Desiderata for Representation Learning: A Causal Perspective

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Abstract

Representation learning constructs low-dimensional representations to summarize essential features of high-dimensional data. This learning problem is often approached by describing various desiderata associated with learned representations; e.g., that they be non-spurious, efficient, or disentangled. It can be challenging, however, to turn these intuitive desiderata into formal criteria that can be measured and enhanced based on observed data. In this paper, we take a causal perspective on representation learning, formalizing non-spuriousness and efficiency (in supervised representation learning) and disentanglement (in unsupervised representation learning) using counterfactual quantities and observable consequences of causal assertions. This yields computable metrics that can be used to assess the degree to which representations satisfy the desiderata of interest and learn non-spurious and disentangled representations from single observational datasets.

Keywords: Causal inference, representation learning, non-spuriousness, disentanglement, probabilities of causation

1 Introduction

Representation learning constructs low-dimensional representations that summarize essential features of high-dimensional data. For example, one may be interested in learning a low-dimensional representation of MNIST images, where each image is a 784-dimensional vector of pixel values. Alternatively, one may be interested in a product review corpus; each review is a 5,000-dimensional word count vector. Given an m-dimensional data point, \( X = (X_1, \ldots, X_m) \in \mathbb{R}^m \), the goal is to find a d-dimensional representation \( Z = (Z_1, \ldots, Z_d) \triangleq (f_1(X), \ldots, f_d(X)) \) that captures \( d \) important features of the data, where \( f_j : \mathbb{R}^m \to \mathbb{R}, j = 1, \ldots, d \) are \( d \) deterministic functions and \( d \ll m \).
A heuristic approach to the problem has been to fit a neural network that maps from the high-dimensional data to a set of labels, and then take the top layer of the neural network as the representation of the image. When labels are not available, a related heuristic is to fit a latent variable model (e.g., a variational autoencoder (Kingma & Welling, 2014)) and output a low-dimensional representation based on the inferred latent variables. In both cases, the hope is that these low-dimensional representations will be useful in the performance of downstream tasks and also provide an interpretation of the statistical relationships underlying the data.

These heuristic approaches do not, however, always succeed in producing representations with desirable properties. For example, as we will discuss in detail, common failure modes involve capturing spurious features that do not transfer well or finding dimensions that are entangled and are hard to interpret. For example, in fitting a neural network to images of animals, with the goal of producing a labeling of the species found in the images, a network may capture spurious background features (e.g., grass) that are highly correlated with the animal features (e.g., the face of a dog). Such spurious features can often predict the label well. But they are generally not useful for prediction in a different dataset or for performing other downstream tasks. The learned representation may also be entangled—a single dimension of the representation may encode information about multiple features (e.g., animal fur and background lighting). Such entangled representations are hard to interpret. The representation itself does not provide guidance on how to separate learned dimensions into more informative dimensions that describe animal fur and background lighting.

While non-spuriousness or disentanglement are natural desiderata of representations, they are often intuitively defined, challenging to evaluate, and hard to optimize over algorithmically. Lacking formal metrics for these desiderata, and not having access to manually labeled features (e.g., grass, animal fur, or background lighting) that provide empirical guidance, prevents us from developing representation learning algorithms that satisfy natural desiderata.

In this work, we take a causal inference perspective on representation learning. This perspective allows us to formalize representation learning desiderata using causal notions, specifically causal relationships among the label $Y$ and the $d$ features captured by $Z_1, \ldots, Z_d$. This yields calculable metrics obtained from the observable implications of the underlying causal relationships. These metrics then enable representation learning algorithms that target these desiderata. (Figure 1 illustrates this workflow.)

We focus on two sets of desiderata in representation learning: (1) efficiency and non-spuriousness in supervised representation learning—i.e., the representations shall efficiently capture non-spurious features of the data—and (2) disentanglement in unsupervised representation learning—i.e., the representations shall encode different features along separate dimensions.

In the supervised setting, the key idea is to view representations as capturing features that are potential causes of the label. From this perspective, a non-spurious representation should capture features that are sufficient causes of the label. This property ensures that the same representation is still likely to be informative of the label when employed on a new dataset. In the running example of images, a representation that captures the dog-face feature is non-spurious because the presence of a dog face is sufficient to determine the dog label (i.e., whether a dog is present in the image). In contrast, a representation based on the grass feature is spurious because it cannot
causally determine the dog label, even though it is highly correlated. We formalize this connection between non-spurious representations and sufficient causes by defining non-spuriousness using the notion of probability of sufficiency (PS), due to Pearl (2011).

The same causal perspective also allows us to formalize the efficiency of representations, which are those that do not pick up redundant features. Again viewing the representation as a potential cause of the label, its efficiency corresponds to the necessity of the cause. In the image example, a representation of both “dog-face” and “four-leg” is inefficient because it is not necessary to know both “dog-face” and “four-leg” to determine the dog label. As dogs always have four legs (in our simplified world), “four-leg” is a redundant feature given “dog-face.” We therefore formalize efficiency of a representation using the notion of probability of necessity (PN). As we will discuss, PN and PS are aspects of a general notion of probabilities of causation, also due to Pearl (2011).

While these causal definitions formalize efficiency and non-spuriousness, they do not immediately imply calculable metrics for these desiderata, because not all causal quantities are estimable from observational data. To obtain calculable metrics, we have to study observable implications of these causal definitions, a problem known as causal identification (Pearl, 2011). We discuss the unique challenges of causal identification with high-dimensional data—specifically, high-dimensional data are often rank-degenerate (Golub et al., 1976; Stewart, 1984)—and we develop identification strategies to address these challenges. These strategies lead to calculable metrics of efficiency and non-spuriousness in the high-dimensional setting, along with the conditions under which they are valid.

Based on these definitions, we develop an algorithmic framework that we refer to as CAUSAL-REP that formulates representation learning as a task of finding necessary and sufficient causes. In a range of empirical studies, we find that CAUSAL-REP is more successful at finding non-spurious representations in both images and text compared to standard benchmarks.

We now turn to the unsupervised setting where our focus will be the desideratum of disentangled representations. Intuitively, disentanglement requires that different dimensions of the learned representation correspond to independent degrees of freedom of objects; that is, we can view disentangled representations as making it possible to generate new examples of objects by separately manipulating the values of the features encoded by such representations.
To develop metrics for disentanglement, we begin with a causal definition of disentanglement due to Suter et al. (2019): different dimensions of the learned representation should correspond to different features that do not causally affect each other. (These features may still be correlated.) While a useful starting place, this definition unfortunately leaves us short of the goal of measuring and optimizing for disentanglement. Naively, it requires one to assess whether causal connections exist among different features, which is a generally impossible task without substantial knowledge about the underlying causal structure (Imbens & Rubin, 2015; Pearl, 2011). In particular, we may observe correlation among variables, with or without causal connections. Thanks to this difficulty, existing disentanglement metrics often rely on ground-truth features or auxiliary labels (Suter et al., 2019; Thomas et al., 2018), which we often do not have access to in practice.

To obtain an operational measure of disentanglement, we again turn to studying the observable implications of its causal definition. The key observation is that the observable implications can exist in the support of the representation, i.e., the set of values the representation can take. Specifically, we find that disentanglement—the absence of causal connections among the features captured by different dimensions of the representation—implies that their support must be independent in observational data (under a standard positivity condition). Visually, features with independent support must have a scatter plot that occupies a hyperrectangular region: the set of possible values each feature can take does not depend on the values of other features. See Figures 8a to 8c for examples.

Building on this connection between disentanglement and independent support, we develop the independence-of-support score (IOSS) as an unsupervised disentanglement metric; it evaluates whether different dimensions of the representation have independent support. We establish the sufficiency of this metric by showing that representations with independent support are identifiable (under suitable conditions). This fundamental identifiability result enables us to enforce disentanglement in representation learning via an IOSS penalty.

Through the study of non-spuriousness/efficiency and disentanglement, we illustrate how causality can provide a fruitful perspective for representation learning. Many intuitively defined desiderata in representation learning can be formalized using causal notions. These causal notions can further lead to metrics that quantify these desiderata and representation learning algorithms that enforce these desiderata.

Related work. There has been a flurry of recent work at the intersection of representation learning and causal inference. We provide a brief review, highlighting the contrast between this literature and our work.

Learning non-spurious representations via invariance. Many works have considered causal formulations of representation learning given datasets from multiple environments (Arjovsky et al., 2019; Cheng & Lu, 2017; Creager et al., 2021; Khasanova & Frossard, 2017; Lu et al., 2021; Mitrovic et al., 2020; Moraffah et al., 2019; Moyer et al., 2018; Puli et al., 2021; Veitch et al., 2021; Zhao et al., 2019). The basic idea is to enforce the invariance of the mapping between the learned representation and the outcome label, thereby encouraging non-spurious representations and enabling out-of-distribution prediction. Similar to these works, our CAUSAL-REP algorithm
targets non-spurious representations. However, it differs in its focus on the setting in which only a single observational dataset is available. That is, we do not assume access to datasets from multiple environments, nor do we leverage the invariance principle.

**Reverse causal inference.** Our work is also related to a body of work on reverse causal inference, a task that aims to find “causes of effects” (Chalupka et al., 2017, 2015; Galhotra et al., 2021; Gelman & Imbens, 2013; Janzing & Schölkopf, 2015; Kilbertus et al., 2018; Kommiya Mothilal et al., 2021; Paul, 2017; Schölkopf et al., 2012, 2013; Wang & Culotta, 2020, 2021; Watson et al., 2021; Weichwald et al., 2014). Existing approaches formulate the search for causes as causal hypotheses generation and testing. In contrast, our CAUSAL-REP algorithm formulates this search as maximizing probabilities of causation (roc) (Mueller et al., 2021; Pearl, 2011, 2019b; Tian & Pearl, 2000), specifically in the context of representation learning. We develop identification conditions for these roc for high-dimensional data such as images and text (Fong & Grimmer, 2016; Grimmer & Fong, 2021; Nabi et al., 2020; Pryzant et al., 2020; Puli et al., 2020; Ranganath & Perotte, 2018; Wang & Blei, 2021, 2019a, 2020; Wang & Culotta, 2020, 2021). Our results build on existing identification results around multiple causes with shared unobserved confounding (Puli et al., 2020; Wang & Blei, 2019a, 2020) and its positivity issues (D’Amour, 2019a,b; D’Amour et al., 2020a; Imai & Jiang, 2019), but are tailored to the representation learning setting where no unobserved confounding is present.

**Disentangled representation learning.** Our independence-of-support score (ioss) relates to the broad literature on disentangled representation learning. Early approaches to disentanglement often enforce statistical independence among different dimensions of the representation (Achille & Soatto, 2018; Bengio et al., 2013; Burgess et al., 2018; Chen et al., 2018; Higgins et al., 2017; Kim & Mnih, 2018; Kumar et al., 2018). However, Locatello et al. (2019a) show that this inductive bias of statistical independence is insufficient for disentanglement due to its non-identifiability. Many recent approaches to disentanglement thus incorporate auxiliary information to enable identifiability in disentanglement (Bouchacourt et al., 2018; Hosoya, 2019; Khemakhem et al., 2020; Locatello et al., 2020, 2019b; Shen et al., 2020; Shu et al., 2019; Träuble et al., 2020; Yang et al., 2020). In contrast to these works, ioss relies on the causal perspectives of Suter et al. (2019) and establishes identifiability in disentanglement without the need for supervision. Focusing on compactly supported representations, we show that representations with independent support are identifiable under suitable conditions.

**Causal structure learning.** Our causal approach to disentanglement also relates to causal structure learning; both involve assessing the existence of causal connections between variables. Traditional approaches to causal structure learning relies on independence tests or score-based methods, often assuming all variables are observed; see Heinze-Deml et al. (2018) for a review. ioss differs from these works in that we allow for unobserved common causes among the observed variables; we also rely on a different observable implication of the lack of causal connections.

**Representation learning for causal inference.** Representation learning and dimensionality reduction have significantly improved the estimation efficiency of causal inference with high-dimensional covariates and treatments (Johansson et al., 2020, 2019; Nabi et al., 2020; Shi et al., 2020; Veitch et al., 2020, 2019; Wu & Fukumizu, 2021). The focus in these work is on how representation learning can help causal inference with high-dimensional covariates. In contrast, we focus on how causal inference can help produce useful representations.
This paper. The rest of the paper is organized as follows. Section 2 develops a causal approach to efficiency and non-spuriousness in supervised representation learning. Section 2.1 formalizes efficiency and non-spuriousness using counterfactual notions, namely the probability of necessity and sufficiency in causal inference. Section 2.2 discusses the identification of these counterfactuals, which leads to metrics for efficiency and non-spuriousness. Section 2.3 formulates representation learning as a task of finding necessary and sufficient causes of the label and develops CAUSAL-REP, an algorithm that targets efficient and non-spurious features. Section 2.4 provides an empirical study of CAUSAL-REP, showing that it captures non-spurious features and improves downstream out-of-distribution prediction. Section 3 switches gears to develop a causal approach to disentanglement in unsupervised representation learning. Section 3.1 reviews a causal definition of disentanglement. Section 3.2 studies its observable implications, showing that causal disentanglement implies independent support of the representation. This leads to i.o.s.s.s, an unsupervised disentanglement metric. Section 3.3 establishes the identifiability of representations with independent support, enabling disentangled representation learning with an i.o.s.s.s penalty. Section 3.4 demonstrates the effectiveness of i.o.s.s.s in empirical studies. Finally, Section 4 concludes the paper with a discussion of causal perspectives on representation learning.

2 Supervised Representation Learning: Efficiency and Non-spuriousness

We begin by formulating a set of desiderata for supervised representation learning. As a running example, we consider a dataset of images and their labels. We focus on settings where the labels are obtained by annotators viewing each image and then labeling; they describe the perceived features of the image, as opposed to the intended features of the image. The goal is to construct a low-dimensional image representation that efficiently captures its essential features.

Formally, we consider a dataset that contains \( n \) \( m \)-dimensional data points and their labels, \( \{x_i, y_i\}_{i=1}^n \in \mathcal{X}^m \times \mathcal{Y} \), assumed to be sampled i.i.d., with \( x_i = (x_{i1}, \ldots, x_{im}) \). The variable \( x_{il} \) might be the value of the \( l \)th pixel of an image, with \( \mathcal{X} = [0, 255] \); it might also be the number of times the \( l \)th word in the vocabulary occur in the document, with \( \mathcal{X} = \mathbb{Z}^+ \). The goal of representation learning is to find a deterministic function \( f : \mathcal{X}^m \to \mathbb{R}^d \) mapping the \( m \)-dimensional data into a \( d \)-dimensional space (\( d \ll m \)). Thus each data point \( x_i \) has a \( d \)-dimensional representation \( z_i \overset{\Delta}{=} f(x_i) \), or equivalently \( z_i = (z_{i1}, \ldots, z_{id}) \overset{\Delta}{=} (f_1(x_i), \ldots, f_d(x_i)) \), with \( f_j : \mathcal{X}^m \to \mathbb{R} \).

Ideally, such a representation should be efficient and non-spurious. It shall efficiently capture features of the image that are essential for determining the label. Below we formalize the desiderata of efficiency and non-spuriousness using counterfactual notions.

2.1 Defining Efficiency and Non-spuriousness using Counterfactuals

To formalize the desiderata of efficiency and non-spuriousness, we posit a structural causal model (scm) that describes the data generating process. This scm will enable us to define these desiderata through counterfactual quantities defined via this model.
2.1.1 A structural causal model of supervised representation learning

We describe a structural causal model (scm) for supervised representation learning and the counterfactual quantities therein.

The structural causal model (scm) of the labeled dataset. We posit Figure 2 as the scm of a data-generating process. (For notation simplicity, we suppress the data point index i and focus on categorical Y.) The figure embodies the assumption that the high-dimensional object \( X = (X_1, \ldots, X_m) \) causally affects its label \( Y \). Moreover, there is no confounder between the two; the label \( Y \) can be fully determined by the object \( X \). These assumptions hold because we assume the labels are obtained by annotators viewing the images; the labeling is solely based on the images.

Figure 2 also posits that different dimensions of the high-dimensional object, \( X_1, \ldots, X_m \), are correlated due to some unobserved common cause \( C \), which can potentially be multi-dimensional. For example, \( C \) can represent the design of an image; nearby pixels tend to be highly correlated if an image has a smooth design. This assumption holds because pixel values of images are not independent; they are determined by common latent variables (Goodfellow et al., 2014; Kingma & Welling, 2014).

The representation \( f(X) \) and its interventions. The representation \( Z = f(X) \) captures features of the high-dimensional data \( X \). For example, \( Z = (Z_1, Z_2) = (\mathbb{I}\{X_{125} \times X_{38} < 0.01\}, X_{164} + 0.3 \cdot X_{76}^2) \) is a two-dimensional representation of \( X \), whose first dimension represents whether pixels 125 and 38 are close to black. More complex representations may capture more meaningful features; e.g., whether a dog face is present in the image.

The representation \( Z = f(X) \) is not included in the scm as a separate causal variable in addition to \( X \). The reason is that the representation \( Z = f(X) \) is a deterministic function of the image pixels \( X \) and the causal variables of a scm must be related by functional relationships perturbed by random disturbances (Pearl, 1995, 2011). In particular, \( Z \) is not a descendant of \( X \) because a value change in \( Z \) would imply a value change in \( X \) (unless \( f \) is a constant function). Nor is \( Z \) an ancestor of \( X \) because a change in the value of \( X \) can also imply a change in \( Z \).

Despite \( Z \) not being explicitly included in the scm, Figure 2 does imply that \( Z = f(X) \) is an ancestor of the label \( Y \). This is because \( X \) is an ancestor of \( Y \). Moreover, interventions on the representation \( Z = f(X) \) are also well defined; they are functional interventions (Correa & Bareinboim, 2020; Eberhardt & Scheines, 2007; Puli et al., 2020), also referred to as stochastic policies (Pearl, 2011, Ch. 4.2). (We will define them rigorously in Section 2.2.) Roughly, suppose a one-dimensional representation \( Z = f(X) \) captures the univariate binary feature of whether the grass is present in the image. Then the intervention \( \text{do}(Z = 1) \)—equivalently \( \text{do}(f(X) = 1) \)—means “turning on” the grass feature in the image. That is, holding the design of the image \( C \) fixed, we change the pixels of the image \( X \) such that the grass is present; \( Z = 1 \). Accordingly, the intervention \( \text{do}(Z = 0) \) means “turning off” the grass feature. Holding the design \( C \) fixed, we change the pixels \( X \) such that the grass is absent, \( Z = 0 \).
**Figure 2**: A scm for supervised representation learning. $X = (X_1, \ldots, X_m)$ represents the high-dimensional object (e.g., pixels of an image), $Y$ represents the outcome label, and $C$ denotes the unobserved common cause of $X_1, \ldots, X_m$.

**Counterfactuals in the scm.** The scm (Figure 2) results in a family of counterfactuals, “what would the value of a variable be if we intervene on some variables of the causal model.” Here we focus on the counterfactuals obtained when we intervene on the image features captured by the representation $Z$. These counterfactuals will allow us to define the efficiency and non-spuriousness of the representations.

We denote $Y(Z = z)$ as the counterfactual label of an image if we force its representation $Z$ to take value $z$.\(^1\) For example, if a one-dimensional representation $Z$ captures the feature “whether grass is present in the image,” then $Y(Z = 0)$ is the counterfactual label had the image had no grass. Accordingly, $Y(Z = 1)$ is the counterfactual label had the image had grass being present.

The key idea here is to evaluate the quality of the representation $Z$ by reasoning about the counterfactual labels: What would the label $Y$ be had $Z$ taken different values? We will show how the desiderata of non-spuriousness and efficiency can be formalized via these counterfactuals. This connection will allow us to develop metrics and algorithms for these desiderata.

### 2.1.2 Counterfactual definitions of non-spuriousness and efficiency

We next discuss how these counterfactual notions can allow us to formalize the non-spuriousness and efficiency of a representation $f(x)$, for a single observed data point $(X = x, Y = y)$.

**Non-spuriousness and its counterfactual definition.** A non-spurious representation captures features that can (causally) determine the label. In the causal model, we say that a representation captures non-spurious features if, given an image without the feature, including this feature would change its label.

\(^1\)The counterfactual label $Y(Z = z)$ is also commonly written as $Y_z$ (Pearl, 2011). Here we employ the parenthesis notation $Y(Z = z)$ to avoid subscripts.
As an example, suppose the label $Y$ indicates whether the image contains a dog. Then a representation $Z$ capturing the presence of dog-face is a non-spurious feature. Given a non-dog image without a dog face ($Y = 0, Z = 0$), adding a dog face to the image turns on the dog label, i.e., $Y(Z = 1) = 1$. In other words, dog-face is a sufficient case of a dog label. In contrast, a representation $Z$ that captures the presence of grass is a spurious feature. Though the presence of grass and the dog label may be highly correlated, adding grass to a non-dog image does not turn on the dog label.

