 2015) augments the VAE by conditioning the generative model on additional inputs \( c \), i.e. the VAE generative model \( y_i = f^\theta(z_i) + \varepsilon_i \) is now replaced by \( y_i = f^\theta(z_i, c_i) + \varepsilon_i \) which requires a minor alteration to the ELBO

\[
\mathcal{L}^\phi = \sum_{i=1}^N \mathbb{E}_{q_\theta(z_i|y_i,c_i)}[\log p_\theta(y_i|z_i, c_i)] - \text{KL}(q_\theta(z_i|y_i,c_i)||p(z_i)).
\]

Conditioning on the extra inputs can greatly increase the expressiveness of the VAE when applied to multimodal data where the indicator driving the multimodality is measurable. For the genomics applications we consider, we use this mechanism to incorporate known covariates as part of the model.

### 3 Neural Decomposition

Our goal is to equip (C)VAEs with feature-level interpretability that would let us characterise the sources of variation for individual features. We aim to achieve this by directly embed decomposable structure within this latent variable model. Specifically, we propose to perform functional ANOVA decomposition as part of the decoder network within the (C)VAE, in order to simultaneously perform dimensionality reduction as well as obtain a feature-level variance decomposition.

For illustration purposes, we consider the special case \( z, c \in \mathbb{R} \). Here we want to decompose the decoder network \( f^\theta(z_i, c_i) \) to extract additive marginal and interaction effects as follows

\[
f^\theta(z_i, c_i) = f^\theta_0 + f^\theta_z(z_i) + f^\theta_c(c_i) + f^\theta_{zc}(z_i, c_i)
\]

as illustrated in Figure 1. However, note that for this functional decomposition to have a meaningful interpretation, identifiability must be enforced.

Traditionally, in functional ANOVA the functions \( f_{ij}, f_{ik}, f_{zc} \) would be represented by either linear mappings, smoothing splines (Gu, 2013) or Gaussian Processes (Kaufman and Sain, 2010). Here, we propose to use deep neural networks instead – a Neural Decomposition (ND). This would enable separating main effects from complex interactions while otherwise maintaining the flexibility of deep generative models, and fast, approximate inference via variational methods. This decoding structure has been illustrated in Fig 3(C).

Next, we formulate the conditions for the decompositions to be unique, and discuss how such constraints can be fulfilled in practice.

#### 3.1 Identifiable Neural Decomposition

For a more general formulation, let us denote the latent and fixed inputs collectively by \( x := (z, c) \) with dimensionality \( D := \dim(x_i) \). We would like to decompose as follows

\[
f^\theta_0 + \sum_k f_k^\theta(x_{ik}) + \sum_{k,l} f_{kl}^\theta(x_{ik}, x_{il}) + \ldots + f^\theta_{1,...,D}(x) \tag{2}
\]

where all functions \( f_k^\theta, f_{kl}^\theta, \ldots, f^\theta_{1,...,D} \) are parameterised by neural networks. We note that without any additional constraints on the neural networks the above decomposition (2) is unidentifiable. This is because the functional subspaces \( f_{ij} \) corresponding to different index sets \( I \) can all be seen as functions defined on the same input space \( \mathbb{R}^D \), being constant in the rest of coordinates, and these subspaces are overlapping:

**Proposition 1.** Let \( f^\theta_0, f^\theta_1, \ldots, f^\theta_{1,...,D} \) be neural networks with the same architecture, i.e. assuming that the networks only differ in the number of inputs. Then for any two disjoint sets of indices \( I \) and \( J \) the functional subspace \( \{ f^\theta_{iz}(x_z) : \theta \in \mathbb{R}^D \} \) is strictly a subset of the functional subspace \( \{ f^\theta_{iz}(x_z, x_J) : \theta \in \mathbb{R}^D \} \).

**Proof:** weights in the first layer corresponding to inputs \( x_J \) can be set to zero, eliminating the effect of \( x_J \).

As a result, without any additional constraints decomposition (2) is not meaningful: it can be used for predictive purposes, but the relative contribution of different terms has no direct interpretation since higher-order interactions can absorb variability that could be explained by main effects or low-order interactions.

