Deep Structural Causal Models for Tractable Counterfactual Inference

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Abstract

We formulate a general framework for building structural causal models (SCMs) with deep learning components. The proposed approach employs normalising flows and variational inference to enable tractable inference of exogenous noise variables—a crucial step for counterfactual inference that is missing from existing deep causal learning methods. Our framework is validated on a synthetic dataset built on MNIST as well as on a real-world medical dataset of brain MRI scans. Our experimental results indicate that we can successfully train deep SCMs that are capable of all three levels of Pearl’s ladder of causation: association, intervention, and counterfactuals, giving rise to a powerful new approach for answering causal questions in imaging applications and beyond. The code for all our experiments is available at https://github.com/biomedia-mira/deepscm.

1 Introduction

Many questions in everyday life as well as in scientific inquiry are causal in nature: “How would the climate have changed if we’d had less emissions in the ’80s?”, “How fast could I run if I hadn’t been smoking?”, or “Will my headache be gone if I take that pill?”. None of those questions can be answered with statistical tools alone, but require methods from causality to analyse interactions with our environment (interventions) and hypothetical alternate worlds (counterfactuals), going beyond joint, marginal, and conditional probabilities [1]. Even though these are natural lines of reasoning, their mathematical formalisation under a unified theory is relatively recent [2].

In some statistics-based research fields, such as econometrics or epidemiology, the use of causal inference methods has been established for some time [3, 4]. However, causal approaches have been introduced into deep learning (DL) only very recently [5]. For example, research has studied the use of causality for disentanglement [6, 7], causal discovery [8, 9], and for deriving causality-inspired explanations [10, 11] or data augmentations [12]. Causal DL models could be capable of learning relationships from complex high-dimensional data and of providing answers to interventional and counterfactual questions, although current work on deep counterfactuals is limited by modelling only direct cause-effect relationships [11] or instrumental-variable scenarios [13], or by not providing a full recipe for tractable counterfactual inference [14].

The integration of causality into DL research promises to enable novel scientific advances as well as to tackle known shortcomings of DL methods: DL is known to be susceptible to learning spurious correlations and amplifying biases [e.g. 15], and to be exceptionally vulnerable to changes in the input distribution [16]. By explicitly modelling causal relationships and acknowledging the difference between causation and correlation, causality becomes a natural field of study for improving the transparency, fairness, and robustness of DL-based systems [17, 18]. Further, the tractable inference of deep counterfactuals enables novel research avenues that aim to study causal reasoning on a

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per-instance rather than population level, which could lead to advances in personalised medicine as well as in decision-support systems, more generally.

In this context, our work studies the use of DL-based causal mechanisms and establishes effective ways of performing counterfactual inference. Our main contributions are: 1) a unified framework for structural causal models using modular deep mechanisms; 2) an efficient approach to estimating counterfactuals by inferring exogenous noise via variational inference or normalising flows; 3) case studies exemplifying how to apply deep structural causal models and perform counterfactual inference. The paper is organised as follows: we first review structural causal models and discuss how to leverage deep mechanisms and enable tractable counterfactual inference. Second, we compare our work to recent progress in deep causal learning in light of Pearl’s ladder of causation [19]. Finally, we apply deep structural causal models to a synthetic experiment as well as to modelling brain MRI scans, demonstrating the practical utility of our framework in answering counterfactual questions.

2 Deep Structural Causal Models

We consider the problem of modelling a collection of $K$ random variables $x = (x_1, \ldots, x_K)$. By considering causal relationships between them, we aim to build a model that not only is capable of generating convincing novel samples, but also satisfies all three rungs of the causation ladder [19].

The first level, association, describes reasoning about passively observed data. This level deals with correlations in the data and questions of the type “What are the odds that I observe...?”, which relates purely to marginal, joint, and conditional probabilities. Intervention concerns interactions with the environment. It requires knowledge beyond just observations, as it relies on structural assumptions about the underlying data-generating process. Characteristic questions ask about the effects of certain actions: “What happens if I do...?” Lastly, counterfactuals deal with retrospective hypothetical scenarios. Counterfactual queries leverage functional models of the generative processes to imagine alternative outcomes for individual data points, answering “What if I had done A instead of B?”. Arguably, such questions are at the heart of scientific reasoning (and beyond), yet are less well-studied in the field of machine learning. The three levels of causation can be operationalised by employing structural causal models (SCMs)\(^2\), recapitulated in the next section.

2.1 Background on structural causal models

A structural causal model $\mathcal{G} := (\mathcal{S}, P(\epsilon))$ consists of a collection $\mathcal{S} = (f_1, \ldots, f_K)$ of structural assignments $x_k := f_k(\epsilon_k; \text{pa}_k)$ (called mechanisms), where $\text{pa}_k$ is the set of parents of $x_k$ (its direct causes), and a joint distribution $P(\epsilon) = \prod_{k=1}^K P(\epsilon_k)$ over mutually independent exogenous noise variables (i.e. unaccounted sources of variation). As assignments are assumed acyclic, relationships can be represented by a directed acyclic graph (DAG) with edges pointing from causes to effects, called the causal graph induced by $\mathcal{G}$. Every SCM $\mathcal{G}$ entails a unique joint observational distribution $P_\mathcal{G}(x)$, satisfying the causal Markov assumption: each variable is independent of its non-effects given its direct causes. It therefore factorises as $P_\mathcal{G}(x) = \prod_{k=1}^K P(\epsilon_k) P_\mathcal{G}(x_k | \text{pa}_k)$, where each conditional distribution $P_\mathcal{G}(x_k | \text{pa}_k)$ is determined by the corresponding mechanism and noise distribution [1].

Crucially, unlike conventional Bayesian networks, the conditional factors above are imbued with a causal interpretation. This enables $\mathcal{G}$ to be used to predict the effects of interventions, defined as substituting one or multiple of its structural assignments, written as ‘$\text{do}(\cdots)$’. In particular, a constant reassignment of the form $\text{do}(x_k := a)$ is called an atomic intervention, which disconnects $x_k$ from all its parents and represents a direct manipulation disregarding its natural causes.

