NeuralQAAD: An Efficient Differentiable Framework for Compressing High Resolution Consistent Point Clouds Datasets

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Abstract: In this paper, we propose NeuralQAAD, a differentiable point cloud compression framework that is fast, robust to sampling, and applicable to consistent shapes with high detail resolution. Previous work that is able to handle complex and non-smooth topologies is hardly scaleable to more than just a few thousand points. We tackle the task with a novel neural network architecture characterized by weight sharing and autodecoding. Our architecture uses parameters far more efficiently than previous work, allowing it to be deeper and more scalable. We also show that the currently only tractable training criterion for point cloud compression, the Chamfer distance, performances poorly for high resolutions. To overcome this issue, we pair our architecture with a new training procedure based on a quadratic assignment problem. This procedure acts as a surrogate loss and allows to implicitly minimize the more expressive Earth Movers Distance (EMD) even for point clouds with way more than 10⁶ points. As directly evaluating the EMD on high resolution point clouds is intractable, we propose a new divide-and-conquer approach based on k-d trees, which we call EM-kD. The EM-kD is shown to be a scaleable and fast but still reliable upper bound for the EMD. NeuralQAAD demonstrates on three datasets (COMA, D-FAUST and Skulls) that it significantly outperforms the current state-of-the-art both visually and qualitatively in terms of EM-kD.

1 Introduction

In recent years, deep learning has been successfully applied to numerous computer vision and graphics tasks, such as classification, segmentation, or compression. However, most progress has been achieved in the 2D domain based on the regular data structure of images. Unfortunately, the same methods cannot easily be generalized to unstructured three-dimensional data. Memory consumption and computational cost usually increase rapidly in 3D, which limits the applicability of GPU-accelerated algorithms. In particular, gradient descent algorithms suffer from this phenomenon. Especially irregular representations such as point clouds or meshes are only sparely covered so far. However, as the typical output of 3D scanners, high resolution point clouds are widely used in autonomous driving, robotics, and other domains to represent surfaces or volumes. A tool capable of reducing the memory requirements of point clouds that can be added to a differential pipeline is therefore highly desirable. Unfortunately, point clouds pose a particular challenge since points may not only be irregularly but also sparsely scattered in space and, in contrast to meshes, do not provide explicit neighborhood information. Furthermore, the habit of neural networks to generate smooth signals seems to hinder the reconstruction of high frequencies, complex topologies, and discontinuities.

The current state-of-the-art in differentiable point cloud compression mainly focuses on the idea of folding one or multiple fixed input manifolds into a target manifold (Chen et al., 2020; Groueix et al., 2018; Deprelle et al., 2019). Regardless of the respective concept, the size of the point clouds that can be processed by current hardware is very limited, since mainly pointwise neural networks are used. Recently, sampling techniques have been used to overcome this problem (see e.g. (Mescheder et al., 2019; Groueix et al., 2018; Deprelle et al., 2019)). Thereby, the difference between two point clouds is usually measured using the Chamfer distance (see e.g. (Chen et al., 2020; Groueix et al., 2018; Zhao et al., 2019)). The popularity of the Chamfer distance stems from its ability to measure the distance from a point in one point cloud to its nearest neighbor in the other and vice versa. However, using the Chamfer metric on point cloud samples badly misguides gradient descent optimization algorithms. This is due to the fact that not all points are considered when determining the nearest neighbors of point cloud samples. Therefore, with high probability, distances between false matches are minimized. However, although a local optimum can be achieved, high frequencies are usually lost as a result. This problem is particularly severe when a point cloud is densely sampled and contains detailed, fine structures. The most commonly used PointNet encoders (Chen et al., 2020; Groueix et al., 2018; Zhao et al., 2019) suffer from exactly this problem as they rely on lossy global pooling layers that can only be generalized to a limited extent. Even worse, the Chamfer distance is generally known as an unsuitable criterion for the similarity of point clouds (Liu et al., 2020; Achlioptas et al., 2018). The earth movers distance (EMD), that solves a linear assignment problem (LAP) between compared point clouds, is considered to be more appropriate (Liu et al., 2020; Achlioptas et al., 2018). However, solving a linear assignment problem has a cubic running time for an exact solution and even approximations are not efficient enough for training neural networks on point clouds with more than a few thousand points.