To turn this into an identifiable learning problem, we need to introduce functional constraints. As in functional ANOVA, we introduce the integral constraints to constrain the marginal effects of every neural network \( f^\theta_0(x) \) to be zero. This can be formalised as follows:

**Proposition 2.** Let neural networks \( f^\theta_0, f^\theta_1, \ldots, f^\theta_{1,...,D} \) be such that they satisfy the following integral constraints

\[
\int f^\theta_{iz}(x_i)dx_i = 0 \quad \text{for all } i \in I
\]

for all neural networks \( f^\theta_{iz} \) in (2), i.e. for every index set \( I \subset \{1, \ldots, D\} \). Then for any \( I \cap J = \emptyset \) the functional subspaces corresponding to \( f^\theta_{iz} \) and \( f^\theta_{iz,j} \) do not overlap any more (apart from the constant zero function). Furthermore, these functional subspaces are orthogonal in \( L^2 \).

\footnote{Whilst we focus on the CVAE setting in this paper, the decomposition is more generally applicable. Thus in this section, we treat all inputs as fixed.}

\footnote{Assuming a fully connected first layer with \( H \) hidden units, i.e. that the first transformation applied to inputs \( x_I \) is \( Wx_I + b \) where \( W \in \mathbb{R}^{H \times \dim(x_I)} \), \( b \in \mathbb{R}^H \), by “the same architecture” we mean that \( \dim(x_I) \) is the only element that is allowed to vary across \( f^\theta_{iz} \).}
Both of these properties (no overlap and orthogonality) are a direct consequence of the integral constraints. The general proof follows derivations as in previous works, see e.g. (Sobol, 1993). We note that the former (no overlap) is sufficient for identifiability, but the latter (orthogonality) leads to an easily interpretable variance decomposition

\[
\text{Var}(f^\theta_0) + \sum_k \text{Var}(f^\theta_k(x_{ik})) + \\
+ \sum_{k,l} \text{Var}(f^\theta_k(x_i, x_j)) + \ldots + \text{Var}(f^\theta_{1,\ldots,D}(x))
\]

For the special case of a two-dimensional input \((x_1, x_2)\), i.e. when the decomposition consists of \(f^\theta_0, f^\theta_1, f^\theta_2, f^\theta_1\), the above leads to the following integral constraints:

- \(\int f^\theta_1(x_1)dx_1 = 0\) and \(\int f^\theta_2(x_2)dx_2 = 0\)
- \(\int f^\theta_1(x_1, x_2)dx_1 = 0\) for all \(x_2\)
- \(\int f^\theta_2(x_1, x_2)dx_2 = 0\) for all \(x_1\).

### 3.2 Inference under integral constraints

We now return to our original goal which is understanding the sources of variation in (C)VAEs. As in eq (1), our goal is to learn a decomposition as part of the generative model – in the VAE framework it corresponds to decomposing the decoder\(^3\). To obtain an identifiable neural decomposition, we need to enforce the above integral constraints for the decoding networks.

This is a constrained optimisation problem, which is non-trivial to solve in the context of deep learning where state-of-the-art off-the-shelf optimisation techniques, such as Adam (Kingma and Ba, 2014), are typically implemented for unconstrained problems only. Thus alternative strategies have to be considered. We turn to the Augmented Lagrangian method, also known as the method of multipliers (Hestenes, 1969; Powell, 1969), to enforce such constraints. We refer to (Platt and Barr, 1988) for an adaptation to neural networks.

\(^3\)We note that we aim to make the functional decomposition (as opposed to the entire (C)VAE) identifiable.

#### Augmented Lagrangian for a single constraint:

For illustration, we first consider the case where our decoder \(f^\theta(x)\) has a univariate input \(x\), and we want to optimise the ELBO subject to a constraint \(\int f^\theta(x)dx = 0\), i.e. we want to restrict the \(f^\theta\) to a subspace such that \(\int f^\theta(x)dx = 0\). To enforce this constraint, we will augment the ELBO with additional penalty term(s) which will be equal to zero when the integral constraints are fulfilled. The resulting objective function is not necessarily a lower bound, but reduces to the ELBO once the constraints become fulfilled during optimisation.

