EUROPEAN ORGANIZATION FOR NUCLEAR RESEARCH (CERN)



Wasserstein normalized autoencoder for anomaly detection

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Abstract

A novel anomaly detection algorithm is presented. The Wasserstein normalized autoencoder (WNAE) is a normalized probabilistic model that minimizes the Wasserstein distance between the learned probability distribution—a Boltzmann distribution where the energy is the reconstruction error of the autoencoder—and the distribution of the training data. This algorithm has been developed and applied to the identification of semivisible jets—conical sprays of visible standard model particles and invisible dark matter states—with the CMS experiment at the CERN LHC. Trained on jets of particles from simulated standard model processes, the WNAE is shown to learn the probability distribution of the input data in a fully unsupervised fashion, such that it effectively identifies new physics jets as anomalies. The model consistently demonstrates stable, convergent training and achieves strong classification performance across a wide range of signals, improving upon standard normalized autoencoders, while remaining agnostic to the signal. The WNAE directly tackles the problem of outlier reconstruction, a common failure mode of autoencoders in anomaly detection tasks.

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1 Introduction

Unsupervised machine-learning algorithms have proven to be powerful tools in the search for new physics at the LHC [1, 2]. They can effectively separate events from known standard model (SM) processes (background) from potential signal events arising from interactions beyond the SM (BSM), without relying on the simulation of specific signal hypotheses. Autoencoders (AEs) are frequently used for unsupervised learning tasks. One of the main challenges when using AE-based unsupervised machine-learning algorithms for anomaly detection is preventing the reconstruction of outliers—examples that are not part of the training data. A well-known example of outlier reconstruction called complexity bias [3] occurs when outliers are less complex than the training data. In this paper, we introduce the Wasserstein normalized autoencoder (WNAE), an evolution of the normalized autoencoder (NAE) algorithm [4], to solve these challenges. The identification of semivisible jets (SVJs) [5] in the CMS experiment [6] is used as a case study to demonstrate the effectiveness of the WNAE in separating signal from background events.

The CMS apparatus [6, 7] is a multipurpose, nearly hermetic detector, designed to trigger on [8–10] and identify electrons, muons, photons, and (charged and neutral) hadrons [11–13]. A global "particle-flow" algorithm [14] aims to reconstruct all individual particles in an event, combining information provided by the all-silicon inner tracker and by the crystal electromagnetic and brass-scintillator hadron calorimeters, operating inside a 3.8 T superconducting solenoid, with data from the gas-ionization muon detectors embedded in the flux-return yoke outside the solenoid. The reconstructed particles are used to build jets [15]—conical sprays of particles—and missing transverse momentum [16]—the imbalance of visible momentum in the transverse plane. Events of interest are selected using a two-tiered trigger system. The first level, composed of custom hardware processors, uses information from the calorimeters and muon detectors to select events at a rate of around 100 kHz within a fixed latency of 4 μ s [8]. The second level, known as the high-level trigger, consists of a farm of processors running a version of the full event reconstruction software optimized for fast processing, and reduces the event rate to a few kHz before data storage [9, 10].

The paper is organized as follows. Section 2 discusses the use and limitations of AEs as anomaly detection tools, focusing on the outlier reconstruction problem, with a brief review of the NAE algorithm. In Section 3, the AE and NAE algorithms are applied to the search for SVJs, and we introduce a metric based on the Wasserstein distance [17, 18] to quantify the degree of outlier reconstruction without relying on the signal in any form during training. Section 4 presents the major advance in this paper: the use of this metric as the direct objective for the autoencoder training to build a more robust anomaly detection algorithm, the WNAE. The advantages of the WNAE over the NAE are discussed. Finally, Section 5 discusses the implications of the findings and possible future directions.

2 Autoencoder-based anomaly detection

2.1 Autoencoders

Autoencoders [19] are a class of neural networks that are intended to learn a compressed representation of the input data. This is usually achieved by mapping the input feature space $\mathcal{X} \subset \mathbb{R}^n$ to a lower-dimensional latent space $\mathcal{Z} \subset \mathbb{R}^m$ via an encoder network $f_e : \mathbb{R}^n \mapsto \mathbb{R}^m$ (m < n) and mapping this latent space to an output space with the same dimension as the input space via a decoder network $f_d : \mathbb{R}^m \mapsto \mathbb{R}^n$. Autoencoders are typically trained by minimizing a loss function that penalizes the difference between the input and output, referred to as the

reconstruction error of the AE. The latter is denoted l_{θ} in the following, where θ represents the parameters of the AE network.

Crucially, when AEs are used for anomaly detection, the network is only exposed to non-anomalous (background) examples during training. If the network is able to learn a compressed representation of these examples, it can reconstruct them with low error. However, since the anomalous (signal) examples are not part of the training set, the AE is expected not to reconstruct them well, resulting in a larger average reconstruction error.

While this reasoning holds for many practical applications of AEs for anomaly detection, it does not follow in general that achieving low reconstruction error on the background examples is sufficient for an AE to effectively identify anomalies, as discussed below. This shortcoming is addressed by the WNAE algorithm introduced in this paper.

2.2 The problem of outlier reconstruction

Given a data distribution p_{data} , an example datum $x \in \mathcal{X}$ is an outlier if it lies in a region with probability density lower than a given threshold τ : $p_{\text{data}}(x) \leq \tau$. In the extreme case where $\tau = 0$, the outlier lies outside of the support of the data distribution.

Outlier reconstruction occurs when an AE, while learning to reconstruct the training examples, also learns to reconstruct outlier examples. More formally: let \mathcal{B} and \mathcal{S} be the supports of the background and signal distributions in the input feature space \mathcal{X} , respectively. The AE is trained to achieve minimum reconstruction error on background samples $x_b \in \mathcal{B}$. Let \mathcal{E} be the region where the AE achieves reconstruction error below some threshold. Since there is no constraint on the reconstruction error of signal samples $x_s \in \mathcal{S}$, there can be a sizable overlap $\mathcal{S} \cap \mathcal{E}$. This overlap can in general be at least partly disjoint from the training data: $(\mathcal{S} \cap \mathcal{E}) - \mathcal{B} \neq \mathcal{O}$. Examples drawn from this region of phase space are assigned low reconstruction error by the AE, even though they are outside the training data. Thus, these examples and the background both have low reconstruction error, decreasing the performance of the reconstruction error as a discriminator. This is illustrated in Fig. 1.

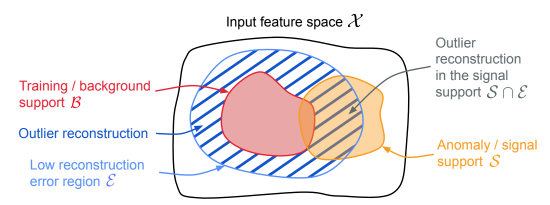


Figure 1: Schematic visualization of the outlier reconstruction failure mode. Signal samples drawn from the hatched area are reconstructed well by the AE, despite not being part of the training set, and thus are not separated from the background. The AE training is assumed to have converged such that the background is reconstructed well.