Viewing the representation $Z$ as a potential cause of the label, a non-spurious representation shall be a sufficient cause of the label. Turning on the feature (captured by) $Z$ should be sufficient to turn on the label. We thus measure the non-spuriousness of a representation by the probability of sufficiency ($\text{ps}$) of the feature $Z$ causing the label $Y$.

**Definition 1 (Non-spuriousness of representations).** Suppose we observe a data point with representation $Z = z$ and label $Y = y$. Then the non-spuriousness of the representation $Z$ for label $Y$ is the probability of sufficiency ($\text{ps}$) of $\mathbb{I}\{Z = z\}$ for $\mathbb{I}\{Y = y\}$:

$$\text{ps}_{Z=z,Y=y} = P(Y(Z = z) = y \mid Z \neq z, Y \neq y). \quad (1)$$

When both the representation $Z$ and the label $Y$ are univariate binary with $z = 1, y = 1$, then Equation (1) coincides with classical definition of $\text{ps}$ (Definition 9.2.2 of Pearl (2011)).

$\text{ps}_{Z=z,Y=y}$ is the probability of the representation $\mathbb{I}\{Z = z\}$ being a sufficient cause of the label $\mathbb{I}\{Y = y\}$. In Pearl’s language, it describes the capacity of the representation to “produce” the label. $\text{ps}$ measures the probability of a positive counterfactual label if we make the feature $\mathbb{I}\{Z = z\}$ be present (and all else equal), conditional on the feature being absent and the label being negative $\{\mathbb{I}\{Z = z\} = 0, \mathbb{I}\{Y = y\} = 0\}$. A non-spurious representation shall have a high $\text{ps}$ for the label of interest.

**Efficiency and its counterfactual definition.** An efficient representation captures only essential features of the data; it does not capture any redundant features. In the causal model, we say that the representation is efficient if, given an image with the feature it captures, removing this feature would change its label.

Returning to the image example with the dog label, a representation that captures “dog-face + four-leg” is inefficient. The reason is that, for an image with both a dog face and four legs, removing one of the dog legs and hence turning off the “dog-face + four-leg” feature does not necessarily turn off the “dog label.” In contrast, the feature “dog-face” is efficient as removing a dog face from an image will turn off the “dog label.”

An efficient representation must capture features that are necessary causes of the label. We can therefore define the efficiency of a representation by the probability of necessity ($\text{pn}$) of the feature causing the label (Pearl, 2011).
Definition 2 (Efficiency of representations). Suppose we observe a data point with representation $Z = z$ and label $Y = y$. Then the efficiency of the representation $Z$ for the label $Y$ is the probability of necessity ($\text{pn}$) of $\mathbb{I}\{Z = z\}$ for $\mathbb{I}\{Y = y\}$:

$$\text{PN}_{Z\mid z, Y\mid y} = P(Y \neq y \mid Z = z, Y = y).$$

When both the representation $Z$ and the label $Y$ are univariate binary with $z = 1, y = 1$, then Equation (2) coincides with classical definition of $\text{pn}$ (Definition 9.2.1 of Pearl (2011)).

$\text{PN}_{Z\mid z, Y\mid y}$ is the probability of the representation $\mathbb{I}\{Z = z\}$ being a necessary cause of the label $\mathbb{I}\{Y = y\}$. It measures the probability of a negative counterfactual label if we remove the feature (i.e., setting $\mathbb{I}\{Z = z\} = 0$ and all else equal), conditional on the feature being present and the label being positive $\{\mathbb{I}\{Z = z\} = 1, \mathbb{I}\{Y = y\} = 1\}$. An efficient representation shall have a high $\text{pn}$ for the label of interest.

2.1.3 Simultaneous assessment of non-spuriousness and efficiency

Representation learning often targets representations that are both non-spurious and efficient. How can one assess non-spuriousness and efficiency simultaneously?

Assessing non-spuriousness and efficiency simultaneously. We invoke the notion of probability of necessity and sufficiency ($\text{pns}$). In more detail, non-spuriousness is intuitively the effectiveness of turning on the label by turning on the feature captured by the representation; efficiency is the effectiveness of turning off the label by turning off this feature. Thus a representation is both non-spurious and efficient if the label responds to the feature in both ways. If the feature is turned on, then the label will be turned on; if the feature is turned off, then the label will be turned off too. $\text{pns}$ calculates exactly this probability.

Definition 3 (Efficiency & non-spuriousness of representations). Suppose we observe a data point with representation $Z = z$ and label $Y = y$. Then the efficiency and non-spuriousness of the representation $Z$ for label $Y$ is the probability of necessity and sufficiency ($\text{pns}$) of $\mathbb{I}\{Z = z\}$ for $\mathbb{I}\{Y = y\}$:

$$\text{PNS}_{Z\mid z, Y\mid y} = P(Y \neq y, Y(Z = z) = y)).$$

When both the representation $Z$ and the label $Y$ are univariate binary with $z = 1, y = 1$, then Equation (3) coincides with classical definition of $\text{pns}$ (Definition 9.2.3 of Pearl (2011)).

Requiring both necessity and sufficiency of the cause is a stronger requirement than requiring only necessity (or only sufficiency). Accordingly, $\text{pns}$ is a weighted combination of $\text{pn}$ and $\text{ps}$,

$$\text{PNS}_{Z\mid z, Y\mid y} = P(Z = z, Y = y) \cdot \text{PN}_{Z\mid z, Y\mid y} + P(Z \neq z, Y \neq y) \cdot \text{PS}_{Z\mid z, Y\mid y},$$

2The distribution of the counterfactual $Y(Z \neq z)$ is defined as that of a soft intervention (Correa & Bareinboim, 2020; Eberhardt & Scheines, 2007); a.k.a. a stochastic policy (Pearl, 2011, Ch. 4.2):

$$P(Y(Z \neq z)) = \int P(Y(Z))P(Z \neq z) \, dZ.$$
as per Lemma 9.2.6 of Pearl (2011). We note that our definitions of \( \text{ps}, \text{pns} \) generalize those in Pearl (2011, Ch. 9) from univariate binary causes to general (continuous, discrete, or multi-dimensional) causes. We next consider two further extensions of the \( \text{pns} \) notion.

**Extension: Efficiency and non-spuriousness over a dataset.** The discussion of efficiency and non-spuriousness (Definitions 1 to 3) has focused on individual data points up to this point, reflecting the fact that probabilities of causation (poc) notions are most commonly discussed with respect to a single occurred event—given that the event \( Z = z \) and \( Y = y \) has occurred, what is the probability that \( Z = z \) is a sufficient and/or necessary cause of \( Y = y \)?

In practice, however, we are often interested in whether a representation, \( f : \mathcal{X} \rightarrow \mathcal{Y} \), can produce an efficient and non-spurious summary for datasets of \( n \) i.i.d. data points, \( \{(x_i, y_i)\}_{i=1}^n \).

We thus extend Definition 3 to this setting:

\[
P_{\text{pns}}(Z, Y) \triangleq \prod_{i=1}^n P_{\text{pns}}(Z_{=z_i}, Y_{=y_i} = \prod_{i=1}^n P(Y(Z \neq z_i) \neq y_i, Y(Z = z_i) = y_i)), \tag{4}
\]

where \( z_i = f(x_i) \) is the representation for data point \( i \). Loosely, this means that a representation is efficient and non-spurious for a dataset if its efficiency and non-spuriousness holds jointly across all the data points. (One can similarly extend Definitions 1 and 2 for \( \text{ps} \) and \( \text{pns} \).)

**Extension: Conditional efficiency and non-spuriousness.** For multi-dimensional representations, one is often interested in the efficiency and non-spuriousness of each of its dimensions. We expect each dimension of the representation to be efficient and non-spurious conditional on all other dimensions.

We thus extend Definition 3 to formalize a notion of conditional efficiency and non-spuriousness. Consider a \( d \)-dimensional representation \( Z = (Z_1, \ldots, Z_d) = (f_1(X), \ldots, f_d(X)) \). The conditional efficiency and non-spuriousness of the \( j \)th dimension \( Z_j \) for data point \( (x_i, y_i) \) is

\[
P_{\text{pns}}(Z_j = z_{ij}, Y_{=y_i} | Z_{-j} = z_{i,-j}) = P(Y(Z_j \neq z_{ij}, Z_{-j} = z_{i,-j}) \neq y_i, Y(Z_j = z_{ij}, Z_{-j} = z_{i,-j}) = y_i), \tag{5}
\]

where \( z_{ij} = f_j(x_i) \) is the \( j \)th dimension of the representation, and \( z_{i,-j} = (z_{ij'})_{j' \in \{1, \ldots, d\} \setminus j} \). Accordingly, the conditional efficiency and non-spuriousness of \( Z_j \) across all \( n \) data points is

\[
P_{\text{pns}}(Z_j, Y | Z_{-j}) \triangleq \prod_{i=1}^n P_{\text{pns}}(Z_{=z_i}, Y_{=y_i} | Z_{-j} = z_{i,-j}), \tag{6}
\]

Conditional efficiency and non-spuriousness describes how the label responds to the \( j \)th feature captured by the representation, holding all other features fixed. Its definition resembles the definition of (unconditional) efficiency and non-spuriousness. The only difference is that the conditional notion contrasts the counterfactual label of \( \{Z_j \neq z_{ij}, Z_{-j} = z_{i,-j}\} \) and \( \{Z_j = z_{ij}, Z_{-j} = z_{i,-j}\} \), while the unconditional notion contrasts that of \( \{Z_j \neq z_{ij}, Z_{-j} \neq z_{i,-j}\} \) and \( \{Z_j = z_{ij}, Z_{-j} = z_{i,-j}\} \).

The conditional efficiency and non-spuriousness across all dimensions of a representation is generally a stronger requirement than the (unconditional) efficiency and non-spuriousness. For example, suppose a dataset of \( (X, Y) \) with \( X = (X_1, X_2, X_3) \) is generated by \( Y = X_1 + \epsilon_y \). Now,
consider a two-dimensional representation, \( Z = (Z_1, Z_2) \), where \( Z_1 = X_1 \) and \( Z_2 = Z_1 + \epsilon, \epsilon \sim \mathcal{N}(0, \sigma^2) \). When \( \sigma^2 \) is small, \( Z_1 \) and \( Z_2 \) are highly correlated. In this case, the (unconditional) efficiency and non-spuriousness of \( Z \) is high because \( (Z_1, Z_2) \) is indeed non-spurious and (nearly) efficient; \( Z_2 \) only introduces a negligible amount of useless information. However, the conditional efficiency and non-spuriousness of \( Z_2 \) given \( Z_1 \) is low because \( Z_2 \) is completely useless given \( Z_1 \).

We will leverage this conditional efficiency and non-spuriousness notion for representation learning in Section 2.3. First, however, we study how to measure (conditional) efficiency and non-spuriousness from observational data.

### 2.2 Measuring Efficiency and Non-spuriousness in Observational Datasets

Definitions 1 to 3 formalize the efficiency and non-spuriousness of representations using POC. These POC are counterfactual quantities; calculating them requires access to a causal structural equation (i.e., the true data generating process), which is rarely available in practice.

In this section, we study the observable implications of Definitions 1 to 3. They lead to strategies to evaluate the efficiency and non-spuriousness of representations with observational data; these strategies are known as causal identification strategies (Pearl, 2011). For simplicity of exposition, we focus on identifying the PNS in Definition 3. (Identification formulas for PN and PS in Definitions 1 and 2 can be similarly derived using Theorem 9.2.15 of Pearl (2011).)

To identify the counterfactual quantity PNS from observational data, we perform two steps of reduction, climbing down the ladder of causation (Pearl, 2019a): (1) connect PNS to interventional distributions, and (2) identify these interventional distributions from observational data. We describe these steps in detail in the next sections. The end product is Theorem 7, which provides an algorithm for calculating a lower bound on the PNS.

#### 2.2.1 From counterfactuals to interventional distributions

To connect the counterfactual quantity PNS with intervention distributions, we generalize the classical identification results for PNS that have been developed for univariate binary causes and outcomes. It turns out that the PNS cannot be point-identified by interventional distributions; for each set of interventional distributions, there may exist many values of PNS that are consistent with the interventional distributions (Pearl, 2011). However, PNS can be bounded by the difference between these two interventional distributions.

**Lemma 4** (A lower bound on PNS). *Assuming the causal graph in Figure 2, the PNS is lower bounded by the difference between two intervention distributions:

\[
P_{\text{PNS}}^{Z = z, Y = y} = P(Y(Z = z) = y, Y(Z \neq z) \neq y) \\
\geq P(Y = y \mid \text{do}(Z = z)) - P(Y = y \mid \text{do}(Z \neq z)).
\]

(7)

The inequality becomes an equality when the outcome \( Y \) is monotone in the representation \( Z \) (in the binary sense); i.e., \( P(Y(Z = z) \neq y, Y(Z \neq z) = y) = 0 \).
Lemma 4 generalizes Theorem 9.2.10 of Pearl (2011) to non-binary $Z$. (The proof is in Appendix A.) It connects the counterfactual quantity $\text{PNS}$ to the intervention distribution $P(Y \mid \text{do}(Z = z))$. An upper bound on $\text{PNS}$ can be similarly obtained by generalizing the upper bound in Theorem 9.2.10 of Pearl (2011). We focus on $\text{PNS}$ here because a larger lower bound on $\text{PNS}$ implies a large $\text{PNS}$. In contrast, a larger upper bound on $\text{PNS}$ does not necessarily imply a large $\text{PNS}$.

Reducing the counterfactual quantity $\text{PNS}$ to the lower bound in Lemma 4 has climbed down one level of the ladder of causation (Pearl, 2019a); we have converted a counterfactual quantity (level three) to one with interventional distributions (level two). Next we will descend one level further, discussing how to identify the interventional distributions $P(Y \mid \text{do}(Z = z))$ from the observational data distribution $P(X, Y)$ (level one); $Z = f(X)$ is the representation of $X$ via some known function $f$. These identification results would also enable the identification of $P(Y \mid \text{do}(Z \neq z)) = \int P(Y \mid \text{do}(Z)) P(Z \mid Z \neq z) \, dZ$.

2.2.2 From interventional distributions to observational data distributions

To calculate the lower bound of $\text{PNS}$ in Equation (7), we need to identify the intervention distribution $P(Y \mid \text{do}(Z = z))$—equivalently $P(Y \mid \text{do}(f(X) = z))$—from observational datasets $(X, Y)$. Below we first state the formal definition of functional interventions $\text{do}(f(X))$. We then discuss the challenges in identifying $P(Y \mid \text{do}(f(X) = z))$ for high-dimensional $X$. To tackle this challenge, we further provide an identification strategy and discuss its theoretical and practical requirements.

The functional intervention $\text{do}(f(X))$. We begin with stating the formal definition of a functional intervention.

**Definition 5** (Functional interventions (Puli et al., 2020)). The intervention distribution under a functional intervention $P(Y \mid \text{do}(f(X) = z))$ is defined as

$$P(Y \mid \text{do}(f(X) = z)) \triangleq \int P(Y \mid \text{do}(X), C) P(X \mid C, f(X) = z) P(C) \, dX \, dC,$$

where $C$ denotes all parents of $X$.

The functional intervention $f(X)$ is a soft intervention on $X$ (Correa & Bareinboim, 2020; Eberhardt & Scheines, 2007) conditional on all its parents $C$, with a stochastic policy $p(X \mid C, f(X) = z)$ (Pearl, 2011, Ch. 4.2). In other words, a functional intervention $\text{do}(f(X) = z)$ considers all interventions on $X$ that are consistent with the functional constraint $f(X) = z$ and its parental structure $C$. Functional interventions recover the standard backdoor adjustment as special cases if we take the function $f$ to be an identity function $f(X) = X$ or one that returns a subset $f(X) = X_S$, $S \subset \{1, \ldots, m\}$; see Appendix B for detailed derivations.

Following this definition, one can write the intervention distribution of interest, $P(Y \mid \text{do}(f(X) = z))$, as follows:

$$P(Y \mid \text{do}(f(X) = z)) = \int P(Y \mid X) \cdot \left[ \int P(X \mid C, f(X) = z) P(C) \, dC \right] \, dX. \quad (9)$$

This equality is due to the $\text{scm}$ in Figure 2: there is no unobserved confounding between $X$ and $Y$, which implies $P(Y \mid \text{do}(X), C) = P(Y \mid X)$. 

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The challenge of identifying $P(X)$ with linear regression when $X$ is high-dimensional is also an example of the rank-degeneracy problem as the data can only inform the underspecification problem in deep learning (D’Amour et al., 2020b). It is akin to the classical identifiability problem (Golub et al., 1976; Stewart, 1984). It is also why, for high-dimensional $X$, even if both $X$ and $Y$ are observed. Intuitively, this non-identifiability problem occurs because the data can only inform $P(Y|X)$ on the low-dimensional manifold in $X^m$ where $P(X)$ is supported. The behavior of $P(Y|X)$ outside this manifold is unconstrained, hence non-identifiable.

Equation (9) provides a way to identify $P(Y|do(f(X) = z))$, provided that one can calculate $P(Y|X)$ and $\int P(X|C, f(X) = z)P(C)\,dC$. At first sight, both quantities seem easy to calculate: one can estimate $P(Y|X)$ from observational sampling of $P(Y,X)$. To calculate $\int P(X|C, f(X) = z)P(C)\,dC$, one may leverage probabilistic factor models (e.g., probabilistic principal component analysis (ppca), Gaussian mixture model (gmm), or variational autoencoder (vae)) because the latent $C$ renders $X_1, \ldots, X_m$ conditionally independent in Figure 2. One can often read off $P(X|C, f(X) = z)$ and $P(C)$ from the probabilistic factor model fit, if the factor model is identifiable.

However, $P(Y|X)$ turns out to be challenging to calculate in practice, especially when $X$ represents high-dimensional objects such as images or text, as we now discuss.

The challenge of identifying $P(Y|X)$ with high-dimensional $X$. The key challenge in identifying $P(Y|X)$ lies in the high-dimensionality of $X$. High-dimensional objects (e.g. images or text) are often only supported on a low-dimensional manifold, i.e., $X_j = g_0(\{X_1, \ldots, X_m\}\setminus X_j)$ for some $j \in \{1, \ldots, m\}$ and some function $g$; see Figure 3 for examples. We refer to this problem as the rank-degeneracy problem (Golub et al., 1976; Stewart, 1984). It is akin to the classical challenge of having highly correlated variables in a linear regression; it is also an example of the underspecification problem in deep learning (D’Amour et al., 2020b).

The rank degeneracy of high-dimensional $X = (X_1, \ldots, X_m)$ challenges the identification of $P(Y|X)$, even if both $X$ and $Y$ are observed. Intuitively, this non-identifiability problem occurs because the data can only inform $P(Y|X)$ on the low-dimensional manifold in $X^m$ where $P(X)$ is supported. The behavior of $P(Y|X)$ outside this manifold is unconstrained, hence non-identifiable.

Figure 3: High-dimensional data such as images (Figure 3a) or text (Figure 3b) are often only supported on a low-dimensional manifold. Low-dimensional data, e.g. the wine dataset (Figure 3c), is often supported in the whole space.

Equation (9) provides a way to identify $P(Y|do(f(X) = z))$, provided that one can calculate $P(Y|X)$ and $\int P(X|C, f(X) = z)P(C)\,dC$. At first sight, both quantities seem easy to calculate: one can estimate $P(Y|X)$ from observational sampling of $P(Y,X)$. To calculate $\int P(X|C, f(X) = z)P(C)\,dC$, one may leverage probabilistic factor models (e.g., probabilistic principal component analysis (ppca), Gaussian mixture model (gmm), or variational autoencoder (vae)) because the latent $C$ renders $X_1, \ldots, X_m$ conditionally independent in Figure 2. One can often read off $P(X|C, f(X) = z)$ and $P(C)$ from the probabilistic factor model fit, if the factor model is identifiable.

However, $P(Y|X)$ turns out to be challenging to calculate in practice, especially when $X$ represents high-dimensional objects such as images or text, as we now discuss.

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The rank degeneracy of high-dimensional $X = (X_1, \ldots, X_m)$ challenges the identification of $P(Y|X)$, even if both $X$ and $Y$ are observed. Intuitively, this non-identifiability problem occurs because the data can only inform $P(Y|X)$ on the low-dimensional manifold in $X^m$ where $P(X)$ is supported. The behavior of $P(Y|X)$ outside this manifold is unconstrained, hence non-identifiable.