One such approach for would be the penalty method, i.e. to incorporate a penalty term \(c \cdot (\int f^\theta(x)dx)^2\) with a fixed penalty \(c\). This approach has the disadvantage that for a fixed value of \(c\) we do not have any guarantees that constraints would be fulfilled exactly.

Alternatively one could introduce a penalty \(\lambda \int f^\theta(x)dx\) where now \(\lambda\) is treated as a parameter. This is analogous to the use of Lagrange multipliers, and following the terminology of (Platt and Barr, 1988), we refer to this as the Basic Differential Multiplier Method (BDMM). Instead of gradient updates \(\lambda^{t+1} = \lambda^t - \eta (\int f^\theta(x)dx)\), BDMM would follow the opposite direction \(\lambda^{t+1} = \lambda^t + \eta (\int f^\theta(x)dx)\) when optimising \(\lambda\). Platt and Barr showed that this corresponds to optimisation behaviour where the system undergoes damped oscillation.

We have empirically compared the behaviour of the penalty method and the BDMM approach on a synthetic problem with two-dimensional inputs, as shown in Figure 2A. Using a fixed penalty does not necessarily lead to fulfilled constraints, whereas the oscillating behaviour of the BDMM leads to the integrals converging towards zero.

Finally, these two penalty terms can be combined, resulting in a hybrid constrained optimisation objective

\[
\min_\theta \left\{ -\mathcal{L}^\theta + \lambda \int f^\theta(x)dx + c \left( \int f^\theta(x)dx \right)^2 \right\}
\]
where $\lambda$ is optimised, $c$ is a fixed constant, and $L_{\theta,\phi}^c$ is our initial ELBO of the (C)VAE. This scheme, the Modified Differential Multiplier Method (MDMM), results in more robust behaviour, both from a theoretical perspective (Platt and Barr, 1988) as well as supported by empirical evidence as shown in our Figure 2. MDMM is used throughout our numerical experiments.

**Enforcing multiple identifiability constraints:**
Next we discuss how to handle multiple constraints, illustrating this on a special case with a two-dimensional input $(x_1, x_2)$. In order to satisfy the constraints $\int f^{\theta}(x_1, x_2)dx_2$ for every value $x_1$ in some interval, we need to introduce a Lagrange multiplier $\lambda_{x_1}(x_1)$ which is now indexed by a continuous-valued $x_1$. The additional penalty corresponding to this term will be $\int \lambda_{x_1}(x_1) \left( \int f^{\theta}_2(x_1, x_2)dx_2 \right) dx_1$. Similarly, the ELBO will also be augmented by $\int \lambda_{x_2}(x_2) \left( \int f^{\theta}_1(x_1, x_2)dx_1 \right) dx_2$ in addition to penalty terms $\lambda_1 \int f^{\theta}_2(x_1)dx_1$ and $\lambda_2 \int f^{\theta}_1(x_2)dx_2$.

In practice, we can choose to estimate these integrals using either quadrature or Monte Carlo estimates.

One might wonder how we know whether the constraints have been (approximately) satisfied. We propose to approach this as follows: we establish a desired tolerance threshold $c$ and evaluate the integrals after optimisation to make sure that all learned NNs have been constrained to the desired functional subspaces within the desired tolerance. We point out that all the additional penalty terms need to be evaluated during training time only. This is important, because once the model has been trained, it can be used for prediction without any additional costs at test time.

### 3.3 Sparse Neural Decomposition

For interpreting what the decomposed (C)VAE has learnt, in addition to obtaining the variance decomposition it is also of practical interest to detect the presence or absence of dependence on certain input coordinates, e.g. detect groups of genes which depend purely on $z$ and not on $c$. This would make it easier to interpret.