Ideally, the reconstruction error l_{θ} should be a function of the probability density of the training data: the lower the probability density of the training data, the higher the reconstruction error. Thus, examples x with $p_{\text{data}}(x) \leq \tau$ must have reconstruction error above a threshold τ_r : $l_{\theta}(x) \geq \tau_r$. The reconstruction error itself can accordingly be used as a summary statistic to find

anomalies. The objective of the autoencoder is therefore to learn the distribution of the training data (the "inlier" distribution), rather than simply to minimize the reconstruction error.

2.3 The normalized autoencoder paradigm

The NAE [4] is based on a proposed mechanism to learn the probability distribution of the training data. In analogy with energy-based models [20–22], often used for generative modeling, the NAE is assigned a normalized probability distribution p_{θ} , parametrized by the AE network weights θ . Following the maximum entropy principle [23], the probability distribution is chosen as the Boltzmann distribution from statistical mechanics:

$$p_{\theta}(x) = \frac{1}{\Omega_{\theta}} \exp\left(-E_{\theta}(x)\right),\tag{1}$$

where E_{θ} is the energy of the system and Ω_{θ} is the partition function:

$$\Omega_{\theta} = \int \mathrm{d}x \exp(-E_{\theta}(x)). \tag{2}$$

The energy of the system is defined as the reconstruction error of the AE:

$$E_{\theta}(x) = l_{\theta}(x). \tag{3}$$

The problem is then recast as a maximum likelihood estimation problem, where the goal is to find the weights θ that maximize the likelihood of the training data, with probability distribution denoted p_{data} . As is common, the negative log likelihood is minimized instead:

$$\mathbb{E}_{x \sim p_{\text{data}}}[-\log p_{\theta}(x)] = \mathbb{E}_{x \sim p_{\text{data}}}[E_{\theta}(x)] + \log \Omega_{\theta}. \tag{4}$$

Calculating the partition function Ω_{θ} is in general intractable, but the problem can be circumvented when using gradient descent to find the optimum, since the gradient of the partition function can be calculated:

$$\nabla_{\theta} \log \Omega_{\theta} = \frac{1}{\Omega_{\theta}} \nabla_{\theta} \Omega_{\theta} = \frac{1}{\Omega_{\theta}} \int_{\mathcal{B}} dx \nabla_{\theta} \exp(-E_{\theta}(x))$$
 (5)

$$= \int dx \frac{1}{\Omega_{\theta}} \exp(-E_{\theta}(x)) \nabla_{\theta}(-E_{\theta}(x))$$
 (6)

$$= \int \mathrm{d}x p_{\theta}(x) \nabla_{\theta}(-E_{\theta}(x)) \tag{7}$$

$$= -\mathbb{E}_{x \sim p_{\theta}} [\nabla_{\theta} E_{\theta}(x)], \tag{8}$$

which is the expectation value of the gradient of the AE reconstruction error over the probability distribution p_{θ} . The gradient of the loss can thus be calculated by sampling from the probability distribution p_{θ} . This can be done via a Markov chain Monte Carlo (MCMC) algorithm. The application of the MCMC in the NAE training procedure is discussed in more detail in Section 3.4 and Appendix C. The loss function to be minimized is then:

$$\mathcal{L}_{\text{NAE}} = \mathbb{E}_{x \sim p_{\text{data}}} [E_{\theta}(x)] - \mathbb{E}_{x' \sim p_{\theta}} [E_{\theta}(x')]. \tag{9}$$

Both terms in Eq. (9) are expectation values of the reconstruction error of the AE. The first term is the expectation value over the training data x (the usual AE loss), referred to as the positive energy. The second term is the expectation value over samples x' drawn from the probability

distribution p_{θ} , referred to as the negative energy. The NAE loss is thus the difference between the positive and negative energy, denoted E_{+} and E_{-} , respectively:

$$\mathcal{L}_{\text{NAE}} \equiv E_{+} - E_{-}.\tag{10}$$

In analogy, the training examples are referred to as positive examples, and examples drawn from p_{θ} are referred to as negative examples. In this way, the NAE is not trained just to reconstruct background examples well, but rather to learn the probability distribution of the training data. The NAE is therefore penalized for having low reconstruction error in regions with low probability density of the training data, hence suppressing the reconstruction of examples outside the background support \mathcal{B} . It is worth emphasizing that the whole procedure is still fully unsupervised. The NAE is trained only on background examples, and the signal is not used in any form during training.

2.4 Markov chain Monte Carlo

The key step in the normalized autoencoder paradigm is to estimate the autoencoder probability p_{θ} . This is achieved by running an MCMC algorithm to obtain a set of examples following the p_{θ} distribution. The Markov chain is only needed to train the network and is not used for inference. The Langevin Monte Carlo algorithm is employed in this work. First, a starting set of points $\{x\}_0$ is drawn from an initial distribution, following the procedure laid out in Section 3.4. The negative samples are obtained by repeating a fixed number of steps, each step proceeding as follows:

$$x_{i+1} = x_i + \lambda \nabla_x \log p_\theta(x_i) + \sigma \epsilon_i, \tag{11}$$

where λ is the step size, σ is the noise coefficient, ϵ_i is an independent draw from a normal distribution on \mathbb{R}^n with mean 0 and covariance matrix equal to the $n \times n$ identity matrix, and n is the number of dimensions of the input feature space. The Markov chain has therefore the following dependence on the reconstruction error of the autoencoder:

$$x_{i+1} = x_i - \lambda \nabla_x E_\theta(x_i) + \sigma \epsilon_i. \tag{12}$$

With ρ_i denoting the probability distribution of the sample $\{x\}_i$, choosing:

$$\sigma = \sqrt{2\lambda},\tag{13}$$

and assuming an infinite number of steps, the probability ρ_i converges towards the target probability p_θ : $\lim_{i\to\infty}\rho_i=p_\theta$. In practice, p_θ needs to be estimated at every gradient descent step, once per batch, while training an NAE. For that reason, the number of Markov chain steps cannot be too large, and the probability p_θ can only be approximated. In order to enhance the gradient term in Eq. (11), the temperature T is introduced:

$$x_{i+1} = x_i + \frac{\lambda}{T} \nabla_x \log p_\theta(x_i) + \sigma \epsilon_i.$$
 (14)

Temperatures T < 1 result in an effective increase of the gradient term.

