Abstract
In representation learning, there has been recent interest in developing algorithms to disentangle the ground-truth generative factors behind a dataset, and metrics to quantify how fully this occurs. However, these algorithms and metrics often assume that both representations and ground-truth factors are flat, continuous, and factorized, whereas many real-world generative processes involve rich hierarchical structure, mixtures of discrete and continuous variables with dependence between them, and even varying intrinsic dimensionality. In this work, we develop benchmarks, algorithms, and metrics for learning such hierarchical representations.

1. Introduction
Autoencoders aim to learn structure in data by compressing it to a lower-dimensional representation with minimal loss of information. Although they have proven useful in many applications (LeCun et al., 2015), the individual dimensions of their representations are often inescrutable, even when the underlying data is generated by simple processes. Motivated by needs for interpretability (Alvarez-Melis & Jaakkola, 2018; Marx et al., 2019), fairness (Creager et al., 2019), and generalizability (Bengio et al., 2013), as well as a basic intuition that representations should model the data correctly, a subfield has emerged which applies representation learning algorithms to synthetic datasets and checks how well representation dimensions “disentangle” the known ground-truth factors behind the dataset.

Perhaps the most common disentanglement approach has been to learn continuous vector representations whose dimensions are statistically independent (and evaluate them using metrics that assume ground-truth factors are also independent), reasoning that factorization is a useful proxy (Ridgeway, 2016; Higgins et al., 2017; Chen et al., 2018; Kim & Mnih, 2018). However, this problem is not identifi-

2. Related Work
In this section, we review work related to our notion of “hierarchical disentangled representations.” However, there are many notions of hierarchy that can be introduced into representations (or into definitions of disentanglement), some of which have little in common except a shift in focus away from flatness or factorization.

Still, the problem of learning a flat, factorized representation has received significant attention over the years. Much of the...
initial work, e.g. from Schmidhuber (1992), Zemel (1994), and Comon (1994), was motivated by classic problems like source separation or biological and information-theoretic arguments about minimum description length (Barlow, 1961). More recently, Ridgeway (2016) argued that factorization was often a useful real-world proxy for disentanglement in the seminal sense of Bengio (2013), which motivated the development of a number of popular methods for training variational autoencoders (VAEs, Kingma & Welling (2013)) to reconstruct data from compressed flat vectors, but with minimal total correlation (TC) between their components (Higgins et al., 2017; Chen et al., 2018; Kim & Mnih, 2018; Dupont, 2018; Kim et al., 2019; Jeong & Song, 2019). We build on these approaches in our work, which we also tie back to some of their original motivating problems like minimum description length (see §8.1).

There are, however, a number of limitations to learning factorized representations. To begin with, the problem was actually shown by Locatello et al. (2018) to be non-identifiable, at least without weak supervision (Locatello et al., 2020a; Klindt et al., 2021). More pressingly, though, factorization sometimes prevents us from learning representations that disentangle independent causal mechanisms with nontrivial structure (Parascandolo et al., 2018; Träuble et al., 2020), which is actually how Bengio (2013) defined the challenge of disentanglement. Our goal in this work is to learn representations that can identify and explicitly model this kind of structure when it exists.

Still, there are a wide variety of ways to incorporate structure into representations or disentanglement. One is simply to change the disentanglement objective, e.g. to encourage different degrees of factorization within and across subgroups (Esmaeili et al., 2018). Another is to change the representation architecture such that “low-level” components are drawn conditionally on “high-level” components from some fixed hierarchy or graphical model (Sønderby et al., 2016; Siddharth et al., 2017; Singh et al., 2019). Others use mixed discrete-continuous representations where continuous representation components are either “global” (marginally independent) or “local” to a specific categorical value (conditionally independent, and sometimes inactive when the categorical takes other values) (Sorrenson et al., 2020; Choi et al., 2020). Typically, though, these approaches only support shallow hierarchies that must be specified by the user in advance, or require instance-level supervision (Yang et al., 2020). Our work is closest to the global-local approach of Choi et al. (2020), but we support arbitrarily deep hierarchies, and also learn them from data.

Other related approaches include relational autoencoders (Wang et al., 2014), which model structure between non-iid flat data, and graph neural networks (Defferrard et al., 2016), which learn flat representations of structured data. In contrast, we model structure within flat inputs. Also relevant are advances in object representations, such as slot attention (Locatello et al., 2020b). While this area has generally not focused on hierarchical nesting, it learns structure and seamlessly handles sets; we view our method as complementary. Finally, our hierarchy detection method is built on work in multiple- and robust manifold learning (Mahapatra & Chandola, 2017; Mahler, 2020). We contribute new robustness innovations and also introduce hierarchy (like Tino & Nabney (2002), but without requiring fixed dimensionality or human feedback).

### 3. Hierarchical Disentanglement Framework

In this section, we outline our framework for modeling hierarchical structure in representations. In our framework, we associate individual data points with paths down a dimension hierarchy (examples in Fig. 2). Dimension hierarchies consist of dimension group nodes (shown as boxes), each of which can have any number of continuous dimensions (shown as ovals) and an optional categorical variable (diamonds) that leads to other groups based on its value. For any data point, we “activate” only the dimensions along its corresponding path. Notation-wise, root(Z) denotes the group at the root of a hierarchy, and child(node) denotes the child groups of a categorical dimension Z_j. In the context of a dataset, for a dimension Z_j or a dimension group g, on(Z_j) or on(g) denotes the subset of the dataset where that Z_j or g is active.

As a potentially more intuitive analogy (as well as a visualization method), we can also understand hierarchical representations in terms of user interfaces with nested groups of sliders and radio buttons (Fig. 1). While traditional representations might consist of a single group of constantly visible sliders (or a mixture of sliders and radio buttons),
4. Hierarchical Disentanglement Benchmarks

In this section, both to clarify our framework and enable testing of our algorithms, we introduce several synthetic benchmark datasets with ground-truth hierarchical structure (see Fig. 2 for instances and dimension hierarchies).

4.1. Spaceshapes

Our first benchmark dataset is Spaceshapes, a binary 64x64 image dataset meant to hierarchically extend dSprites, a shape dataset common in the disentanglement literature (Matthey et al., 2017). Like dSprites, Spaceshapes images have location variables x and y, as well as a categorical shape with three options (in our case, moon, star, and ship). However, depending on shape, we add other continuous variables with qualitatively different effects: moons have a phase; stars have a sharpness to their shine; and ships have an angle. Finally, ships can optionally have a jet, which has a length (jetlen), but this is only defined at the deepest level of the hierarchy. The presence of jetlen alters the intrinsic dimensionality of the representation; it can be either 3D or 4D depending on the path. As in dSprites, variables are sampled from continuous or discrete uniforms. An interactive visualization of a representation trained to model this ground-truth hierarchy can be viewed here.

4.2. Chopsticks

Our second benchmark, Chopsticks, is actually a family of arbitrarily deep timeseries datasets. Chopsticks samples are 64D linear segments, each of which can have a uniform-sampled slope and/or intercept; different dataset variants can have one, the other, both, or either but not both. For all variants, segments initially span the whole interval. However, we then flip a coin to determine whether to chop the segment, in which case we add a uniform offset to the slope and/or intercept of the right half. We repeat this process recursively up to a configurable maximum depth, biasing probabilities so that we have equal probability of stopping at each level; each chop requires increasing local dimensionality to track additional slopes and intercepts. Although the underlying process is quite simple, the structure can be made arbitrarily deep, making it a useful benchmark for testing structure learning. We provide more details in §A.2, and interactive visualizations are also available for the depth-2 either and depth-3 both variants.