The feature-level variance decomposition from ND does not explicitly identify such groups by default. One approach would be to apply an ad hoc thresholding to decide which effects are zero. For a more principled probabilistic approach, in this section we introduce Bernoulli random variables for every decoder network and for every data dimension. For the special case when $z, c \in \mathbb{R}$, we would introduce $s_z^{(j)}, s_c^{(j)}, s_{zc}^{(j)} \sim \text{Bernoulli}(p_0)$ for every feature $j = 1, \ldots, P$, each indicating the presence of $f^{\theta}_z, f^{\theta}_c$ and $f^{\theta}_{zc}$ non-zero effects for feature $j$. I.e. we define the decoder to have the following structure $y_i | z_i, c_i, \theta, s = s_z f^{\theta}_z(c_i) + s_c f^{\theta}_c(z_i) + s_{zc} f^{\theta}_{zc}(z_i, c_i) + \varepsilon_i$ where $s_z, s_c, s_{zc}$ have the same dimensionality as observations $y$. To implement variational inference via the reparameterisation trick, we use the continuous relaxation of Bernoulli random variables (Maddison et al., 2017; Jang et al., 2017). We specify a prior $p(s) = \text{RelaxedBernoulli}(p_0)$ for all $s_z, s_c, s_{zc}$ and approximating distributions $q(s) = \text{RelaxedBernoulli}(u)$ where $u$ are variational parameters. We use $p_0 = 0.1$ throughout.

The integration of the sparsity priors and the neural decomposition model is summarised in Figure 3.

![Figure 3: Diagram of the Neural Decomposition decoding model. For every gene in panel (a), we learn an identifiable decomposition of decoder networks $f^{\theta}_z(z_i), f^{\theta}_c(c_i)$ and $f^{\theta}_{zc}(z_i, c_i)$ as shown in panel (c), which are then elementwise multiplied with the respective Bernoulli sparsity masks (panel (b)) for easier interpretability.](image)

### 4 Related work

From the application perspective, Märtens et al. (2019) have considered a similar problem setting, but they relied on Gaussian Processes (GPs) as opposed to neural networks. While in the GP framework we can enforce functional constraints elegantly in closed form, its scalability properties are different. We have empirically found ND to be much more scalable to high-dimensional data, both in terms of compute and memory (see Supplementary for details).

There has been substantial interest towards explaining black-box models. A common strategy is to approximate the neural network with a simpler, interpretable model. Such approaches can be divided into global (Tan et al., 2018) and local explanations (Ribeiro et al., 2016; Lundberg and Lee, 2017). As opposed to such post hoc explanation methods, ND has decomposable structure built in as part of the model.
5 Experiments

The value of ND lies in scalable and identifiable functional decompositions. In real data, however, the true underlying functional decompositions are unknown, thus we are unable to quantify the correctness of the inferred decompositions on real data. For this reason, we seek to carefully characterise the behaviour of ND in a controlled setting before considering the real-life genomics example. Our implementation of ND is available in https://github.com/kasparmartens/NeuralDecomposition.

5.1 Synthetic data

We first investigate the behaviour of ND in a controlled setting. We generated synthetic data with a goal to mimic patterns in real gene expression data, thus including linear relationships, monotone warpings as well as non-monotone non-linear dependency structures (see Supplementary for details). We applied three variants of ND-CVAE: (i) with a linear decoder, (ii) non-linear decoder without identifiability constraints and (iii) our full implementation.

Figure 4 illustrates the inferred variance decompositions for 25 features with varying dependency structures (e.g. the first five features exhibit a pure additive effect). Neither the linear-CVAE (panel (B)) nor the ND-CVAE without constraints (panel (C)) have been able to identify the correct decomposition: the former is unable to capture non-linearities in the data due to its restrictive modelling assumptions, and the latter suffers from unidentifiability which makes its inferred variance decomposition arbitrary. Supp Table 1 quantifies performance over repeated experiments.

To demonstrate that our approach is not restricted to a univariate z, we have included an experiment using a two-dimensional latent space (see batch correction example in Supplementary).