While the observational distribution relates to statistical associations and interventions can predict causal effects, SCMs further enable reasoning about counterfactuals. These are hypothetical retrospective interventions, given an observed outcome: ‘What would $x_i$ have been if $x_j$ were different, given that we observed $x$?’ This type of question effectively offers explanations of the data, since we can analyse the changes resulting from manipulating each variable. Counterfactual queries can be mathematically formulated as a three-step procedure [2, Ch. 7]:

1. **Abduction:** Predict the ‘state of the world’ (the exogenous noise, $\epsilon$) that is compatible with the observations, $x$, i.e. infer $P_{\mathcal{G}}(\epsilon | x)$.

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\(^2\)SCMs are also known as (nonlinear) structural equation models or functional causal models.
2. Action: Perform an intervention (e.g. \( \text{do}(x_k := \tilde{x}_k) \)) corresponding to the desired manipulation, resulting in a modified SCM \( \tilde{G} = \tilde{G}_{x; \text{do}(\tilde{x}_k)} \) [1, Sec. 6.4].

3. Prediction: Compute the quantity of interest based on the distribution entailed by the counterfactual SCM, \( P_{\tilde{G}}(\tilde{x}) \).

With these operations in mind, the next section explores a few options for building flexible, expressive, and counterfactual-capable functional mechanisms for highly structured data.

### 2.2 Deep mechanisms

In statistical literature (e.g. epidemiology, econometrics, sociology), SCMs are typically employed with simple linear mechanisms (or generalised linear models, involving an output non-linearity). Analysts attach great importance to the regression weights, as under certain conditions these may be readily interpreted as estimates of the causal effects between variables. While this approach generally works well for scalar variables and can be useful for decision-making, it is not flexible enough to model higher-dimensional data such as images. Solutions to this limitation have been proposed by introducing deep-learning techniques into causal inference [8, 14].

We call an SCM that uses deep-learning components to model the structural assignments a **deep structural causal model** (DSCM). In DSCMs, the inference of counterfactual queries becomes more complex due to the potentially intractable abduction step (inferring the posterior noise distribution, as defined above). To overcome this, we propose to use recent advances in normalising flows and variational inference to model mechanisms for composable DSCMs that enable tractable counterfactual inference. While here we focus on continuous data, DSCMs also fully support discrete variables without the need for relaxations (see Appendix C). We consider three types of mechanisms that differ mainly in their invertibility, illustrated in Fig. 1.

**Invertible, explicit:** Normalising flows model complex probability distributions using transformations from simpler base distributions with same dimensionality [20]. For an observed variable \( x \), diffeomorphic transformation \( f \), and base variable \( \epsilon \sim P(\epsilon) \), the output density \( p(x) = p(\epsilon) |\text{det} \nabla f(\epsilon)|^{-1} \), evaluated at \( \epsilon = f^{-1}(x) \) [21, 22]. For judicious choices of \( f \), the Jacobian \( \nabla f \) may take special forms with efficiently computable determinant, providing a flexible and tractable probabilistic model whose parameters can be trained via exact maximum likelihood. Furthermore, flows can be made as expressive as needed by composing sequences of simple transformations. For more information on flow-based models, refer to the comprehensive survey by Papamakarios et al. [22]. Note that this class of models also subsumes the typical location-scale and inverse cumulative distribution function transformations used in the reparametrisation trick [23, 24], as well as the Gumbel trick for discrete variable relaxations [25, 26].

Although normalising flows were originally proposed for unconditional distributions, they have been extended to conditional densities [27], including in high dimensions [28, 29], by parametrising the transformation as \( x = f(\epsilon; \text{pa}_X) \), assumed invertible in the first argument. In particular, conditional
flows can be adopted in DSCMs to represent invertible, explicit-likelihood mechanisms (Fig. 1a):

\[ x_i := f_i(\epsilon_i; \mathbf{p}_a), \quad p(x_i \mid \mathbf{p}_a) = p(\epsilon_i) \cdot |\text{det} \nabla_{\epsilon_i} f_i(\epsilon_i; \mathbf{p}_a)|^{-1} \bigg|_{\epsilon_i = f_i^{-1}(x_i; \mathbf{p}_a)}. \] (1)

**Amortised, explicit:** Such invertible architectures typically come with heavy computational requirements when modelling high-dimensional observations, because all intermediate operations act in the space of the data. Instead, it is possible to use arbitrary functional forms for the structural assignments, at the cost of losing invertibility and tractable likelihoods \( p(x_k \mid \mathbf{p}_k) \). Here, we propose to separate the assignment \( f_k \) into a ‘low-level’, invertible component \( h_k \) and a ‘high-level’, non-invertible part \( g_k \)—with a corresponding noise decomposition \( \epsilon_k = (u_k, z_k) \)—such that

\[ x_k := f_k(\epsilon_k; \mathbf{p}_k) = h_k(u_k; g_k(z_k; \mathbf{p}_k), \mathbf{p}_k), \quad P(\epsilon_k) = P(u_k)P(z_k). \] (2)

In such a decomposition, the invertible transformation \( h_k \) can be made shallower, while the upstream non-invertible \( g_k \) maps from a lower-dimensional space and is expected to capture more of the high-level structure of the data. Indeed, a common implementation of this type of model for images would involve a probabilistic decoder, where \( g_k \) may be a convolutional neural network, predicting the parameters of a simple location-scale transformation performed by \( h_k \) [24].