Although these datasets are designed to have clear hierarchical structure, there are certain ambiguities in how to structure aspects of the dimension hierarchies, which we discuss in §6.1.

5. Hierarchical Disentanglement Algorithms

We next present a method for learning hierarchical disentangled representations from data alone. We split the problem into two brunch-themed algorithms, MIMOSA (which infers hierarchies) and COFHAE (which trains autoencoders).

5.1. Learning Hierarchies with MIMOSA

The goal of our first algorithm, MIMOSA (Multi-manifold
Algorithm 1 MIMOSA(X)

1: Encode the data $X$ using a smooth autoencoder to reduce the initial dimensionality. Store as $Z$.
2: Construct a neighborhood graph on $Z$ using a Ball Tree (Omphuendo, 1989).
3: Run LocalSVD (Algorithm 3) on each point in $Z$ and its neighbors to identify local manifold directions.
4: Run BuildComponent (Algorithm 5) to successively merge neighboring points with similar local manifold directions.
5: Run MergeComponents (Algorithm 6) to combine similar components over longer distances and discard outliers.
6: Run ConstructHierarchy (Algorithm 7) to create a dimension hierarchy based on which components enclose others.
7: return the hierarchy and component assignments.

IsoMap On Smooth Autoencoder, is to learn a hierarchy $\hat{H}$ from data, as well as an assignment vector $A_0$ of data points to hierarchy leaves. MIMOSA consists of the following steps (see Appendix for Algorithms 3-7 and complexity, and Fig. 3 for a detailed example):

Dimensionality Reduction (Algorithm 1, line 1): We start by performing an initial reduction of $X$ to $Z$ using a flat autoencoder. While we could start with $Z = X$, performing this reduction saves computation as later steps (e.g. finding neighbors) scale linearly with $|Z|$. Although this requires choosing $|Z|$, we find the exact value is not critical as long as it exceeds the (max) intrinsic dimensionality of the data. We also find it important to use differentiable activation functions (e.g. Softplus rather than ReLU) to keep latent manifolds smooth; see Fig. A.4 for more.

Manifold Decomposition (Algorithms 3-6): We next decompose $Z$ into a set of manifold “components” by computing SVDs locally around each point and merging neighboring points with sufficiently similar subspaces. We then perform a second merging step over longer length scales, combining equal-dimensional components with similar local SVDs along their nearest boundary points and discarding small outliers, which we found was necessary to handle interstitial gaps when two manifolds intersect. The core of this step is based on a multi-manifold learning method (Mahapatra & Chandola, 2017), but we make efficiency as well as robustness improvements by combining ideas from RANSAC (Fischler & Bolles, 1981) and contagion dynamics (Mahler, 2020). The merging step is a new contribution.

It bears emphasis that manifold decomposition, which groups points based on the similarity of local principal components, is distinct from clustering, which groups points based on proximity. In the examples we consider, even hierarchical iterative clustering methods like OPTICS (Ankerst et al., 1999) will not suffice, as nearby points may lie on different manifolds.

Hierarchy Identification (Algorithm 7): Finally, we construct a tree by drawing edges from low-dimensional components to the higher-dimensional components that best “enclose” them, which we define using a ratio of inter-component to intra-component nearest neighbor distances; we believe this is novel. We use this tree and the component dimensions to construct a dimension hierarchy and a set of assignments from points to paths, which we output.

Hyperparameters: Each of these steps has several hyperparameters, and we provide a full listing and sensitivity study in §A.5. The one we found most critical was the minimum SVD similarity to merge neighboring points.

5.2. Training Autoencoders with COFHAE

Algorithm 2 COFHAE(X)

1: hierarchy, assignments = MIMOSA(X) # Algorithm 1
2: $HAE_\theta = \text{init}_\text{hierarchical}_\text{autoencoder}(\text{hierarchy})$
3: $D_\psi = \text{init}_\text{discriminator}()$
4: for $x, a \sim \text{minibatch}(X, \text{assignments})$ do
5: \hspace{1em} $a', z = HAE_\theta.\text{encode}(x; \tau)$ # Algorithm 8
6: \hspace{1em} $x' = HAE_\theta.\text{decode}(\text{concat}(a', z))$ # normal NN
7: \hspace{1em} $z' = \text{copy}(z)$
8: \hspace{1em} for $i = 1 .. |z_0|$ do
9: \hspace{2em} shuffle $z'_i$, over minibatch indices where on($z_i$)
10: \hspace{2em} end for
11: \hspace{1em} $L_\theta = L_x(x', x) + \lambda_1 L_a(a', a) - \lambda_2 \log \frac{D_\psi(z)}{1 - D_\psi(z)}$
12: \hspace{1em} $L_\psi = -\log D_\psi(z' - \theta) - \log (1 - D_\psi(z))$
13: \hspace{1em} $\theta = \text{descent}_\text{step}(\theta, L_\theta)$
14: \hspace{1em} $\psi = \text{descent}_\text{step}(\psi, L_\psi)$
15: end for
16: return $HAE_\theta$

Our first stage, MIMOSA, gives us the hierarchy and assignments of data down it. In the second stage, COFHAE (COnditionally Factorized Hierarchical AutoEncoder, Algorithms 2 and 8), we learn an autoencoder that respects this hierarchy via (differentiable) masking operations that impose structure on flat representations.

Hierarchical Encoding (Algorithm 8): Instances $x$ pass through a neural network encoder to an initial vector $z_{pre}$, whose dimensions correspond to both continuous and categorical dimensions. We then pass each set of categoricals through a softmax with temperature $\tau$, and use them to recursively mask the entirety of $z_{pre}$ based on the hierarchy. We finally split this masked vector into a continuous vector $z$ and a list of estimated assignments $a'$, outputting both.

Supervising Assignments: Although we lack ground-truth
during training, we do have assignments $a$ from MIMOSA (for at least a subset of the dataset). We add a penalty $L_a(a', a)$, weighted by $\lambda_1$, to make encoded $a'$ match $a$.

**Conditional Factorization:** Kim & Mnih (2018) penalize the total correlation (TC) between dimensions of flat continuous representations $z$ with two tricks. First, noting that TC is the KL divergence between $q(z)$ (the joint distribution of the encoded $z$) and $\bar{q}(z) = \prod_{j=1}^{|z|} q(z_j)$ (the product of its marginals), they approximate samples from $\bar{q}(z)$ by randomly permuting the values of each $z_i$ across batches (Arcones & Gine, 1992). Second, they approximate the KL divergence between the two distributions using the density ratio trick (Sugiyama et al., 2012) on an auxiliary discriminator $D_\psi(z)$, where $KL(q(z)||\bar{q}(z)) \approx \log \frac{D_\psi(z)}{1-D_\psi(z)}$ if $D_\psi(z)$ outputs accurate probabilities of $z$ having been sampled from $\bar{q}$. We adopt a similar approach, except instead of permuting each $z_i$ across the full batch $B$, we only permute it where it is active, i.e. $B \cap \text{on}(z_i)$ (defined using the hardened version of the mask). This approximates a hierarchical version of $\bar{q}(z)$ where each dimension distribution is a mixture of 0 and the product of its active marginals. $D_\psi(z)$ then lets us estimate the KL between this distribution and $q(z)$, which we penalize and weight with $\lambda_2$.