In this paper we present NeuralQAAD, a deep autodecoder for compact representation of high resolution point cloud datasets. NeuralQAAD is based on a specific neural network architecture that overlays multiple foldings defined on learned and shared features to reconstruct point clouds. Our main contributions are

- a new scaleable point cloud autodecoder architecture that is able to efficiently recover high resolution consistent shapes,
- 2. training algorithms that are motivated by a newly defined quadratic assignment problem (QAP),
- 3. a validation that our approach outperforms the current state-of-the-art on high resolution point clouds, and finally
- 4. a novel dataset of detailed skull CT-scans.

2 Related Work

In the following, we give a brief overview of the three main areas of related work: 1. algorithms and data structures to process point clouds using deep learning, 2. neural network architectures for compressing point clouds, and 3. assignment problems.

2.1 Deep Learning for 3D Point clouds

Various data structures for representing 3D data in the context of deep learning have been examined, including meshes (Bruna et al., 2014), voxels (Wang et al., 2018; Maturana and Scherer, 2015), multi-view (Su et al., 2015), classifier space (Mescheder et al., 2019) and point clouds (Qi et al., 2017a). Point clouds are often the raw output of 3D scanners and are characterized by irregularity, sparsity, missing neighborhood information, and permutation invariance. To cope with these challenges, many approaches convert point clouds into other representations such as discrete grids or polygonal meshes.

Nonetheless, each data structure comes with its downsides. Voxelization of point clouds (Girdhar et al., 2016; Sharma et al., 2016; Wu et al., 2016; Roveri et al., 2018) enables the use of convolutional neural networks but sacrifices sparsity. This leads to significantly increased memory consumption and decreased resolution. Utilizing meshes requires to reconstruct neighborhood information. This can either be done handcrafted, with the risk of wrongly introducing a bias, or learned for a particular task (Chen et al., 2020). In both cases, again, memory consumption increases significantly due to the storage of graph structures. Related to the transformation into meshes are approaches that define convolution directly on point clouds (Tatarchenko et al., 2018; Xu et al., 2018). Another approach is to embed point clouds into a classifier space via fully connected neural networks (Mescheder et al., 2019; Park et al., 2019). However, this approach does not take into account the advantages of point clouds, as neural networks tend to smooth the irregularity of point clouds and thus may miss detailed structures. Also, these methods are not trainable out of the box, as a countable finite (point) set must be defined on an uncountable space.

PointNet (Qi et al., 2017a) was the first neural network to directly work on point clouds. It applies a pointwise fully connected neural network to extract pointwise features that are combined by max-pooling to create global features. Thus, all the previously mentioned disadvantages apply to PointNet. There is numerous work on extending PointNet to be sensitive to local structures (see e.g. (Qi et al., 2017b; Zhou and Tuzel, 2018)). In this paper, we focus on unsupervised learning. Most of the aforementioned methods are only applicable in a supervised setting. We deploy PointNet to produce intermediate encodings but discard it for the final compression.

2.2 Deep Point Cloud Compression

An autoencoder is the standard technique when it comes to compressing data using deep learning. However, recent work (Park et al., 2019) demonstrates that solely utilizing a decoder can be sufficient, at least in the 3D domain. Here, the encoder is replaced by a trainable latent tensor. Deep generative models like generative adversarial networks (GANs) (Goodfellow et al., 2014) or variational autoencoders (VAEs) (Kingma and Welling, 2014) can also be applied to reduce dimensionality. But they add overhead complexity and are rarely used for this purpose, although for VAEs (Zadeh et al., 2019) it may be sufficient to train only one decoding part, just like for standard autoencoders.