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3This non-identifiability of $P(Y|X)$ is also why, for high-dimensional $X$, directly fitting a neural network for $P(Y|X)$ can pick up spurious correlations and fail in out-of-distribution prediction. Similar failure can also occur with linear regression when $X$ is approximately low rank; e.g., when different dimensions of $X$ are highly correlated.
In more detail, suppose \( p(x, y), p(x), p(y \mid x) \) denote the relevant densities (and assume they exist). Any \( h_0(x, y) \) function that satisfies

\[
h_0(x, y) \cdot p(x) = p(x, y) \quad \forall x \in \mathcal{X}^m
\]

is a valid density for \( p(y \mid x) \). Under rank degeneracy, it turns out that there exist many different functions \( h_0(x, y) \) that satisfy Equation (10). Hence \( p(y \mid x) \) is non-identifiable. The reason is that rank degeneracy implies \( p(x) = 0 \) for \( x \in \tilde{X} \subseteq \mathcal{X}^m \), where \( \tilde{X} \) is a set with positive measure. If some function \( h'_0 \) satisfies Equation (10), then the function \( h''_0 \) would also satisfy Equation (10) if they differ only on the set \( (x, y) \in \tilde{X} \times \mathcal{Y} \). Note that \( h'_0 \) and \( h''_0 \) are not almost surely equal; they differ on a set \( \tilde{X} \) with positive measure. Thus, \( h'_0 \) and \( h''_0 \) are two different densities both valid for \( p(y \mid x) \), which implies the non-identifiability of \( p(y \mid x) \).

As a more concrete example, consider a high-dimensional vector of image pixels \( X \) that lives on a low-dimensional manifold; i.e., such that \( X_j - g_0(\{X_1, \ldots, X_m\} \setminus X_j) \) is identically zero in the observational data (Goodfellow et al., 2014; Kingma & Welling, 2014). This rank degeneracy implies that for any \( p(y \mid x) = h_0(x, y) \) compatible with the observational data distribution, the conditional \( p(y \mid x) = h_0(x, y) + \alpha \cdot (x_j - g_0(\{x_1, \ldots, x_m\} \setminus x_j)), \forall \alpha \in \mathbb{R} \) is also compatible with the observational data.

This fundamental non-identifiability of \( P(Y \mid X) \) with high-dimensional \( X \) suggests that we cannot hope to identify functional interventions \( P(Y \mid \text{do}(f(X) = z)) \) where the function \( f \) non-trivially depends on all \( X_1, \ldots, X_m \), or any of its subset that are also rank degenerate.

**Causal identification of \( P(Y \mid f(X)) \) for a restricted set of \( f \).** Given the fundamental non-identifiability of \( P(Y \mid X) \) with high-dimensional \( X = (X_1, \ldots, X_m) \), we restrict our attention to representations that only nontrivially depends on a “full-rank” subset; i.e., \( Z = f(X) = f((X_j)_{j \in S}), \) for some function \( f : \mathcal{X}^{|S|} \rightarrow \mathbb{R}^d \), and a set \( S \subseteq \{1, \ldots, m\} \), where \( p((x_j)_{j \in S}) > 0 \) for all values \( (x_j)_{j \in S} \in \mathcal{X}^{|S|} \). We term this requirement “observability.”

Focusing on such representations \( f(X) = \tilde{f}((X_j)_{j \in S}) \), we calculate its intervention distributions by returning to the definition of functional interventions (Definition 5),

\[
P(Y \mid \text{do}(f(X) = z)) = \int P(Y \mid (X_j)_{j \in S}, C) P((X_j)_{j \in S} \mid C, f(X) = z) P(C) \ d(X_j)_{j \in S} \ dC.
\]

Equation (11) implies that \( P(Y \mid \text{do}(f(X) = z)) \) is identifiable as long as one can identify the unobserved common cause \( C \) from data. This condition is often called the “pinpointability” (or “effective observability”) of \( C \) (Wang & Blei, 2019a, 2020). The following lemma summarizes the identification of \( P(Y \mid \text{do}(f(X) = z)) \) under these conditions.

**Lemma 6** (Identification of \( P(Y \mid \text{do}(f(X) = z)) \)). Assume the causal graph in Figure 2. Suppose the representation only effectively depends on a subset \( (X_j)_{j \in S} \) of \( (X_1, \ldots, X_m) \); i.e., \( f(X) = \tilde{f}((X_j)_{j \in S}) \) for some function \( f : \mathcal{X}^{|S|} \rightarrow \mathbb{R}^d \) and some set \( S \subseteq \{1, \ldots, m\} \). Then the intervention distribution \( P(Y \mid \text{do}(f(X) = z)) \) is identifiable by

\[
P(Y \mid \text{do}(f(X) = z)) = \int P(Y \mid f(X) = z, h(X)) \cdot P(h(X)) \ dh(X),
\]

---

4https://www.kaggle.com/crowdflower/twitter-airline-sentiment, https://archive.ics.uci.edu/ml/datasets/wine
if the following conditions are satisfied:

1. (pinpointability) the unobserved common cause $C$ is pinpointable; i.e., $P(C \mid X) = \delta_{h(X)}$ for a deterministic function $h$ known up to bijective transformations,

2. (positivity) $(X_j)_{j \in S}$ satisfies the positivity condition given $C$; i.e., $P((X_j)_{j \in S} \in \tilde{X} \mid C) > 0$ for any set $\tilde{X} \subset X^{\mid S \mid}$ such that $P((X_j)_{j \in S} \in \tilde{X}) > 0$,

3. (observability) $P((X_j)_{j \in S} \in \tilde{X}) > 0$ for all subsets $\tilde{X} \subset X^{\mid S \mid}$ with a positive measure.

Lemma 6 describes a set of representations $Z = f(X)$ for which can we evaluate their intervention distribution, and hence the lower bound of their RNS for $Y$. (The proof is in Appendix C.)

Lemma 6 is based on three conditions. The pinpointability ensures that the terms in Equation (11) involving $C$ (e.g. $P(C)$, $P((X_j)_{j \in S} \mid C, f(X) = z)$) are estimable from observational data. We note that pinpointability serves a different purpose here than in Wang & Blei (2019a, 2020) and Puli et al. (2020). We invoke pinpointability for Lemma 6 because functional interventions require the knowledge of $C$; there is no unobserved confounding issue. In contrast, Wang & Blei (2019a, 2020) and Puli et al. (2020) invoke pinpointability to handle unobserved confounders.

Pinpointability is often approximately satisfied when the high-dimensional $X$ is driven by a low-dimensional factor $C$. In this case, $P(C \mid X)$ is often close to a point mass (Chen et al., 2020b). For example, when the $m$-dimensional data $X$ is generated by a known $d$-dimensional probabilistic factor model, then the latent factor $C$ is approximately pinpointable when $m \gg d$. In these probabilistic factor models, $P(C \mid X)$ becomes increasingly closer to a point mass as $m \to \infty$ but $d$ fixed (Bai & Li, 2016; Chen et al., 2020b; Wang & Blei, 2019b). In practice, one assesses pinpointability by fitting probabilistic factor models to $X$. We will discuss these practical details in the next section.

The second condition of Lemma 6 is the positivity of $(X_j)_{j \in S}$ given $C = h(X)$, also known as the overlap condition (Imbens & Rubin, 2015). This condition ensures that $P(Y \mid (X_j)_{j \in S}, C)$ in Equation (11) is estimable from observational data. Positivity loosely requires that all values of $(X_j)_{j \in S}$ are possible conditional on $C = h(X)$. For example, it is violated when $S = \{1, \ldots, n\}$ because it is impossible to observe $X = x$ and $C = c$ simultaneously when $h(x) \neq c$. In practice, positivity is more likely to be satisfied when $(X_j)_{j \in S}$ is a low-dimensional vector; e.g., the image representation only depends on a few image pixels that are far away from each other.

Finally, the observability condition requires that it must be possible to observe all possible values of $(X_j)_{j \in S}$. Together with the positivity condition, this implies that $P((X_j)_{j \in S} \mid C) > 0$ for all $(X_j)_{j \in S}$ and, therefore, $P(Y \mid (X_j)_{j \in S}, C)$ in Equation (12) is estimable from observational data. This condition is violated if the observed $(X_j)_{j \in S}$ is rank-degenerate; e.g., $(X_j)_{j \in S}$ with $S = \{1, \ldots, m\}$ represents the whole high-dimensional image vector supported only on a low-dimensional manifold. This rank degeneracy would render $P(Y \mid (X_j)_{j \in S}, C)$ non-identifiable, for the same reason as why $P(Y \mid X)$ is non-identifiable with rank-degenerate $X$. In practice, this observability condition is more likely to hold when $(X_j)_{j \in S}$ is low-dimensional.

Lemma 6 describes a class of representations $Z = f(X)$ whose intervention distributions are identifiable from observational data. Combining Lemmas 4 and 6 allows us to lower bound the RNS of these representations and evaluate their efficiency and non-spuriousness.
Theorem 7 (Evaluating efficiency and non-spuriousness with observational data). Under the assumptions of Lemma 6, the efficiency and non-spuriousness of the representation $Z = f(X) = \hat{f}((X_j)_{j\in S})$ in the dataset $\{x_i, y_i\}_{i=1}^n$ is lower bounded by

$$PNS_a(f(X), Y) \geq PNS_n(f(X), Y)$$

$$\triangleq \prod_{i=1}^n \int [P(Y = y_i | f(X) = f(x_i), C) - P(Y = y_i | f(X) \neq f(x_i), C)] \cdot P(C) \, dC,$$

where $C = h(X)$ is the unobserved common cause pinpointed by the observational data $X$ in Lemma 6. Similarly, the conditional efficiency and non-spuriousness of the $j$th dimension of $(f_1(X), \ldots, f_d(X))$ is lower bounded by

$$PNS_n(f_j(X), Y | f_{-j}(X)) \geq PNS_n(f_j(X), Y | f_{-j}(X))$$

$$\triangleq \prod_{i=1}^n \int [P(Y = y_i | f_j(X) = f_j(x_i), f_{-j}(X) = f_{-j}(x_i), C) - P(Y = y_i | f_j(X) \neq f_j(x_i), f_{-j}(X) = f_{-j}(x_i), C)] \cdot P(C) \, dC.$$

Theorem 7 is an immediate consequence of Lemmas 4 and 6. It suggests that we evaluate efficiency and non-spuriousness using Equations (13) and (14) in practice. We operationalize this result in the next section.

2.2.3 Measuring efficiency and non-spuriousness in practice

We operationalize Theorem 7 to measure efficiency and non-spuriousness for a representation $f(X)$. The full algorithm is in Algorithm 1. The algorithm involves three steps: (1) Pinpoint the unobserved common cause $C$; (2) Estimate the conditional $P(Y | f(X), C)$; (3) Calculate the lower bounds in Theorem 7 (Equations (13) and (14)) to measure efficiency and non-spuriousness. We discuss each step below.

Pinpointing the unobserved common cause $C$. Theorem 7 requires that the unobserved common cause $C$ must be pinpointable by the observational data: $P(C | X) = \delta_{h(X)}$ for some deterministic function $h$. Moreover, this pinpointing function $h$ needs to be known up to bijective transformations. How can we assess pinpointability and extract $h$ in practice?

One can assess pinpointability by fitting a probabilistic factor model, viewed as a causal generative model; e.g., probabilistic principal component analysis (PPCA) (Tipping & Bishop, 1999), variational autoencoder (VAE) (Kingma & Welling, 2014; Rezende et al., 2014), mixture models (McLachlan & Basford, 1988), or mixed membership models (Airoldi et al., 2008; Blei et al., 2003; Erosheva & Fienberg, 2005; Pritchard et al., 2000).

Specifically, suppose the dataset of $n$ i.i.d. data points $(x_i)_{i=1}^n$ is assumed to be generated by a PPCA model, then one can assess pinpointability by first fitting a PPCA model,

$$c_i \overset{i.i.d.}{\sim} \mathcal{N}(0, I_K),$$

$$x_l | c_i \overset{i.i.d.}{\sim} \mathcal{N}(c_i^\top \theta_l, \sigma^2),$$

where

$$c_i \overset{i.i.d.}{\sim} \mathcal{N}(0, I_K),$$

$$x_l | c_i \overset{i.i.d.}{\sim} \mathcal{N}(c_i^\top \theta_l, \sigma^2),$$

for $i = 1, \ldots, n$, and $l = 1, \ldots, m$. (15) (16)
where \( c_i \) is the latent variable for each data point \( x_i = (x_{i1}, \ldots, x_{im}) \), and \( \theta = (\theta_1, \ldots, \theta_m) \) is the set of parameters. One then infers the parameters \( \theta \) and the posterior \( p(c_i \mid x_i) \) for each \( i \), using standard posterior inference algorithms like variational inference (Blei et al., 2017) or Markov chain Monte Carlo methods (Gilks et al., 1995). Pinpointability then approximately holds if \( p(c_i \mid x_i) \) is close to a point mass for all \( i \)’s.

The choice of the latent dimensionality \( K \) is a challenging task. We choose \( K \) in practice by searching for a factor model that can both fit data well (e.g., via a posterior predictive check (Gelman et al., 1996)) and approximately satisfy pinpointability. A larger \( d \) tends to have a better fit to the data, while a smaller \( d \) is more likely to satisfy pinpointability. We proceed if pinpointability holds, along with the other conditions—observability and positivity—from Theorem 7.

Given a fitted factor model, one can extract the pinpointing function \( h \) by computing the posterior mean, 
\[
h(x) \approx \mathbb{E}[c_i \mid x_i = x].
\]

The same procedure applies if the data is generated by other probabilistic factor models.

A reader might ask: Why probabilistic factor models? The reason is that the causal graph Figure 2 encodes a conditional independence structure that coincides with the defining feature of probabilistic factor models. In more detail, Figure 2 assumes that each dimension of the data \( X = (X_1, \ldots, X_m) \) is rendered conditionally independent given an unobserved common cause \( C \). This conditional independence is precisely the defining structure of probabilistic factor models,

\[
ce_i \sim p(\cdot \mid \lambda_C), \quad i = 1, \ldots, n,
\]

\[
x_{il} \mid c_i \sim p(\cdot \mid c_i \mid \theta_l), \quad l = 1, \ldots, m,
\]

with i.i.d. data \( x_i = (x_{i1}, \ldots, x_{im}) \) and latent \( c_i \). We therefore use probabilistic factor models to assess the pinpointability of \( C \) in Figure 2.

**Calculate the conditional** \( P(Y \mid f(X), C) \). After pinpointing \( C \), we still need the conditional \( P(Y \mid f(X), C) \) to calculate the lower bound in Theorem 7.

To estimate \( P(Y \mid f(X), C) \), we can fit a model to the observational data \((x_i, y_i)_{i=1}^n\), with \( f(\cdot) \) being the representation function of interest and \( c_i = h(x_i) \) estimated from the first step. As an example, we may posit a linear model,

\[
P(Y \mid f(X), C) = \mathcal{N}((\beta_0 + \beta^T f(X) + \gamma^T C), \sigma^2),
\]

and estimate the parameters \( \beta = (\beta_1, \ldots, \beta_d) \) and \( \gamma \) via maximum likelihood. One may also posit other more flexible models like a partially linear model (with a nonlinear \( f_C \)) and target a categorical outcome:

\[
P(Y \mid f(X), C) = \text{Categorical}(\text{softmax}(\beta_0 + \beta^T f(X) + \gamma^T f_C(C))).
\]

**Calculate the lower bounds in Theorem 7.** We finally calculate the lower bounds in Equations (13) and (14) using the pinpointed \( C \) and the estimated conditional \( P(Y \mid f(X), C) \) from the previous two steps.
Algorithm 1: Calculating the (lower bound of) efficiency and non-spuriousness of a representation

**input**: The observational data and its label \( \{ x_i, y_i \}_{i=1}^n \); representation function \( f(\cdot) \);
the probabilistic factor model that generates the data \( P(X, C) \).

**output**: The (lower bound of) efficiency and non-spuriousness of representation \( f(X) \) for label \( Y \)

Fit a probabilistic factor model (Equations (17) and (18)) and infer \( p(c_i \mid x_i) \);
if Pinpointability holds, i.e., \( p(c_i \mid x_i) \) is close to a point mass for all \( i \) then

**foreach** datapoint \( i \) do
- Pinpoint the unobserved common cause: \( \hat{c}_i = h(x_i) \approx \mathbb{E} \left[ c_i \mid x_i \right] \);
- Calculate the conditional \( P(Y \mid f(X), C) \) by fitting a model to \( \{ x_i, \hat{c}_i, y_i \}_{i=1}^n \) (e.g., Equations (19) and (20));

end

Calculate the (lower bound of) efficiency and non-spuriousness \( \text{PNS}_n(f(X), Y) \) and \( \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)), j = 1, \ldots, d \) (Equations (13) and (14));

end

As an example, we calculate the lower bound of the conditional efficiency and non-spuriousness \( \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)) \) with the linear model (see Appendix E for the detailed derivation):

\[
\log \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)) \\
\approx \left( \frac{1}{2\sigma^2} \sum_{i=1}^{n} (\beta_j \cdot (f_j(x_i) - \mathbb{E} \left[ f_j(x_i) \right]))^2 \\
+ 2 \cdot \sum_{i=1}^{n} \beta_j \cdot (f_j(x_i) - \mathbb{E} \left[ f_j(x_i) \right]) \cdot \gamma^\top (c_i - \mathbb{E} \left[ c_i \right]) \right) + \text{constant}. \tag{21}
\]

Similarly, we can obtain the lower bound of the (unconditional) efficiency and non-spuriousness,

\[
\log \text{PNS}_n(f(X), Y) \\
\approx \left( \frac{1}{2\sigma^2} \sum_{i=1}^{n} \sum_{j=1}^{d} (\beta_j \cdot (f_j(x_i) - \mathbb{E} \left[ f_j(x_i) \right]))^2 \\
+ 2 \cdot \sum_{j=1}^{d} \beta_j \cdot (f_j(x_i) - \mathbb{E} \left[ f_j(x_i) \right]) \cdot \gamma^\top (c_i - \mathbb{E} \left[ c_i \right]) \right) + \text{constant}. \tag{22}
\]

Equations (21) and (22) illustrate how conditional efficiency and non-spuriousness differ from the unconditional version. The conditional notion considers each \( \beta_j \cdot f_j(X) \) at a time. The unconditional notion lumps together all \( d \) dimensions of the representation and considers \( \sum_{j=1}^{d} \beta_j \cdot f_j(X) \) as a whole. In this sense, the conditional PNS is finer-grained notion than the (unconditional) PNS.
A Linear example. We illustrate Algorithm 1 on a toy rank-degenerate image dataset. (We analyze more complex and higher-dimensional data in Section 2.4.) Imagine we collect a dataset of $n = 1000$ images. Each image is characterized by its values on $m = 5$ chosen representative pixels, $\mathbf{X} = (X_1, \ldots, X_5)$, accompanied by a label about the image brightness $Y$. Both the pixel values $\mathbf{X}$ and the labels $Y$ are real-valued.

We simulate such a dataset of image pixels and labels. As the pixel values are often highly correlated, we generate $\mathbf{x}_i = (x_{i1}, \ldots, x_{i5})$ from a multivariate Gaussian distribution with strong (positive or negative) correlations—all pairwise correlations are $> 0.8$. As the brightness label only depends on a small number of pixels, we simulate $y_i$ from a linear model that only uses two of the five pixels

$$y_i = \beta^*_1 x_{i1} + \beta^*_2 x_{i2} + \epsilon_i, \quad \epsilon_i \sim \mathcal{N}(0, \sigma^2),$$

where $\beta^*_1 = 0.5, \beta^*_2 = 1.0$.

The goal is to evaluate the efficiency and non-spuriousness of a representation $f(\mathbf{X}) = (X_2, 0.5 \cdot X_1 + X_4)$. We apply Algorithm 1: pinpoint the unobserved common cause $C$, estimate $P(Y \mid f(\mathbf{X}), C)$, and finally calculate the lower bound of efficiency and non-spuriousness $\text{PNS}_n(f(\mathbf{X}), Y)$ and $\text{PNS}_n(f_j(\mathbf{X}), Y \mid f_{-j}(\mathbf{X})), j = 1, \ldots, d$.