2.5 Related work

There has long been interest in applying autoencoders to model a data distribution, mostly for generative tasks. Variational autoencoders (VAE) [24] make use of the Kullback-Leibler divergence to impose a Gaussian prior in the latent space \mathcal{Z} and the decoder f_d learns to model the data distribution from this Gaussian prior. Adversarial autoencoders (AAE) [25] minimize the

2.5 Related work 5

reconstruction error and enforce a Gaussian prior in the latent space by training the network in an adversarial way, similarly to generative adversarial networks (GANs) [26]. The encoder f_e plays the role of the generator network and a discriminator network estimates the probability that a latent sample arises from the prior distribution rather than being the output of the generator $f_e(x)$, where x is a training datum. In Wasserstein autoencoders (WAE) [27], the loss function is composed of two terms: a divergence between the latent space $\mathcal Z$ and a (Gaussian) prior, and the Wasserstein distance between the training data distribution and that learned by the AE (the output of the decoder f_d). The Wasserstein distance between two probability distributions p_1 and p_2 is defined as:

$$W(p_1, p_2) = \inf_{\gamma \in \Pi(p_1, p_2)} \mathbb{E}_{(x, x') \sim \gamma} [\|x - x'\|], \qquad (15)$$

where $\Pi(p_1, p_2)$ denotes the set of all joint distributions whose marginal distributions are p_1 and p_2 and inf indicates the infimum over this set. This distance measures the cost of morphing p_1 into p_2 . Using the Kantorovich-Rubinstein duality [28], this expression can be written in the dual form as:

$$W(p_1, p_2) = \sup_{f \in \mathcal{F}_{\mathcal{L}}} \mathbb{E}_{x \sim p_1} \left[f(x) \right] - \mathbb{E}_{x \sim p_2} \left[f(x) \right], \tag{16}$$

where the supremum is over all 1-Lipschitz functions $\mathcal{F}_{\mathcal{L}}$. The Wasserstein distance in Eqs. (15) and (16) is typically approximated by relaxing the constraint on the coupling distribution γ or the function f and replacing it with a simpler constraint or regularizing the loss function. In Wasserstein GANs [29], the 1-Lipschitz constraint is approximated by implementing f as a neural network and clipping its weights. In WAEs, the constraint on the coupling distribution γ is relaxed and the loss function is regularized by adding this term to the loss function: $\lambda[D_f(\gamma_1, p_1) + D_f(\gamma_2, p_2)]$, where λ is a positive constant, γ_1 and γ_2 are the marginal distributions of γ , and D_f denotes an f-divergence [27]. An exact solution to the optimal transport problem in Eq. (15) is implemented in Ref. [30], but does not scale well to large data sets. Nevertheless, computing the Wasserstein distance on minibatches of the entire data set is efficient and was found to provide an unbiased estimator of the gradients [31].

As explained in Section 2.2, an ideal outlier detection algorithm must learn the training data distribution. For that reason, and as further highlighted in Section 3.3, the vanilla autoencoder defined in Section 2.1 is not a suitable algorithm for anomaly detection. While the reconstruction error of VAEs can directly be used as an anomaly score, VAEs have been shown to be prone to outlier reconstruction [32, 33]. On the other hand, algorithms based on GANs, often constructed similarly to AAEs, need to be modified to provide an anomaly score. Generative probabilistic novelty detection [34], GANomaly [35], and AnoGAN [36] follow different approaches, but all are challenging to train because of the iterative adversarial procedure [26]. The NAE, described in Section 2.3, is based on a different principle: it relies on an MCMC algorithm to sample data points from the probability distribution learned by the network. Though it is a normalized probabilistic model, its loss function takes the form of a minimax rule, which leads to numerical instability and therefore requires ad-hoc regularization [4, 37]. In addition, as detailed in Section 3.4, the NAE has a failure mode that can cause outlier reconstruction.

Since the first results using AEs to search for new physics at the LHC [1, 2], there has been active research on this topic. The CMS experiment has implemented two realtime anomaly detection algorithms in the level-1 trigger to select new physics signatures: one based on a VAE using global trigger information, and another based on a convolutional AE using calorimeter trigger information [38]. The CMS experiment has recently conducted a search for new physics in dijet resonances with anomalous jet substructure using five different anomaly detection algorithms, two of which are based on AEs [39]. The ATLAS experiment looked for new

physics in the invariant mass distribution of pairs of SM objects consisting of one light-flavor or b jet, and a lepton, a photon, or a second light-flavor or b jet, using an AE to select anomalous data [40]. NAEs were first used in high-energy physics in the context of jet tagging for new physics searches [41].

3 Case studies

3.1 Data set generation

Astronomical observations [42–46] provide compelling evidence for dark matter (DM) in the universe. However, particles that could constitute the DM are not found in the SM of particle physics. If DM particles can be produced in high-energy collisions at the LHC by new physics processes, their existence can be inferred from the signature they leave in detectors. This work considers a specific class of models of new physics, described below.

Hidden valley models [47] are new physics scenarios in which the SM is expanded with a dark sector that has its own particles and forces. The dark sector particles are not charged under any SM gauge group and hence are not directly detectable. Similarly, SM particles are not charged under the dark gauge group. Hidden valley theories may, under certain conditions, result in SVJs as an experimental signature [5], where some of the particles in the final state are not detectable. If the dark sector has a new non-Abelian SU(N) confining interaction, it would exhibit showering and hadronization qualitatively similar to the SM quantum chromodynamics (QCD). In analogy with the SM sector, the new confining force and the new particles charged under it are referred to as dark QCD and dark quarks, respectively. More specifically, we consider a dark QCD interaction $SU(N_c^{\text{dark}})$, where N_c^{dark} denotes the number of dark colors, with a confinement scale $\Lambda_{\rm dark}$ and $N_{\rm f}^{\rm dark}$ flavors of dark quark χ with a mass m_{χ} , which communicates with the SM via a mediator Φ with mass m_{Φ} . In such a model, QCD-like showers occur when $m_{\Phi} \gg \Lambda_{\rm dark}$ and $m_{\chi} \sim \Lambda_{\rm dark}$. The dark quarks therefore form bound states called dark hadrons. The unstable dark hadrons, π_{dark} and ρ_{dark} , decay back to SM particles to produce a visible signature, depending on the symmetries of the theory. The stable dark hadrons, $\pi_{\mathrm{dark}}^{\mathrm{DM}}$ and $\rho_{\rm dark}^{\rm DM}$, are invisible and escape detection, giving rise to missing transverse momentum. The ensemble of stable dark hadrons and the decay products of unstable dark hadrons from the dark QCD shower constitutes an SVJ. The fraction of stable dark hadrons in the dark QCD shower can be parametrized by the invisible fraction r_{inv} :

$$r_{\rm inv} = \left\langle \frac{\text{Number of stable dark hadrons}}{\text{Total number of dark hadrons}} \right\rangle.$$
 (17)

Figure 2 illustrates the production of SVJs. The CMS and ATLAS Collaborations have published searches for resonant production of pairs of SVJs [48, 49]. The production of SVJs via *t*-channel interactions through a bifundamental scalar mediator, charged under both SM QCD and dark QCD, is also possible [50], with one published search from ATLAS [49]. The latter production mode is considered in this paper.