Most commonly, deep learning based compression methods for 3D point clouds either rely on voxelization (Girdhar et al., 2016; Sharma et al., 2016; Wu et al., 2016) or have one neuron per point in a fully connected output layer (Fan et al., 2017; Achlioptas et al., 2018). The latter approaches are hardly trainable for large-scale point clouds and require a huge number of parameters even for small point clouds. In contrast, FoldingNet (Yang et al., 2018) and FoldingNet++ (Chen et al., 2020) rely on PointNet for encoding and use pointwise decoders that are based on the folding concept. Thereby folding describes the process of transforming a fixed input point cloud to a target point cloud by a pointwise neural network. FoldingNet++ additionally learns graph structures for localized transformations partly avoiding the problem of the smooth signal bias. Another approach based on the concept of folding is AtlasNet (Groueix et al., 2018), which folds and translates multiple 2D patches to cover a target surface. The extension AtlasNetV2 (Deprelle et al., 2019) makes these patches trainable. AtlasNet and AtlasNetV2 suffer from the problem that they are hardly scalable and sometimes fail to glue all patches together without artifacts. Other pointwise autoencoders include 3D Point Capsule Networks (Zhao et al., 2019), which use multiple latent vectors per instance to capture different basis functions while using a dynamic routing scheme (Sabour et al., 2017).

All of the state-of-the-art point cloud autoencoders (Chen et al., 2020; Groueix et al., 2018; Zhao et al., 2019) are trained by optimizing the Chamfer distance. Older approaches such as (Fan et al., 2017; Achlioptas et al., 2018) trained on the earth mover distance, as well, but are restricted in point cloud size. In this work, we provide a *surrogate loss* for minimizing the EMD even for high resolution point clouds.

2.3 Assignment Problems

In its most general form, an assignment problem consists of several agents and several tasks. The goal is to find an assignment from the agents to the tasks that minimizes some cost function. The term assignment problem is often used as a synonym for the linear assignment problem (LAP). LAPs have the same number of agents and tasks. The cost function is defined on all assignment pairs. Kuhn proposed a polynomial time algorithm, known as the Hungarian algorithm (Kuhn, 1955), that can solve a LAP in quartic runtime. Later algorithms such as (Edmonds and Karp, 1972) or (Jonker and Volgenant, 1986) improved to cubic runtime. A fast and parallelizable *ɛ*-approximation algorithm (the auction algorithm) to determine an almost optimal assignment was introduced in (Bertsekas, 1988) and used in (Fan et al., 2017) to calculate the EMD. For NeuralQAAD, we use a GPUaccelerated implementation of the Auction algorithm (Liu et al., 2020). Although being fast in comparison to previous algorithms, it is by no means fast enough to act as a gradient descent criterion. Other assignment approximations exist such as the Sinkhorn divergence (Feydy et al., 2019), but to the best of our knowledge none of them can be implemented efficiently enough on GPUs for gradient-based optimization.

Besides the LAPs, there are nonlinear assignment problems which differ in the construction of the underlying cost function. For nonlinear assignment problems, the cost function can have more than two dimensions. In this work, we utilize a four dimensional quadratic assignment problem (QAP) to model registration tasks. More precisely, we use the formulation of (Beckman and Koopmans, 1957), in which the general objective is to distribute a set of facilities to an equally large set of locations. In contrast to the LAP, there is no cost directly defined on pairs between facilities and locations. Instead, costs are modeled by a flow function between facilities and a distance function between locations. Given two facilities, the cost of their assignment is calculated by multiplying the flow between them with the distance between the locations to which they are assigned. This definition ensures that facilities that are linked by a high flow are placed close together. However, finding an exact solution or even an ϵ -approximative solution for a QAP is NP-hard (Burkard, 1984). Hence, even for small instances, it is almost impossible to solve a QAP in a reasonable time. In this paper, we propose algorithms that can be efficiently executed on GPUs and allow our architecture to follow quadratic assignments.



Figure 1: The topology and algorithms dedicated to NeuralQAAD. The initial assignment between the input point cloud Q and a target point cloud P is determined by QAAD_GREEDY. P is described by a latent vector \mathbf{l}_j and stored in the lookup table L. A point \mathbf{q}_i is processed along with \mathbf{l}_j by a shared MLP to create common features. Subsequently, the common features and \mathbf{q}_i are folded by a patch-specific MLP. The optimizer updates the input point \mathbf{q}_i , the latent code \mathbf{l}_i , and all MLP parameters while improving the initial assignment through QAAD_REASSIGNMENT.