This approach presumes ground-truth continuous variables should be conditionally independent given categorical values, which is a major assumption. However, it is less strict than the assumption taken by many disentanglement methods, i.e. that continuous variables are independent marginally, and it may remain useful as an inductive bias.

### 6. Hierarchical Disentanglement Metrics

In this section, we develop metrics for quantifying how well learned representations and hierarchies match ground-truth.

![Initial 4D AE representation (projected to 3D, color = true assignment)](image1)

**Figure 3.** Breakdown of MIMOSA for the depth-2 either version of Chopsticks, colored by ground-truth assignments. MIMOSA learns an initial 4D softplus AE representation (left), decomposes it into lower-dimensional components by grouping together neighboring points with similar local SVDs (second from left), merges them over longer distances while discarding outliers (second from right), and finally uses enclosure relationships to infer a hierarchy (right). In this case, correspondence between the assignment of points to learned components vs. ground-truth is very close (99.8% purity, covering 93.7% of the training set, and with no $H$-error—see §6.2 for definitions of these metrics). Similar examples are shown for other datasets in Figs. A.11-A.16 of the Appendix.

#### 6.1. Desiderata and Invariances

Our goal in designing metrics is to measure whether we have learned the “right representation,” both in terms of global structure and specific variable correspondences. In an ideal world, we would measure whether a learned representation $Z$ is identically equal to a ground-truth $V$. However, most existing disentanglement metrics are invariant to permutations, so that dimensions $V_i$ can be reordered to match different $Z_j$, as well as univariate transformations, so that the values of $Z_j$ do not need to be identical to $V_i$. In the case of methods like the SAP score (Kumar et al., 2017), these univariate transformations must be linear, but as the uniformity of scaling can be arbitrary, we permit general nonlinear transformations, as long as they are 1:1, or invertible.

Hierarchical representations have an additional ambiguity about the right “vertical” placement of continuous variables. For example, on Spaceshapes, the phase, shine, and angle variables could all be “merged up” to a single top-level variable whose effect changes based on shape. Alternatively, x and y position could be “pushed down” and duplicated for each shape despite their analogous effects (see Fig. A.10 for an illustration). In terms of our user interface analogy from Fig. 1 (or our specific implementation), “merge up” and “push down” transformations correspond to moving sliders into or out of outlined groups, but keeping their effects on the outputs the same, as well as preserving the structure of nested radio buttons. To a user interacting with such representations, they would appear almost identical, except some slider labels might change with radio button settings. Because of this functional near-equivalence, we defer the problem of deciding the most natural vertical placement of continuous variables to future work, and make our main metrics invariant to them.
6.2. MIMOSA Metrics: \textit{H}-error, Purity, Coverage

The first metric we use to evaluate MIMOSA is \textit{H}-error, which measures whether learned hierarchy \( \hat{H} \) has the same essential structure as the ground-truth hierarchy \( H \). We define \( H \)-error in terms of the tree edit distance of Zhang & Shasha (1989) (i.e., minimum number of insertions, edits, or deletions to transform \( H \) into \( \hat{H} \)), but between normalized “merged up” representations of each hierarchy; details are in §A.3. This metric is 0 if and only if both hierarchies are the same up to the transformations described in §6.1.

The second MIMOSA metric is \textit{purity}, which measures whether the assignments output by MIMOSA match ground-truth. To compute it, we iterate over each leaf in \( H \), find the leaf in \( \hat{H} \) to which most of its assigned points belong, and then compute the fraction that belong to the majority. Then we average these scores across \( \hat{H} \), weighting by the number of points in each leaf. This metric only falls below 1 if leaves contain points with different ground-truth assignments.

The final metric we use to evaluate MIMOSA is \textit{coverage}. Since MIMOSA discards small outlier components, it is possible that the final set of assignments will not cover the full training set. If almost all points are discarded this way, the other metrics may not be meaningful. As such, we measure coverage as the fraction of the training set which is not discarded. We note that hyperparameters can be tuned to ensure high coverage without knowing ground-truth assignments.

6.3. COFHAE Metrics: \( R^4 \) and \( \tilde{R}^4 \) Scores

Per our desiderata, we seek to check whether every ground-truth variable \( V_i \) can be mapped invertibly to some learned dimension \( Z_j \). As a preliminary definition, we say that a learned \( Z_j \) \textit{corresponds} to a ground-truth \( V_i \) over some set \( S \subseteq \mathbb{R} \) if a bijection between them exists; that is,

\[
\exists f(\cdot) : S \rightarrow \mathbb{R} \text{ s.t. } f(V_i) = Z_j \text{ and } f^{-1}(Z_j) = V_i \quad (1)
\]

We say that \( Z \) \textit{disentangles} \( V \) if all \( V_i \) have a corresponding \( Z_j \). To measure the extent to which bijections exist, we can simply try to learn them (over random splits of many paired samples of \( V_i \) and \( Z_j \)). Concretely, for each pair of learned and true dimensions, we train univariate models to map in both directions, compute their coefficients of determination \( (R^2) \), and take their geometric mean:

\[
f \equiv \min_{f \in F} \mathbb{E}_{\text{train}}[ (f(X) - Y)^2 ]
\]

\[
R^2(X \rightarrow Y) \equiv \mathbb{E}_{\text{test}} \left[ 1 - \frac{\sum (f(X) - Y)^2}{\sum (Y - Y)^2} \right]
\]

\[
R^2(X \leftrightarrow Y) \equiv \sqrt{R^2(X \rightarrow Y) \cdot R^2(Y \rightarrow X)},
\]

where we average over train/test splits (we use 5), assume \( F \) is sufficiently flexible to contain the optimal bijection (we use gradient-boosted decision trees), and assume our dataset is large enough to reliably identify \( f \in F \). In the limit, \( R^2(X \leftrightarrow Y) \) can only be 1 if a bijection exists, as any region of non-zero mass in the joint distribution of \( X \) and \( Y \) where this is false would imply \( \mathbb{E}[(f(X) - Y)^2] > 0 \) or \( \mathbb{E}[(f(Y) - X)^2] > 0 \). In the special case that \( Y \) is discrete rather than continuous, we use classifiers for \( f \) and accuracy instead of \( R^2 \), but the same argument holds.

To measure whether a set of variables \( Z \) disentangles another set of variables \( V \), we check if, for each \( V_i \), there is at least one \( Z_j \) for which \( R^2(V_i \leftrightarrow Z_j) = 1 \):

\[
R^4(V, Z) \equiv \frac{1}{|V|} \sum_i \max R^2(V_i \leftrightarrow Z_j),
\]

We call this the “right-representation” \( R^2 \), or \( R^4 \) score. Note that this metric is related to the existing SAP score (Kumar et al., 2017), except we allow for nonlinearity, require high \( R^2 \) in both directions, and take the maximum over each score column rather than the difference between the top two entries (which avoids assuming ground-truth is factorized).