The SVJs may have different radiation patterns than SM jets, which can be exploited to identify them. The details depend on the exact spectrum of the dark hadrons, which in turn depends on $N_{\rm c}^{\rm dark}$, $N_{\rm f}^{\rm dark}$, $\Lambda_{\rm dark}$, the masses m_{χ} , and the nonperturbative physics in the dark sector, all of which are unknown. Therefore, anomaly detection, which does not require a detailed simulation of every target signal hypothesis in this large parameter space, is a powerful complement to traditional, supervised searches for SVJs, such as Ref. [48].

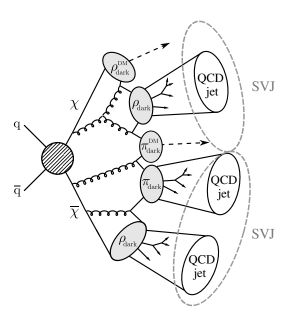


Figure 2: An illustration of collider SVJ production. The dashed black arrows indicate stable, undetectable DM candidate particles. Figure adapted from Ref. [51].

Simulated events are used to train and evaluate the performance of the machine-learning models discussed in this paper. The background considered in this study consists of top-antitop quark production ($t\bar{t}$) with additional jets, where each top quark decays either hadronically or leptonically. Background and SVJ signal events are generated at the parton level with MADGRAPH5_aMC@NLO 2.6.5 [52], final-state partons are hadronized with PYTHIA 8.240 [53], and the interactions of the resulting particles with the CMS detector [6] are simulated using GEANT4 [54]. The NNPDF3.1 parton distribution functions are used for all processes [55]. The hadronization in the dark sector during the signal generation follows the procedure described in Ref. [48], and all dark hadrons have a fixed mass, 20 GeV. Because of the double hadronization, first in the dark sector and then in the SM sector, the substructure of SVJs is expected to be different from that of SM jets. In particular, SVJs are expected to be wide, and therefore jets are reconstructed using the anti- $k_{\rm T}$ algorithm [56, 57] with radius parameter R=0.8. Each jet is represented by the following features, describing its substructure:

- the major and minor axes [58], which characterize the elliptical shape of the jet in the η - ϕ plane (where η is the pseudorapidity and ϕ is the azimuthal angle);
- the first energy flow polynomial (EFP1) [59], a multiparticle energy correlator that directly results from infrared and collinear safety;
- the $C_2^{(0.5)}$ energy correlation function [60], a ratio of two ratios of energy correlation functions that can be used to determine if a jet has two hard subjets;
- the transverse momentum dispersion p_T^D [58], which describes the spread of transverse momenta of the jet constituents;
- the soft-drop mass [61], the mass of the jet after soft wide-angle radiation has been removed; and
- the 2- and 3-subjettiness τ_2 and τ_3 [62], describing the compatibility of the jet with a two- or three-prong structure, respectively;

for a total of n=8 inputs to the AE. These input features are chosen to exploit potential differences between SM jets and BSM jets in shape (major and minor axes), energy distribution

 $(p_{\rm T}^{\rm D})$ and EFP1), prong structure $(C_2^{(0.5)}, \tau_2, \tau_3)$, and mass of the high-momentum core of the jet (softdrop mass). These features represent a space of physical variables that facilitates the classification task by providing some separation between signal and background, even though a single boundary in this space, unbiased across all possible signal models, may be difficult to define. For each event, the two jets with highest transverse momenta are used for training and evaluation. In addition, among the two selected jets, only jets identified via generator-level information as originating from dark-sector particles are considered as anomalous signal jets when evaluating the network performance. The inputs are preprocessed to follow a normal distribution using quantile normalization, as implemented in the SCIKIT-LEARN package [63].

3.2 Training setup

The architecture is a fully connected network with five hidden layers. All layers have ten nodes, except for the third one, which, as the bottleneck, has six. The performance of the different models discussed in this paper was found to have little dependence on either the number of layers or the number of nodes per layer; the listed choices are found to produce a sufficiently expressive network for this anomaly detection task. The reconstruction of the AE is parametrized by the network weights θ . The reconstruction error l_{θ} for an example x is defined as the mean squared error between the features x_i of x and the features $\hat{x}_{\theta,i}$ of the reconstructed output:

$$l_{\theta}(x) = \frac{1}{n} \sum_{i=1}^{n} (x_i - \hat{x}_{\theta,i})^2.$$
 (18)

The loss function of the AE is the reconstruction error averaged over the batch:

$$\mathcal{L}_{AE} = \frac{1}{N} \sum_{i=1}^{N} l_{\theta}(x^{(j)}), \tag{19}$$

where $x^{(j)}$ denotes the example in position j in the batch and N is the number of training examples in the batch. The background data set is split into three independent parts: a training set on which the loss is minimized (70%), a validation set used to monitor the training (15%), and a test set used to evaluate the performance of the network (15%). The background data set has a total of 47000 jets and the number of trainable parameters is 534. The AE is trained until the loss function evaluated on the validation set ceases to improve. The AE is then tested using multiple SVJ signal models, as well as the background test sample.

3.3 Autoencoder training

The issue of outlier reconstruction is found to be especially impactful in the case of the SVJ search. While previous studies have shown that AEs are an effective tool for the task of discriminating between the signal and the SM background from light-flavor jets [64], the performance is found to degrade significantly when jets originating from t quarks in the tt̄ process are included. This is shown by training an AE solely on the tt̄ background. The cause is understood to be outlier reconstruction, based on the nature of tt̄ events. Boosted jets from high-momentum top quarks, which contain multiple prongs and genuine missing transverse momentum from neutrinos, are the closest SM analog to SVJs. These events span multiple topologies, as they can also include boosted W boson jets, light-flavor quark jets, and gluon jets from initial- or final-state radiation.

The receiver operating characteristic curve compares the true- and false-positive rates for the AE reconstruction error used as a discriminator, and the area under the curve (AUC) is used

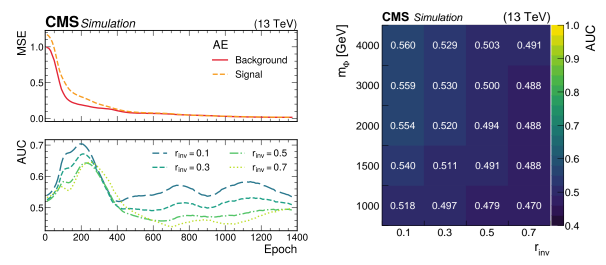


Figure 3: Left: the reconstruction error (upper panel) and the AUC scores (lower panel) for the AE trained on the $t\bar{t}$ background, evaluated during each training epoch on $t\bar{t}$ background jets and signal models with $m_{\Phi}=2000\,\text{GeV}$ and $r_{\text{inv}}=0.3$ (upper) or $r_{\text{inv}}=0.1,0.3,0.5,0.7$ (lower). Right: The AUC scores for the same AE, evaluated for the epoch with the minimal background reconstruction error, for the classification of several SVJ signal hypotheses against the $t\bar{t}$ background. The AUC scores are close to 0.5, indicating that the AE is unable to discriminate between the SVJ signal and the $t\bar{t}$ background.

to quantify the performance of this discriminator. As shown in Fig. 3, when achieving minimum reconstruction error on the $t\bar{t}$ background, the AE is unable to discriminate between the SVJ signal and the $t\bar{t}$ background, and the AUC score is close to 0.5. In order to check that this behavior does not occur because the bottleneck was too large, the same experiment was repeated with a bottleneck of only five, four, three, or two nodes. The results were found to be consistent with a bottleneck size of six, indicating that the issue does not arise from insufficient compression of the data.