As the conditional likelihood \( p(x_k \mid \mathbf{p}_k) \) in this class of models is no longer tractable because \( z_k \) cannot be marginalised out, it may alternatively be trained with amortised variational inference. Specifically, we can introduce a variational distribution \( Q(z_k \mid x_k, \mathbf{p}_k) \) to formulate a lower bound on the true marginal conditional log-likelihood, which will be maximised instead:

\[ \log p(x_k \mid \mathbf{p}_k) \geq \mathbb{E}_{Q(z_k \mid x_k, \mathbf{p}_k)}[\log p(x_k \mid z_k, \mathbf{p}_k)] - D_{\text{KL}}[Q(z_k \mid x_k, \mathbf{p}_k) \mid \mid P(z_k)]. \] (3)

The argument of the expectation in this lower bound can be calculated similarly to Eq. (1):

\[ p(x_k \mid z_k, \mathbf{p}_k) = p(u_k) \cdot |\det \nabla_{u_k} h_k(u_k; g_k(z_k, \mathbf{p}_k), \mathbf{p}_k)|^{-1} \bigg|_{u_k = h_k^{-1}(x_k; g_k(z_k, \mathbf{p}_k), \mathbf{p}_k)}. \] (4)

The approximate posterior distribution \( Q(z_k \mid x_k, \mathbf{p}_k) \) can for example be realised by an encoder function, \( e_k(x_k, \mathbf{p}_k) \), that outputs the parameters of a simple distribution over \( z_k \) (Fig. 1b), as in the auto-encoding variational Bayes (AEVB) framework [24].

**Amortised, implicit:** While the models above rely on (approximate) maximum-likelihood as training objective, it is admissible to train a non-invertible mechanism as a conditional implicit-likelihood model (Fig. 1c), optimising an adversarial objective [30–32]. Specifically, a deterministic encoder \( e_j \) would strive to fool a discriminator function attempting to tell apart tuples of encoded real data \((x_j, e_j(x_j); \mathbf{p}_j), \mathbf{p}_j)\) and generated samples \((f_j(\epsilon_j; \mathbf{p}_j), \epsilon_j, \mathbf{p}_j)\).

### 2.3 Deep counterfactual inference

Now equipped with effective deep models for representing mechanisms in DSCMs, we discuss the inference procedure allowing us to compute answers to counterfactual questions.

**Abduction:** As presented in Section 2.1, the first step in computing counterfactuals is abduction, i.e. to predict the exogenous noise, \( \epsilon \), based on the available evidence, \( x \). Because each noise variable is assumed to affect only the respective observed variable, \( (\epsilon_k)_{k=1}^K \) are conditionally independent given \( x \), therefore this posterior distribution factorises as \( P_\Theta(\epsilon \mid x) = \prod_{k=1}^K P_\Theta(\epsilon_k \mid x_k, \mathbf{p}_k) \). In other words, it suffices to infer the noise independently for each mechanism, given the observed values of the variable and of its parents\(^3\).

For invertible mechanisms, the noise variable can be obtained deterministically and exactly by just inverting the mechanism: \( \epsilon_i = f_i^{-1}(x_i; \mathbf{p}_a) \). Similarly, implicit-likelihood mechanisms can be approximately inverted by using the trained encoder function: \( \epsilon_j \approx e_j(x_j; \mathbf{p}_j) \).

Some care must be taken in the case of amortised, explicit-likelihood mechanisms, as the ‘high-level’ noise \( z_k \) and ‘low-level’ noise \( u_k \) are not independent given \( x_k \). Recalling that this mechanism is

\(^3\)Note that here we assume full observability, i.e. no variables are missing when predicting counterfactuals. We discuss challenges of handling partial evidence in Section 6.
trained along with a conditional probabilistic encoder, \(Q(z_k | x_k; \mathbf{pa}_k)\), the noise posterior can be approximated as follows, where \(\delta_x(\cdot)\) denotes the Dirac delta distribution centred at \(w\):

\[
P_{\theta}(\epsilon_k | x_k, \mathbf{pa}_k) = P_{\theta}(z_k | x_k, \mathbf{pa}_k) P_{\theta}(u_k | z_k, x_k, \mathbf{pa}_k) \\
\approx Q(z_k | e_k(x_k; \mathbf{pa}_k)) \delta_{h_k^{-1}(x_k; g_k(z_k; \mathbf{pa}_k), \mathbf{pa}_k)}(u_k).
\]  

(5)

**Action:** The causal graph is then modified according to the desired hypothetical intervention(s), as in the general case (Section 2.1). For each intervened variable \(x_k\), its structural assignment is replaced either by a constant, \(x_k := \bar{x}_k\)—making it independent of its former parents (direct causes, \(\mathbf{pa}_k\)) and of its exogenous noise (\(\epsilon_k\))—or by a surrogate mechanism \(x_k := f_k(\epsilon_k; \mathbf{\overline{pa}}_k)\), forming a set of counterfactual assignments, \(\mathbf{\overline{S}}\). This then defines a counterfactual SCM \(\overline{S} = (\mathbf{S}, P_{\theta}(\epsilon | \mathbf{x}))\).

**Prediction:** Finally, we can sample from \(\mathbf{\overline{S}}\): Noise variables that were deterministically inverted (either exactly or approximately) can simply be plugged back into the respective forward mechanism to determine the new output value. Notice that this step is redundant for observed variables that are not descendants of the ones being intervened upon, as they will be unaffected by the changes.

As mentioned above, the posterior distribution over \((z_k, u_k)\) for an amortised, explicit-likelihood mechanism does not factorise (Eq. (5)), and the resulting distribution over the counterfactual \(x_k\) cannot be characterised explicitly. However, sampling from it is straightforward, such that we can approximate the counterfactual distribution via Monte Carlo as follows, for each sample \(s\):

\[
\tilde{z}_k^{(s)} \sim Q(z_k | e_k(x_k; \mathbf{pa}_k)) \\
u_k^{(s)} = h_k^{-1}(x_k; g_k(z_k^{(s)}; \mathbf{pa}_k), \mathbf{pa}_k) \\
\tilde{x}_k^{(s)} = \tilde{h}_k(u_k^{(s)}; \tilde{g}_k(z_k^{(s)}; \mathbf{\overline{pa}}_k), \mathbf{\overline{pa}}_k).
\]  

(6)