Although \( R^4 \) is useful for measuring correspondence between sets of variables that are both always active, it does not immediately apply to hierarchical representations unless inactive variables are represented somehow, e.g. as 0 (an arbitrary implementation decision that affects \( R^2 \) by changing \( \mathbb{E}[Y] \)). It also lacks invariance to merge-up and push-down operations. Instead, we seek \textit{conditional correspondence} between \( V_i \) and a set of dimensions in \( Z \), defined as

\[
\forall V_i \in \text{on}(V_i) \exists Z_i = \{Z_j, Z_k, \ldots\} \text{ s.t.}
\]

\[
(a) \ V_i \text{ corresponds to } Z_j \text{ over } \text{on}(V_i) \cap \text{on}(Z_j), \]

\[
(b) \ \text{on}(Z_j) \cap \text{on}(Z_k) = \emptyset \text{ for all } j \neq k, \text{ and}
\]

\[
(c) \bigcup_{z \in Z} \text{on}(z) = \text{on}(V_i), \tag{4}
\]

or rather that we can find some tiling of \( \text{on}(V_i) \) into regions where it corresponds 1:1 with different \( Z_j \) which are never active simultaneously. This allows for one \( Z_j \) to correspond to non-overlapping elements of \( V \) (e.g. merging up), as well as for one \( V_i \) to be modeled by non-overlapping elements of \( Z \) (e.g. pushing down).

We can then formulate a conditional \( \tilde{R}^4 \) score which quantifies how closely conditional correspondence holds:

\[
\tilde{R}^2(V_i, g) \equiv \max_{g' \in \text{children}(Z_j)} \left( \frac{\max_{j \in g} \left( R^2(V_i \leftrightarrow Z_j \mid \text{on}(V_i) \cap \text{on}(g)) \right)}{\text{|on}(V_i)|} \right),
\]

for a dimension group \( g \); the overall disentanglement is:

\[
\tilde{R}^4(V \leftrightarrow Z) \equiv \frac{1}{|V|} \sum_{i=1}^{\text{|V|}} \tilde{R}^2(V_i, \text{root}(Z)).
\]
In the special case that $V$ and $Z$ are flat, $R^4_z$ reduces to $R^4$. We note that even for flat representations, the $R^4$ score may be a useful measure of disentanglement when ground-truth variables are not factorized.

7. Experimental Setup

**Benchmarks:** We ran experiments on nine benchmark datasets: Spaceshapes, and eight variants of Chopsticks (varying slope, intercept, both, and either at recursion depths of 2 and 3). See §4 for more details, and Fig. A.1 for preliminary experiments on noisy data.

**Algorithms:** In addition to COFHAE with MIMOSA, we trained a number of flat baselines. As fully continuous baselines, we trained autoencoders (AE), variational autoencoders (Kingma & Welling, 2013) (VAE), the $\beta$-total correlation autoencoder (Chen et al., 2018) (TCVAE), and FactorVAE (Kim & Mnih, 2018). As mixed discrete-continuous baselines, we trained JointVAE (Dupont, 2018) and CascadeVAE (Jeong & Song, 2019), providing them with the ground-truth structure of discrete variables. Finally, we ran COFHAE ablations using the ground-truth hierarchy and assignments, testing all possible combinations of loss terms and comparing conditional vs. marginal TC penalties; results are in Fig. 5. See §A.1 for training and model details.

**Metrics:** To evaluate hierarchies, we computed purity, coverage, and $H$-error (§6.2). Results are in Table 1. To measure disentanglement, we primarily use $R^4_z$ (§6.3); results for all datasets and models are in Fig. 4. We also compute the following baseline metrics: the SAP score (Kumar et al., 2017) (SAP), the mutual information gap (Chen et al., 2018) (MIG, estimated using 2D histograms), the FactorVAE score (Kim & Mnih, 2018) (FV AE), and the DCI disentanglement score (Eastwood & Williams, 2018) (DCI). Most implementations were adapted from disentanglement_lib (Locatello et al., 2018). We also compute our marginal $R^4$ score. Results across metrics are shown for a subset of datasets and models in Fig. 6.

**Hyperparameters:** COFHAE is only given instances $X$, which complicates cross-validation. However, we can still tune its hyperparameters to ensure assignments $a'$ match MIMOSA outputs $a$ and reconstruction loss for $x$ is low (which fail to happen if the adversarial term dominates). Over a grid of $\tau$ in $\{\frac{1}{2}, 5, 1\}$, $\lambda_1$ in $\{10, 100, 1000\}$, and $\lambda_2$ in $\{1, 10, 100\}$, we select the model with the lowest training reconstruction loss $L_z$ from the $\frac{1}{3}$ with the lowest assignment loss $L_a$. For MIMOSA, hyperparameters can be tuned to ensure high coverage (purity and $H$-error require side-information); see §A.5 for more details.

For our baselines, we show results at $\beta=5$ for TCVAE, $\gamma=10$ for FactorVAE, $\beta=1$, $C_s=C_r=10$ for JointVAE, and $\beta=2$ for CascadeVAE (with other hyperparameters set to the same settings as the original paper). However, we tested each method across a variety of strength and capacity hyperparameters, and show more complete results in Fig. A.7.

![Disentanglement results over 5 restarts](image)

**Figure 4.** Hierarchical disentanglement results for representation learning methods (baselines and COFHAE + MIMOSA) over all nine datasets. COFHAE almost perfectly disentangles ground-truth on the six simplest versions of Chopsticks, with some degradations on the two most complex versions (with very deep hierarchies) and on Spaceshapes (with a shallower hierarchy, but higher-dimensional inputs). Baseline methods were generally much more entangled, though JointVAE, $\beta$-TCVAE, and CascadeVAE are competitive in certain cases.

![Figure 5.](image)

**Figure 5.** Ablation study for COFHAE on the depth-2 both version of Chopsticks (over 5 restarts). Hierarchical disentanglement is low for flat AEs (Flat); adding the ground-truth hierarchy $H$ improves it (Hier $H$), as does also adding supervision for ground-truth assignments $A$ (H+A). Adding a FactorVAE-style marginal TC penalty $(H+A+TC(Z))$ does not appear to help disentanglement, but making that TC penalty conditional $(H+A+TC(Z|on))$, i.e. COFHAE) brings it close to the near-optimal disentanglement of a hierarchical model whose latent representation is fully supervised $(H+A+Z)$. However, the hierarchical conditional TC penalty fails to produce this same disentanglement without any supervision over assignments $(H+TC(Z))$.

8. Results and Discussion

MIMOSA consistently recovered the right hierarchies. Per Table 1, we consistently found the right hierarchy for all
sampling to ensure sufficient coverage). Finally, alongside non-identifiability and optimization issues, MIMOSA errors (e.g., merge-up/push-down differences for Spaceshapes and suboptimal purity and coverage for Chopsticks) also may play a role, as evidenced by performance improvements in our full COFHAE ablations in Fig. A.6. Despite all of these issues, COFHAE is still closer to optimal, at best and on average, than any of our baseline algorithms (even on Spaceshapes, where it is possible for a flat representation to disentangle all features except jet length). We note also that our baselines often performed worse with increasing disentanglement penalty strength (Fig. A.7), with the closest COFHAE competitor, JointVAE, achieving its best results at its minimal tested value $\gamma=1$ (i.e., equivalent to a normal VAE). These results are consistent with the fact that minimizing marginal rather than conditional TC on these datasets prevents models from learning the right representation.