The AUC scores evolve in nontrivial ways throughout the training. As shown in Fig. 3, the AUC score improves during the first stages of the training, indicating that the AE is performing as desired. However, after further training, the AUC score starts to degrade, and eventually converges to a value close to 0.5. This is a clear indication that the requirement to minimize the reconstruction error on the $t\bar{t}$ background is too weak to achieve good discrimination; it does not prevent the AE from also reconstructing signal events.

3.4 Normalized autoencoder training

The most delicate part of the NAE training is the tuning of the MCMC. A suboptimal MCMC would not accurately sample the distribution p_{θ} , and in turn, the NAE would not be able to learn the training data probability distribution p_{data} . In theory, the convergence of an MCMC algorithm does not depend on the random initial set of points from which the Markov chain starts. However, there are two motivations for using a nonrandom initial set of points. First, the Markov chain is not infinitely long, and so there is no guarantee of convergence starting from any initial distribution. Second, from one batch to another, the neural network weights and p_{θ} both change only slightly, so a good initialization for batch k usually also works for batch k+1. In addition, tuning the hyperparameters of the Markov chain is a complex task since the p_{θ} distribution is unknown. The procedure to initialize the MCMC and tune its associated hyperparameters, which accounts for these considerations, is detailed in Appendix C.

After tuning the MCMC algorithm, the evolution of the positive and negative energies of an NAE with the loss function defined in Eq. (10) is presented in Fig. 4. It shows two failure modes: negative loss values, indicating the negative energy is higher than positive energy; and divergence of both energies.

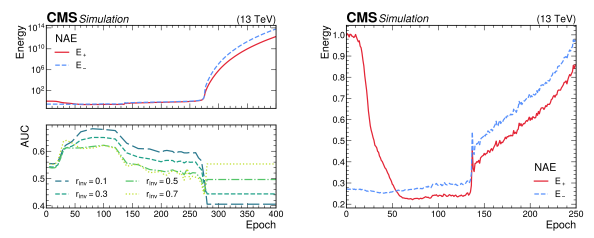


Figure 4: Left: NAE training showing the divergence of the loss function, in terms of positive and negative energy (upper panel), and the AUC for several signal hypotheses with fixed $m_{\Phi} = 2000\,\text{GeV}$ and varying r_{inv} values (lower panel). Right: the positive and negative energies from the upper panel of the left plot, shown for epoch < 250—before the divergence—and 0.18 < energy < 1.4 on a linear scale, to illustrate their differences.

Negative values of the energy difference occur when the network did not learn the background probability distribution, such that p_{data} is higher than p_{θ} in regions with low reconstruction error. This is a severe failure mode, illustrating that energy difference minimization is not sufficient to guarantee the intended behavior.

The divergence of the negative energy was also reported in previous work [4, 41]. As a consequence of the form of the loss function, after the positive energy has reached a minimal value, further minimization of the loss is achieved by increasing the negative energy, effectively leading to a divergence of both energy terms and the loss itself, as shown in Fig. 4. The distributions of a selected input feature τ_3 for the positive, negative, and signal samples before and after the start of the divergence are provided in Fig. 5. The large reconstruction error after the divergence leads to very large gradient values and thus large steps in the MCMC. This results in the negative samples reaching the boundaries of the sampling phase space and no longer being representative of p_{θ} .

A simple modification of the NAE loss can mitigate these issues:

$$\mathcal{L} = \log \cosh(E_+ - E_-). \tag{20}$$

Because the log cosh function is positive definite, this loss removes the incentive to favor negative values of the energy difference and leads to a more stable training procedure. The evolution of the energies, loss value, and AUC score for a training with this loss function are shown in Fig. 6. Their behavior in this figure can be qualitatively understood as follows. There is a first stage in which the energy difference is minimized, giving the largest improvement in AUC. Once the difference between positive and negative energies is approximately zero, there is a second stage in which the network fine-tunes its knowledge of p_{θ} , further increasing the AUC score until it reaches a plateau around epoch 4000. There is subsequently a third stage

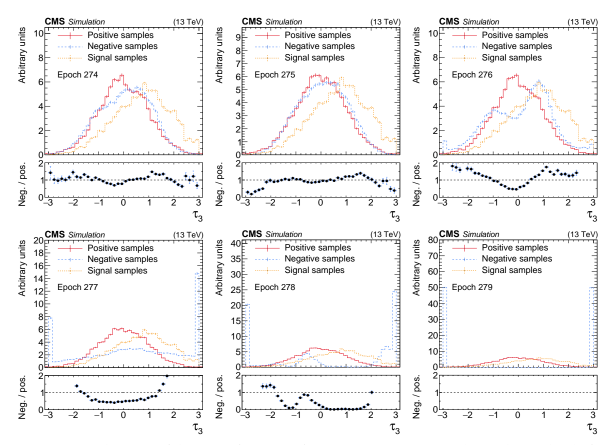


Figure 5: Distributions of the input feature τ_3 for positive, negative, and signal samples, before (epochs 274) and after (epochs 275–279) the start of the divergence of the NAE loss. The signal distributions are overlaid for illustration; signal samples are not used during the training. All distributions are normalized such that their integral is 100.

in which a new form of mode collapse is observed: the energy difference is still approximately zero and the positive energy is further minimized, but the AUC score begins to decrease. Crucially, it is not possible to avoid this degradation by relying solely on the positive and negative energies. On the other hand, making use of the AUC score as a stopping condition for the training would be in contradiction with the unsupervised nature of the NAE. The evolution of the Wasserstein distance, also shown in this figure, is discussed in Section 4.

This collapse arises as a form of overtraining of the network, leading to outlier reconstruction. While keeping the reconstruction error low, the network eventually achieves low reconstruction in a region $\mathcal E$ that contains, but is larger than, the support of the training data $\mathcal B$, while keeping $\mathbb E_{x\sim\mathcal E}[E(x)]$ comparable to $\mathbb E_{x\sim\mathcal B}[E(x)]$. As the overlap of $\mathcal E$ with the support of the signal $\mathcal S$ becomes larger, the AUC score decreases. This behavior is shown schematically in Fig. 7. Before the mode collapse, $\mathcal E$ and $\mathcal B$ overlap, while $\mathcal E$ and $\mathcal S$ do not. After the mode collapse, $\mathcal E$ expands and can partially include $\mathcal S$, reducing the difference in reconstruction error between background and signal and correspondingly lowering the AUC score. In both cases, the difference between the positive and the negative energies is zero.