We first pinpoint $C$. Suppose it is known that the data $\mathbf{X}$ is generated by a PPCA. We fit a one-dimensional PPCA (Equation (15)) to $\{\mathbf{x}_i\}_{i=1}^n$; the latent variable $c_i$ is thus a scalar. The fit leads to the posterior of the unobserved common cause, $p(c_i \mid \mathbf{x}_i)$. We assess the pinpointability of $C$ by calculating $\text{Var}(c_i \mid \mathbf{x}_i)$ for all $i$. We find the variance is smaller than 0.01 for all $i$, implying that $p(c_i \mid \mathbf{x}_i)$ is fairly close to a point mass and pinpointability is approximately satisfied. (The threshold of 0.01 is a subjective choice.) We then calculate $\hat{c}_i = \mathbb{E}[c_i \mid \mathbf{x}_i]$ for all $i$.

We next estimate $P(Y \mid f(\mathbf{X}), C)$ by fitting a linear model to the dataset $\{(y_i, f(\mathbf{x}_i), \hat{c}_i)\}_{i=1}^n$ with $f(\mathbf{x}_i) = (x_{i2}, 0.5 \cdot x_{i1} + x_{i4}),$

$$y_i \sim \mathcal{N}(\beta_0 + \beta_1 \cdot x_{i2} + \beta_2 \cdot (0.5 \cdot x_{i1} + x_{i4}) + \gamma \cdot \hat{c}_i, \sigma^2).$$

Fitting this linear model returns the regression coefficients $\{\hat{\beta}_0, \hat{\beta}_1, \hat{\beta}_2, \hat{\gamma}\}$.

Finally, plugging in these regression coefficients to Equations (21) and (22) gives the efficiency and non-spuriousness, with $f_1(\mathbf{x}_i) = x_{i2}$ and $f_2(\mathbf{x}_i) = 0.5 \cdot x_{i1} + x_{i4}$.

### 2.3 CAUSAL-REP: Learning Efficient and Non-spurious Representations

We have developed a strategy to evaluate the efficiency and non-spuriousness of a given representation in Section 2.2. Here we utilize this strategy to improve representation learning in both supervised and unsupervised settings.

#### 2.3.1 Representation learning as finding necessary and sufficient causes

As representation learning requires efficient and non-spurious representations, we formulate representation learning as a task of finding necessary and sufficient causes of the label.
Representation learning as finding necessary and sufficient causes. To operationalize this formulation, we search for a representation that maximizes the conditional efficiency and non-spuriousness for a given dataset, following Definition 3 and Equation (6). Thus we view an ideal representation as one in which each dimension of the representation captures features that are essential and non-spurious given all other dimensions.

We perform representation learning by maximizing the sum of log \( \text{PNS} \) across all dimensions of the representation,

\[
\max_f \sum_{j=1}^d \log \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)),
\]

where \( \text{PNS}_n(\cdot, Y \mid \cdot) \) measures the conditional efficiency and non-spuriousness of \( f_j(X) \) as in Equation (6).

Classes of representation functions. To perform this maximization over representations in practice, we consider parameterized classes of representation functions \( f(\cdot) \). One option is to consider a class of neural network functions with a fixed architecture, subject to the constraint that each output dimension has zero mean and unit standard deviation, e.g.,

\[
\{ f : \text{multilayer perceptrons } \mathbb{R}^m \to \mathbb{R}^d \text{ with two hidden layers of size 512} \\
\text{s.t. } \{f(x_i)\}_{i=1}^n \text{ has sample mean 0 and standard error 1} \}.
\]

Another option is to consider representations that are convex combinations of the \( m \)-dimensional data,

\[
\{ f : f(X) = X_1 \times_m W_{m \times d}, \sum_{l=1}^m W_{lj} = 1, W_{lj} \geq 0, \forall l \in \{1, \ldots, m\}, j \in \{1, \ldots, d\} \).
\]

Each dimension of such representations is a convex combination of \( X_1, \ldots, X_m \), that is, \( f_j(X) = \sum_{l=1}^m W_{lj} X_l, j = 1, \ldots, d \).

A third option is to consider representations that select relevant subsets of the \( m \)-dimensional data,

\[
\{ f : f(X) = X_1 \times_m W_{m \times d}, W_{lj} \in \{0, 1\}, \sum_{l=1}^m W_{lj} \leq 1, \forall l \in \{1, \ldots, m\}, j \in \{1, \ldots, d\} \}.
\]

Each dimension of such representations selects one (or none) of \( X_1, \ldots, X_m \). The \( d \)-dimensional representation \( f(X) \) thus selects at most \( d \) features from \( X_1, \ldots, X_m \).

CAUSAL-REP: Maximizing \( \text{PNS} \) in practice. Solving Equation (25) involves calculating counterfactual quantities such as conditional \( \text{PNS} \). As is discussed in Section 2.2, these quantities are not directly calculable without access to the causal structural equations.
To employ Equation (25) in practice, we employ the lower bound of \( \text{PNS} \) derived in Equation (7). Specifically, we restrict our attention to representations whose lower bound of \( \text{PNS} \) is identifiable via Theorem 7, and find the representation that maximizes this lower bound:

\[
(\text{CAUSAL-REP objective}) \quad \max_f \sum_{j=1}^{d} \log \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)) + \lambda \cdot R(f ; \{x_i, y_i, c_i\}_{i=1}^{n}),
\]

(29)

where \( \text{PNS}_n(f_j(X), Y \mid f_{-j}(X)) \) is the \( \text{PNS} \) lower bound (Equation (14)) in Theorem 7. The parameter \( \lambda \geq 0 \) indicates regularization strength. The term \( R(f ; \{x_i, y_i, c_i\}_{i=1}^{n}) \) is a regularization penalty,

\[
R(f ; \{x_i, y_i, c_i\}_{i=1}^{n}) = \frac{1}{d} \sum_{j=1}^{d} \log(1 - \text{Rsq}(f_j(x_i) ; c_i)) - \alpha \cdot ||W(f)||_2^2,
\]

(30)

where \( \text{Rsq}(f_j(x_i) ; c_i) \) is the R-squared of regressing the \( j \)-th dimension of the representation \( f_j(x_i) \) against the unobserved common cause \( c_i \). The quantity \( ||W(f)||_2 \) represents the \( L_2 \)-norm of the \( f \) function’s parameters, e.g., all the weight parameters of a neural network \( f \); \( \alpha \) is the relative weight of the two regularization penalties.

The first part of the objective (Equation (29)) is the lower bound of conditional \( \text{PNS} \) developed in Theorem 7. Following the practical discussions in Section 2.2, its calculation requires pinpointing the unobserved common cause \( C = h(X) \) and fitting a model to \( P(Y \mid f(X), C) \) (e.g., Equation (19)).

The second part of the objective is the regularization penalty. It encourages representations whose lower bound of \( \text{PNS} \) is identifiable from observational data. Specifically, these regularization penalties aim to enforce the positivity and observability conditions in Lemma 6. (The pinpointability condition is enforced in a separate step.)

In more detail, the first penalty \( \frac{1}{d} \sum_{j=1}^{d} \log(1 - \text{Rsq}(f_j(x_i) ; c_i)) \) in Equation (30) encourages the positivity of \( f(X) \) given \( C \) in Lemma 6. The sample R-squared \( \text{Rsq}(f_j(x_i) ; c_i) \) evaluates the variation of \( f_j(X) \) explainable by \( C \). When the R-squared is equal to one, then the positivity condition is violated. Accordingly, a close-to-one value of R-squared implies that \( f_j(X) \) nearly violates the positivity condition. The penalty \( \frac{1}{d} \sum_{j=1}^{d} \log(1 - \text{Rsq}(f_j(x_i) ; c_i)) \) takes a large negative value if any dimension of the representation \( f(X) \) nearly violates the positivity condition.

The second penalty \( -\alpha \cdot ||W(f)||_2^2 \) encourages representations that satisfy the observability condition. Specifically, it penalizes the coefficients in front of \( (X_1, \ldots, X_m) \) in the representation function \( f \). Imposing a large regularization parameter \( \alpha > 0 \) leads to representations \( f(X) \) that only effectively depend on a small subset of \( (X_1, \ldots, X_m) \); i.e., \( f(X) = \tilde{f}(X_{j \in S}) \) for some function \( \tilde{f} \), with \( S \subseteq \{1, \ldots, m\} \) being a set with only a few elements. Such representations are more likely to satisfy the observability condition—namely, \( P((X_j)_{j \in S}) > 0 \) for all values of \( (X_j)_{j \in S} \)—because lower-dimensional \( (X_j)_{j \in S} \) are more likely to have full rank (Udell & Townsend, 2019).
Out-of-distribution prediction with CAUSAL-REP. Given a representation returned by CAUSAL-REP, how do we make predictions, especially on out-of-distribution data?

To make predictions, we train a prediction function that maps the CAUSAL-REP representation to the labels. We note that the predictions are only made using the CAUSAL-REP representation; it does not involve the unobserved common cause $C$. This prediction function is different from the prediction model fitted for $P(Y \mid f(X), C)$ (cf. Equations (19) and (20)). The rationale is that CAUSAL-REP encourages non-spurious representations; it implies that the relationship between the representation $f(X)$ and the label $Y$ should generalize to out-of-distribution data. However, CAUSAL-REP places no constraints on the relationship between $C$ and $Y$.

Specifically, suppose CAUSAL-REP returns $\hat{f}(\cdot)$ as the representation function; i.e., it maximizes Equation (29). Then we posit a model for $P(Y \mid \hat{f}(X))$; e.g., a linear model:

$$P(Y \mid \hat{f}(X)) = \mathcal{N}(\beta_0^{\text{pred}} + (\beta^{\text{pred}})^\top \hat{f}(X), (\sigma^{\text{pred}})^2),$$

or a flexible exponential family model:

$$P(Y \mid \hat{f}(X)) = \text{EF}(g^{\text{pred}}(\hat{f}(X))),$$

where $\text{EF}$ indicates an exponential family distribution and $g^{\text{pred}}$ indicates a link function. We fit this model to the training data $\{(\hat{f}(x_i), y_i)_{i=1}^n\}$ using maximum likelihood estimation, and then use this fitted model to predict on out-of-distribution data; e.g., we calculate $\mathbb{E}[Y \mid \hat{f}(X_{\text{test}})]$.

CAUSAL-REP with perfectly correlated spurious and non-spurious features. We have described CAUSAL-REP, a representation learning algorithm that targets non-spurious and efficient representations. A reader might ask: What if spurious and non-spurious features are perfectly correlated in the training dataset? That is, what if the spurious feature is on if and only if the non-spurious feature is also on? Intuitively, we can not hope to tease apart spurious and non-spurious representations in this case. How would CAUSAL-REP fare?

In such a perfect correlation case, the CAUSAL-REP algorithm would capture neither the spurious feature nor the non-spurious one. The reason is that both features are excluded from the feasible set of representations considered by CAUSAL-REP. Their $\text{pns}$ lower bounds are both not identifiable, but CAUSAL-REP only considers representations whose $\text{pns}$ lower bound is identifiable.

In more detail, suppose $f_s((X_j)_{j \in T_s})$ captures the spurious feature and $f_n((X_j)_{j \in T_n})$ captures the non-spurious features, where $T_s, T_n \subset \{1, \ldots, m\}$. They depend on disjoint subsets of $X$, $T_s \cap T_n = \emptyset$, but they are perfectly correlated $f_s((X_j)_{j \in T_s}) = f_n((X_j)_{j \in T_n})$. Then both features must be measurable with respect to the unobserved common cause $C$: $f_s((X_j)_{j \in T_s}) = f_n((X_j)_{j \in T_n}) = q(C)$ for some function $q$. This measurability makes their $\text{pns}$ lower bounds non-identifiable; it violates the positivity assumption required by Theorem 7.

---

$^5$The reason is that $C$ must render all $X_j$’s conditionally independent due to the causal graph (Figure 2). It must also render the spurious and non-spurious features conditionally independent, because they depend on disjoint subsets of $X$. As the two features are perfectly correlated, the only way to render them conditionally independent is to make them measurable with respect to $C$. 

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Algorithm 2: CAUSAL-REP (Supervised)

input : The training data and its label \( \{x_{i\text{train}}, y_{i\text{train}}\}_{i=1}^{n_{\text{train}}} \), the (out-of-distribution) test data \( \{x_{i\text{test}}, y_{i\text{test}}\}_{i=1}^{n_{\text{test}}} \), and the probabilistic factor model that generates the training data \( P(X, C) \)

output : CAUSAL-REP representation function \( \hat{f}(\cdot) \); predictions on the test data \( \hat{y}_i, i = 1, \ldots, n_{\text{test}} \)

# Representation learning with CAUSAL-REP
Fit a probabilistic factor model (Equations (17) and (18)) to the training data and infer \( p(c_i | x_{i\text{train}}), i = 1, \ldots, n_{\text{train}} \);

if Pinpointability holds, i.e. \( p(c_i | x_{i\text{train}}) \) is close to a point mass for all \( i \) then

foreach training datapoint \( i \) do

Pinpoint the unobserved common cause \( C : c_i = h(x_{i\text{train}}) \triangleq \mathbb{E}[c_i | x_{i\text{train}}] \) for \( i = 1, \ldots, n_{\text{train}} \);

end

Maximize Equation (29) to obtain the CAUSAL-REP representation \( \hat{f} \);

end

# Out-of-distribution prediction with the CAUSAL-REP representation
Estimate \( P(Y | \hat{f}(X)) \) by fitting a model (e.g. Equations (31) and (32)) to \( \{\hat{f}(x_{i\text{train}}), y_{i\text{train}}\}_{i=1}^{n_{\text{train}}} \)

Predict on test data using the fitted model: \( \hat{y}_i = \mathbb{E}[Y | \hat{f}(x_{i\text{test}})], i = 1, \ldots, n_{\text{test}} \)

2.3.2 CAUSAL-REP and the linear example continued

We described each step of CAUSAL-REP in the previous section. Algorithm 2 summarizes these steps. The key step is to maximize the CAUSAL-REP objective (Equation (29)). This step involves a nested loop of optimization: an outer loop with respect to the representation parameters \( W(f) \), and an inner loop to estimate \( P(Y | f(X), C) \). This nesting is required because calculating the CAUSAL-REP objective involves estimating \( P(Y | f(X), C) \) by maximum likelihood.

To maximize the CAUSAL-REP objective, one can use standard gradient-based algorithms to handle this nested loop. When the inner loop—the maximum likelihood estimation of \( P(Y | f(X), C) \)—has a closed-form solution (e.g., under the linear model in Equation (19)), we can plug in the solution to the CAUSAL-REP objective and directly apply a gradient-based method for optimization. When the inner loop does not admit a closed form, we can alternate between the gradient updates of the two optimizations, e.g., alternating between (1) one gradient update step to maximize the CAUSAL-REP objective, and (2) multiple gradient update steps until convergence to estimate \( P(Y | f(X), C) \) with maximum likelihood.

The linear example continued. We illustrate CAUSAL-REP (Algorithm 2) with the linear example in Section 2.2.3: pinpoint the unobserved common cause \( C \) from the training data \( \{x_{i\text{train}}, y_{i\text{train}}\}_{i=1}^{n_{\text{train}}} \), obtain the CAUSAL-REP objective \( \hat{f} \), and predict on some out-of-distribution test data \( \{x_{i\text{test}}\}_{i=1}^{n_{\text{test}}} \). We focus on learning two-dimensional representations, \( f(X) = (f_1(X), f_2(X)) \).

We first perform the same pinpointability step to obtain \( \hat{c}_i \) as in Section 2.2.3.
We next maximize the CAUSAL-REP objective (Equation (29)). We calculate the CAUSAL-REP objective by plugging in the calculation of \( p_{n_{\text{sc}}} (f_j (X), Y | f_{-j} (X)) \) in Equation (21), where we adopt a linear model for \( P(Y | f(X), C) \). In more detail, to optimize the CAUSAL-REP objective, we need to write (the parameters of) the fitted model of \( P(Y | f(X), C) \) as a function of the representation function \( f \). We use the closed-form estimates of these linear model parameters when fitted to the training data \( \{x_i^{\text{train}}, y_i^{\text{train}}\}_{i=1}^{n_{\text{train}}} \) :

\[
(\hat{\beta}_0, \hat{\beta}_1, \hat{\beta}_2, \hat{\gamma}) = \arg\min \sum_{i=1}^{n} (y_i^{\text{train}} - \beta_0 - \beta_1 \cdot f_1(x_i^{\text{train}}) - \beta_2 \cdot f_2(x_i^{\text{train}}) - \gamma \cdot \hat{c}_i)^2
\]

\[
= (\hat{X}^\top \hat{X})^{-1}(\hat{X}^\top \hat{Y}),
\]

where \( \hat{X} \) is an \( n_{\text{train}} \times 4 \) matrix with \( \hat{X}_{i1} = 1, \hat{X}_{i2} = f_1(x_i^{\text{train}}), \hat{X}_{i3} = f_2(x_i^{\text{train}}), \hat{X}_{i4} = \hat{c}_i \), and \( \hat{Y} \) is an \( n \times 1 \) vector with \( \hat{Y}_{i1} = y_i^{\text{train}}, i = 1, \ldots, n \). (When closed-form solutions are not available, we need to perform gradient-based optimization to obtain the fitted model parameters.)

Plugging in these terms, we maximize the CAUSAL-REP objective via gradient descent with respect to the parameters of the representation function \( \hat{W}(f) \). The optimal \( \hat{W}(f) \) give the CAUSAL-REP representation function \( \hat{f} \).

Finally, we perform out-of-distribution prediction with the CAUSAL-REP representation \( \hat{f} \). We train a prediction function by fitting a linear model to \( P(Y | \hat{f}(X)) \),

\[
y_i^{\text{train}} = \beta_0^{\text{pred}} + \beta_1^{\text{pred}} \cdot \hat{f}_1(x_i^{\text{train}}) + \beta_2^{\text{pred}} \cdot \hat{f}_2(x_i^{\text{train}}) + \epsilon, \epsilon \sim \mathcal{N}(0, (\sigma_{\text{pred}})^2).
\]

We obtain the estimated regression coefficients \( \{\hat{\beta}_0^{\text{pred}}, \hat{\beta}_1^{\text{pred}}, \hat{\beta}_2^{\text{pred}}, \sigma_{\text{pred}}\} \) by maximum likelihood. They allow us to make predictions on the test data \( \{x_i^{\text{test}}\}_{i=1}^{n_{\text{test}}} \),

\[
\hat{y}_i = \mathbb{E}[Y | \hat{f}(x_i^{\text{test}})] = \hat{\beta}_0^{\text{pred}} + \hat{\beta}_1^{\text{pred}} \cdot \hat{f}_1(x_i^{\text{test}}) + \hat{\beta}_2^{\text{pred}} \cdot \hat{f}_2(x_i^{\text{test}}).
\]

### 2.3.3 Extending CAUSAL-REP to unsupervised settings

We extend CAUSAL-REP to unsupervised settings where labels are not available. We focus on the task of instance discrimination in unsupervised representation learning (Hadsell et al., 2006), where the goal is to find representations that can distinguish the different subjects.

**The unsupervised CAUSAL-REP.** We focus on the unsupervised setting where some form of data augmentation is available. That is, the dataset contains one raw observation for each subject; this raw observation is then augmented multiple times, which leads to multiple observations per subject. Given this unsupervised dataset, which attach the subject ID to each observation as a supervised label. We can thereby formulate the instance discrimination problem as a supervised task, one that finds representations that are informative of the subject IDs and extend CAUSAL-REP to an unsupervised setting. The resulting algorithm turns out to be closely related to contrastive learning (Chen et al., 2020a).

In more detail, we begin with a dataset with \( n \) i.i.d. samples, \( \{x_1, \ldots, x_n\} \). Assume that each data point is associated with a different subject \( i \). Next, we augment each data point with \( U - 1 \) augmentations, producing an augmented dataset, \( \{\{x_i^{u}\}_{u=1}^{U}\}_{i=1}^{n}, U \geq 2 \). We then create an \( n\)-
dimensional label, \(y^u_i = (y^u_{i1}, \ldots, y^u_{im})\), for each data point in the augmented dataset, where \(y^u_{is} = \mathbb{I}\{x^u_i \text{ belongs to } s\}\), s = 1, \ldots, n. \]

This labeling turns the unsupervised problem into a supervised one with respect to the augmented labeled dataset \(\{\{x^u_i, y^u_i\}^U_{u=1}\}^n_{i=1}\). We then employ the CAUSAL-REP objective to find representations that are necessary and sufficient causes of the n outcomes \((Y_1, \ldots, Y_n)\):

\[
\max_f \sum^n_{s=1} \sum^m_{j=1} \log \text{PNS}_{n,U}(f_j(X), Y_s \mid f_{-j}(X)) + \lambda \cdot R(f ; \{\{x^u_i, y^u_i; c^u_i\}^U_{u=1}\}^n_{i=1}),
\]

where \(\text{PNS}_{n,U}(f_j(X), Y_s \mid f_{-j}(X))\) is the conditional efficiency and spuriousness (Equation (6)) for the sth outcome; it is calculated on the augmented dataset \(\{\{x^u_i, y^u_i\}^U_{u=1}\}^n_{i=1}\). The penalty term \(R(\cdot)\) is the same penalty as in the CAUSAL-REP objective (Equations (29) and (30)). Solving Equation (36) produces non-spurious and efficient representations of \(X\) for instance discrimination. (We summarize the algorithm in Algorithm 4.)