3 Method

In the following, we give a formal description of the addressed problem and derive constraints for a corresponding deep learning solution. We then construct a neural network architecture based on the concept of folding (Yang et al., 2018) that satisfies these constraints. Finally, we demonstrate how to train this architecture by designing and efficiently solving a quadratic assignment problem. Figure 1 shows an overview of the proposed NeuralQAAD architecture.

3.1 Problem Formulation

Given a set $S = \{P_i \mid 1 \le i \le N\}$ of 3D point clouds

$$P_i = \left\{ \mathbf{p}_{ij} \in \mathbb{R}^3 \mid 1 \le j \le M \right\} \tag{1}$$

all having the same number of points, we seek for a pair of functions f, g where $f : \mathbb{R}^{3 \cdot M} \mapsto \mathbb{R}^{l}, l \ll M$ encodes a point cloud into a lower dimensional feature vector

$$\mathbf{l}_i = f(P_i) \in \mathbb{R}^l, \tag{2}$$

and $g : \mathbb{R}^l \mapsto \mathbb{R}^{3 \cdot M}$ decodes the feature vector \mathbf{l}_i back into a point cloud

$$\overline{P}_i = g(\mathbf{l}_i) \tag{3}$$

such that the error

$$E(f,g) = \frac{1}{N} \sum_{i=1}^{N} d(\overline{P}_i, P_i), \qquad (4)$$

is minimal with respect to some function d which measures the similarity of two point clouds. As discussed in Section 2.2, folding transforms equation (3) into

$$\overline{P}_i = g(\mathbf{l}_i, Q), \tag{5}$$

where $Q = \{\mathbf{q}_j \mid 1 \le j \le M\}$ is a single fixed input point cloud.

As point clouds possess no explicit neighborhood information and are invariant to permutations, the encoder function f should also have these properties. Therefore, we further divide f into a pointwise local feature descriptor \hat{f} transforming the pointset P_i into a feature set

$$\hat{F}_i = \left\{ \hat{f}(\mathbf{p}_{ij}) \mid 1 \le j \le M \right\} \tag{6}$$

and a permutation-invariant global feature descriptor \tilde{f} which converts the feature set into a lower dimensional feature vector

$$\tilde{l}_i = \tilde{f}(\hat{F}_i) \in \mathbb{R}^l.$$
(7)

Likewise, substituting the decoder function g with a pointwise reconstruction function \hat{g} changes (5) into

$$\hat{P}_i = \left\{ \hat{\mathbf{p}}_{ij} \mid 1 \le j \le M \right\} \text{ with } \hat{\mathbf{p}}_{ij} = \hat{g}(\tilde{l}_i, \mathbf{q}_j).$$
(8)

Thus, instead of finding two functions f, g that minimizes (4), our objective is now to find three functions \hat{f}, \hat{g} and \tilde{f} that minimize

$$E(\hat{f}, \hat{g}, \tilde{f}) = \frac{1}{N} \sum_{i=1}^{N} d(\hat{P}_i, P_i).$$
(9)

Minimizing (9) using deep learning algorithms, in general, is very memory and computational intensive. For example, a comparably small batch of 32 point clouds each consisting of M = 65000 points is equal to processing a rather large batch of around 2653 MNIST¹ images. Therefore the realization of \hat{f}, \hat{g} and \tilde{f} through deep neural networks requires computationally efficient and memory-optimized methods as presented in the following.

¹http://yann.lecun.com/exdb/mnist/

3.2 EM-kD

The measure d for the similarity of point clouds is typically chosen to be either the Chamfer distance (Groueix et al., 2018; Zhao et al., 2019)

$$d_{C}(\hat{P}, P) = \sum_{\mathbf{p} \in P} \min_{\hat{\mathbf{p}} \in \hat{P}} \|\mathbf{p} - \hat{\mathbf{p}}\|_{2} + \sum_{\hat{\mathbf{p}} \in \hat{P}} \min_{\mathbf{p} \in P} \|\mathbf{p} - \hat{\mathbf{p}}\|_{2}$$
(10)

or the augmented Chamfer distance (Chen et al., 2020)

$$d_A(\hat{P}, P) = \max\left\{\sum_{\mathbf{p}\in P}\min_{\hat{\mathbf{p}}\in\hat{P}}\|\mathbf{p}-\hat{\mathbf{p}}\|_2, \sum_{\hat{\mathbf{p}}\in\hat{P}}\min_{\mathbf{p}\in P}\|\mathbf{p}-\hat{\mathbf{p}}\|_2\right\}$$
(11)

with previously detected weaknesses (Liu et al., 2020; Achlioptas et al., 2018).