Consider an uncorrelated Gaussian decoder for images as a concrete example, predicting vectors of means and variances for each pixel of \(x_k\): \(g_k(z_k; \mathbf{pa}_k) = (\mu_k(z_k; \mathbf{pa}_k), \sigma^2_k(z_k; \mathbf{pa}_k))\). Exploiting the reparametrisation trick, counterfactuals that preserve \(x_k\)’s mechanism can be computed simply as

\[
u_k^{(s)} = (x_k - \mu_k(z_k^{(s)}; \mathbf{pa}_k)) \odot \sigma_k(z_k^{(s)}; \mathbf{pa}_k), \\
\tilde{x}_k^{(s)} = x_k + [\mu_k(z_k^{(s)}; \mathbf{\overline{pa}}_k) - \mu_k(z_k^{(s)}; \mathbf{pa}_k)].
\]

This showcases how true image counterfactuals are able to retain pixel-level details. Typical conditional generative models would output only \(\mu_k(z_k; \mathbf{\overline{pa}}_k)\) (which is often blurry in vanilla variational auto-encoders [33]), or would in addition have to sample \(P(u_k)\) (resulting in noisy images).

### 3 Related Work

Deep generative modelling has seen a wide range of contributions since the popularisation of variational auto-encoders (VAEs) [24], generative adversarial networks (GANs) [34], and normalising flows [21]. These models have since been employed to capture conditional distributions [27, 29, 32, 35], and VAEs and GANs were also extended to model structured data by incorporating probabilistic graphical models [36–38]. In addition, deep generative models have been heavily used for (unsupervised) representation learning with an emphasis on disentanglement [39–42]. However, even when these methods faithfully capture the distribution of observed data, they are capable of fulfilling only the association rung of the ladder of causation.

Interventions build on the associative capabilities of probabilistic models to enable queries related to changes in causal mechanisms. By integrating a causal graph into the connectivity of a deep model, it is possible to perform interventions with GANs [14] and causal generative NNs [8]. VAEs can also express causal links using specific covariance matrices between latent variables, which however restrict the dependences to be linear [6]. Despite reaching the second rung of the causal ladder, these methods lack tractable abduction capabilities and therefore cannot generate counterfactuals.
We quantitatively compare the associative capabilities of all models by evaluating their evidence. We use this setup to study the capabilities of our framework in comparison to models with less explicitly model associations, interventions, nor causal structure. Potentially closest to our work is a method for counterfactual explainability of visual models, which extends CausalGANs [14] to predict reparametrised distributions over image attributes following an assumed causal graph [10]. However, this approach performs no abduction step, instead resampling the noise of attributes downstream from the intervention(s), and does not include a generative model of imaging data. To the best of our knowledge, the proposed DSCM framework is the first flexible approach enabling end-to-end training and tractable inference on all three levels of the ladder of causation for high-dimensional data.

4 Case Study 1: Morpho-MNIST

We consider the problem of modelling the causal model of a synthetic dataset based on MNIST digits [47], where stroke thickness causes the brightness of the digit: thicker digits are brighter whereas thinner digits are dimmer. This simple dataset allows for examining the three levels of causation in a controlled and measurable environment. We use morphological transformations on MNIST [48] to generate a dataset with known causal structure and access to the ‘true’ process of generating counterfactuals. The SCM for this synthetic dataset is as follows:

\[
\begin{align*}
t &:= f_T(\epsilon_T^*) = 0.5 + \epsilon_T^*, & \epsilon_T^* &\sim \Gamma(10, 5) , \\
i &:= f_i(\epsilon_i^*; t) = 191 \cdot \sigma(0.5 \cdot \epsilon_i^* + 2 \cdot t - 5) + 64 , & \epsilon_i^* &\sim \mathcal{N}(0, 1) , \\
x &:= f_X(\epsilon_X^*; i, t) = \text{SetIntensity}(\text{SetThickness}(\epsilon_X^*; i); i) , & \epsilon_X^* &\sim \text{MNIST} ,
\end{align*}
\]

where SetIntensity(\cdot; i) and SetThickness(\cdot; t) refer to the operations that act on an image of a digit and set its intensity to i and thickness to t (see Appendix A.1 for details), x is the resulting image, \(\epsilon^*\) is the exogenous noise for each variable and \(\sigma(\cdot)\) is the logistic sigmoid.

We use this setup to study the capabilities of our framework in comparison to models with less causal structure. We adapt the true causal graph from Eq. (7) and model thickness and intensity using (conditional) normalising flows and employ a conditional VAE for modelling the image. In particular, we adopt the causal graphs shown in Fig. 2 and test a fully independent model (Fig. 2a), a conditional decoder model (Fig. 2b), as well as our full causal model (Fig. 2c). All our experiments were implemented within PyTorch [49] using the Pyro probabilistic programming framework [50], and implementation details can be found in Appendices A.2 and B.2.

We quantitatively compare the associative capabilities of all models by evaluating their evidence lower bound (Eq. (3)), log-likelihoods and reconstruction errors as shown in Table 1. We find that performance improves consistently with the model’s capabilities: enabling conditional image generation improves \(p(x|t, i)\), and adding a causal dependency between \(t\) and \(i\) improves \(p(i|t)\). Further, we examine samples of the conditional and unconditional distributions in Appendix A.3.1.