The optimization objective in Equation (36) considers the n outcomes \(\{Y_s\}\) one at a time and then averages over them. It focuses on the contrast of one-vs-all, finding representations that distinguish each subject from the rest. This objective does not consider the n outcomes as a single n-dimensional outcome \((Y_1, \ldots, Y_n)\) and calculate its \(\text{PNS}\).

**Unsupervised CAUSAL-REP and contrastive learning.** The unsupervised CAUSAL-REP objective (Equation (36)) is closely related to contrastive learning (Chen et al., 2020a), whose learning objective is to maximize

\[
\sum^U_{u_1=1} \sum^U_{u_2=1} - \log \frac{\exp(d(f(x^{u_1}_i), f(x^{u_2}_i)))}{\sum_{i' \neq i} \sum^U_{u_1=1} \exp(d(f(x^{u_1}_i), f(x^{u_1}_{i'})))},
\]

where \(d(\cdot, \cdot)\) is some distance function. This contrastive learning objective targets the same instance discrimination task as the unsupervised CAUSAL-REP does; it aims for representations that can determine whether two data points come from the same subject. Contrastive learning thus is also able to predict whether each data point belongs to subject s, for s = 1, \ldots, n, as is the unsupervised CAUSAL-REP.

That said, the two algorithms only coincide when different dimensions of \(X\) are independent—they do not share an unobserved common cause as in Figure 2. The two algorithms behave very differently when such a common cause exists, especially when the common cause induces some spurious feature (e.g., image color). Concretely, there may be a subset of \((X_1, \ldots, X_m)\) that is highly correlated with the label \(Y\) but cannot causally determine \(Y\). In such cases, contrastive learning may pick up these spurious features if no augmentations are performed for these features (e.g., there is no augmentation that randomly changes the color of images). In contrast, the unsupervised CAUSAL-REP would not capture these spurious features.

This difference illustrates how CAUSAL-REP learns non-spurious representations in a different way than data-augmentation-based methods (including contrastive learning). CAUSAL-REP capitalizes on a causal perspective and its treatment of high-dimensional \(X\). Data-augmentation-based methods instead wipe out the correlation between spurious features and the label by performing data augmentation. We will illustrate this difference further in Section 2.4.5.
2.4 Empirical Studies of CAUSAL-REP

We study CAUSAL-REP in both image and text datasets. We find that probabilities of causation ( poc ) are effective in distinguishing efficient/inefficient and non-spurious/spurious representations. Moreover, CAUSAL-REP finds non-spurious features in both supervised and unsupervised settings; it also outperforms existing unsupervised representation learning algorithms in downstream out-of-distribution prediction.

2.4.1 How well do probabilities of causation measure efficiency and non-spuriousness of features?

We first study the correspondence between probabilities of causation ( ps, pn, pns ) and the efficiency/non-spuriousness of representations. We generate features with known efficiency and non-spuriousness properties; we find that the (lower bound of) probabilities of causation in Lemma 4 are consistent with these properties.

The simulated data. We simulate two binary features Z_1, Z_2 and two binary outcomes Y_1, Y_2:

\[
\begin{align*}
Z_1 & \sim \text{Bernoulli}(0.4), \\
Z_2 & = Z_1 \oplus \text{Bernoulli}(p), \ p \in \{0, 0.1, \ldots, 0.9, 1\}, \\
Y_1 & = Z_1 \oplus \text{Bernoulli}(0.2), \\
Y_2 & = (Z_1 \& Z_2) \oplus \text{Bernoulli}(0.2),
\end{align*}
\]

where \( \oplus \) indicates the XOR operator. We vary the parameter \( p \); a small \( p \) implies a high correlation between \( Z_1 \) and \( Z_2 \). The generative process of \( Y_1, Y_2 \) implies that (1) \( Z_1 \) is necessary (a.k.a. efficient) and sufficient (a.k.a. non-spurious) for \( Y_1 \), but is necessary and insufficient for \( Y_2 \); (2) \( Z_2 \) is neither necessary nor sufficient for \( Y_1 \), but is necessary and insufficient for \( Y_2 \); (3) \( Z_1 \& Z_2 \) is necessary and sufficient for \( Y_2 \). Moreover, when the correlation between \( Z_1 \) and \( Z_2 \) increases, then using both features as a representation become increasingly inefficient; the conditional necessity decreases. We calculate the lower bound of \( \text{ps, pn, pns} \) of \( Z_1, Z_2 \) for both outcomes \( Y_1, Y_2 \); the exact value is not identifiable from data.

Results. Figures 4, 12 and 13 present the probabilities of causation of the features. Figure 4a shows that the (lower bound of) conditional \( \text{pns} \) can correctly signal the non-spuriousness of features: \( Z_2 \) is spurious for \( Y_1 \), but \( Z_1 \) is non-spurious for \( Y_1 \), and both \( Z_1, Z_2 \) are non-spurious for \( Y_2 \). Moreover, we study how probabilities of causation ( poc ) fare when the correlation between \( Z_1 \) and \( Z_2 \) increases. Figure 4 shows that the (lower bounds of) \( \text{poc} \) of \( Z_1 \) for \( Y_2 \) increase as the correlation increases, which is consistent with the intuition that \( Z_2 \) becomes less necessary for \( Y_2 \) given \( Z_1 \) given increasingly higher correlations. Similarly, Figures 12 and 13 show that (the lower bounds of) the unconditional \( \text{poc} \) of \( Z_2 \) for \( Y_1 \) also increase. It is also consistent with the intuition: \( Z_2 \) is an increasingly better surrogate of \( Z_1 \) for \( Y_1 \) under higher correlations between \( Z_1 \) and \( Z_2 \).
We next study the supervised CAUSAL-REP in image datasets: Colored MNIST (Arjovsky et al., 2019) and CelebA (Liu et al., 2015). We consider the version of supervised CAUSAL-REP algorithm which adopts a VAE with 64 latent dimensions as the probabilistic factor model for pinpointing. For

2.4.2 Does the supervised CAUSAL-REP pick up spurious features in synthetic data?

We next study the performance of the supervised CAUSAL-REP in a toy synthetic dataset. We use PPCA as the pinpointing factor model and linear functions as the class of representation functions.

The simulated data. We generate a dataset of core and spurious features \( (X_1, \ldots, X_5) \), who are highly correlated in the training set but not so in the test set,

\[
\tilde{X}_1^{\text{train}}, \ldots, \tilde{X}_5^{\text{train}} \sim \mathcal{N}(0, 0.05 \cdot I_5 + 0.95 \cdot J_5),
\]

\[
\tilde{X}_1^{\text{test}}, \ldots, \tilde{X}_5^{\text{test}} \sim \mathcal{N}(0, 0.05 \cdot I_5 + 0.05 \cdot J_5),
\]

where \( I_5 \) is a \( 5 \times 5 \) identity matrix and \( J_5 \) is a \( 5 \times 5 \) all-ones matrix. For both training and test sets, we inject noise to features to lower the signal-to-noise ratio of the problem: \( X_j \sim \mathcal{N}(\tilde{X}_j, 0.4^2), j = 1, 2; X_j \sim \mathcal{N}(\tilde{X}_j, 0.3^2), j = 3, 4, 5. \) Finally, we generate an outcome \( Y \) that only depends on the core features \( Y \sim \mathcal{N}(\beta_1 X_1 + \beta_2 X_2, 1) \), where the coefficients are drawn from a uniform \( \beta_j \sim \text{Unif}[0, 10], j = 1, 2. \)

Results. We compare the performance of CAUSAL-REP, linear regression, and an oracle algorithm; the oracle is equipped with the knowledge of which features are spurious and only performs linear regression against the non-spurious features. We study whether these algorithms pick up spurious features by evaluating their the out-of-distribution (ood) predictive performances. If an algorithm picks up spurious features, then its ood predictions will suffer, because spurious features are much less predictive in test data than in training data. Figure 5a presents the result: CAUSAL-REP outperforms linear regression in ood predictive R-squared; its predictive performance is not far from the oracle algorithm.

2.4.3 Does the supervised CAUSAL-REP produce non-spurious representations for image data? A study on Colored MNIST and CelebA

We next study the supervised CAUSAL-REP in image datasets: Colored MNIST (Arjovsky et al., 2019) and CelebA (Liu et al., 2015). We consider the version of supervised CAUSAL-REP algorithm which adopts a VAE with 64 latent dimensions as the probabilistic factor model for pinpointing. For
Figure 5: CAUSAL-REP learns non-spurious representations in both supervised and unsupervised settings. (a) CAUSAL-REP outperforms linear regression in OOD prediction in toy synthetic data. (b) CAUSAL-REP outperforms baseline representation learning algorithms (e.g. directly fitting neural networks, or performing contrastive learning, or adopting VAE representations) in the colored and shifted MNIST dataset. The dashed yellow line indicates the theoretical maximum of OOD predictive accuracy. (Higher is better.)

representation functions, we consider a two-layer neural network with 20-dimensional outputs. We again evaluate the non-spuriousness of the representations by the OOD predictive accuracy; if the learned representation captures spurious features, then it will suffer in OOD prediction.

Competing methods. We compare CAUSAL-REP with a baseline representation learning algorithm that fits a neural network to the labels and extracts its penultimate layer as the representation (Bengio et al., 2013). As we use VAE for pinpointing in CAUSAL-REP in this study, we also compare with directly adopting the VAE representation for OOD prediction.

The Colored MNIST study. We focus on the colored MNIST data with the digits ‘3’ and ‘8’ and colors ‘red’ and ‘green’; see Appendix G.2 for the detailed experimental setup.

Figure 6a presents the OOD predictive accuracy of CAUSAL-REP across different levels of spurious correlations. CAUSAL-REP outperforms other baseline methods in OOD predictive accuracy when the spurious correlation is larger than 0.5. Its performance remains close to the theoretical maximum (the yellow dashed line) across different levels of the spurious correlation, suggesting that CAUSAL-REP does not pick up spurious features. In contrast, the OOD performance of representation learning by directly fitting neural networks quickly degrades as the spurious correlation increases; so does OOD prediction with the VAE representations.

Figure 6b evaluates the robustness of CAUSAL-REP against the latent dimensionality choice of the pinpointing factor model, fixing the spurious correlation at 0.9. We find that the performance of CAUSAL-REP does not change much as we vary the latent dimensionality of pinpointing VAE. The relative performance of CAUSAL-REP and the competing methods also stay stable.
Figure 6: (a) The supervised CAUSAL-REP learns non-spurious representations in colored MNIST and outperforms baseline representation learning algorithms (e.g. directly fitting neural networks, or adopting VAE representations) in OOD prediction. (b) The performance of CAUSAL-REP is robust to the choice of the latent dimensionality of probabilistic factor models. The dashed yellow line indicates the theoretical maximum of OOD predictive accuracy. (Higher is better.)

The CelebA study. We next study CAUSAL-REP on the CelebA dataset (Liu et al., 2015). We create training and test sets, each containing 5,000 data points, by subsampling the CelebA datasets. We focus on face attributes with a relatively balanced distribution in the raw CelebA dataset. We designate pairs of target attributes and spurious attributes, and subsample such that the two are highly correlated in the training set and not as correlated in the test set. We then perform representation learning and OOD prediction for the target labels, using CAUSAL-REP and other competing methods.

Table 1 presents the results for different pairs of target and spurious face attributes. Though the spurious label and the target label are highly correlated, CAUSAL-REP can pick up non-spurious features that inform the target label and outperform the baseline algorithm that directly fits a neural network. In most settings, CAUSAL-REP also outperforms OOD prediction with VAE representations. The exception is that VAE representations can outperform CAUSAL-REP representations when the spurious correlations climb up to 0.9, though the performance of CAUSAL-REP is still competitive. The spurious features in these settings violate the positivity condition: Mustache and goatee exclusively appear on males in the subsampled dataset.
Table 1: CAUSAL-REP learns non-spurious representations in the CelebA dataset and outperforms baseline representation learning algorithms in ood prediction.

2.4.4 Does the supervised CAUSAL-REP produce non-spurious representations for text data? A study on reviews corpora and sentiment analysis

We next study CAUSAL-REP on text datasets: the Amazon (Wang et al., 2010, 2011), Tripadvisor, and Yelp reviews corpora, and the IMDB-L, IMDB-S, and Kindle dataset as is processed in Wang & Culotta (2020, 2021). In these studies, we convert these corpora into bags of words. We use PPCA as a pinpointing factor model for CAUSAL-REP and consider representations whose each dimension is a convex combination of words.

The reviews corpora study. We begin with the raw reviews datasets from Amazon, Tripadvisor, and Yelp. We create a binary label for each review by converting 4 and 5 stars to a positive label and 1 and 2 stars to a negative label. We then inject irrelevant words to the training dataset as spurious features by randomly adding in “as”, “also”, “am”, “an” to reviews with positive labels; the resulting spurious correlation is around 0.9. We create two test datasets: one is in-distribution with the spurious words present as in the training set; the other is out-of-distribution without the randomly added spurious words.

Figure 7 presents the predictive accuracy of CAUSAL-REP in both test sets. We find that the predictive performance of CAUSAL-REP is stable across in-distribution and out-of-distribution test sets, suggesting that it learns non-spurious representations. In contrast, logistic regression predicts well in the in-distribution test set but not in the out-of-distribution test set, suggesting that it picks up spurious features that only exist in the training set.

Table 4 presents the most informative words of the (positive or negative) ratings, suggested by the CAUSAL-REP representation and the logistic regression coefficients. Across three reviews corpora, logistic regression returns the spurious words “as”, “also”, “am”, “an” as the top words. In contrast, CAUSAL-REP extracts words that are more relevant for the ratings.

\[\text{http://times.cs.uiuc.edu/wang296/Data/}\]
\[\text{https://www.yelp.com/dataset/documentation/main}\]
Figure 7: CAUSAL-REP learns non-spurious representations across reviews text corpora; its predictive performance is stable across in-distribution and out-of-distribution test sets.

Table 2: CAUSAL-REP outperforms naive representation learning algorithms in predicting on counterfactual test sets.

|                      | Observational test set | Counterfactual test set |
|----------------------|------------------------|-------------------------|
|                      | Logistic Regression    | CAUSAL-REP              | Logistic Regression | CAUSAL-REP              |
| IMDB-L               | 0.669                  | 0.645                   | 0.591              | 0.642                   |
| IMDB-S               | 0.836                  | 0.682                   | 0.570              | 0.621                   |
| Kindle               | 0.850                  | 0.618                   | 0.468              | 0.572                   |

The sentiment analysis study. We next employ CAUSAL-REP for sentiment classification on the IMDB-L, IMDB-S, and Kindle datasets (Wang & Culotta, 2020, 2021). We evaluate CAUSAL-REP on their raw observational and counterfactual test sets, without employing additional data augmentations as in Wang & Culotta (2020, 2021). The observational test sets are in-distribution test sets, and the counterfactual test sets are out-of-distribution test sets.

Table 2 presents the predictive performance of CAUSAL-REP compared with logistic regression. While logistic regression outperforms in in-distribution predictive accuracy, CAUSAL-REP outperforms in out-of-distribution prediction. Moreover, the predictive performance of CAUSAL-REP is similar across in-distribution and out-of-distribution test sets, suggesting that CAUSAL-REP produces non-spurious representations.

2.4.5 How well does unsupervised CAUSAL-REP perform on instance discrimination? A study on colored and shifted MNIST

Finally, we study CAUSAL-REP in the unsupervised setting. We focus on image datasets in the unsupervised setting because non-spurious features that distinguish different images are more readily defined in the image domain than in the text domain. We evaluate the non-spuriousness of the unsupervised CAUSAL-REP again by ood prediction. Given the unsupervised CAUSAL-REP representation, we fit a prediction model to the target label and test its predictive performance on ood test sets.
**Competing methods.** We compare CAUSAL-REP with baseline representation learning algorithms that fit a neural network to the subject ID label in Section 2.4.5. We also compare with contrastive learning (Chen et al., 2020a) and the VAE representation.

**The colored and shifted MNIST study.** We construct the colored and shifted MNIST dataset by coloring and shifting digits in a way that is highly correlated with the digit labels; see Appendix G.4 for details. (Representation learning with the unsupervised CAUSAL-REP does not make use of the digit labels; these labels are only used in producing a prediction function from the representations to the labels.)

Figure 5b presents the ood predictive accuracy of the unsupervised CAUSAL-REP on colored and shifted MNIST. CAUSAL-REP outperforms baseline representation learning algorithms when the spurious correlation is high, and their predictive accuracy stays close to the theoretical maximum (the yellow dashed line). Comparing CAUSAL-REP with contrastive learning with shift augmentation, we find that the ood predictive accuracy of contrastive learning degrades when the spurious correlations increase over 0.7. It is because the shift augmentation can not rule out the spurious color feature; it can only avoid picking up the spurious shift feature. This observation echoes the discussion in Section 2.4.5, illustrating how the unsupervised CAUSAL-REP relies on a different mechanism to produce non-spurious representations than contrastive learning and data augmentation.

3 **Unsupervised Representation Learning:**

**Disentanglement**

In this section we study the desideratum of disentanglement in unsupervised representation learning. Unsupervised representation learning aims to find a low-dimensional representation for high-dimensional objects without the help of labels. Given an \( m \)-dimensional object, \( X = (X_1, \ldots, X_m) \), the goal is again to find a \( d \)-dimensional representation, \( Z = (Z_1, \ldots, Z_d) \triangleq (f_1(X), \ldots, f_d(X)) \).

We focus on a causal definition of disentanglement: different dimensions of the representation shall encode features that do not causally affect each other (Suter et al., 2019). An absence of causal relationships among different dimensions of the representation can be viewed as independent manipulability of each dimension. Other related criteria include the notion of independent mechanisms (Parascandolo et al., 2018; Schölkopf et al., 2012; Suter et al., 2019) and the concept of independently controllable factors (Thomas et al., 2018, 2017). Based on this causal definition, we will develop unsupervised metrics and algorithms for unsupervised disentanglement.

3.1 **The Causal Definition of Disentanglement**

We begin with reviewing the causal definition of disentanglement. A representation is (causally) disentangled if (1) it represents features that can generate the original data and (2) its dimensions correspond to features that do not causally affect each other (Suter et al., 2019). Definition 8 casts these requirements in terms of a structural causal model (scm) (Pearl, 2011).
Figure 8: Disentangled features have independent support even though they may be correlated. Moreover, the independence of support can distinguish disentangled and entangled features. This figure illustrates how entangled and disentangled features differ using pairwise scatter plots. Figure 8a considers the ground truth features (shape, scale, orientation) of the dSprites dataset. These features are disentangled. They also have independent support; e.g., conditional on ‘scale’, the set of values that ‘orientation’ can take does not change with ‘scale.’ Visually, these disentangled features have scatter plots that occupy rectangular (or hyperrectangular) regions. Figure 8b considers the same features but in a subset of the dSprites dataset where the features are correlated. These features, though correlated, are still disentangled; they also have independent support. Figure 8c considers three entangled features, each of which is a nonlinear transformation of the three ground-truth features. These features are not disentangled. Their supports are also not independent. Conditional on ‘entangle1’, the possible values ‘entangle2’ can take depends on the value of ‘entangle1.’

**Definition 8** (Causal Disentanglement (Suter et al., 2019)). A $d$-dimensional representation, $Z = (Z_1, \ldots, Z_d)$, is disentangled if it represents features that generate the object of interest $X = (X_1, \ldots, X_m)$ according to the following causal SCM:

\[
\begin{align*}
C & \leftarrow u_C, \\
Z_j & \leftarrow f^z_j(C, u_{z,j}), & j = 1, \ldots, d, \\
X_l & \leftarrow f^x_l(Z, u_X), & l = 1, \ldots, m,
\end{align*}
\]
where $C = (C_1, \ldots, C_K)$ denotes an $K$-dimensional unobserved confounder that can affect $Z$ and hence induce correlations among its components $(Z_1, \ldots, Z_d)$. Figure 2 describes this $\text{scm}$ using a causal DAG.\(^8\)

As an example to illustrate this definition, we return to the dataset of animal images in Section 1. If we had access to the $d$ ground-truth features that generated the dataset (e.g., fur color, number of legs), we could create a $d$-dimensional representation that concatenates them. Such a representation would be disentangled, as the ground-truth features do not causally affect each other. That said, they may still be correlated due to some unobserved confounders (a.k.a., common causes). For example, the animal’s genetic information may be an unobserved confounder; the genes can causally affect these features, and thus induce correlations among them.