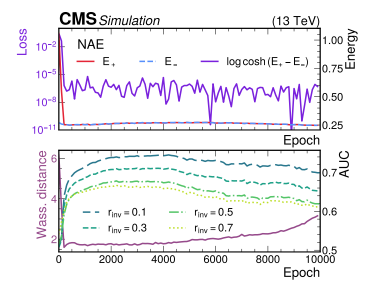


Figure 6: As a function of epoch during the training of an NAE with the loss function from Eq. (20): the positive and negative energies and the value of the loss function (upper panel); the Wasserstein distance between negative and positive samples and the AUC for several signal hypotheses with fixed $m_{\Phi} = 2000\,\text{GeV}$ and varying r_{inv} (lower panel).

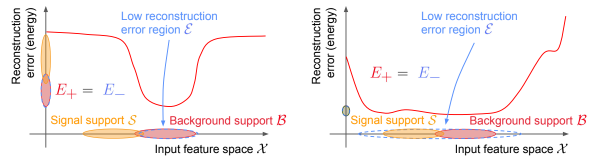


Figure 7: Schematic representation of the mode collapse when using the loss function described in Eq. (20). The energy landscape is shown before (left) and after (right) the mode collapse. On the right, the reconstruction errors for the signal and background supports are completely overlapping on the vertical axis. The symbols E_+ and E_- denote the positive and negative energies, respectively.

4 The Wasserstein normalized autoencoder

4.1 The Wasserstein distance as a measure of outlier reconstruction

The failure mode described at the end of Section 3.4 is a consequence of the fact that the energy difference is unable to distinguish the two cases shown in Fig. 7. In fact, no stopping condition based solely on combinations of positive and negative energies was found to be a good indicator of the start of this mode collapse. In other words, without explicitly monitoring the AUC during training, there is no clear stopping condition; choosing a stopping condition based on the AUC, however, violates the principle of unsupervised learning. In order to overcome this problem, a different metric is used: the Wasserstein distance W (also known as the earth mover's distance) from Eq. (15) between the probability distribution of the training data $p_{\rm data}$ and the probability distribution learned by the network p_{θ} , in analogy to what was applied successfully to generative adversarial networks [29]:

$$W(p_{\text{data}}, p_{\theta}) = \inf_{\gamma \in \Pi(p_{\text{data}}, p_{\theta})} \mathbb{E}_{(x, x') \sim \gamma} \left[\|x - x'\| \right]. \tag{21}$$

The Wasserstein distance is found to be a good indicator of the start of the mode collapse, as seen in Fig. 6 around epoch 4000: the Wasserstein distance increases significantly when this collapse occurs and the AUC decreases. The kink in the Wasserstein distance, quantified as the point in which its derivative changes by more than a given threshold, is then identified as a stopping condition for the training of the NAE. It is worth stressing that the Wasserstein distance, calculated from the positive and negative samples, is completely agnostic to the signal, and yet it is found to be closely correlated with the performance of the NAE as an outlier detector. With this prescription, the NAE is able to successfully learn the tt distribution and effectively detect the signal, as shown in Fig. 8.

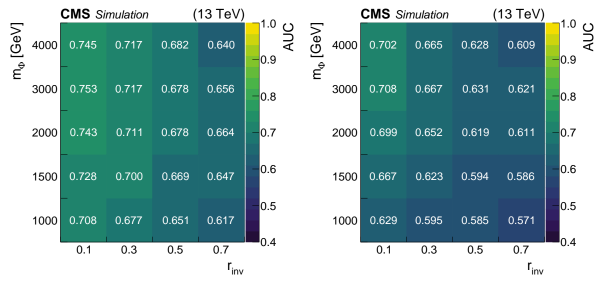


Figure 8: Left: the AUC scores for an NAE trained on the tt background and tested against a grid of possible SVJ signals, before the increase of the Wasserstein distance (at epoch 3000). Right: the AUC scores for the same NAE after the increase in Wasserstein distance (at epoch 10000).

4.2 The Wasserstein distance as a loss function

Having shown that the Wasserstein distance is a good indicator of the performance of the NAE as an outlier detection algorithm, it is natural to use it directly as a loss function. In this case, for

an anomaly-detecting autoencoder, the dependence of the Wasserstein distance on the network weights θ is nontrivial and requires a specific setup to ensure gradients can propagate. The procedure to obtain a differentiable Wasserstein distance for use as a loss function is detailed in Appendix A. The resulting model is called the Wasserstein normalized autoencoder (WNAE), with source code publicly available online [65]. The flowchart for training the WNAE is shown in Fig. 9.

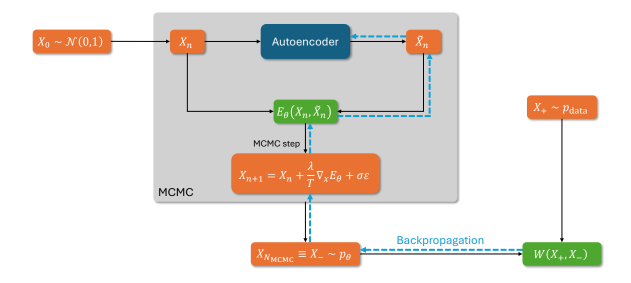


Figure 9: Flowchart of the Wasserstein normalized autoencoder training. The negative examples are generated via MCMC using the energy function of the model, which is the Boltzmann-distributed reconstruction error. The energy function is computed from random input feature values X_n and the corresponding reconstructed feature values \widetilde{X}_n , obtained by passing the inputs through the autoencoder. The positive examples are compared to the negative examples through the Wasserstein distance. The gradients are backpropagated through the entire MCMC chain.

4.3 Training the Wasserstein normalized autoencoder

The training is found to qualitatively proceed in two phases: first, a coarse adjustment in which the Wasserstein distance decreases sharply while the model adapts p_{θ} to produce negative samples that resemble physical examples, and second, a fine-tuning phase in which the model learns the specifics of the training sample. The use of a learning rate scheduler was found to decrease the training time while preserving the stability of the training. In the case of a constant learning rate, if the learning rate is too small, the time needed for the first phase increases significantly, while if it is too large, the training becomes unstable in the second phase. Technical details about the learning rate scheduler can be found in Appendix B.

The computational bottleneck for training the WNAE is the calculation of the Wasserstein distance between the positive and negative samples, which takes approximately 90% of the computation time for each training batch. The second most impactful operation is the backpropagation through the MCMC chain, which takes approximately 7% of the computation time. The time needed for the calculation of the Wasserstein distance heavily depends on the batch size, which was set to 4096 to allow for a robust estimation of the distance between positive and negative samples. Lowering the batch size reduces the computation time at the expense of decreasing the stability of the loss function. Depending on the batch size, the training of

the WNAE can thus be significantly slower than that of the NAE. However, the Wasserstein distance is found to be a much more reliable and stable stopping condition. All of these computational considerations only impact training, not inference.