Compared to the Chamfer distance, the Earth Mover Distance (EMD)

$$d_{EMD}(\hat{P}, P) = \min_{\phi: P \to \hat{P}} \sum_{\mathbf{p} \in P} \|\mathbf{p} - \phi(\mathbf{p})\|_2 \qquad (12)$$

is considered to be more representative for visual varieties (Liu et al., 2020; Achlioptas et al., 2018) in case that ϕ is (almost) a bijection. We define

$$\arg d_{EMD}(\hat{P}, P) = \{(\mathbf{p}, \hat{\mathbf{p}} = \phi(\mathbf{p}))\}_{\mathbf{p} \in P}$$
(13)

as the linear assignment problem (LAP) imposed by the EMD.

Since the use of the EMD even for moderately large point clouds leads to impracticable runtimes, we introduce a divide-and-conquer approximation of the real EMD, which we call EM-kD. The EM-kD is defined as

$$d_{EM-kD}(\hat{P},P) = \sum_{(\hat{\mathcal{P}},\mathcal{P})\in k_d(\hat{P})\nabla k_d(P)} d_{EMD}(\hat{\mathcal{P}},\mathcal{P}), \quad (14)$$

where the k_d function constructs k-d tree subspaces and ∇ defines the diagonal product that enumerates the canonical subspace pairs. Afterwards, the (approximated) EMD between each subspace pair is measured. In all our experiments we use the auction algorithm as an EMD approximation.

The EM-kD closely approximates the EMD as can be seen by considering the following two cases. First, we assume that the low frequencies of both point clouds are similar, i.e., the k-d subspaces will also be similar. Here, the EM-kD may slightly overestimate the EMD at most at the subspace boundaries. The second case considers the situation where the low frequencies of both point clouds are different and the k-d subspaces are different as well. In this situation, the EM-kD may significantly overestimate the EMD. Overall, EM-kD judges similar to the EMD, but penalizes very dissimilar point clouds even more. As a side benefit, the EM-kD may also be well suited for other tasks where the EMD is typically used, such as point cloud completion. While being a reasonable upper bound of the true EMD, the EMkD is greatly scaleable as the subtask size of the auction algorithm can be controlled through the depth of the k-d trees. In our experiments, using the EM-kD reduces the runtime by more than an order of magnitude, more precisely by a factor of 36.

In all our experiments, we used the EM-kD to evaluate NeuralQAAD. In Section 3.4 we present a training procedure to implicitly optimizing the EMkD and hence the EMD.

3.3 Network Architecture

In the following, we discuss the three key components that we have identified for a successful implementation of a scalable autodecoder for point clouds based on a deep neural network. Since our approach is mainly motivated by a Quadratic Assignment Problem and realizes an Auto Decoder, we call it NeuralQAAD.

First, NeuralQAAD follows the idea of folding. In contrast to previous work that folds a fixed (Yang et al., 2018) or trainable (Deprelle et al., 2019) point cloud by uniformly sampling a specific manifold such as a square, we choose to deform a random sample of the target data set as this is very likely to be closer to any other sample than an arbitrary primitive geometry and thus is better suited for initialization. For a dataset with strong low frequency changes, clustering of multiple random examples may be even more advantageous, but we leave this for future research as it turned out to be not beneficial in our experiments.

Second, we found that splitting the input point cloud into K many patches that are independently folded (Deprelle et al., 2019), i.e. K independent multilayer perceptrons (MLPs), is essential but also uses resources highly inefficiently. Each patch has its own set of parameters and thus its own set of extracted features. Increasing the number of patches and thus the number of MLPs, quickly limits their potential depth. We observe that weight sharing across the first layers of each MLP does not decrease the performance, but enables the realization of deeper and/or wider architectures with the same number of parameters. Similarly, low-level feature sharing can be used to scale the number of patches.