Notably, the causal definition of disentanglement opens up the possibility to consider correlated features that are disentangled. We work with this disentanglement definition in the rest of this section.

### 3.2 Measuring Disentanglement with Observational Data

How do we assess the disentanglement of a representation? In the do notation, a representation $Z$ is disentangled if, for $j = 1, \ldots, d$,

$$P(Z_j \mid \text{do}(Z_{-j} = z_{-j})) = P(Z_j), \forall z_{-j} \in Z_{-j},$$

where $P(\cdot \mid \text{do}(\cdot))$ denotes the intervention distribution (Pearl, 2011). We cannot, however, estimate $P(Z_j \mid \text{do}(Z_{-j}))$ in practice. The reason is that we only have access to an observational dataset of the representation $\{z_i\}_{i=1}^n = \{f(x_i)\}_{i=1}^n$. This dataset is generally insufficient for identifying $P(Z_j \mid \text{do}(Z_{-j}))$ due to the unobserved common cause $C$ (Imbens & Rubin, 2015; Pearl, 2011). This difficulty of estimating $P(Z_j \mid \text{do}(Z_{-j}))$ is a core challenge in assessing disentanglement.

\(^8\)This causal $\text{scm}$ allows $Z_j$ to depend on only a subset of the $K$ confounder dimensions because the constraint imposed by a $\text{scm}$ resides in the absence of causal connections.
To tackle this challenge, we consider the possibility of finding observable implications of disentanglement. These observable implications will be necessary conditions associated with Equation (40). Though possibly insufficient, they can still help us reject representations that are not disentangled. Specifically, when a representation violates these observable implications, it cannot be disentangled.

### 3.2.1 Observable implications of disentanglement: The independent support condition

We turn to the specific observable implications that underpin our approach to disentanglement. We will prove that if a representation is disentangled, then its different dimensions must have independent support under a standard positivity condition.

**Theorem 9** (Disentanglement ⇒ Independent support). Assume the unobserved common cause $C$ satisfies a positivity condition: for all $j$, we have $P(Z_j \mid C) > 0$ iff $P(Z_j) > 0$. Then the support of the interventional distribution coincides with that of the observational distribution:

$$
\text{supp}(Z_j \mid \text{do}(Z_{j'} = z_{j'})) = \text{supp}(Z_j \mid Z_{j'} = z_{j'}),
$$

(41)

where $j, j' \in \{1, \ldots, d\}, j \neq j'$, and the density at $z_{j'}$ is nonzero, $p(z_{j'}) > 0$. As a consequence, different dimensions of a disentangled representation $Z = (Z_1, \ldots, Z_d)$ must have independent support:

$$
\text{supp}(Z_1, \ldots, Z_d) = \text{supp}(Z_1) \times \cdots \times \text{supp}(Z_d),
$$

(42)

$$
\text{supp}(Z_j \mid Z_S) = \text{supp}(Z_j) \text{ for all } S \subseteq \{1, \ldots, d\} \setminus j.
$$

**Proof.** We focus on proving Equation (41). (The remainder of the proof is in Appendix I.)

First notice that the positivity condition on $C$ guarantees that $\text{supp}(C) = \text{supp}(C \mid Z_j = z_j)$ for all $p(z_j) > 0$:

$$
\mathbb{I}\{P(C \mid Z_j) > 0\} = \mathbb{I}\left\{ \frac{P(C)P(Z_j \mid C)}{P(Z_j)} > 0 \right\} = \mathbb{I}\{P(C) > 0\} \mathbb{I}\{P(Z_j \mid C) > 0\}
$$

$$
= \mathbb{I}\{P(C) > 0\} \mathbb{I}\{P(Z_j) > 0\}
$$

$$
= \mathbb{I}\{P(C) > 0\},
$$

(43)
where the third equality is due to the positivity condition. Therefore, Equation (41) holds because

\[
\mathbb{I}\{P(Z_j \mid \text{do}(Z_{j'})) > 0\} = \mathbb{I}\left\{ \int P(Z_j \mid Z_{j'}, C)P(C)\,dC > 0 \right\} = \int \mathbb{I}\{P(Z_j \mid Z_{j'}, C) > 0\} \mathbb{I}\{P(C) > 0\} \,dC = \int \mathbb{I}\{P(Z_j \mid Z_{j'}, C, Z_{j'}) > 0\} \mathbb{I}\{P(C \mid Z_{j'}) > 0\} \,dC = \mathbb{I}\{P(Z_j \mid Z_{j'}) > 0\},
\]

where the third equality is due to Equation (43).

The intuition behind Theorem 9 is that two variables can have their support depend on each other in only two ways: (1) the causal connection between them, and (2) their common cause induces a dependence among their supports. Therefore, two causally disconnected variables must have independent support under the positivity condition, which guarantees the common cause does not causally affect their support. Positivity is also known as the overlap condition and it is a standard assumption in causal inference with observational data (Imbens & Rubin, 2015; Pearl, 1995). It implies that the unobserved common cause $C$ can induce correlations among $Z_j$’s, though it can not affect their support. Any combination of $Z_j$’s values must be possible.

Theorem 9 shows that the absence of causal arrows between two variables implies that their support must be independent, even in the presence of unobserved confounders. In other words, the relationship between the support of the variables is robust to unobserved confounding. It implies that any disentangled $Z$ must satisfy the independent support condition despite of the unobserved confounder $C$.

Consider compactly supported representations: each dimension of the representations must be supported on a closed and bounded interval. Then “independent support” implies a hyperrectangular joint support. As an example, Figures 8a and 8b illustrate the scatterplots of two disentangled discrete features; they occupy rectangular regions, regardless of their correlations. Figure 8c illustrate the scatter plots of two entangled features, which occupy curvilinear, non-rectangular regions.

Theorem 9 is most suitable for detecting entangled representations whose different dimensions causally affect each other’s support. For example, in a dataset of animals, the feature “the animal being a rabbit” can affect the support of the feature “the background being the sea” because rabbits can not swim. These two features are entangled; they also violate the independent support condition. In the next sections, we will leverage these observations stemming from Theorem 9 to develop a disentanglement metric.
3.2.2 Assessing disentanglement with the IOSS

To assess disentanglement, we build on Theorem 9 to develop the independence-of-support score (iOSS). It quantifies the extent to which a representation violates the independent support condition.

**Definition 10** (Independence-of-support score (iOSS)). Suppose a representation $Z$ has bounded support and $\sup Z_j - \inf Z_j > 0$, $j = 1, \ldots, d$. Then the iOSS of $Z$ is the Hausdorff distance between the joint support of $(Z_1, \ldots, Z_d)$ and the product of each individual’s support:

$$\text{iOSS}(Z_1, \ldots, Z_d)$$

$$\triangleq d_H(\text{supp}(Z_1, \ldots, Z_d), \text{supp}(Z_1) \times \cdots \times \text{supp}(Z_d))$$

$$= d(\text{supp}(Z_1) \times \cdots \times \text{supp}(Z_d), \text{supp}(Z_1, \ldots, Z_d)),$$

where $\tilde{Z}_j = (Z_j - \inf Z_j)/(\sup Z_j - \inf Z_j)$ is the standardized $Z_j$, and $d_H(\cdot, \cdot)$ is the Hausdorff distance. The second equality is due to $\text{supp}(Z_1, \ldots, Z_d) \subseteq \text{supp}(Z_1) \times \cdots \times \text{supp}(Z_d)$.

iOSS calculates the distance between the current support of the representation $Z$ and the (fictitious) support of $Z$ if it were disentangled. The larger the iOSS is, the more entangled the representation is. When $Z$ is disentangled, $\text{iOSS}(Z) = 0$.

iOSS focuses on representations with bounded support, which allows us to standardize each $Z_j$ to have $\inf \tilde{Z}_j = 0$ and $\sup \tilde{Z}_j = 1$. This standardization step makes iOSS scale-invariant; it also makes the different $Z_j$ equivalent in their importance in iOSS.

To compute iOSS in practice, we can directly compute the support of discrete-valued representations and hence their iOSS. For continuous-valued representations, we compute the sample iOSS. Given $n$ samples, $\{Z_i\}_{i=1}^n = \{(Z_{i1}, \ldots, Z_{id})\}_{i=1}^n$, we rescale the representation to the unit hypercube,

$$\tilde{Z}_{ij} = \frac{Z_{ij} - \min_{i=1,\ldots,n} Z_{ij}}{\max_{i=1,\ldots,n} Z_{ij} - \min_{i=1,\ldots,n} Z_{ij}},$$

and compute the sample iOSS as follows:

$$\widehat{\text{iOSS}}(\{Z_i\}_{i=1}^n) = \max_{k=1,\ldots,K} \min_{i=1,\ldots,n} (U_k - Z_{ij})^2,$$

where $U_k \overset{iid}{\sim} \text{Unif}[0, 1]^d$, $k = 1, \ldots, K$, are $K$ uniform draws from $\prod_j \text{supp}(Z_{ij})$. To improve the robustness of iOSS in practice, one may also replace the $\max, \min$ with the $(100 - \alpha)$th and $\alpha$th quantile, where $\alpha$ is a small positive number.

---

9The Hausdorff distance between sets $X$ and $Y$ is

$$d_H(X, Y) = \max\{d(X, Y), d(Y, X)\},$$

where

$$d(x, Y) = \inf\{d(x, y) \mid y \in Y\},$$

$$d(X, Y) = \sup\{d(x, Y) \mid x \in X\}.$$
The sample loss is comparable for representations with a fixed sample size $n$, dimension $d$, and uniform draws $K$. As $n \to \infty$ and $K/n \to \infty$, the sample loss approaches loss almost surely, $\hat{\text{loss}}\left(\{Z_i\}_{i=1}^n\right) \to \text{loss}(Z)$.

To see loss in action, we compute the sample loss for the three representations in Figure 8 with $K = 10^d \cdot n$. The sample loss of the entangled features in Figure 8c is substantially higher than that of the disentangled features in Figures 8a and 8b. In Section 3.4, we will show that loss better distinguishes entangled and disentangled representations than existing disentanglement metrics.

### 3.3 Learning Disentangled Representations with loss

To learn disentangled representations, we propose to use the sample loss as a regularizer in unsupervised representation learning.

Before presenting the algorithm, we first establish the identifiability of representations with independent support. This result will license the use of loss as a regularizer for unsupervised disentanglement.

**Theorem 11** (Identifiability of representations with independent support). For any two $d$-dimensional representations, $Z = f(X) = (Z_1, \ldots, Z_d)$ and $Z' = f'(X) = (Z'_1, \ldots, Z'_d)$, such that

1. The representations $Z, Z'$ both satisfy the independent support condition and have compact support in $\mathbb{R}^d$, i.e.

   $$\text{supp}(Z_1, \ldots, Z_d) = [a_1, b_1] \times \cdots \times [a_d, b_d], \quad (44)$$
   $$\text{supp}(Z'_1, \ldots, Z'_d) = [a'_1, b'_1] \times \cdots \times [a'_d, b'_d], \quad (45)$$

   where $a_j \leq b_j, a'_j \leq b'_j, j = 1, \ldots, d$,

2. The functions $f, f'$ are differentiable, and $f = h \circ f'$ with $h : \mathbb{R}^d \to \mathbb{R}^d$ for some bijective differentiable $h = (h_1, \ldots, h_d)$. Further, the dependency graph of $h_i(Z_1, \ldots, Z_d)$ on $Z_j$ (i.e. whether $h_i(Z_1, \ldots, Z_d)$ nontrivially depends on $Z_j$) remains the same across all values of $Z_{-j} = (Z_1, \ldots, Z_{j-1}, Z_{j+1}, \ldots, Z_d)$ within the support, i.e.

   $$\{\partial h_i(Z_1, \ldots, Z_d) \over \partial Z_j = 0\} \text{ is a constant function in } Z_{-j} \quad \forall i, j \in \{1, \ldots, d\}. \quad (46)$$

3. All components of $h$ being invertible, i.e. $h_j(Z'_1, \ldots, Z'_d)$ is an invertible function with respect to each of $Z'_k, k \in \{1, \ldots, d\}$, for all values $Z'_{-k}$ within the support and for all $j$.

4. For each $i \in \{1, \ldots, d\}$, there exist an open set $\overline{Z}_{-i} \subset \text{supp}(Z'_{-i})$, such that

   $$\cap_{j \neq i} \{Z'_i : h_j(Z'_1, \ldots, Z'_{-i} = z'_{-i}) \in [a_j, b_j]\} \cap \{Z'_i : h_i(Z'_1, \ldots, Z'_{-i} = z'_{-i}) \in [a_i, b_i]\}, \forall z_{-i} \in \overline{Z}_{-i}, \quad (47)$$

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10We are grateful to Shubhangi Ghosh, Luigi Gresele, Julius von Kugelgen, Michel Besserve, and Bernhard Schölkopf for pointing out a counterexample to a previous version of the theorem.
then we have

$$Z_1, \ldots, Z_d = \text{perm}(q_1(Z'_1), \ldots, q_d(Z'_d)),$$

where the $q_j$ are continuous bijective function with a compact domain in $\mathbb{R}$. (The proof is in Appendix J.)

To understand the intuition behind Theorem 11, we consider a toy example of a two-dimensional compactly supported representation $(Z_1, Z_2)$ with independent support: $Z_1 \in [1, 2]$, $Z_2 \in [0, 2]$. Next consider an entanglement of this representation $(Z'_1, Z'_2)$, which is a bijective transformation of $(Z_1, Z_2)$:

$$Z'_1 = Z_1 + Z_2, \quad Z'_2 = Z_1 - Z_2.$$

We will show that $(Z'_1, Z'_2)$ does not have independent support, then the support of $Z_1 - Z_2$ depends on the value of $Z_1 + Z_2$. To see why, consider the case when $Z_1 + Z_2 = 4$, then we must have $Z_1 = Z_2 = 2$ due to the support constraints on $Z_1, Z_2$. Hence $Z_1 - Z_2 = 0$, thus the support of $Z_1 - Z_2$ is $\{0\}$. Following a similar argument, the support of $Z_1 - Z_2$ is $\{1\}$ when $Z_1 + Z_2 = 1$. Therefore, the support of $Z_1 - Z_2$ depends on values of $Z_1 + Z_2$, and hence they have dependent support.

The proof of Theorem 11 relies on a similar argument as in the toy example. It builds on the following observation. Suppose a variable $W \in \mathbb{R}$ has support $[U_W, L_W]$. Then the support of $g(W) \in \mathbb{R}$ is $[\min\{g(U_W), g(L_W)\}, \max\{g(U_W), g(L_W)\}]$ if $g$ is continuous and bijective (subject to other assumptions of Theorem 11). In other words, the end points of $g(W)$’s support are functions of end points of $W$’s support. Therefore, any function $Z_{W'}$ that non-trivially depends on a second variable $W'$ will violate the independent support condition; the support of $Z_{W'}(W)$ will depend on $W'$. In this case, the mappings between two representations can only involve coordinate-wise mappings (e.g., permutation and coordinate-wise bijective mappings); the mapping cannot involve multiple dimensions of the representation.

This argument requires several assumptions in addition to independent support. Assumption 2 posit that the dependency graph of $h$ does not change across values. To study the dependency structure of $h$, it is sufficient to study the boundary of the support of $Z, Z'$. Loosely, this assumption guarantees that “$h_i$ nontrivially depends on two or more of its arguments” implies that “$h_i$ nontrivially depends on two or more of its arguments at the boundary of the support.” Assumption 3 assumes the invertibility of the $h$ functions, which facilitates the discussion on the dependency structure of $h$, namely how $Z'_j$ depends on $Z_{j'}$ for $j, j' \in \{1, \ldots, d\}$. Assumption 4 ensures that the dependency structure of $h(\cdot)$ has observable implications on the support boundary of $Z, Z'$; no support boundary is vacuous in the sense that it does not depend on any components of the $h(\cdot)$ function.

Theorem 11 establishes the identifiability of representations with independent support. It focuses on representations that generate the same $\sigma$-algebra; two representations $Z, Z'$ satisfy $\sigma(Z) = \sigma(Z')$ if there exists a bijective function $L$ such that $Z = L(Z')$. Representations with the same $\sigma$-algebra are “information-equivalent”; they capture the same information about the raw data $X$. Among these information-equivalent representations, there is a unique representation (up to coordinate-wise bijective transformations) that has independent support. Together with “disentanglement $\Rightarrow$ independent support” (Theorem 9), this unique representation must also be disentangled (if it exists).
We now illustrate disentangled representation learning with 1oss. Begin with an unsupervised representation learning objective \( L(\{x_i\}_{i=1}^n; \{z_i\}_{i=1}^n) \), and use the sample 1oss as a regularizer:

\[
L_{\text{1oss}}(\{x_i\}_{i=1}^n; \{z_i\}_{i=1}^n) = -L(\{x_i\}_{i=1}^n; \{z_i\}_{i=1}^n) + \lambda \cdot \text{1oss}(\{z_i\}_{i=1}^n),
\]

(48)

As a concrete example, the representation learning objective can be the VAE objective, a popular generative modeling approach to unsupervised representation learning (Kingma & Welling, 2014):

\[
L(\{x_i\}_{i=1}^n; \{z_i\}_{i=1}^n) = -\mathbb{E}_{q_\theta(z_i | x_i)} [\log p_\theta(z_i, x_i) - \log q_\theta(z_i | x_i)]
\]

where we adopt the (approximate) posterior mean of the latent variables as the representation \( z_i = \mathbb{E}_{q_\theta(z_i | x_i)} [z_i | x_i] \), and where \( \lambda > 0 \) is a regularization parameter.

To perform stochastic optimization on the \( L_{\text{1oss}} \) objective, we subsample batches \( \{z_i\}_{i \in B} \) of the data and update the parameters with the gradients of \( L_{\text{1oss}}(\{z_i\}_{i \in B}) \). Algorithm 3 summarizes this algorithm for learning disentangled representations with 1oss. In Section 3.4, we will show that it can produce representations that are more disentangled.

**Algorithm 3: Disentangled representation learning with 1oss**

**input**: Training data \( \{x_i\}_{i=1}^n \); Test data \( \{\tilde{x}_i\}_{i=1}^n \)

**output**: Representations of the test data \( \{\tilde{z}_i\}_{i=1}^n \)

Minimize Equation (48) to obtain \( q_\theta \);

Compute representations for the test data: \( \tilde{z}_i = \mathbb{E}_{q_\theta(z_i | x_i)} [z_i | x_i] \).

### 3.4 Empirical Studies of IOSS

We study 1oss in unsupervised image datasets. We find that 1oss is more effective at distinguishing between disentangled and entangled representations than other unsupervised disentanglement metrics. Unsupervised representation learning with the 1oss penalty results in representations with better disentanglement.

#### 3.4.1 Can 1oss distinguish entangled and disentangled representations?

We first study whether 1oss can distinguish entangled and disentangled representations.

**Generating entangled/disentangled representations.** Focusing on datasets with correlated ground truth features, we subsample each dataset so that the correlation among the ground-truth features is \( > 0.8 \). Given the subsampled dataset, we take the ground-truth features as disentangled representations. We then generate entangled representations by applying nonlinear transformations to the ground-truth features. Given ground-truth features \( \tilde{Z}_1, \ldots, \tilde{Z}_d \), we construct entangled representations by setting \( Z_i = f(\tilde{Z}_1, \ldots, \tilde{Z}_d; \theta_i) + Z_i \) with \( \theta_i \sim \text{Unif}([-2.5, 2.5]) \), where \( f(\cdot; \theta) \) is a third-order polynomial with coefficients \( \theta \).

**Evaluation metrics.** For each pair of generated entangled and disentangled representations, we calculate different disentanglement scores for each. We report the percentage of entangled and disentangled pairs that are correctly labeled by the metric. For example, if 1oss returns a smaller
IOSS (this paper) & 0.998(0.045) & 0.968(0.176) & 0.980(0.140) & 0.892(0.311) \\
Total Correlation & 0.858(0.349) & 0.070(0.255) & 0.162(0.369) & 0.090(0.286) \\
Wasserstein Dependency & 0.956(0.205) & 0.478(0.500) & 0.310(0.463) & 0.964(0.186) \\
Intervention Robustness (oracle) & 1.000(0.000) & 1.000(0.000) & 1.000(0.000) & 1.000(0.000) \\

| Table 3: IOSS outperforms existing disentanglement metrics in distinguishing entangled and disentangled features. The table presents the proportion of disentangled/entangled pairs that the metric correctly distinguish. Intervention Robustness Score is an oracle metric as it makes use of ground-truth features. (Higher is better.)