5.2 CelebA

The motivation behind ND is understanding tabular (non-image) data – this is the reason why we learn the decomposition on the level of observed features. However, the ND methodology is generally applicable to a variety of large-scale problems, and in this section we demonstrate ND on CelebA data (Liu et al., 2015).4

5.2.1 Pixel-level decomposition

To be able to capture subtle effects like “glasses” and “beard” which are typically not captured in the inferred latent space of the VAE (see e.g. empirical results in (Kim and Mnih, 2018)), we include these as fixed covariates. Thus, we consider a CVAE with four covariates $c_1=$ Gender, $c_2=$ Smiling, $c_3=$ Glasses, $c_4=$ Beard and a 1D latent variable $z$. By performing a pixel-level decomposition with our ND with a functional form $f_θ^z(z) + \sum_k f_θ^{cz}(z, c_k)$ where $k \in \{1, 2, 3, 4\}$, thus capturing the marginal additive effects of $z$ and $\{c_k\}$ as well as their interactions as part

4Our interest is not to tune ND to develop a state-of-the-art model for images (for this purpose, we would e.g. use CNNs within our CVAE), but instead to demonstrate ND on an easy-to-visualise high-dimensional use case. 

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Figure 4: The true variance decomposition $\text{Var}(f_z)$, $\text{Var}(f_c)$, $\text{Var}(f_{zc})$ on a synthetic data set shown in (A1-A3) for all 25 features (which have been grouped in blocks of 5 for visual aid), compared to the inferred decompositions by (B) linear-CVAE, (C) unidentifiable ND, and (D) identifiable ND.
of the decoding structure. The inferred pixel-level sparsity masks in Figure 5 help us interpret what each of the decoder networks has learnt. Note that \( z \) captures how dark/light the background is, whereas the additive effects of covariates affect only relevant parts of the faces (e.g. glasses or beard). The interaction effects between \( z \) and \( c_k \) are relatively small: they mostly affect pixels that interact with the overall background darkness, e.g. long hair for women, or the outer borders of glasses and beard.

However, the additive structure in ND has led to visually better quality images. The same is indicated by the loglikelihoods on held-out images of smiling men: for CVAE (-3181 ± 630) these are lower than for ND (-2497 ± 319).

5.3 Gene expression

We next examine a publicly available time-series single-cell RNA-seq (scRNA-seq) data set of bone marrow derived dendritic cells responding to particular stimuli (Shalek et al., 2014). In this experiment, cells were exposed to either LPS (a component of Gram-negative bacteria) or PAM (a synthetic mimic of bacterial lipopeptides), and scRNA-seq was performed at 1, 2, 4 and 6 h after stimulation. With the capture time information, the original study studied single-cell gene expression dynamics under the two exposures, however, the cells behave asynchronously and heterogeneity exists within the cellular populations at each time point (i.e. after 1 hour of stimulation, the cells do not reach exactly the same biochemical state). Previous analyses have suggested this data set is more suited to a latent variable (so called pseudotime) analysis (Campbell and Yau, 2018) where the capture times are treated as unreliable and the latent variable is used instead to capture a continuous measure of how the cells are continually changing in response to each stimuli (Figure 7).

Figure 5: Inferred sparsity masks on CelebA data.

5.2.2 Improved extrapolation

Despite the empirical success of deep generative models, their generalization (e.g. to under-represented subpopulations) in the presence of sampling bias is still an active area of research (Zhao et al., 2018). Here, we demonstrate how additive structure in the decoder can improve the extrapolation properties of the CVAE. We use CelebA data to mimic sampling bias as follows: we exclude all images of all smiling men from the training set, and at test time, we visualise predictions, both for CVAE and ND-CVAE, as shown in Figure 6,

- First, using \( z = 0 \), Gender=Male, Smiling=False, Glasses=False, Beard=False, i.e. a scenario that was observed during training
- New scenarios Smiling=True (2nd panel), with additionally Glasses (3rd) and Beard (4th panel)

Figure 6: Generalization to unseen subgroups.

It is not surprising that CVAE has not been able to generate clear images for unseen subgroups: the decoder has not been trained on these particular inputs.