\(R^4\) provides more insight into disentanglement than baselines. One way to evaluate an evaluation metric is to test it against a precisely known quantity. In this case, we know the \(H+A+Z\) model, whose encoder is supervised to match ground-truth, should receive a near-perfect score. The only metrics to do this consistently are \(R^4\) and \(R^4\). Note that the DCI disentanglement score, based on the entropy of normalized feature importances from an estimator predicting single ground-truth factors from all learned dimensions, comes close. Intuitively, this metric could behave similarly to \(R^4\) if its estimator was trained to be sparse (placing importance on as few dimensions as possible). Using \(R^4\)s of univariate estimators is more direct, and also incorporates information from the DCI informativeness score.

Another way to evaluate an evaluation metric is to test whether quantitative differences capture salient qualitative differences. To this point, specifically to compare \(R^4\) and \(R^4\), we consider several examples in Fig. A.8 and Fig. A.9. First, we see that for the Spaceshapes COFHAE model in Fig. A.8c (or here), its \(R^4\) score (0.89) is higher than its \(R^4\) (0.75). This increase is due to the fact that \(R^4\) penalizes “push-down” differences (§6.1) between the learned and true factors representing \(x\) and \(y\) position, while \(R^4\) is invariant to them. However, the overall increase is less dramatic than one might expect due to moderate decreases in corre-
spondence scores for other dimensions (e.g. 0.98→0.89 for jetlen), which occur because $R_4^i$ is not biased by spurious equality between dimensions which are both inactive. Another example of a difference between $R^4$ and $R^4_i$ (illustrating invariance to “merging up” rather than “pushing down”) is for the Spaceshapes $\beta$-TCVAE in Fig. A.8b. In this case, histograms show that one $\beta$-TCVAE variable ($Z_i$) corresponds closely to both moon phase and star shine (and to a lesser extent, jetlen), only one of which is active at a time. The $R^4$ score (0.47) assigns low scores to these correspondences, but $R^4_i$ (0.69) properly factors them in.

**COFHAE and MIMOSA subcomponents improve performance.** Though COFHAE contains many moving parts, results in Fig. 5 and Fig. A.6 suggest they all count. Autoencoders only achieve optimal disentanglement if provided with the hierarchy, assignments, and a conditional (not marginal) penalty on the TC of continuous variables; no partial subset suffices. In the Appendix, Fig. A.5 shows ablations and sensitivity analyses for MIMOSA that validate its subcomponents are important as well.

### 8.1. Remark on Identifiability and Parsimony

From Locatello et al. (2018), we know all forms of TC minimization permit multiple solutions (though they often improve disentanglement empirically, especially when ground-truth factors are non-Gaussian). However, what about the other components of our method, such as MIMOSA?

MIMOSA does not minimize an objective function, so questions of identifiability might seem moot. However, we could reformulate it as trying to find a small set of low-dimensional and bounded-curvature manifolds that approximately contain a large fraction of the data. More concretely, we could place penalties or constraints on, e.g., the cardinality, dimensionality, and mean or percentiles of error and principal curvature magnitudes over the set. Such a problem might well be identifiable (up to the transformations discussed in §6.1), though analyzing it is beyond the scope of this work.

However, perhaps a better-motivated formulation that covers both MIMOSA and COFHAE would be to return to minimum description length (MDL)—the same problem that motivated much of the initial research into factorized representations (Barlow, 1961; Zemel, 1994). As an example, assume we are given a dataset of $N$ instances, $\frac{7}{8}$ of which lie on a 1D manifold, and $\frac{1}{8}$ of which lie on an 8D manifold. If we must encode instances as flat vectors of 32-bit floats, those vectors will need to be at least 8D for accurate reconstruction, meaning the dataset’s description length will be $8 \times 32 \times N = 256N$ bits (plus the size of the model, which is negligible for sufficiently large $N$). However, if we use a disentangled hierarchical representation, we need either 1 or 8 floats to represent each instance (plus a single bit to distinguish between them). In that case, the description length would be $(\frac{1}{8} \times 8 \times 32) + (\frac{7}{8} \times 1 \times 32) + 1) N = 61N$ bits, which is minimal (assuming the model is not much larger). The problem of learning factorized representations within each manifold might remain non-identifiable, but the MDL argument for doing so remains the same as in Zemel (1994). This example suggests that (disentangled) hierarchical representations might spontaneously emerge as the (partially identifiable) solution to MDL objectives, at least for datasets that lie on multiple manifolds.

### 9. Conclusion

In this work, we introduced a novel formulation of hierarchical disentanglement, where ground-truth representation dimensions are organized into a tree and activated or deactivated based on the values of categorical dimensions. We presented benchmarks, algorithms, and metrics for learning and evaluating such hierarchical representations.

There are a number of promising avenues for future work. One is extending our methods to handle a wider variety of underlying structures, e.g. dimension DAGs, or integrating our methods with object representation techniques to better model generative processes involving ordinal variables or unordered sets (Locatello et al., 2020b). Another is to better solve or understand hierarchical disentanglement as we have already formulated it, e.g. by improving robustness to noise (Fig. A.1) or providing a better theoretical understanding of identifiability, perhaps through the lens of description length. Finally, there are ample opportunities to apply these techniques to real-world data that we expect to have hierarchical multiple-manifold structure, such as patient phenotype or population genetics datasets.

More generally, we feel it is important for representation learning to move beyond flat vectors, and work towards explicitly modeling the rich structure contained in the real world. Symbolic AI and cognitive science researchers have made compelling arguments that future AI progress should be evaluated not by improvements in accuracy or reconstruction error, but by how well models build their own interpretable models of the world (Lake et al., 2017). Our work takes steps in this direction.

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A. Appendix

A.1. Training and Architecture Details

For Chopsticks, our encoders and decoders used two hidden layers of width 256, and our loss function $L_x$ was defined as a zero-centered Gaussian negative log likelihood with $\sigma = 0.1$. For Spaceshapes, encoders and decoders used the 7-layer convolutional architecture from Burgess et al. (2018), and our loss function $L_x$ was Bernoulli negative log likelihood. All models were implemented in Tensorflow; code is available at https://github.com/detak/hierarchical-disentanglement.

For both models, the assignment loss $L_{a}$ was set to mean-squared error, but only for assignments that were defined: we set undefined assignment components to -1, then let $L_a(a,a') = \sum_i 1[a_i' \geq 0](a_i - a_i')^2$.

All activation functions were set to ReLU (max(0,x)) or Softplus (ln(1+e^x)), e.g. for the initial smooth autoencoder, which was also trained with dimensionality equal to one plus the maximum intrinsic dimensionality of the dataset. We investigate varying this parameter in Fig. A.5 and find it can be much larger, and perhaps would have produced better results (though nearest neighbor calculation and local SVD computations would have been slower).

All models were trained for 50 epochs with a batch size of 256 on a dataset of size 100,000, split 90%/10% into train/test. We used the Adam optimizer with a learning rate starting at 0.001 and decaying by $\frac{1}{10}$ halfway and three-quarters of the way through training.

For COFHAE, we selected softmax temperature $\tau$, the assignment penalty strength $\lambda_1$, and the adversarial penalty strength $\lambda_2$ based on training set reconstruction error and MIMOSA assignment accuracy. Splitting off a separate validation set was not necessary, as the most common problem we faced was poor convergence, not overfitting; the adversarial penalty would dominate and prevent the procedure from learning a model that could reconstruct $X$ or $A$.