Figure 10: IOSS can better distinguish entangled and disentangled representations than existing unsupervised disentanglement metrics on the mpi3d dataset.

value for the disentangled representation than the entangled one in the pair, then it is deemed correct labeling. Similar procedures apply to other unsupervised disentangled metrics. A metric can effectively distinguish entanglement and disentanglement if this percentage is high.

**Competing methods.** We compare IOSS with existing unsupervised disentanglement metrics that do not rely on ground truth features: total correlation (Chen et al., 2018) and Wasserstein dependency (Xiao & Wang, 2019). We also compared with an oracle supervised disentanglement metric: the intervention robustness score (Suter et al., 2019); it targets the same causal disentanglement definition as IOSS but relies on ground truth features.

**Results.** Table 3 presents the results. The IOSS outperforms baseline unsupervised disentanglement metrics in distinguishing disentanglement from entanglement. Figures 10 and 15 to 17 also show that IOSS can better separate disentangled and entangled representations than existing unsupervised disentanglement metrics.

3.4.2 Does the IOSS regularizer encourage disentangled representations?

We next apply the IOSS penalty to learn disentangled representations via VAEs across all four datasets: mpi3d, smallnorb, dsprites, and cars3d. We work with subsampled datasets such that the ground truth features are highly correlated (with correlation ≈ 0.8) following a similar process as in Section 3.4.1. We then fit VAEs with an increasingly strong regularization with the IOSS
penalty. We evaluate the disentanglement of the learned representation using the supervised disentanglement score of intervention robustness score (Suter et al., 2019); a higher score implies better disentanglement.

**Competing methods.** We compare vae+iooss with other unsupervised disentanglement algorithms including classical vae (Kingma & Welling, 2014), FactorVAE (Kim & Mnih, 2018), betaVAE (Higgins et al., 2017), and betaTCVAE (Chen et al., 2018).

**Disentanglement of vae+iooss-learned representations.** Figure 11a shows that vae+iooss produce more causally disentangled representations than existing unsupervised disentanglement algorithms when ground truth features are highly correlated. These results corroborate Figures 8b and 8c. When features are highly correlated, they may still be disentangled. Hence enforcing statistical independence like beta-TCVAE and FactorVAE in this setting does not perform disentanglement. This setting violates the core assumption of beta-TCVAE and FactorVAE that the disentangled factors are independent; i.e., their joint distribution is a product of their marginals. Thus beta-TCVAE and FactorVAE do not perform as well in disentanglement with correlated features. In contrast, VAE with iooss regularization can accommodate correlated factors, as the causal graph Figure 2 allows an unobserved common cause between the factors.

Moreover, Figure 11b shows that, across datasets, the vae+iooss-learned representation becomes increasingly disentangled as we regularize with the iooss penalty. It implies that the iooss penalty indeed encourages learning causally disentangled representations. Figure 18 also illustrates that regularizing for iooss indeed encourages independent support across different dimensions of the learned representations.

**Tradeoff between informativeness and disentanglement.** We finally evaluate the tradeoff between informativeness and disentanglement in representation learning. We increase the regularization strengths of iooss and evaluate the log-likelihood of the fitted vae. Figure 11c shows that the log-likelihood stays stable despite the increasing regularization for disentanglement. While increasing the regularization strengths of iooss encourages disentanglement, it does not compromise the informativeness of the learned representations in vae. This result suggests the orthogonality between the disentanglement desiderata and other informativeness-related desiderata.

4 Discussion

We have shown that desiderata for representation learning, often discussed informally but rarely defined formally, can be usefully formalized from the perspective of causal inference. By studying the observable implications of our causal definitions, we develop metrics that evaluate representational desiderata and algorithms that enforce these desiderata.

We have focused on two examples of representation learning desiderata: (1) efficiency and non-spuriousness in supervised representation learning, and (2) disentanglement in unsupervised representation learning. Our overall workflow can be summarized as follows: “desiderata” → “causal definitions” → “observable implications” → “metrics and algorithms.” We have shown that this workflow can lead to practical evaluation metrics and representation learning algorithms.
Figure 11: (a) VAE+IOSS outperforms existing unsupervised disentanglement algorithms in producing representations that are more causally disentangled. (b) Regularizing for IOSS encourages learning disentangled representations. The intervention robustness score of the representation increases as we increase the regularization strengths of IOSS. (c) There is no obvious tradeoff between informativeness and disentanglement. The model fit stays stable despite we increase the regularization strengths of IOSS.

Targeting the desiderata of efficiency and non-spuriousness in the supervised setting, we view the representation as a cause of the label and formalize these desiderata as the probability of necessity and sufficiency (PNS) of the cause. Studying the observable implications of PNS enables us to measure efficiency and non-spuriousness with observational data. It also allows us to formulate representation learning as a task of finding necessary and sufficient causes. We operationalize this task by developing the CAUSAL-REP algorithm. En route, we develop identification results for the PNS given high-dimensional (rank-degenerate) data.

To encourage unsupervised disentanglement, we again begin with a causal definition, and we obtain metrics for disentanglement that again rely on studying its observable implications. We capitalize on the observation that causally disentangled variables—which have no causal connections among each other—must have independent support under a standard positivity condition. Based on this insight, we develop the independence-of-support score (IOSS) for assessing causal disentanglement. We further establish the identifiability of representations with independent support, which enables representation learning with a IOSS penalty.

For future work, one promising direction is to identify other desiderata in representation learning (or machine learning tasks in general) that may be formalized using causal notions. One may follow the workflow to obtain evaluation metrics and learning algorithms for the desiderata.

Another direction is to extend the metrics and algorithms developed in this work. One may extend the CAUSAL-REP algorithm to general anti-causal learning, without requiring the assumptions (e.g., pinpointability) required in Lemma 6. Such an extension would enable the treatment of reverse causal inference via probabilities of causation. On the disentanglement side, IOSS can
also be extended to learn primitives for compositional generalization. Primitives share similar properties as disentangled features; they cannot causally affect each other. In this sense, ioss-type ideas can potentially offer new approaches to compositional generalization.

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Supplementary Materials

A Proof of Lemma 4

Proof. The proof generalizes the proof of Theorem 9.2.10 in Pearl (2011).

We first notice the following:
\[ P(Y(Z = z) = y) = P(Y(Z = z) = y, Y(Z \neq z) = y) + P(Y(Z = z) = y, Y(Z \neq z) \neq y) \]  
(49)

Next denote \( \epsilon \triangleq P(Y(Z = z) \neq y, Y(Z \neq z) = y) \geq 0 \). Then we have
\[ P(Y(Z \neq z) = y) = P(Y(Z = z) = y, Y(Z \neq z) = y) + P(Y(Z = z) \neq y, Y(Z \neq z) = y) \]  
(50)
\[ = P(Y(Z = z) = y, Y(Z \neq z) = y) + \epsilon. \]  
(51)

Substituting Equation (51) into Equation (49) implies that
\[ P(Y(Z = z) = y) = P(Y(Z \neq z) = y) + P(Y(Z = z) = y, Y(Z \neq z) \neq y) - \epsilon, \]  
(52)
which implies
\[ P(Y(Z = z) = y, Y(Z \neq z) \neq y) = P(Y(Z = z) = y) - P(Y(Z \neq z) = y) + \epsilon \]  
(53)
\[ = P(Y = y \mid do(Z = z)) - P(Y = y \mid do(Z \neq z)) + \epsilon \]  
(54)
\[ \geq P(Y = y \mid do(Z = z)) - P(Y = y \mid do(Z \neq z)). \]  
(55)

The inequality becomes equality when \( \epsilon = 0 \) under the monotonicity condition in Lemma 4.

B The definition of functional interventions recovers backdoor adjustment

The definition of functional interventions (Definition 5) recovers the standard backdoor adjustment as special cases when the function \( f(X) \) returns \( X \) or its subset (assuming the causal graph Figure 2).

For example, when \( f(X) = X \), then
\[ P(Y \mid do(f(X) = z)) = P(Y \mid do(X = z)) \]
\[ = \int P(Y \mid do(X), C)P(X \mid C, X = z)P(C) \, dX \, dC \]
\[ = \int P(Y \mid X) \cdot \delta_{X=z} \cdot P(C) \, dX \, dC \]
\[ = P(Y \mid X = z), \]

53
where the third inequality is due to no unobserved confounding between $X$ and $Y$ in Figure 2.

As another special case, suppose the function $f(X)$ returns the subset of $X = (X_1, \ldots, X_m)$ except $X_1$, i.e., $f(X) = (X_2, \ldots, X_m)$. Then Equation (8) recovers backdoor adjustment:

$$P(Y \mid \text{do}(f(X) = z)) = P(Y \mid \text{do}((X_2, \ldots, X_m) = z))$$

$$= \int P(Y \mid \text{do}(X), C)P(X \mid C, (X_2, \ldots, X_m) = z)P(C) \, dX \, dC$$

$$= \int P(Y \mid X, (X_2, \ldots, X_m) = z) \cdot P(X_1 \mid C)P(C) \, dX \, dC$$

$$= \int P(Y \mid X_1, (X_2, \ldots, X_m) = z) \cdot P(X_1 \mid (X_2, \ldots, X_m) = z, C)P(C) \, dX_1 \, dC$$

$$= \int P(Y, X_1 \mid (X_2, \ldots, X_m) = z, C)P(C) \, dX_1 \, dC$$

$$= \int P(Y \mid (X_2, \ldots, X_m) = z, C)P(C) \, dC,$$

where the third inequality is due to no unobserved confounding between $X$ and $Y$ in Figure 2, and the fifth inequality is due to the conditional independence of $X_1, \ldots, X_m$ given $C$ in Figure 2.

### C Proof of Lemma 6

**Proof.** We calculate the intervention distribution for functional interventions $f(X) = \tilde{f}((X_j)_{j \in S})$

$$P(Y \mid \text{do}(f(X)))$$

$$= \int P(Y \mid (X_j)_{j \in S}, C)P((X_j)_{j \in S} \mid C, f(X))P(C) \, d(X_j)_{j \in S} \, dC,$$  \hspace{1cm} (56)

$$= \int P(Y \mid (X_j)_{j \in S}, C, f(X))P((X_j)_{j \in S} \mid C, f(X))P(C) \, d(X_j)_{j \in S} \, dC,$$  \hspace{1cm} (57)

$$= \int P(Y \mid C, f(X))P(C) \, dC,$$  \hspace{1cm} (58)

$$= \int P(Y \mid f(X), h(X)) \cdot P(h(X)) \, dh(X).$$  \hspace{1cm} (59)

The first equation is due to the definition of functional interventions on $f(X)$ (Puli et al., 2020). It is a soft intervention on $X$ with a stochastic policy conditional on the parents of $X$. The stochastic policy is $P(X \mid f(X), PA(X))$ where $PA(X)$ denotes the parents of $X$.

The second equation is due to $P(Y \mid (X_j)_{j \in S}, C, f(X)) = P(Y \mid (X_j)_{j \in S}, C)$. The second equation is due to $P(Y \mid (X_j)_{j \in S}, C, f(X)) = P(Y \mid (X_j)_{j \in S}, C)$. The second equation is due to $P(Y \mid (X_j)_{j \in S}, C, f(X)) = P(Y \mid (X_j)_{j \in S}, C)$.
The third equation is due to the observability and positivity condition. Observability and positivity guarantee that $P((X_j)_{j \in S}, C) > 0$ for all $(X_j)_{j \in S}, C$. Thus we can calculate $P(Y|(X_j)_{j \in S}, C, f(X)) = P(Y|(X_j)_{j \in S}, C)$. Moreover, the two conditions imply that $P(f(X), C) > 0$ for all $f(X), C$, and hence we can calculate $P((X_j)_{j \in S} | C, f(X))$.

The fourth equation is due to the pinpointability condition.

\[\square\]

## D Pinpointing the unobserved common cause $C$

The CAUSAL-REP algorithm begins with a step of pinpointing the unobserved common cause $C$. In this step, we infer $C$ from the observational data $X$, when $C$ is low-dimensional and $X$ is high-dimensional. Operationally, as $C$ renders $X = (X_1, \ldots, X_m)$ conditionally independent, we infer $C$ by fitting a probabilistic factor model to $X$,

\[p(x, c; \phi) = p(x; c; \phi) = p(c) \prod_{j=1}^{m} p(x_{ij} | c; \phi). \tag{60}\]

Specifically, we consider VAE, a flexible probabilistic factor model (Kingma & Welling, 2014),

\[C_i \sim p(c_i), \tag{61}\]
\[X_i | C_i \sim p(x_i | c_i; \theta) = \text{EF}(x_i | f_{\theta}(c_i); \lambda_{\theta}), \tag{62}\]

where EF is an exponential family distribution, $\theta = (f_{\theta}, \lambda_{\theta})$ are the parameters, and $f_{\theta} : C \rightarrow X$ is a flexible neural network.

Next we infer $p(c_i | x_i)$ using variational approximation $p(c_i | x_i) \approx q_{\phi^*}(c_i | x_i)$ where $q_{\phi^*}(c | x)$ maximizes the evidence lower bound (ELBO) objective of VAE (Blei et al., 2017),

\[q_{\phi^*} = \arg \max_{q_{\phi}} \sum_{i=1}^{n} \left[\log p(x_i, c_i) - \log q_{\phi}(c_i | x_i)\right],\]

and $q_{\phi}(\cdot | x)$ parametrizes a flexible family of distributions with parameters $\phi$. For example, we can have $q_{\phi}(\cdot | x) = \mathcal{N}(Z_1, \phi(x), Z_2, \phi(x) \cdot I)$ or even more flexible non-Gaussian distributions via normalizing flows.

Finally, we set

\[C_i = h(X_i) = \mathbb{E}_{q_{\phi^*}(c_i | x_i)} [C_i | X_i], i = 1, \ldots, n. \tag{63}\]

Though we use variational approximation for $p(c_i | x_i)$, Equation (63) can often give a good approximation of $C_i$ when $\dim(X) \gg \dim(C)$, or more precisely when the pinpointability condition holds (Chen et al., 2020b; Wang & Blei, 2019b).
E Calculating PNS lower bounds with linear models

We calculate the lower bound of the conditional efficiency and non-spuriousness
\(\text{PNS}_n(f_j(X), Y | f_{-j}(X))\) with the linear model (Equation (19)):

\[
\text{PNS}_n(f_j(X), Y | f_{-j}(X)) = \prod_{i=1}^n \left[ \int \left( P(Y = y_i | f_j(X) = f_j(x_i), f_{-j}(X) = f_{-j}(x_i), C) 
\right. \\
- P(Y = y_i | f_j(X) \neq f_j(x_i), f_{-j}(X) = f_{-j}(x_i), C) \right] \cdot P(C) \, dC \\
= \prod_{i=1}^n \int \left[ \mathcal{N}(y_i; \beta_0 + \beta_j f_j(x_i) + \sum_{j' \neq j} \beta_{j'} f_{j'}(x_i) + \gamma^\top C, \sigma^2) 
\right. \\
- \mathcal{N}(y_i; \beta_0 + \beta_j \mathbb{E}[f_j(x_i)] + \sum_{j' \neq j} \beta_{j'} f_{j'}(x_i) + \gamma^\top C, \sigma^2) \right] \cdot P(C) \, dC \\
= \prod_{i=1}^n \int \left[ \exp \left( -\frac{(\gamma^\top (c_i - C) + \epsilon_i)^2}{2\sigma^2} \right) 
\right. \\
- \exp \left( -\frac{(\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) + \gamma^\top (c_i - C) + \epsilon_i)^2}{2\sigma^2} \right) \right] \cdot P(C) \, dC \\
\approx \prod_{i=1}^n \int \left[ \left( 1 - \frac{(\gamma^\top (c_i - C) + \epsilon_i)^2}{2\sigma^2} \right) 
\right. \\
- \left. \left( 1 - \frac{(\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) + \gamma^\top (c_i - C) + \epsilon_i)^2}{2\sigma^2} \right) \right] \cdot P(C) \, dC \\
= \prod_{i=1}^n \int \left( \frac{(\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2 + 2 \beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) \cdot (\gamma^\top (c_i - C) + \epsilon_i)}{2\sigma^2} \right) \\
\cdot P(C) \, dC \times (2\pi\sigma^2)^{-\frac{d}{2}} \\
= \prod_{i=1}^n \left( \frac{(\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2 + 2 \beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) \cdot (\gamma^\top (c_i - \mathbb{E}[C]) + \epsilon_i)}{2\sigma^2} \right) \\
\times (2\pi\sigma^2)^{-\frac{d}{2}} \\
= \prod_{i=1}^n \exp \left( \frac{(\beta_j f_j(x_i) - \mathbb{E}[f_j(x_i)])^2 + 2 \beta_j (f_j(x_i) - \mathbb{E}[f_j(x_i)]) (\gamma^\top (c_i - \mathbb{E}[C]) + \epsilon_i)}{2\sigma^2} - 1 \right) \\
\times (2\pi\sigma^2)^{-\frac{d}{2}} \\
= \exp \left( \sum_{i=1}^n \frac{(\beta_j (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2 + 2 \beta_j (f_j(x_i) - \mathbb{E}[f_j(x_i)]) (\gamma^\top (c_i - \mathbb{E}[C]))}{2\sigma^2} - n \right) \\
\times (2\pi\sigma^2)^{-\frac{d}{2}},
\]
where $\mathcal{N}(\cdot)$ denotes the Gaussian density, $\epsilon_i = y_i - (\beta_0 + \beta^T f(x_i) + \gamma^T c_i)$ is the residual of the regression in Equation (19). Equations (65) and (68) make use of Taylor approximation $\exp(x) \approx 1 + x$. Equation (69) makes use of $\sum_{i=1}^{n} \epsilon_i \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) \approx \mathbb{E}[\epsilon \cdot (f_j(X) - \mathbb{E}[f_j(X)])] = 0$ in the regression model.

Finally, in CAUSAL-REP Algorithms 2 and 4, we often impose an $R^2$ penalty to encourage the positivity of $f_j(X)$ given $C$. In these cases, we often have $\sum_{i=1}^{n} \beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) \cdot (\gamma^T (c_i - \mathbb{E}[C])) \approx \beta_j \text{Cov}(f_j(X), \gamma^T C) \approx 0$. Thus, we can further approximate the $\text{PNS}$ by

$$\text{PNS}_n(f_j(X), Y | f_{-j}(X)) \approx \exp \left( \frac{\sum_{i=1}^{n} (\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2}{2\sigma^2} - n \right) \times (2\pi\sigma^2)^{-\frac{n}{2}},$$

and thus,

$$\log \text{PNS}_n(f_j(X), Y | f_{-j}(X)) \approx \left( \frac{\sum_{i=1}^{n} (\beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2}{2\sigma^2} \right) + \text{constant},$$

Similarly, we can obtain the lower bound of the (unconditional) efficiency and non-spuriousness, similar to Equation (69),

$$\log \text{PNS}_n(f(X), Y) \approx \left( \frac{1}{2\sigma^2} \sum_{i=1}^{n} \left[ \sum_{j=1}^{d} \beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]))^2 \right] + 2 \cdot \sum_{j=1}^{d} \beta_j \cdot (f_j(x_i) - \mathbb{E}[f_j(x_i)]) \cdot \gamma^T (c_i - \mathbb{E}[C]) \right) + \text{constant}.$$

## F Details of the unsupervised CAUSAL-REP algorithm

We summarize the unsupervised CAUSAL-REP in Algorithm 4.

## G Details of the empirical studies for CAUSAL-REP and additional empirical results

### G.1 Details of Section 2.4.1

Figures 12 and 13 present additional results for Section 2.4.1. As $Z_1$ and $Z_2$ become increasingly correlated, the (lower bounds of) unconditional roc of $Z_2$ for $Y_1$ also increase. It is also consistent with the intuition: $Z_2$ is an increasingly better surrogate of $Z_1$ for $Y_1$ give higher correlations between $Z_1$ and $Z_2$. 
Algorithm 4: CAUSAL-REP (Unsupervised)

**input**: The observational training data (without labels) \( \{x_i\}_{i=1}^n \); the probabilistic factor model that generates the training data \( P(X, C) \)

**output**: CAUSAL-REP representation function \( \hat{f}(\cdot) \)

Augment the unsupervised training dataset into a supervised one \( \{\{x_u^i, y_u^i\}_{u=1}^U\}_{i=1}^n \) following Equation (35);

Fit a probabilistic factor model (Equations (17) and (18)) and infer \( \{p(c_u^i | x_u^i)\}_{u=1}^U \}_{i=1}^n ;

if Pinpointability holds, i.e. \( p(c_u^i | x_u^i) \) is close to a point mass for all \( i, u \) then

- Pinpoint the unobserved common cause \( C: c_u^i = h(x_u^i) \triangleq \mathbb{E}[c_u^i | x_u^i] \) for all \( u = 1, \ldots, U \);

end

Maximize Equation (36) to obtain the CAUSAL-REP representation \( \hat{f} \);

end

G.2 Details of the colored MNIST study

We study CAUSAL-REP on the colored MNIST dataset (Arjovsky et al., 2019). The dataset builds on the original MNIST data but color the image in a way that is highly correlated with the digits label. We focus on the digits ‘3’ and ‘8’ and colors ‘red’ and ‘green.’