We conducted an analysis using ND where we encoded the stimulant to which the cells were exposed as a binary covariate \( (c \in \{\text{LPS}, \text{PAM}\}) \) and used a single dimension for the latent variable \( z \). Each gene was therefore modelled as a combination of main effects due to 1) LPS/PAM exposure, 2) temporal effects (independent of stimulant type), and 3) temporal interaction effects that were modulated by the stimulant.

Figure 7: Above: Schema illustrating pseudotime analysis approach to mouse dendritic cells. Below: Inferred \( z \) (pseudotime) correlates with capture time.
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Figure 8: ND on the large-scale scRNA-seq data of dendritic cells, using \( c \in \{ \text{LPS, PAM} \} \), lets us decompose gene-level variation (shown for selected 3 genes) into the marginal \( z \) effect, the marginal \( c \) effect, and the non-linear interaction between the two, together with the fraction of variance attributed to each component. As opposed to earlier linear approaches (Campbell and Yau, 2018), the decomposition provided by ND is non-linear.

We applied ND to 820 cells using the 7,500 most variable genes, following the previously published linear analysis (Campbell and Yau, 2018). Our inferred latent dimension was indeed correlated with capture time and therefore is indeed a measure of the continuous progression of these cells under stimulation (Figure 7).

Furthermore, we were able to decompose the expression of each gene into components that were dependent on non-linear additive effects of pseudotime \( z \), covariate \( c \), and interactions \((z, c)\). In Figure 8, we highlight three genes previously identified to exhibit interaction effects. However, a limitation of the previous analysis was the reliance on linear models, which leaves open the possibility of model misspecification driving false positives in this very noisy single cell expression data. Here we have used a more flexible, non-linear CVAE-based approach. The genes \( \text{Tnf, Rasgefb1} \) and \( \text{Tnfaip3} \) all exhibited strong interaction effects and the dependence of the gene expression on \((z, c)\) does follow a near-linear behaviour even under a flexible non-linear model, thus confirming the findings of (Campbell and Yau, 2018) under a more flexible class of models, while providing a more accurate feature-level variance decomposition (rightmost panels in Figure 8).

We next identified the top 50 genes with the strongest 1) additive \( z \), 2) additive \( c \), and 3) interactions effects as identified by ND, and applied UMAP, a popular dimensionality reduction and visualisation algorithm, to examine these subsets in a two-dimensional representation (Supp Fig 1). As expected, the set of genes with strong additive \( z \) effects but smaller additive \( c \) and interaction effects, showed considerable intermixing between cells stimulated under the two conditions. These genes exhibit behaviours that are largely independent of stimulus so we would not expect segregation of the genes by stimulus type. In contrast, the UMAP visualisation of the gene sets with strong additive \( c \) or interaction effects shows considerable separation between the LPS and PAM stimulated cells. These are genes whose behaviour is heavily influenced by the type of stimulation used as well as the latent variable.

Overall, this suggests that ND-CVAE is identifying relevant structure in the single cell data and correctly attributing the appropriate feature behaviours to the relevant subsets of genes. Furthermore, unlike previous analyses, our flexible non-linear models permits greater robustness to model misspecification.

6 Discussion

We have proposed a VAE-framework where we embed functional ANOVA decompositions within the decoding structure. This specification allows us to associate variation in the latent space and other covariates to feature-level variability, leading to interpretability that is not present in existing (C)VAE models. Our work brings together ideas from classical statistics and constrained optimisation, while leveraging modern deep learning software to develop comparatively fast, scalable, and interpretable models for dimensionality reduction.

The ND construction makes the functional decomposition (with fixed inputs) identifiable. In principle, the entire model could be made identifiable, when combining ND with monotonicity constraints on the neural networks (Pierson et al., 2019), however, this would considerably restrict the flexibility of the model.
Acknowledgements

KM was supported by a UK Engineering and Physical Sciences Research Council Doctoral Studentship. CY is supported by a UK Medical Research Council Research Grant (Ref: MR/P02646X/1) and by The Alan Turing Institute under the EPSRC grant EP/N510129/1.

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