The interventional distributions can be directly compared to the true generative process. Figure 3 shows that the densities predicted by our full model after intervening on \(t\) closely resemble the
Table 1: Comparison of the associative abilities of the models shown in Fig. 2. The image is denoted by $x$, thickness by $t$, and intensity by $i$. Quantities with $\geq$ are lower bounds. MAE refers to the mean absolute error between pixels of the original image and of its reconstruction.

| Model     | $\log p(x, t, i)$ $\geq$ | $\log p(x | t, i)$ $\geq$ | $\log p(t)$ $\geq$ | $\log p(i | t)$ | $\text{MAE}(x, x')$ |
|-----------|---------------------------|---------------------------|-------------------|----------------|----------------------|
| Independent | -5925.26                  | -5919.14                 | -0.93            | -5.19         | 4.50                 |
| Conditional | -5526.50                  | -5520.37                 | -0.93            | -5.19         | 4.26                 |
| Full       | -5692.94                  | -5687.71                 | -0.93            | -4.30         | 4.43                 |

Figure 3: Distributions of thickness and intensity in the true data (left), and learned by the full (centre) and conditional (right) models. Contours depict the observational (red, shaded) and interventional joint densities for $\text{do}(t := f_T(\epsilon_T) + 1)$ (blue, solid) and $\text{do}(t := f_T(\epsilon_T) - 0.5)$ (green, dashed).

Figure 4: Counterfactuals generated by the full model. (left) Counterfactual ‘trajectories’ of two original samples, A and B, as their thickness and intensity are modified, overlaid on the learned joint density $p(t, i)$. (right) Original and counterfactual images corresponding to samples A and B.

true behaviour. The conditional and independent models operate equivalently and are incapable of modelling the relationship between $t$ and $i$, capturing only their marginal distributions.

Lastly, we examine the full model’s ability to generate counterfactuals. The other two models were omitted as they are incapable of accomplishing interventions, a prerequisite for counterfactual inference. Examples of previously unseen images and generated counterfactuals are shown in Fig. 4. We see that our model is capable of generating convincing counterfactuals that preserve the digit identity while changing thickness and intensity consistently with the underlying causal model.

5 Case Study 2: Brain Imaging

Our real-world application touches upon fundamental scientific questions in the context of medical imaging: how would a person’s anatomy change if particular traits were different? We illustrate with a (simplified) example that our DSCM framework may provide the means to answer such counterfactual
queries, which may enable entirely new research into better understanding the physical manifestation of lifestyle, demographics, and disease. Here, we model the appearance of brain MRI scans given the person’s age and biological sex, as well as brain and ventricle volumes, using population data from the UK Biobank [51]. Ventricle and total brain volumes are two quantities that are closely related to brain age [52] and can be observed relatively easily. We adopt the causal graph shown in Fig. 5a and otherwise follow the same training procedure as for the MNIST experiments.

The learned DSCM is capable of all three levels of the causal hierarchy. We present the analysis of lower levels in Appendix B.3.1 and focus here on counterfactuals, shown in Fig. 5b (more examples in Appendix B.3.2). The difference maps show plausible counterfactual changes: increasing age causes slightly larger ventricles while decreasing the overall brain volume (first column). In contrast, directly changing brain volume has an opposite effect on the ventricles compared to changing age (second column). Intervening on ventricle volume has a much more localised effect (third column), while intervening on the categorical variable of biological sex has smaller yet more diffuse effects. Note how the anatomical ‘identity’ (such as the cortical folding) is well preserved after each intervention.

6 Conclusion

We introduce a novel general framework for fitting SCMs with deep mechanisms. Our deep SCM (DSCM) framework fulfils all three rungs of Pearl’s ladder of causation—in particular, it is the first to enable efficient abduction of exogenous noise, permitting principled counterfactual inference. We demonstrate the potential of DSCMs with two case studies: a synthetic task of modelling Morpho-MNIST digits with a known causal structure and a real-world example with brain MRI.

The ability to correctly generate plausible counterfactuals could greatly benefit a wide variety of possible applications, e.g.: explainability, where differences between observed and counterfactual data can suggest causal explanations of outcomes; data augmentation, as counterfactuals can extrapolate beyond the range of observed data (e.g. novel combinations of attributes); and domain adaptation, since including the source of the data as an indicator variable in the causal model could enable generating counterfactual examples in a relevant target domain.

The proposed method does not come without limitations to be investigated in future work. Like the related approaches, the current setup requires all variables to be observed when computing a counterfactual, which may limit its applicability in certain scenarios. This could be alleviated by imputing the missing data via MCMC or learning auxiliary distributions. Further work should study more closely the dynamic behaviour of deep mechanisms in SCMs. While not observed in our experiments, neural networks may not learn to cleanly separate the roles of its inputs on the output.

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4 Ventricles are fluid-filled cavities identified as the dark areas in the centre of the brain.
as expected—which could require custom counterfactual regularisation similar to losses used in image-to-image translation [46] and explainability work [11]. The use of such flexible models also raises questions about the identifiability of the ‘true’ mechanism, as counterfactuals may not be uniquely defined. Lastly, it would be interesting to examine whether this framework can be applied to causal discovery, attempting to uncover plausible causal structures from data.

**Broader Impact**

Causal inference can be applied to a wide range of applications, promising to provide a deeper understanding of the observed data and prevent the fitting of spurious correlations. Our research presents a methodological contribution to the causal literature proposing a framework that combines causal models and deep learning to facilitate modelling high-dimensional data.

Because of the general applicability of deep learning and causal inference, our framework could have a broad impact of enabling fairer machine learning models explicitly modelling causal mechanisms, reducing spurious correlations and tackling statistical and societal biases. The resulting models offer better interpretability due to counterfactual explanations and could yield novel understanding through causal discovery.

However, causal modelling relies on strong assumptions and cannot always unambiguously determine the true causal structure of observational data. It therefore is necessary to carefully consider and communicate the assumptions being made by the analyst. In this light, our methodology is susceptible to being used to wrongly claim the discovery of causal structures due to careless application or intentional misuse. Particularly, the use of ‘black-box’ components as causal mechanisms may exacerbate concerns about identifiability, already present even for simple linear models. Whereas deep causal models can be useful for deriving insights from data, we must be cautious about their use in consequential decision-making, such as in informing policies or in the context of healthcare.

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A Synthetic Morpho-MNIST Experiment

A.1 Data Generation

We use the original MNIST dataset [47] together with the morphometric measurements introduced with Morpho-MNIST [48] to add functionality to measure intensity as well as set the intensity and thickness to a given value.