Third, as discussed in Section 3.1, a batch of large point clouds may not be processed at once. To cope with this problem, NeuralQAAD only *folds* a sample from each point cloud of a batch during a single forward pass. Since the latent code \tilde{l}_i of an instance

depends on all its related points, the encoding function \tilde{f} needs to be adapted. We realize \tilde{f} as a trainable lookup table and thus renounce the explicit calculation of the pointwise function \hat{f} . In addition to our motivation for autodecoders to minimize memory usage, (Park et al., 2019) claims that this topology utilizes computational resources more effectively.

Considering the three constituents just discussed, the problem of finding functions \hat{f}, \hat{g} and \tilde{f} that minimize (9) can now be rephrased into the problem of training several multilayer perceptrons MLP_k , one for each input patch Q_k , and a lookup table $L \in \mathbb{R}^{N \times l}$ such that

$$E(MLP, L, Q) = \frac{1}{N} \sum_{i=1}^{N} d(P_i, \hat{P}_i), \qquad (15)$$

is minimized, where the elements of \hat{P}_i are given as

$$\hat{\mathbf{p}}_{i,j} = MLP_k(\mathbf{l}_i, \mathbf{q}_{k,j}), k \in K.$$
(16)

Without loss of generality, we do not explicitly state the split into various patches to simplify notation in the following. The inverse assignment from each predicted point to its source point is given by $\mathbf{q}_{k,j} = MLP_k^+(\mathbf{l}_i, \hat{\mathbf{p}}_{i,j}), k \in K$.

3.4 Training

Optimizing NeuralQAAD only on subsets of the input point cloud Q directly affects the construction of the loss function. The common pattern for calculating a distance between points clouds is similar to a registration task that can be split into the two steps

- 1. Identifying a suitable matching between both point clouds.
- 2. Determine the actual deformation of one point cloud to the other.

Gradient descent is only done for the second step whereas the first step characterizes how appropriate a metric is. Apart from the fact that the Chamfer distance is not reliable and the EMD is not tractable, another general consideration comes into play: The matching that steers the folding process has to comply with the predominant bias of neural networks, which is smoothing. Therefore, we choose a matching that allows the optimizer to follow the quadratic assignment problem described next.

Given an input point cloud Q, a target point cloud P, a weight function $w(\mathbf{p}, \hat{\mathbf{p}}) = \|\mathbf{p} - \hat{\mathbf{p}}\|_2$, and a flow function $f(\mathbf{q}, \hat{\mathbf{q}}) = \frac{1}{w(\mathbf{q}, \hat{\mathbf{q}})}$ we want to find a bijective mapping $A : Q \mapsto P$ such that

$$E(A) = \sum_{\mathbf{q}, \hat{\mathbf{q}} \in Q} f(\mathbf{q}, \hat{\mathbf{q}}) \cdot w(A(\mathbf{q}), A(\hat{\mathbf{q}}))$$
(17)

is minimized. Intuitively speaking, this means that by solving this quadratic assignment problem, we assure

that points close to each other in Q are matched with points close to each other in P. Note that LAPs do not exhibit this property and, hence, might inhibit gradient descent. By construction, minimizing the distances between all QAP matches optimizes any criterion like the EMD, EM-kD or Chamfer distances, too.

Algorithm 1 Calculate an initial QAP solution A between a target point cloud P and the image \hat{P} of an input point cloud Q. The image \hat{P} is calculated by the NeuralQAAD *MLP* given a latent vector **I**.

procedure QAAD_GREEDY(l, Q, P)// Predict input point cloud $\hat{P} = MLP(\mathbf{l}, Q)$ // Set of initial QAP solution $A = \emptyset$ // Assignment for each canonical k-d tree subspace pair for $(\hat{\mathcal{P}}, \mathcal{P}) \in k_d(\hat{P}) \nabla k_d(P)$ do // Matching of EMD $\mathcal{A} = \arg d_{EMD}(\hat{\mathcal{P}}, \mathcal{P})$ // If duplicates exist, use the smallest distance only for $(\boldsymbol{\hat{p}},\boldsymbol{p})\in\mathcal{A}$ do if $\nexists(\bar{\mathbf{p}}, \mathbf{p}) \in \mathcal{A} : ||(\bar{\mathbf{p}} - \mathbf{p})|| < ||(\hat{\mathbf{p}} - \mathbf{p})||$ then $A = A \cup \{(MLP^+(\mathbf{l}, \hat{\mathbf{p}}), \mathbf{p})\}$ // Randomly assign the remaining points for $(\hat{\mathbf{p}}, \mathbf{p}) \in \mathcal{A}$ do if $\nexists(MLP^+(\mathbf{l}, \hat{\mathbf{p}}), \bar{\mathbf{p}}) \in A$ then $\vec{\mathbf{p}} \sim uniform(\{\vec{\mathbf{p}} \in \mathcal{P} | \nexists(\mathbf{q}, \vec{\mathbf{p}}) \in A\})$ $A = A \cup \{(MLP^+(\mathbf{l}, \hat{\mathbf{p}}), \vec{\mathbf{p}}))\}$ // Return the initial assignment return A