To create a training set, we color the ‘3’ images in red with probability \( p \) and in green with probability \( 1 - p \). Next, we color the ‘8’ images in red with probability \( 1 - p \) and in green with probability \( p \). When \( p \in [0, 1] \) is large, then the color of the image is highly correlated with the digit label in such a training set. We further add noise to the ground truth digit label by randomly flipping the labels with a probability of 0.25. The best possible predictive accuracy is thus 0.75 (the yellow dashed line in Figure 6).

The color of the images is a spurious feature in the training set; it has a high correlation with the digit label but does not causally determine the label. In contrast, the features of the digits themselves are non-spurious features; they are highly correlated with the digit label and can causally determine the label.

To create a test set, we color the images such that the color and images are correlated oppositely. We color the ‘3’ images in red with probability \( 1 - p \) and the ‘8’ images in red with probability \( p \). As the color-image relationship is very different, a representation learning algorithm will predict poorly in the test set if it only captures color as a feature in the training set.

G.3 Details of the reviews corpora study

Table 4 presents the most informative words of the (positive or negative) ratings, suggested by the CAUSAL-REP representation and the logistic regression coefficients. Across three reviews corpora, logistic regression returns the spurious words “as”, “also”, “am”, “an” as the top words. In contrast, CAUSAL-REP extracts words that are more relevant for the ratings.
G.4 Details of the colored and shifted MNIST study

The colored and shifted MNIST dataset is created similarly as in the colored MNIST. We consider four shifts: \((dx, dy) = (0, 1), (1, 0), (0, 1), (1, 1)\)—labeled 0, 1, 2, 3—and shift the images such that the shift label is highly correlated with the digit label in the training set.

We evaluate the non-spuriousness of the representations using a downstream prediction task with domain shift. Given a labeled training set where the digit features are no longer correlated with the spurious features, we learn a mapping from the representations to the digit label. A representation can only predict the digit label well if it captured the non-spurious digit features in unsupervised representation learning.

H Details of Figure 8

Here we include the full pairwise scatter plot (Figure 14) of the three sets of features in Figure 8.

To generate the factors for Figure 14a, we randomly sample 7,000 data points from the dsprites dataset.

To generate the factors for Figure 14b, we first sort the data points of the dsprites dataset in descending order by columns ['shape', 'scale', 'orientation', 'positionX', 'positionY']. Next we create the correlated factors dataset by taking the top 5000 data points and then randomly draw 700 data points from the rest.

To generate the factors for Figure 14c, we randomly subsample 7,000 data points and then generate the entangled factors as follows:

\[
\begin{align*}
\text{entangle1} &= 6 \cdot \text{shape}^1 + 8 \cdot \left(\frac{\text{scale}}{\text{sd(scale)}}\right)^3 + 1 \cdot \left(\frac{\text{orientation}}{\text{sd(orientation)}}\right)^3 + 0.2 \cdot \mathcal{N}(0, 1), \\
\text{entangle2} &= 12 \cdot \text{shape}^2 + 1 \cdot \left(\frac{\text{scale}}{\text{sd(scale)}}\right)^2 + 8 \cdot \left(\frac{\text{orientation}}{\text{sd(orientation)}}\right)^1 + 0.2 \cdot \mathcal{N}(0, 1), \\
\text{entangle3} &= 0 \cdot \text{shape}^3 + 4 \cdot \left(\frac{\text{scale}}{\text{sd(scale)}}\right)^1 + 4 \cdot \left(\frac{\text{orientation}}{\text{sd(orientation)}}\right)^2 + 0.2 \cdot \mathcal{N}(0, 1).
\end{align*}
\]

I Proof of Theorem 9

**Proof.** We first prove \(\text{supp}(Z_1, \ldots, Z_d) = \text{supp}(Z_1) \times \cdots \times \text{supp}(Z_d)\). Notice that the causal disentanglement of \(Z_1, \ldots, Z_d\) (Figure 9) implies that

\[
P(Z_1, \ldots, Z_d) = \int P(Z_1, \ldots, Z_d | C) P(C) \, dC = \int P(Z_1 | C) \cdots P(Z_d | C) P(C) \, dC. \tag{73}
\]

Therefore, we have

\[
P(Z_1, \ldots, Z_d) > 0 \iff P(Z_j | C \in C) > 0 \quad \forall j \in \{1, \ldots, d\} \text{ for some set } P(C \in C) > 0. \tag{74}
\]

Together with the positivity condition, which requires

\[
P(Z_j | C \in C) > 0 \text{ for any set } P(C \in C) > 0 \iff P(Z_j) > 0, \tag{75}
\]

59
we have
\[ P(Z_1, \ldots, Z_d) > 0 \iff P(Z_j) > 0 \quad \forall j \in \{1, \ldots, d\}. \quad (76) \]
As \( \text{supp}(G) = \mathbb{I}\{P(G) > 0\} \), we can rewrite it as
\[ \text{supp}(Z_1, \ldots, Z_d) = \text{supp}(Z_1) \times \ldots \times \text{supp}(Z_d), \quad (77) \]
i.e. \( Z_1, \ldots, Z_d \) satisfy the full support condition.

Next we show that \( \text{supp}(Z_i \mid Z_S) = \text{supp}(Z_i) \).

First notice that
\[ \mathbb{I}\{P(Z_i, Z_S) > 0\} = \mathbb{I}\{P(Z_i \mid Z_S)P(Z_S) > 0\} \]
\[ = \mathbb{I}\{P(Z_i \mid Z_S) > 0\} \times \mathbb{I}\{P(Z_S) > 0\} \] \quad (78)
Therefore the full support condition \( \mathbb{I}\{P(Z_i, Z_S) > 0\} = \mathbb{I}\{P(Z_i) > 0\} \times \mathbb{I}\{P(Z_S) > 0\} \) implies that
\[ \mathbb{I}\{P(Z_i) > 0\} = \mathbb{I}\{P(Z_i \mid Z_S) > 0\} \text{ for } Z_S \text{ s.t. } \mathbb{I}\{P(Z_S) > 0\}. \quad (80) \]
This implies \( \text{supp}(Z_i \mid Z_S) = \text{supp}(Z_i) \) for \( Z_S \in \text{supp}(Z_S) \).

Finally, we consider the support of do interventions.
\[ P(Z_S \mid \text{do}(Z_i = Z_i)) = \int P(Z_S \mid Z_i, C)P(C)\,dC, \quad (81) \]
\[ P(Z_S \mid Z_i = Z_i) = \int P(Z_S \mid Z_i, C)P(C \mid Z_i)\,dC. \quad (82) \]
Positivity guarantees that the support of \( P(C) \) must be the same as \( P(C \mid Z_i) \). (Because \( \text{supp}(C \mid Z_i) \subseteq \text{supp}(C) \) and positivity guarantees the other direction.) Therefore, the support of \( P(Z_S \mid \text{do}(Z_i = Z_i)) \) and \( P(Z_S \mid Z_i = Z_i) \) are the same.

Therefore, if \( \text{supp}(Z_S \mid Z_i) \neq \text{supp}(Z_S) \), then \( \text{supp}(P(Z_S \mid \text{do}(Z_i = Z_i))) \neq \text{supp}(P(Z_S)) \) and hence \( P(Z_S \mid \text{do}(Z_i = Z_i)) \neq P(Z_S) \).

\[ \square \]

### J  Proof of Theorem 11

**Proof.** We follow a proof-by-contradiction argument. Suppose the opposite: \( Z = \mathbf{h}(Z') \) where \( \mathbf{h} \) nontrivially depends on two or more of its arguments.

The first two assumptions imply that \( \mathbf{h} : \mathbb{R}^d \to \mathbb{R}^d \) is differentiable and invertible.

Moreover, Equation (46) ensures that the (nontrivial) dependency graph (whether \( h_i \) nontrivially depends on \( Z_j \) of \( h_i(Z_1, \ldots, Z_d), i \in \{1, \ldots, d\} \) on \( Z_1, \ldots, Z_d \) remains constant over all values of \( Z_1, \ldots, Z_d \) within the support. This assumption ensures that it is sufficient to study the
We next consider a different mapping between \( Z, Z' \) to conclude about the dependency structure of \( h(\cdot) \) and thus disentanglement. In other words, “\( h \) nontrivially depends on two or more of its arguments” implies that “\( h \) nontrivially depends on two or more of its arguments at the boundary of the support.”

Finally, we will try to argue that if (1) \( Z' \) satisfies the independent support condition (2) the function \( h_i \) (for some \( i \)) nontrivially depends on two or more of \( Z'_j, Z'_k, j, k \in \{1, \ldots, d\} \), then \( Z \) will violate the independent support condition. (The fourth assumption is to exclude the corner cases where some support constraints are vacuous.)

Begin with two examples that illustrate this argument.

**Warm-up example 1.** Consider two two-dimensional representations \( Z' = (Z'_1, Z'_2) \) and \( Z = (Z_1, Z_2) \). They satisfy \( Z = (Z_1, Z_2) = h(Z') \) where

\[
\begin{align*}
Z_1 &= h_1(Z'_1, Z'_2) = Z'_1 + Z'_2, \\
Z_2 &= h_2(Z'_1, Z'_2) = Z'_1 - Z'_2.
\end{align*}
\]

Moreover, assume the support of \( Z \) is \( \text{supp}(Z_1, Z_2) = [a_1, b_1] \times [a_2, b_2] \), which satisfies the independent support condition.

Next we show that \( h(Z'_1, Z'_2) \) cannot satisfy the independent support condition.

Recall that that \( \text{supp}(Z_1, Z_2) = [a_1, b_1] \times [a_2, b_2] \) implies that \( \text{supp}(Z'_1 + Z'_2) = [a_1, b_1] \) and \( \text{supp}(Z'_1 - Z'_2) = [a_2, b_2] \). It implies that \( \text{supp}(Z'_1 \mid Z'_2) = [a_1 - Z'_2, b_1 - Z'_2] \cap [a_2 + Z'_2, b_2 + Z'_2] \). All end points of this intersection involves \( Z'_2 \), so the support of \( \text{supp}(Z'_1 \mid Z'_2) \) depends on \( Z'_2 \) (for values of \( Z'_2 \) s.t. \( \text{supp}(Z'_1 \mid Z'_2) \) is non-empty); it violates the independent support condition.

More broadly, \( Z' \) does not satisfy the independent support condition because both \( h_1(Z'_1, Z'_2) \) and \( h_2(Z'_1, Z'_2) \) non-trivially depend on both \( Z'_1, Z'_2 \) across all values of \( Z'_1, Z'_2 \) within the support.

Visually, independent support condition ensures that the support is rectangular. If we apply a rotation-like transformation, then it can no longer be rectangular in the original coordinate system. More generally, if we apply any transformations that will “entangle” the coordinates of the original representation at its support boundary, then the independent support condition cannot hold for both the original representation and the post-transformation representation. That said, we still need Equation (46) to extend argument about the boundary to one about the whole function being entangling.

**Warm-up wexample 2.** We next consider a different mapping between \( Z \) and \( Z' \),

\[
\begin{align*}
Z_1 &= h_1(Z'_1, Z'_2) = Z'_1(Z'_1 - Z'_2), \\
Z_2 &= h_2(Z'_1, Z'_2) = Z'_1 - Z'_2.
\end{align*}
\]

Again assume independent support for \( Z \): \( \text{supp}(Z_1, Z_2) = [a_1, b_1] \times [a_2, b_2] \).

In this case, \( \text{supp}(Z'_1 \mid Z'_2) = \{Z'_1 : Z'_1(Z'_1 - Z'_2) \in [a_1, b_1]\} \cap [a_2 + Z'_2, b_2 + Z'_2] \). Again the boundary of the support depends on \( Z'_2 \).

The boundary of the support \( \{Z'_1 : h_1(Z'_1, Z'_2) \in [a_1, b_1]\} \) does not depend on \( Z'_2 \) only when \([a_1, b_2]\) is a vacuous bound for \( h_1 \), e.g. \( a_1 = 0, b_1 = 1 \), \( h_1(Z'_1, Z'_2) = \frac{\exp(-Z'_1 + Z'_2)}{1 + \exp(-Z'_1 + Z'_2)} \).
**General argument.** Suppose some $h_l$ nontrivially depend on two or more of its arguments, say $Z'_j, Z'_k, j, k \in \{1, \ldots, d\}$. Then we argue that the support of $Z'_j, Z'_k$ will not be independent.

Recall that $Z$ satisfies the independent support condition, $\text{supp}(Z_1, \ldots, Z_d) = [a_1, b_1] \times \cdots \times [a_d, b_d]$. We consider the support $\text{supp}(Z'_j \mid Z'_k ; Z'_{-j,-k}) = \cap_{l=1}^d \{Z'_j : h_l(Z'_1, \ldots, Z'_d) \in [a_l, b_l]\}$, fixing $Z'_{-j,-k}$ at some value within the support. We argue that (1) $\{Z'_j : h_l(Z'_1, \ldots, Z'_d) \in [a_l, b_l]\}$ implies the support of $Z'_j$ is dependent on $Z'_k$, and (2) $\text{supp}(Z'_j \mid Z'_k ; Z'_{-j,-k})$ also depends on $Z'_k$.

The first argument is via the invertibility of $h_l$ for all values of $Z'_{-j}$. Fix $Z'_{-j,-k}$ at some value within the support. Then $\text{supp}(Z'_j \mid Z'_k ; Z'_{-j,-k}) = [\min\{B^1_l, B^2_l\}, \max\{B^1_l, B^2_l\}]$, where $B^1_l = h^{-1}(a_l; Z'_k, Z'_{-j,-k}), B^2_l = h^{-1}(b_l; Z'_k, Z'_{-j,-k})$, where $h^{-1}(h_l; Z'_k, Z'_{-j,-k})$ nontrivially depends on $Z'_k$ because of Equation (46) and that $h_l$ nontrivially depends on $Z'_k$.

The second argument is via the assumption Equation (47), which ensures that the support constraint imposed by $h_l$ is not subsumed by other $h_j, j \neq l$, and its non-trivial dependence on $Z'_k$ will make the support $\text{supp}(Z'_j \mid Z'_k ; Z'_{-j,-k})$ also non-trivially depend on $Z'_k$ for some values of $Z'_{-j,-k}$ within the support.

\[\square\]

**K  Details of empirical studies of IOSS and additional empirical results**

**K.1  Details of Section 3.4.1**

Figures 10 and 15 to 17 also show that ioss can better separate disentangled and entangled representations than existing unsupervised disentanglement metrics.

**K.2  Details of Section 3.4.2**

Figure 18 illustrates that regularizing for ioss indeed encourages independent support across different dimensions of the learned representations.
Figure 12: Lower bounds of probabilities of causation are consistent with intuitive notions of feature necessity and sufficiency. As $Z_1$ and $Z_2$ become increasingly highly correlated, (a) the roc of $Z_1$ for $Y_1$ stays high, (b) the roc of $Z_2$ for $Y_1$ starts to increase when the correlation turns positive, and (c) the roc of $Z_1$&$Z_2$ increases.
Figure 13: Lower bounds of probabilities of causation are consistent with intuitive notions of feature necessity and sufficiency. As $Z_1$ and $Z_2$ become increasingly highly correlated, (a) the poc of $Z_1$ for $Y_2$ increases, (b) the poc of $Z_2$ for $Y_2$ increases, and (c) the poc of $Z_1 \& Z_2$ stays high.
| Amazon | CAUSAL-REP | Logistic Regression |
|--------|------------|---------------------|
| 1      | love_this_camera, recommend_this_camera, my_first_digital, great, best_camera, camera_if_you, this_camera_and, camera_have, excellent_camera, camera_bought_this; | love_my, the_tracfone, am, an, also, as, it_real, which_is, too, so_much, is_so_much, which_is_pretty, faster, must_for, when_use |
| 2      | this_camera, camera, camera_is, pictures, picture, the_camera, digital, camera_for, this_camera_is, digital_camera; |  |
| 3      | really_nice, hold_the, excellent_it, this_one_it, easy_it, is_superb, nice_if, returning, too_low, you_need_more; |  |
| 4      | with_this, aa, took, came, yet, pictures_of, camera_in, computer, pictures_in, for_those; |  |
| 5      | camera_was, expect, the_photos, by, camera_are, blurry, sony, have_an, had_some, wife; |  |

| Tripadvisor | CAUSAL-REP | Logistic Regression |
|-------------|------------|---------------------|
| 1           | front_door, typical, lady, room_is, sized, yogurt, of_italian, in_we, was_at, front_desk_to, was_easy; | am, as, an, also, we_have, consideration, and_nice, real, stay_here_again, good |
| 2           | lobby, man, directions, door, open, tried, seemed, with_the, by_the, almost; |  |
| 3           | man_at, the_price, is_in, above_and_beyond, we_checked_out, in_central_rome, coliseum, great_for, covered, place_to_sit; |  |
| 4           | the_front, front, front_desk, desk, the_front_desk, at_the_front, desk_staff, front_desk_staff, desk_was, front_of; |  |
| 5           | the_people_at, the_people, stayed_nights, people_at, was_very_helpful, would_not_recommend, lovely_and, great_location, staff_at, of_my, pricey; |  |

| Yelp       | CAUSAL-REP | Logistic Regression |
|------------|------------|---------------------|
| 1          | but_if, looking, what_you, you_go, but_if_you, that_you, to_do, lot, youll, try, own, do_not; | an, as, am, also, japanese_food, because_its, sure_to_get, ive, ive Been, at_night, up_for_it |
| 2          | even_if, your_place, this_restaurant, for_you, you_should, thank_you, or_if, thank, lamb, fabulous, is_awesome; |  |
| 3          | want_the, then_you, ahead, hollywood, dont_want, suggest, with, please, check_this_place, all_that; |  |
| 4          | are, you_like, if_you_like, here_if, are_looking, here, are_in, is_the_place, you_need, are_looking_for; |  |
| 5          | if_you, if, you, want, you_want, you_are, if_you_are, want_to, if_you_want, your, you_want_to, you_dont, dont; |  |

Table 4: Across the Amazon, Tripadvisor, and Yelp reviews corpora, CAUSAL-REP learns representation that does not rely on the injected words that are spuriously correlated with the sentiment. The table shows the top 12 words identified by the five-dimensional representation from CAUSAL-REP; the five dimensions are not ordered. The representation obtained from logistic regression relies heavily on the spurious words “am”, “an”, “also”, “as.”
Figure 14: Disentangled features have independent support even though they may be correlated. Moreover, IOSS can distinguish disentangled and entangled features. This figure illustrates how entangled and disentangled features differ using pairwise scatter plots. Figure 14a considers the ground truth features (shape, scale, orientation) of the dsprites dataset. These features are disentangled. They also have independent support, e.g., conditional on ‘scale’, the set of values that ‘orientation’ can take does not change with ‘scale’. Visually, these causally disentangled features have scatter plots that occupy rectangular (or hyperrectangular) region. Figure 14b considers the same features but in a subset of the dsprites dataset where the features are correlated. These features, though correlated, are still disentangled; they also have independent support. Figure 14c considers three entangled features, each of which is a nonlinear transformation of the three ground-truth features. These features are not disentangled. Their support are also not independent. Conditional on ‘entangle1’, the possible values ‘entangle2’ can take depends on the value of ‘entangle1.’

Figure 15: IOSS can better distinguish entangled and disentangled representations than existing unsupervised disentanglement metrics on the dsprites dataset.
Figure 16: IOSS can better distinguish entangled and disentangled representations than existing unsupervised disentanglement metrics on the smallnorb dataset.

Figure 17: IOSS is competitive in distinguishing entangled and disentangled representations compared with existing unsupervised disentanglement metrics on the cars3d dataset.
Figure 18: Increasing regularization strengths of IOSS encourages the learned representation to have independent (i.e. hyperrectangular) support. (a) The pairwise scatter plot of the learned representations without IOSS regularization; some dimensions of the representation do not have independent support. (b) The pairwise scatter plot of the learned representations with IOSS regularization; most dimensions of the representation have independent support.