We implement \texttt{MeasureIntensity} by following the processing steps proposed by Castro et al. [48], and measure the intensity $i$ of an image as the median intensity of pixels within the extracted binary mask. Once the intensity is measured, the entire image is rescaled to match the target intensity, with values clamped between 0 and 255 (images are assumed to be in unsigned 8-bit format).

Originally, Morpho-MNIST only proposed relative thinning and thickening operations. We expand those operations to absolute values by calculating the amount of dilation or erosion based on the ratio between target thickness and measured thickness.

Finally, we follow Eq. (7) to modify each image within the MNIST dataset and randomly split the original training set into a training and validation set. We show random samples from the resulting test set in Fig. A.1.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{example.png}
\caption{Random exemplars from the synthetically generated Morpho-MNIST test dataset}
\end{figure}

\subsection*{A.2 Experimental Setup}

We use (conditional) normalising flows for all variables apart from the images, which we model using (conditional) deep encoder-decoder architectures. The flows consist of components that constrain the support of the output distribution (where applicable) and components relevant for fitting the distribution. We use unit Gaussians as base distributions for all exogenous noise distributions $P(\epsilon)$ and, if available, we use the implementations in PyTorch [49] or Pyro [50] for all transformations. Otherwise, we adapt the available implementations, referring to [53] for details. We indicate with $\theta$ the modules with learnable parameters.
We train all models for 1000 epochs and report the results of the model with the best validation loss.

We implement the decoder network as a CNN:

\[ t := f_T(\epsilon_T) = (\exp \odot \text{AffineNormalisation} \odot \text{Spline}_{\theta}(\epsilon_T), \quad (A.1) \]
\[ i := f_I(\epsilon_I; t) = (\text{AffineNormalisation} \circ \text{sigmoid} \circ \text{ConditionalAffine}_{\theta}(t)) \circ (\epsilon_I). \quad (A.2) \]

In the independent model, where \( i \) is not conditioned on \( t \), we use instead

\[ i := f_I(\epsilon_I) = (\text{AffineNormalisation} \circ \text{sigmoid} \circ \text{Spline}_{\theta} \circ \text{Affine}_{\theta})(\epsilon_I). \quad (A.3) \]

We found that including normalisation layers help learning dynamics\(^5\) and therefore include flows to perform commonly used normalisation transformations. For doubly bounded variable \( y \) we learn the flows in unconstrained space and then constrain them by a sigmoid transform and rescale to the original range using fixed affine transformations with bias \( \min(Y) \) and scale \( [\max(Y) - \min(Y)] \). We constrain singly bounded values by applying an exponential transform to the unbounded values and using an affine normalisation equivalent to a whitening operation in unbounded log-space. We denote those fixed normalisation transforms as AffineNormalisation and use a hat to refer to the unconstrained, normalised values (e.g. \( \widehat{\text{pa}}_k \)). The Spline\( \theta \) transformation refers to first-order neural spline flows \( [53] \). Affine\( \theta \) is an element-wise affine transformation, and sigmoid refers to the logistic function. ConditionalAffine\( \theta(\cdot) \) is a regular affine transform whose transformation parameters are predicted by a context neural network taking \( \cdot \) as input. In the case of \( f_I(\epsilon_I; t) \), the context network is represented by a simple linear transform. Further, we model \( x \) using a low-level flow:

\[ h_x(u_X; \text{pa}_X) = \left[ \text{Preprocessing} \circ \text{ConditionalAffine}_{\theta}(\widehat{\text{pa}}_X) \right](u_X), \quad (A.4) \]

where the ConditionalAffine transform practically reparametris the noise distribution into another Gaussian distribution and Preprocessing describes a fixed preprocessing transformation. We follow the same preprocessing as used with RealNVP \([54]\). The context network for the conditional affine transformation is the high-level mechanism \( g_X(z_X; \text{pa}_X) \) and is implemented as a decoder network that outputs the bias for of the affine transformation, while the log-variance is fixed to \( \log \sigma^2 = -5 \). We implement the decoder network as a CNN:

\[
g_X(z_X; \text{pa}_X) = \left( \text{Conv}_\theta(1; 1; 1; 0) \circ \text{ConvTranspose}_\theta(1; 4; 2; 1) \circ \text{ReLU} \circ \text{BN}_\theta \circ \text{Conv}_\theta(64; 4; 2; 1) \circ \text{ReLU} \circ \text{BN}_\theta \circ \text{Linear}_\theta(1024) \circ \text{ReLU} \circ \text{BN}_\theta \circ \text{Linear}_\theta(1024) \right)(z_X, \widehat{\text{pa}}_X), \quad (A.5)\]

where the operators describe neural network layers as follows: BN is batch normalisation; ReLU the ReLU activation function; Conv\((c; k; s; p)\) and ConvTranspose\((c; k; s; p)\) are a convolution or transposed convolution using a kernel with size \( k \), a stride of \( s \), a padding of \( p \) and outputting \( c \) channels; Linear\((h)\) is a linear layer with \( h \) output neurons; and Reshape\((\cdot)\) reshapes its inputs into the given shape \( \cdot \). Lastly, \([z_X, \text{pa}_X]\) denotes the concatenation of \( z_X \) and \( \text{pa}_X \), and \( z_X \in \mathbb{R}^{106} \).

Equivalently, we implement the the encoder function as a simple CNN that outputs mean and log-variance of a independent Gaussian:

\[
e_X(x; \text{pa}_X) = \left( [\text{Linear}_\theta(16), \text{Linear}_\theta(16)] \circ \text{LeakyReLU}(0.1), \widehat{\text{pa}}_X \right)
\circ \text{BN}_\theta \circ \text{Linear}_\theta(100) \circ \text{Reshape}(128 \cdot 7 \cdot 7)
\circ \text{LeakyReLU}(0.1) \circ \text{BN}_\theta \circ \text{Conv}_\theta(128; 4; 2; 1)
\circ \text{LeakyReLU}(0.1) \circ \text{BN}_\theta \circ \text{Conv}_\theta(64; 4; 2; 1))(x), \quad (A.6)\]

where LeakyReLU\((\ell)\) is the leaky ReLU activation function with a leakiness of \( \ell \).