Specifically, for each restart, we ran COFHAE with $\tau$ in $\{2, 3, 4\}$, $\lambda_1$ in $\{10, 100, 1000\}$, and $\lambda_2$ in $\{1, 10, 100\}$. We then selected the model with the lowest training MSE $\sum_n ||x_n - x_n'||_2^2$, but restricting ourselves to the 33.3% of models with the lowest assignment loss $\sum_n L_a(a_n, a_n')$.

For evaluating $R^4$ and $R^d$, we used gradient boosted decision trees, which were faster to train than neural networks.

A.2. Additional Chopsticks Details

In this section, we clarify the generative process behind the different variants of Chopsticks, and discuss alternatives.

Chopsticks can be generated by the following Python code (the exact code we used is slightly different due to the need to save ground-truth factors):

```python
def Bern(p):
    return int(np.random.uniform() < p)

def Unif(a,b):
    return a + np.random.uniform() * (b-a)

def stick_segment(variant, T):
    slope = Unif(-0.01, 0.01) * np.arange(T)
    inter = Unif(-0.2, 0.2) + np.zeros(T)

    if variant == 'slope':
        return slope
    elif variant == 'inter':
        return inter
    elif variant == 'both':
        return slope+inter
    elif variant == 'either':
        return slope if Bern(0.5) else inter

def chopsticks(depth, variant, T=64):
    stick = stick_segment(variant, T)
    chop = Bern(1-np.power(2.0,-(depth-1)))

    if chop:
        stick2 = chopsticks(depth-1, variant, T//2)
    else:
        stick[T//2:] += stick2
    return stick
```

For all variants, at depth $d \geq 1$, we sample a linear “stick”, and then “chop” it with probability $1 - 2^{-(d-1)}$. If we chop the stick, then we recursively generate a new stick of half the length, which we add to the second half of the current stick. We choose chop probabilities in this way so that, on average, we have equal counts of samples at each depth.

Although this framework already gives us a wide diversity of datasets, we could consider others. One option is to add noise, which hurts MIMOSA, though it depends on the dataset (Fig. A.1). Another option is to sample slopes and intercepts non-uniformly or even over non-convex sets; see e.g. Fig. A.2a, where we set slope/intercept magnitudes at least a threshold away from 0, which introduces gaps into the initial representation. In general, MIMOSA continues to return the right hierarchy for all such slope/intercept distributions, though COFHAE disentanglement tends to drop when variables are sampled from Gaussians, likely because of symmetry (with proper rescaling, we can rotate factorized Gaussians in any direction and preserve factorization; the same is not true for uniforms, though Locatello et al.
In preliminary experiments, we find that noise poses the greatest problem for identifying the lowest-dimensional components, e.g., the 1D components in (b) that end up being classified as 2D or 3D. Tuning parameters would help, but we lack labels to cross-validate. (2018) show there must exist analogous, if more complex, transformations. Yet another possibility is to overwrite the slope and/or intercept when recursing, rather than offsetting them (Fig. A.2b). Overwriting slope does not affect MIMOSA performance, though it changes the orientation of lower-dimensional components within higher-dimensional manifolds, which can affect COFHAEE, but overwriting the intercept can break the geometrical nesting of manifolds at large slopes. Although we considered many of these options, we ultimately decided it was pedagogically best for our benchmark to distribute instances over maximally simple (but still arbitrarily deep) underlying manifold structures.

A.3. Computing $H$-error

Our $H$-error metric is meant to quantify the “edit distance” between two dimension hierarchies $H$ and $\hat{H}$, but in a way that is invariant to merge-up and push-down operations, as well as reorderings of child groups. To implement it, we first convert $H$ and $\hat{H}$ to a canonical form where each dimension group is labeled by the minimum downstream dimension of its leaves (which equals the dimension of the manifold component at the matching location in the original enclosure hierarchy), which renders us invariant to merge-up and push-down operations. We then reorder children in terms of the (sorted) concatenation of their downstream labels, which renders us invariant to child ordering in most cases. Finally, we apply the Zhang-Shasha algorithm for tree edit distance between ordered, labeled trees (Zhang & Shasha, 1989; Paassen et al., 2015) to get our final $H$-error.

A.4. Complexity and Runtimes

Per Fig. A.3, the total runtime of our method is dominated by COFHAEE, an adversarial autoencoder method which has the same complexity as FactorVAE (Kim & Mnih, 2018) (linear in dataset size $N$ and number of training epochs, and strongly affected by GPU speed).

MIMOSA could theoretically take more time, however, as the complexity of constructing a ball tree (Omohundro, 1989) for nearest neighbor queries is $O(|Z|N \log N)$. As such, initial dimensionality reduction is critical; in our
Spaceshapes experiments, \(|Z|\) is 7, whereas \(|X|\) is 4096.

Other MIMOSA steps can also take time. With a num_nearest_neighbors of \(k\), the complexity of running local SVD on every point in the dataset is \(O(N(|Z|^2k + |Z|k^2 + k^3))\), providing another reason to reduce initial dimensionality and keep neighborhood size manageable (though ideally \(k\) should increase with \(|Z|\) to robustly learn local manifold directions). Iterating over the dataset in BuildComponent and computing cosine similarity will also have complexity at least \(O(Nkd(d + |Z|))\) for components of local dimensionality \(d\), and detecting component boundaries can actually have complexity \(O(Nke^d)\) (if this is implemented, as in our experiments, by checking if projected points are contained in their neighbors’ convex hulls—though we also explored a much cheaper \(O(Nkd^2d)\) strategy of checking for the presence of neighbors in all principal component directions that worked almost as well).

Although these scaling issues are worth noting, MIMOSA was still relatively fast in our experiments, where runtimes were dominated by neural network training (Fig. A.3).

**A.5. MIMOSA Hyperparameters**

In this section, we list and describe all hyperparameters for MIMOSA, along with values that we used for our main results. We also present sensitivity analyses in Fig. A.5.

**MIMOSA initial autoencoder (Algorithm 1, line 1)**

- initial_dim - the dimensionality of the initial smooth autoencoder. As the sensitivity analysis in Fig. A.5 shows, this does not need to be as low as the intrinsic dimensionality of the data, which MIMOSA will estimate, and ideally should be a little larger. We defaulted to using the maximum intrinsic dimensionality plus 1; in a real-world context where this information is not available, it can be estimated by starting at initial_dim = \(|X|\) and reducing until initial autoencoder reconstruction error starts increasing.

- Training and architectural details appropriate for the data modality (e.g. convolutional layers for images). See §A.1 for our choices.

**LocalSVD (Algorithm 3)**

- num_nearest_neighbors - neighborhood size for LocalSVD and traversal; we used 40. Must exceed initial_dim; could replace with a search radius.

- ransac_frac - the fraction of neighbors to refit SVD. We used 2/3. Note that we do not run traditional, multi-step RANSAC (Fischler & Bolles, 1981), but a more efficient two-step approximation, where we define the loss term based an aggregation of reconstruction errors across dimensions. Another (less efficient but potentially more robust) option would be to iteratively re-fit SVD using the points with lowest reconstruction error at each dimension, and check if the resulting eigenvalues meet our cutoff criteria.

- eig_cumsum_thresh - the minimum fraction of variance SVD dimensions must explain to determine local dimensionality. We used 0.95. For noisy or sparse data, it might be useful to reduce this parameter.