4.4 Performance of the Wasserstein normalized autoencoder

The WNAE is found to be significantly more stable in training, not showing any of the forms of divergence described previously. It is able to achieve strong anomaly detection performance while using the minimum of the validation loss, the Wasserstein distance evaluated on the validation set, as a clear stopping condition. Figure 10 shows the evolution of the Wasserstein distance and the AUC score for several SVJ signal hypotheses. The minimum of the Wasserstein distance corresponds to the maximum discrimination power. Figure 11 shows the AUC score evaluated for a grid of SVJ signals, showing good discrimination power across the parameter space, on par with or better than any of the NAE models trained above. The WNAE is thus found to be a more reliable, stable, and effective tool for anomaly detection in the context of the SVJ search. In addition, this algorithm can achieve the same performance with all hidden layers having higher dimension than the input feature space. The requirement to minimize the Wasserstein distance controls the behavior of the network without separately requiring an architecture with a lower-dimensional bottleneck.

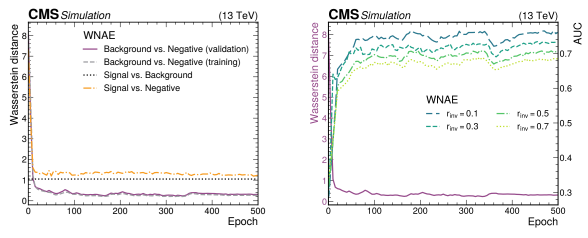


Figure 10: Left: the Wasserstein distance between pairs of the positive, negative, and signal samples during the WNAE training. Right: the Wasserstein distance between negative and positive samples and the AUC scores from the same WNAE for several signal hypotheses with fixed $m_{\Phi} = 2000\,\text{GeV}$ and varying r_{inv} .

The fact that the WNAE behaves as desired can be seen in Figs. 12 and 13, where the distributions of the input features for the positive, negative, and signal samples are shown at the start and end of the training. Starting from an essentially multivariate distribution, induced by the random weight initialization of the network, the WNAE is able to learn the distribution of the training data, as evidenced by the distribution of negative samples eventually converging toward that of the positive samples.

4.5 Complexity bias in the Wasserstein normalized autoencoder

A common weakness of plain AEs is the so-called complexity bias, in which the AE tends to be able to discriminate effectively only against anomalous examples whose distribution in input space is more complex than that of the training data. This can be demonstrated by inverting the background and signal samples and observing that, in this case, an AE trained on SM jets can

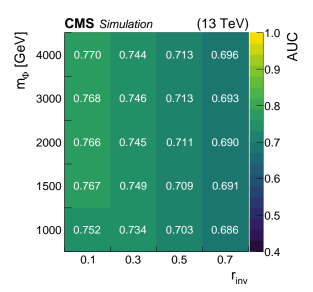


Figure 11: The AUC scores for a WNAE trained on the tt background and tested on a grid of possible SVJ signal models.

tag SVJs as anomalous, but the reverse is not true. Because the WNAE learns the probability distribution of the training data in input space, this issue is mitigated, as shown in Fig. 14. The high AUC at the start of the training is caused by the random uniform initialization of the network weights and the different correlations between the input features of the signal and background jets. The AUC then decreases as the network learns a distribution closer to the physical input data, before increasing again as the network starts to more precisely learn the distribution of SVJ examples and thus becomes able to tag SM jets as anomalous.

5 Conclusions

Anomaly detection using autoencoders (AEs) relies on learning a reconstruction function that gives high reconstruction error to phase space regions with low probability density, such that they can be identified as anomalous. However, standard AEs are prone to learn to reconstruct outliers because they are free to minimize the reconstruction error outside the training phase space. In addition, they may exhibit complexity bias, learning to identify examples as anomalous only if their feature distribution is more complex than the training data. The normalized autoencoder (NAE) paradigm promotes the AE reconstruction error to an energy function in the framework of energy-based models, in order to define a normalized probabilistic model. This is achieved by minimizing the negative log-likelihood of the training data from the energy-based model probability. In practice, this method presents a number of failure modes, such as divergence of the loss function and phase space degeneracy, leading to low reconstruction error for phase space regions distinct from the training data.

The Wasserstein normalized autoencoder (WNAE), an improvement over the NAE, is introduced to solve these failure modes. This is achieved by directly minimizing the Wasserstein distance between the probability distribution of the training data and the Boltzmann distribution of the energy function of the model. This Wasserstein distance is found to be highly correlated with signal identification performance while still being fully signal-agnostic, preserving the unsupervised nature of the approach. The performance is studied in the context of a search for new physics with the CMS experiment, using top-antitop quark production as the standard model background and nonresonant semivisible jet production from a strongly

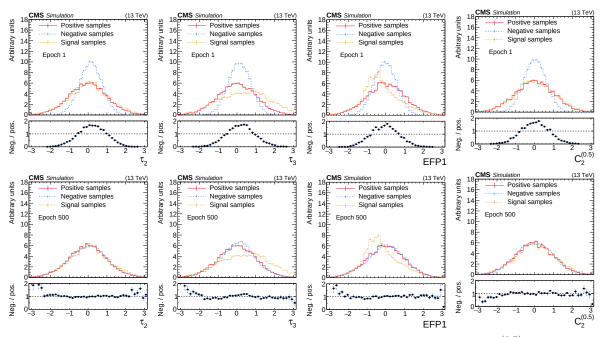


Figure 12: The distributions of half of the input variables, τ_2 , τ_3 , EFP1, and $C_2^{(0.5)}$, for the positive, negative, and signal samples, at the start (upper) and at the end (lower) of the WNAE training. The signal distributions are overlaid for illustration; signal samples are not used during the training. All distributions are normalized such that their integral is 100.

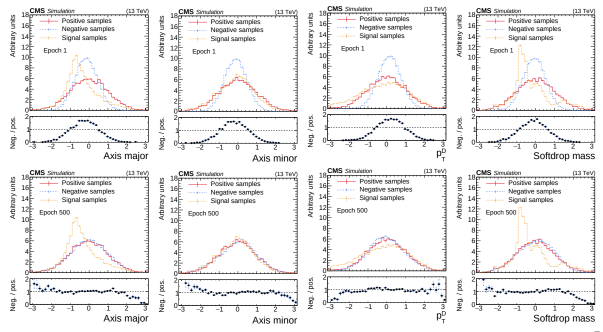


Figure 13: The distributions of the other half of the input variables, axis major, axis minor, $p_{\rm T}^{\rm D}$, and softdrop mass, for the positive, negative, and signal samples, at the start (upper) and at the end (lower) of the WNAE training. The signal distributions are overlaid for illustration; signal samples are not used during the training. All distributions are normalized such that their integral is 100.