We use Adam \([55]\) for optimisation with batch size of 256 and a learning rate of \( 10^{-4} \) for the encoder-decoder and 0.005 for the covariate flows. We set the number of particles (MC samples) for estimating the ELBO to 4. We use 32 MC samples for estimating reconstruction and counterfactuals. We train all models for 1000 epochs and report the results of the model with the best validation loss.

### A.3 Additional Results

Here we further illustrate the associative, interventional, and counterfactual capabilities of the trained independent, conditional, and full models. (Continued on the next page.)

\(^5\)We observed that not normalising the inputs can lead to the deep models prioritising learning the dependence on the variable with largest magnitude. This phenomenon should be investigated further.
A.3.1 Association

Figure A.2: Random samples generated by the independent, conditional and full model. Note how all models appear to have the same unconditional generation capacity.

Figure A.3: Conditional samples generated by the independent, conditional, and full model. The high-level noise, $z_X$, is shared for all samples from each model, ensuring the same ‘style’ of the generated digit. The independent model generates images independent of the thickness and intensity values, resulting in identical samples. For the conditional and full models, thickness and intensity change consistently along each column and row, respectively.

Figure A.4: Reconstructions. These are computed as Monte Carlo averages approximating $\mathbb{E}_{Q(z_X|e_X(x_{pa_X}))}[g_X(z_X; pa_X)]$, where $e_X$ and $g_X$ are the image encoder and decoder networks. All models seem capable of producing faithful reconstructions.
Figure A.5: Comparison of the target covariates and the corresponding values measured from the generated images. The leftmost column refers to the accuracy of the SetThickness and SetIntensity transforms used in generating the synthetic dataset, and the remaining three columns describe the fidelity of samples generated by each of the learned models. While images sampled from the independent model are trivially inconsistent with the sampled covariates, the conditional and full models show comparable conditioning performance.

A.3.2 Intervention

Figure A.6: Difference between conditioning and intervening, based on the trained full model. The joint density $p(t, i)$ is shown as contours in the background, for reference, and the ‘violin’ shapes represent the density of one variable when conditioning or intervening on three different values of the other variable. Since $t$ causes $i$, notice how $p(t|i)$ (left) is markedly different from $p(t|do(i))$ (middle), which collapses to $p(t)$. On the other hand, $p(i|do(t))$ and $p(i|t)$ (right) are identical.
Figure A.7: Original samples and counterfactuals from the full model. The first column shows the original image and true values of the non-imaging data. The even rows show the difference maps between the original image and the corresponding counterfactual image. We observe that all counterfactuals preserve the digits’ identity and style. Our model even generates sensible counterfactual images (with some artefacts) in very low-density regions, e.g. ’0’ with $do(i = 64)$ (thick but dim), and very far from the original, e.g. ’2’ with $do(t = 5.0)$.

B Brain Modelling

B.1 Data Generation

The original three-dimensional (3D) T1-weighted brain MRI scans have been pre-processed by the data providers of the UK Biobank Imaging study using the FSL neuroimaging toolkit [56]. The pre-processing involves skull removal, bias field correction, and automatic segmentation of brain structures. In addition, we have rigidly registered all scans to the standard MNI atlas space using an in-house image registration tool, which enabled us to extract anatomically corresponding mid-axial 2D slices that were used for the experiments presented in this paper. The 2D slices were normalised in intensity by mapping the minimum and maximum values inside the brain mask to the range $[0, 255]$. Background pixels outside the brain were set to zero. Age and biological sex for each subject were retrieved from the UK Biobank database along with the pre-computed brain and ventricle volumes. These volumes are derived from the 3D segmentation maps obtained with FSL, and although these
are image-derived measurements, they may serve as reasonable proxies of the true measurements within our (simplified yet plausible) causal model of the physical manifestation of the brain anatomy.

Figure B.1: Random examplars from the test set of the adopted UK Biobank dataset

B.2 Experimental Setup

The setup for the brain imaging experiment closely follows the MNIST example as described in Appendix A.2. We randomly split the available 13,750 brain images into train, validation and test sets with the respective ratios 70%, 15% and 15%. During training, we randomly crop the brain slices from their original size of 233 px × 197 px to 192 px × 192 px and use center crops during validation and testing. The cropped images are downsampled by a factor of 3 to a size of 64 px × 64 px.

We use the same low-level mechanism for the image \( x \) as with MNIST images but change the encoder and decoder functions to a deeper architecture with 5 scales consisting of 3 blocks of \((\text{LeakyReLU}(0.1) \circ \text{BN} \circ \text{Conv})\) each as well as a linear layer that converts to and from the latent space with 100 dimensions. We directly learn the binary probability of the sex \( s \) and use the following invertible transforms to model the age \( a \), brain volume \( b \), and ventricle volume \( v \) as

\[
a := f_A(\epsilon_A) = (\exp \circ \text{AffineNormalisation} \circ \text{Spline}_A)(\epsilon_A),
\]
\[
b := f_B(\epsilon_B; s, a) = (\exp \circ \text{AffineNormalisation} \circ \text{ConditionalAffine}(s, \hat{a}))(\epsilon_B),
\]
\[
v := f_V(\epsilon_V; a, b) = (\exp \circ \text{AffineNormalisation} \circ \text{ConditionalAffine}(\hat{b}, \hat{a}))(\epsilon_V),
\]

where the context networks are implemented as a fully-connected network with 8 and 16 hidden units, and a \(\text{LeakyReLU}(0.1)\) nonlinearity.