- eig_decay_thresh - the minimum multiplicative factor by which SVD eigenvalues must decay to determine local dimensionality. We used 4. It might also be useful to reduce this parameter for sparse data.

Note that our LocalSVD algorithm can be seen as a faster version of Multiscale SVD (Little et al., 2009), which is used in an analogous way by Mahapatra & Chandola (2017), but would require repeatedly computing singular value decompositions over different search radii for each point.

**BuildComponent (Algorithm 5)**

- cos_simil_thresh - neighbors’ local SVDs must be this similar to add to the component. This corresponds to the \(\epsilon\) parameter from Mahapatra & Chandola (2017). We used 0.99 for Chopsticks and 0.95 for Spaceshapes; in general, we feel this is one of the most important parameters to tune, and should generally be reduced in the presence of noise or data scarcity.

- contagion_num - only add similar points to a manifold component when a threshold fraction of their neighbors have already been added. This is useful for robustness, and corresponds to the \(T\) parameter from Mahler (2020) (but expressed as a number rather than a fraction). We used 5 for Chopsticks and 3 for Spaceshapes. Values above 20% of
Algorithm 3 LocalSVD(Z)

1: Run SVD on Z (a design matrix of dimension num_nearest_neighbors by initial_dim)
2: if ransac_frac < 1 then
3:   for each dimension d from 1 to initial_dim − 1 do
4:     for each point z_n do
5:       Compute the reconstruction error for z_n using the only first d SVD dimensions
6:     end for
7:   end for
8:   Take the norm of reconstruction errors across dimensions, giving a vector of length num_nearest_neighbors
9:   Re-fit SVD on points whose error-norms are less than the 100 × ransac_frac percentile value.
10: end if
11: for each dimension d from 1 to initial_dim − 1 do
12:   Check if the cumulative sum of the first d eigenvalues is at least eig_cumsum_thresh
13:   Check if the ratio of the dth to the d + 1st eigenvalue is at least eig_decay_thresh
14: if both of these conditions are true then
15:   return only the first d SVD components
16: end if
17: end for
18: return the full set of SVD components otherwise

Algorithm 4 TangentPlaneCos(U, V)

1: if U and V are equal-dimensional then
2:   return |det(U · V^T)|
3: else
4:   return 0
5: end if

Algorithm 5 BuildComponent(z_i, neighbors, svds)

1: Initialize component to z_i and neighbors z_j not already in other components where TangentPlaneCos(svds_i, svds_j) ≥ cos_simil_thresh (Algorithm 4).
2: while the component is still growing do
3:   Add all points z_k for which at least contagion_num of their neighbors z_ℓ are already in the component with TangentPlaneCos(svds_k, svds_ℓ) ≥ cos_simil_thresh.
4:   Skip adding any z_k already in another component.
5: end while
6: return the set of points in the component

Algorithm 6 MergeComponents(components, svds)

1: Discard components smaller than min_size_init.
2: for each component c_i do
3:   Construct a local ball tree for the points in c_i.
4:   Set c_i.edges to points not contained in the convex hull of their neighbors in local SVD space.
5: end for
6: Initialize edge overlap matrix M of size |components| by |components| to 0.
7: for each ordered pair of equal-dimensional components (c_i, c_j) do
8:   Set M_ij to the fraction of points in c_i.edges for which the closest point in c_j.edges has local SVD tangent plane similarity above cos_simil_thresh.
9: end for
10: Average M with its transpose to symmetrize.
11: Merge all components c_i ≠ c_j of equal dimensionality d where M_ij ≥ min_common_edge_frac(d).
12: Discard components smaller than min_size_merged.
13: return the merged set of components
Algorithm 7 ConstructHierarchy(components)

1: for each component $c_i$ do
2: Set $c_i$.neighbor_lengthscale to the average distance of points to their nearest neighbors inside the component (computed using the local ball tree from Algorithm 6)
3: end for
4: for each pair of different-dimensional components $(c_i, c_j)$, $c_i$ higher-dimensional do
5: Compute the average distance from points in $c_i$ to their nearest neighbors in $c_j$ (via ball tree).
6: Divide this average distance by $c_i$.neighbor_lengthscale.
7: if the resulting ratio $\leq$ neighbor_lengthscale_mult then
8: Set $c_j \in c_i$ ($c_j$ is enclosed by $c_i$)
9: end if
10: end for
11: Create a root node with edges to all components which do not enclose others.
12: Transform the component enclosure DAG into a tree (where enclosing components are children of enclosed components) by deleting edges which:
   1. are redundant because an intermediate edge exists, e.g. if $c_1 \in c_2 \in c_3$, we delete the edge between $c_1$ and $c_3$.
   2. are ambiguous because a higher-dimensional component encloses multiple lower-dimensional components (i.e. has multiple parents). In that case, preserve only the edge with the lowest distance ratio.
13: Convert the resulting component enclosure tree into a dimension hierarchy:
   1. If the root node has only one child, set it to be the root. Otherwise, begin with a dimension group with a single categorical dimension whose options point to groups containing each child.
   2. For the rest of the component tree, add continuous dimensions until the total number of continuous dimensions up to the root equals the component’s dimensionality.
   3. If a component has children, add a categorical dimension that includes those child groups as options (recursing down the tree), along with an empty group ($\emptyset$) option.
14: return the dimension hierarchy

Algorithm 8 HAE$_\theta$.encode($x; \tau$)

1: Encode $x$ using any neural network architecture as a flat vector $z_{pre}$, with size equal to the number of continuous variables plus the number of categorical options in HAE$_\theta$.hierarchy.
2: Associate each group of dimensions in the flat vector with variables in the hierarchy.
3: For all of the categorical variables, pass their options through a softmax with temperature $\tau$.
4: Use the softmax outputs to recursively mask all components of $z_{pre}$ corresponding to variables below each option in HAE$_\theta$.hierarchy.
5: return the masked representation, separated into discrete $a'$, continuous $z$, as well as the mask $m$ (for determining active dimensions later).

Figure A.4. Comparison of the latent spaces learned by MIMOSA initial autoencoders with ReLU (left) vs. Softplus (right) activations on three versions of Chopsticks (depth=1 either, depth=2 either, and depth=3 slope). Each plot shows encoded data samples colored by their ground-truth location in the dimension hierarchy. Because ReLU activations are non-differentiable at 0, the resulting latent manifolds contain sharp corners where local SVD directions change discontinuously, causing issues for BuildComponent and MergeComponents within MIMOSA (Algorithms 5 and 6). Representations learned by autoencoders with smooth activation functions work much better.
Figure A.5. Effect of varying different hyperparameters (and ablating different robustness techniques) on MIMOSA. Default values are shown with vertical gray dotted lines, and for each hyperparameter (top to bottom), average coverage (left), purity (middle), and $H$ error (right) when deviating from defaults are shown for three versions of the Chopsticks dataset. Results suggest both a degree of robustness to changes (degradations tend not to be severe for small changes), but also the usefulness of various components; for example, results markedly improve on some datasets with $\text{contagion_num} > 1$ and $\text{ransac_frac} < 1$ (implying contagion dynamics and RANSAC both help). Many parameters exhibit tradeoffs between component purity and dataset coverage.
num_nearest_neighbors will likely produce poor results, and we found the greatest increases in robustness just going from 1 (or no contagion dynamics) to 2.