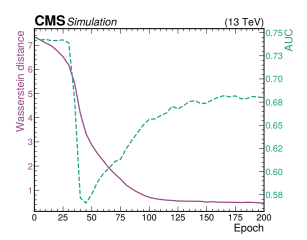


Figure 14: The Wasserstein distance between the positive and negative samples and the AUC score during the training of a WNAE on an SVJ signal ($m_{\Phi} = 2000 \,\text{GeV}$, $r_{\text{inv}} = 0.3$), with the $t\bar{t}$ background used for testing.

coupled dark sector as the proposed signal model. The classification of the signal events as outliers by the WNAE is shown to be on par with or better than that of the NAE. Further, the WNAE approach is found to mitigate complexity bias, as it can effectively identify top quark jets as anomalous when trained on semivisible jet signal events.

Though simulated samples were used to develop the WNAE, in practice it may be preferable to use observed data directly for training, in order to limit biases arising from differences between simulation and observation. In this case, the training data may contain anomalies, which would reduce the anomaly detection performance of the WNAE. The WNAE can be straightforwardly trained using observed data from a control region with no anomalous examples, if such a region can be defined and follows the same probability distribution as the observed data where the WNAE will be applied. When no assumption at all can be made about the nature of the anomalies, such as the case of triggering at a high-energy physics experiment, alternative solutions may exist. The WNAE associates low probability density regions with high reconstruction error; because anomalies necessarily have low probability density, they still tend to have relatively high reconstruction error even when included in the training data set. Therefore, the training data set can be iteratively refined by selecting a given fraction of examples with the lowest reconstruction error, in order to reduce the proportion of anomalous data. This would result in a self-supervised training for the WNAE. We leave the development of such a procedure for future work.

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References 19

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A Gradient dependence between MCMC and AE

During the MCMC process, the gradient with respect to the feature space needs to be calculated to compute each step. At the same time, the dependency on the weights of the neural network must be retained in order to compute the gradient with respect to the weights for backpropagation.

For generative applications, a model g_{θ} , parametrized by weights θ , may be trained to minimize the Wasserstein distance W between the training examples x that it is trying to replicate and the samples $g_{\theta}(z)$ that it generates from random noise z with prior distribution p_z :

$$\mathcal{L}_{\text{generative}} = \inf_{\gamma \in \Pi(p_{\text{data}}, p_z)} \mathbb{E}_{(x, z) \sim \gamma} \left[\| x - g_{\theta}(z) \| \right]. \tag{22}$$

In this case, the Wasserstein distance depends explicitly on the network weights θ via $g_{\theta}(z)$.

However, this is not the case for an AE with the loss function defined in Eq. (21), since neither the positive nor the negative examples pass through the network explicitly. The dependency is in the generation of the negative examples, which is done via the MCMC described in Section 2.3. Equation (14) shows that the dependency on the network weights is in the gradient of the energy function with respect to the input features. As such, in order to use the Wasserstein distance as a loss function, the dependency of each step of the MCMC on the weights of the network must be retained:

$$\nabla_{\theta} x_{i+1,\theta} = \nabla_{\theta} x_{i,\theta} - \lambda \nabla_{\theta} \nabla_{x} E_{\theta}(x_{i,\theta}). \tag{23}$$

The differentiable implementation of the Wasserstein distance metric is provided in the POT package [30]. In the PYTORCH package [66], the dependency retention in Eq. (23) is accomplished by explicitly enabling the <code>create_graph</code> flag when calling the <code>autograd</code> method on the AE energy in each MCMC step. This ensures that the computational graph, which PYTORCH uses to perform backpropagation, includes the dependency of the negative examples on the network weights throughout the MCMC.

B Learning rate scheduler

The torch.optim.lr_scheduler.ReduceLROnPlateau method [66] is used to reduce the learning rate by a factor of 0.8 when the Wasserstein distance does not decrease for 20 epochs. The values for all hyperparameters of the learning rate scheduler can be found in Table B.1. The evolution of the learning rate during the training of the WNAE is shown in Fig. B.1.

Hyperparameter	Value
Initial learning rate	2×10^{-4}
Mode	"min"
Factor	0.8
Patience	20
Cooldown	10
Threshold	0.001
Threshold mode	"rel"
Minimum learning rate	0

Table B.1: The hyperparameters of the learning rate scheduler.

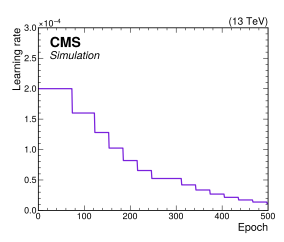


Figure B.1: The learning rate during the training of the WNAE from Section 4.3.

C Tuning the MCMC

Markov chains initialized with persistent contrastive divergence (PCD) [67] are found to accurately sample from the p_{θ} distribution. Therefore, PCD is used to initialize the Markov chain for each batch. In PCD, the Markov chain in batch k starts not from a new random set of points, but rather from the final set of points obtained from the Markov chain in batch k-1. This procedure can be tuned by adding a restart fraction r, meaning that at every batch, a fraction r of the points sampled for the previous batch are discarded and replaced by new, random points, while the remaining points from the previous batch are reused [37]. The restart fraction r is a hyperparameter of the MCMC algorithm and is set to 5%.

The tuning of the Markov chain hyperparameters—the coefficients of Eq. (11), the step size λ and the noise σ —is performed as follows. The number of MCMC steps $N_{\rm MCMC}$ is fixed to 10 as a compromise between training time and sampling accuracy. The MCMC hyperparameters λ and σ are chosen to satisfy Eq. (13), leaving one degree of freedom, and the temperature T was constrained to the interval (0,1]. These parameters are optimized to maximize the AUC

for discrimination between SVJs and top quark jets, using Eq. (20) as the loss function, with two constraints to mitigate bias toward the signal hypothesis. First, at the maximum AUC, the input feature distributions of the negative and positive samples should match, as discussed in Section 3.4. Second, the Wasserstein distance between the positive and negative samples should be minimal, as discussed in Section 4.1. If the MCMC parameters are correctly set and the network is sufficiently trained, then the network learns $p_{\theta} = p_{\text{data}}$, so the AUC is greater than 0.5, assuming the distribution of the signal is not also equal to p_{data} , in which case no discrimination would be possible. In this case, the distribution of the negative samples is representative of the p_{data} distribution. The values of the Markov chain hyperparameters obtained from this optimization are provided in Table C.1.

Trainings with a considerably larger gradient coefficient λ compared to the optimum were found to reach larger AUCs; however, these failed to achieve a good match between the distributions of negative and positive samples. Such MCMC configurations, though appealing if only the AUC is considered, result in an inadequate NAE that has not learned $p_{\theta} = p_{\text{data}}$: the AUC is large only because of the specific location of the signal hypotheses. This choice would therefore bias the performance of the NAE towards the signal hypotheses used for tuning, spoiling the unsupervised nature of the approach.

Table C.1: The hyperparameters of the MCMC.

Hyperparameter	Value
Number of steps N_{MCMC}	10
Restart fraction <i>r</i>	0.05
Temperature <i>T</i>	0.063
Noise σ	0.22
Step size λ	0.0242

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