B.3 Additional Results

Likewise, we present more detailed analyses of the model trained on UK Biobank brain images and covariates, in terms of modelling the observational distribution and computing various counterfactual queries. (Continued on the next page.)
B.3.1 Association

Figure B.2: Random samples from the model trained on the UK Biobank dataset

\[ b = 800 \text{ ml} \]
\[ v = 10 \text{ ml} \]
\[ v = 100 \text{ ml} \]
\[ v = 1000 \text{ ml} \]
\[ b = 1200 \text{ ml} \]
\[ b = 1600 \text{ ml} \]
\[ v = 10 \text{ ml} \]
\[ v = 100 \text{ ml} \]
\[ v = 1000 \text{ ml} \]

Figure B.3: Conditional samples from the model trained on the UK Biobank dataset. Images in each 3×3 block share the same the high-level noise vector, \( z_X \). Each row consistently changes the brain size, whereas each column changes the ventricle volume.

Figure B.4: Original samples and reconstructions from the model trained on the UK Biobank dataset
(a) Age vs. brain volume: $p(a, b|s)$. Here we see differences in head size across biological sexes (reflected in brain volume), as well as a downward trend in brain volume as age progresses.

(b) Age vs. ventricle volume: $p(a, v|b \in \cdot)$. As expected from the literature [52], we observe a consistent increase in ventricle volume with age, in addition to a proportionality relationship with the overall brain volume.

Figure B.5: Densities for the true data (KDE) and for the learned model. The overall trends and interactions present in the true data distribution seem faithfully captured by the model.
### B.3.2 Counterfactual

| Original | do(s = male) | do(a = 40 y) | do(a = 80 y) | do(b = 800 ml) | do(b = 1600 ml) | do(v = 11 ml) | do(v = 110 ml) |
|----------|--------------|--------------|--------------|----------------|----------------|---------------|---------------|
| ![Image](image1.png) | ![Image](image2.png) | ![Image](image3.png) | ![Image](image4.png) | ![Image](image5.png) | ![Image](image6.png) | ![Image](image7.png) | ![Image](image8.png) |
| s = female | a = 49 y | b = 1153 ml | v = 26.62 ml |
| ![Image](image9.png) | ![Image](image10.png) | ![Image](image11.png) | ![Image](image12.png) | ![Image](image13.png) | ![Image](image14.png) | ![Image](image15.png) | ![Image](image16.png) |
| s = male | a = 59 y | b = 1078 ml | v = 19.99 ml |
| ![Image](image17.png) | ![Image](image18.png) | ![Image](image19.png) | ![Image](image20.png) | ![Image](image21.png) | ![Image](image22.png) | ![Image](image23.png) | ![Image](image24.png) |
| s = female | a = 50 y | b = 1095 ml | v = 46.84 ml |
| ![Image](image25.png) | ![Image](image26.png) | ![Image](image27.png) | ![Image](image28.png) | ![Image](image29.png) | ![Image](image30.png) | ![Image](image31.png) | ![Image](image32.png) |
| s = female | a = 60 y | b = 1035 ml | v = 24.29 ml |
| ![Image](image33.png) | ![Image](image34.png) | ![Image](image35.png) | ![Image](image36.png) | ![Image](image37.png) | ![Image](image38.png) | ![Image](image39.png) | ![Image](image40.png) |
| s = male | a = 70 y | b = 1062 ml | v = 34.87 ml |

Figure B.6: Original samples and counterfactuals from the model trained on the UK Biobank dataset. The first column shows the original image and true values of the non-imaging data. The even rows show the difference maps between the original image and the corresponding counterfactual image.
C  Discrete counterfactuals

As mentioned in the main text, the DSCM framework supports not only low- and high-dimensional continuous data, but also discrete variables. In particular, discrete mechanisms with a Gumbel–max parametrisation have been shown to lead to counterfactuals satisfying desirable properties \cite{57}. For example, they are invariant to category permutations and are stable, such that increasing the odds only of the observed outcome cannot produce a different counterfactual outcome. More computational details and properties of the Gumbel distribution are found in Maddison and Tarlow \cite{58}.

Consider a discrete random variable over $K$ categories, $y$, with a conditional likelihood described by logits $\lambda$, assumed to be a function $g_Y$ of its parents, $pa_Y$:

$$P(y = k | pa_Y) = \frac{e^{\lambda_k}}{\sum_{l=1}^{K} e^{\lambda_l}}, \quad \lambda = g_Y(pa_Y). \tag{C.1}$$

Under the Gumbel–max parametrisation, the mechanism generating $y$ can be described as

$$y := f_Y(\epsilon_Y; pa_Y) = \arg \max_{1 \leq l \leq K} (\epsilon_{Yl} + \lambda_l), \quad \epsilon_{Yl} \sim \text{Gumbel}(0, 1). \tag{C.2}$$

Samples from the Gumbel(0, 1) distribution can be generated by computing $-\log(-\log U)$, where $U \sim \text{Unif}(0, 1)$.

The Gumbel distribution has certain special properties \cite{58} that enable tractable abduction. Given that we observed $y = k$, samples can be generated from the exact posterior $P(\epsilon_Y | y = k, pa_Y)$:

$$\epsilon_Y^k = G_k + \log \sum_l e^{\lambda_l} - \lambda_k, \quad G_k \sim \text{Gumbel}(0, 1),$$

$$\epsilon_{Yl} = -\log(e^{-G_l - \lambda_l} + e^{-\epsilon_Y^k - \lambda_k}) - \lambda_l, \quad G_l \sim \text{Gumbel}(0, 1), \quad \forall l \neq k. \tag{C.3}$$

Finally, given an upstream counterfactual intervention such that $\tilde{X} = \tilde{g}_Y(\tilde{pa}_Y)$, the counterfactual outcome for $y$ can be determined simply as

$$y = f_Y(\epsilon_Y; \tilde{pa}_Y) = \arg \max_{1 \leq l \leq K} (\epsilon_{Yl} + \tilde{\lambda}_l). \tag{C.4}$$

Note that this entire derivation applies to a truly discrete variable, without the need for continuous relaxations as commonly used in deep generative models \cite{25, 26}, as the likelihood is given in closed form and no gradients of expectations are necessary.