**MergeComponents (Algorithm 6)**

- **min_size_init** - discard initial components smaller than this, which helps speed up the algorithm (by reducing the number of pairwise comparisons) and avoid merges through single-point components. We used 0.02% of the dataset size, or 20 points.
- **min_size_merged** - discard merged components smaller than this, which helps exclude spurious interstitial points at boundaries where low-dimensional components intersect. We used 2% of the dataset size, or 2000 points.
- **min_common_edge_frac(d)** - the minimum fraction of edges that two manifold components must share in common to merge, as a function of dimensionality $d$. We used $2^{−d−1} + 2^{−d−2}$; this is based on the idea that two neighboring (possibly distorted) hypercubes of dimension $d$ should match on one of their sides; since they have $2^d$ sides, the fraction of matching edge points would be $2^{−d}$. However, for robustness (as not all manifold segments will be hypercubes, and even then some edge points may not match), we average that fraction with the smaller fraction that would need for a $d + 1$ dimensional hypercube, or $2^{−d−1}$, for our resulting $2^{−d−1} + 2^{−d−2}$. In general, we found that this choice was not critical in the noiseless data case, as matches were common for separated components with the same true assignments and rare for others, but it did help in cases with many intersecting components.

**ConstructHierarchy (Algorithm 7)**

- **neighbor_lengthscale_mult** - the threshold for deciding whether a higher-dimensional component “encloses” a lower-dimensional component, expressed as a ratio of (1) the average distance from lower-dimensional component points to their nearest neighbors in the higher-dimensional component (inter-component distance), to (2) the average distance of points in the higher-dimensional component to their nearest neighbors in that same component (intra-component distance). We used 10, which we found was robust for our benchmarks, though it may need to be increased if ground-truth components are higher-dimensional than those in our benchmarks.
Figure A.6. A fuller version of main paper Fig. 5 showing COFHA ablations on all datasets. Hierarchical disentanglement tends to be low for flat AEs (Flat), better with ground-truth hierarchy \( H \) (Hier \( H \)), and even better after adding supervision for ground-truth assignments \( A \) (\( H + A \)). Adding a FactorVAE-style marginal TC penalty (\( H + A + TC(Z) \)) sometimes helps disentanglement, but making that TC penalty conditional (\( H + A + TC(Z|on) \)), i.e. COFHA, tends to help more, bringing it close to the near-optimal disentanglement of a hierarchical model whose latent representation is fully supervised (\( H + A + Z \)). Partial exceptions include the hardest three datasets (Spaceshapes and depth-3 compound Chopsticks), where disentanglement is not consistently near 1; this may be due to non-identifiability or adversarial optimization difficulties.

Figure A.7. Varying disentanglement penalty hyperparameters for baseline algorithms (TCVAE, FactorVAE, CascadeVAE, and JointVAE). Markers indicate mean \( R^2 \) over 5 trials, with standard deviation errorbars. In contrast to COFHA (mean performance in red, with standard deviation in pink), no setting produces near-optimal disentanglement, and disentanglement often decreases with increasing disentanglement penalty strength.
Figure A.8. Pairwise histograms of ground-truth vs. learned variables for a flat autoencoder (top left), $\beta$-TCVAE (top right), and the best-performing run of COFHAE (bottom) on Spaceshapes. Histograms are conditioned on both variables being active, and dimension-wise components of the $R^4_c$ score are shown on the right. $\beta$-TCVAE does a markedly better job disentangling certain components than the flat autoencoder, but in this case, COFHAE is able to fully disentangle the ground-truth by modeling the discrete hierarchical structure. See Fig. A.9 for a hierarchical latent traversal, or https://hreps.s3.amazonaws.com/viz/index.html?dataset=spaceshapes&model=cofhae for an interactive visualization.
Figure A.9. Hierarchical latent traversal plot for the Spaceshapes COFHAE model shown in Fig. A.8c. Individual latent traversals show the effects of linearly sweeping each active dimension from its 1st to 99th percentile value (center column shows the same input with intermediate values for all active dimensions). Consistent with Fig. A.8c, the model is not perfectly disentangled, though primary correspondences are clear: star shine is modeled by \( Z_5 \), moon phase is modeled by \( Z_8 \), ship angle is modeled by \( Z_{10} \), ship jetlen is modeled by \( Z_{12} \), and \((x, y)\) are modeled by \((Z_3, Z_4)\), \((Z_6, Z_7)\), and \((Z_{11}, Z_9)\) respectively for each shape. See also an interactive visualization.

Figure A.10. Three different potential hierarchies for Spaceshapes which all have the same structure of variable groups and dimensionalities, but with different distributions of continuous variables across groups. The ambiguity in this case is that the continuous variable that modifies each shape (phase, shine, angle) could either be a child of the corresponding shape category, or be “merged up” and combined into a single top-level continuous variable that controls the shape in different ways based on the category. Alternatively, the location variables \( x \) and \( y \) could instead be “pushed down” from the top level and duplicated across each shape category. In each of these cases, the learned representation still arguably disentangles the ground-truth factors—in the sense that for any fixed categorical assignment, there is still 1:1 correspondence between all learned and ground-truth continuous factors. We deliberately design our \( R^4_c \) and \( H \)-error metrics in §6 to be invariant to these transformations, leaving this specific disambiguation to future work.

Figure A.11. MIMOSA-learned initial encoding (left), components (middle), and hierarchy (right) for Spaceshapes. Initial points are in 7 dimensions and projected to 3D for plotting. Three identified components are 3D and one is 4D. Analogue of Fig. 3 in the main text.
Figure A.12. MIMOSA-learned initial encoding (left), 2D and 1D components (middle), and hierarchy (right) for depth-2 Chopsticks varying the slope. Analogue of Fig. 3 in the main text.

Figure A.13. MIMOSA-learned initial encoding (left), 2D, 1D, and 3D components (middle), and hierarchy (right) for depth-3 Chopsticks varying the slope. Initial points are in 4 dimensions and projected to 3D for plotting. Analogue of Fig. 3 in the main text.

Figure A.14. MIMOSA-learned initial encoding (left), 2D and 4D components (middle), and hierarchy (right) for depth-2 Chopsticks varying both slope and intercept. Initial points are in 5 dimensions and projected to 3D for plotting. Analogue of Fig. 3 in the main text.

Figure A.15. MIMOSA-learned initial encoding (left), 2D, 4D, and 6D components (middle), and hierarchy (right) for depth-2 Chopsticks varying both slope and intercept. Initial points are in 7 dimensions and projected to 3D for plotting. Analogue of Fig. 3 in the main text.

Figure A.16. MIMOSA-learned initial encoding (left), 1D-3D components (middle), and hierarchy (right) for depth-3 Chopsticks varying either slope or intercept. Note that the learned hierarchy is not quite correct (two nodes at the deepest level are missing). Initial points are in 5 dimensions and projected to 3D. Analogue of Fig. 3.
Figure A.17. Pairwise histograms of ground-truth vs. learned variables for COFHAE on the most complicated hierarchical benchmark (Chopsticks at a recursion depth of 3 varying either slope or intercept). Histograms are conditioned on both variables being active, and dimension-wise components of the $R^2_c$ score are shown on the right. Despite the depth of the hierarchy, COFHAE representations model it fairly well.