On first sight, QAPs seem to be harder to solve than LAPs. However, we do not need to solve them explicitly, but remove the limitation of the optimizer of following a linear assignment. The first step is a greedy algorithm called QAAD_GREEDY that generates an initial matching before gradient-descent. The second step is a dynamic reassignment algorithm called QAAD_REASSIGNMENT, that monotonically improves equation (15). During our experiments it became evident that neither of the two steps alone lead to a superior solution. The novel shared NeuralQAAD architecture can also be trained on the augmented Chamfer Distance training procedure which already results in a significant performance improvement and does not produce any additional assignment overhead at all, but still is not as good as our QAAD training scheme (see Section 4).

QAAD_GREEDY (see Algorithm 1) is conceptually similar to calculating the EM-kD. Instead of using the calculated distance between the input and the target point cloud the determined LAP matches are used as the initial assignment solution. To establish a perfect matching, which is not guaranteed by the Algorithm 2 Refining the initial QAP solution A for a query of points $Q \subset Q$ of the input point cloud Q. At this, the image \hat{P} of the query given a latent code I is constructed by the current state of the NeuralQAAD *MLP* and evaluated against the target point cloud P. However, for efficiency only a subset $\mathcal{P} \subset P$ of the target is considered for reassignment.

procedure QAAD_REASSIGNMENT($\mathbf{l}, Q, \mathcal{P}, A$) for $a \in O$ do // Predict input subset and assign their nearest // neighbors in target subset $\mathcal{A}(q) = \operatorname{argmin}_{\mathbf{p} \in \mathcal{P}} \left\| \mathit{MLP}(\mathbf{l}, \mathbf{q}) - \mathbf{p} \right\|$ // Check current loss $L_B = \|MLP(\mathbf{l}, \mathbf{q}) - A(\mathbf{q})\|$ $+ \| MLP(\mathbf{l}, A^{-1}(\mathcal{A}(\mathbf{q}))) - \mathcal{A}(\mathbf{q}) \|$ // Check loss after hypothetical reassignment $L_A = \|MLP(\mathbf{l},\mathbf{q}) - \mathcal{A}(\mathbf{q})\|$ + $\|MLP(\mathbf{l}, A^{-1}(\mathcal{A}(\mathbf{q}))) - A(\mathbf{q}))\|$ // Decide where reassignment is beneficial if $L_A \leq L_B$ then // Delete old assignment $A = A \setminus \{(\mathbf{q}, A(\mathbf{q}))\}$ $A = A \setminus \{ (A^{-1}(\mathcal{A}(\mathbf{q})), \mathcal{A}(\mathbf{q})) \}$ // Add new assignment $A = A \cup \{(\mathbf{q}, \mathcal{A}(\mathbf{q}))\}$ $A = A \cup \{ (A^{-1}(\mathcal{A}(\mathbf{q})), A(\mathbf{q})) \}$ // Return the refined assignment return A

auction algorithm, the input point closest to a repeatedly chosen target point is matched. Any remaining points are randomly matched. Like for the EMkD, QAAD_GREEDY can be efficiently parallelized on a GPU. Even for the largest tested dataset D-Faust (Bogo et al., 2017) QAAD_GREEDY runs only minutes. Additionally, since it only runs once, the runtime is negligible considering the common training time of neural networks.

Since the input point cloud is trainable, QAAD_GREEDY may make assignment errors at subspace borders and the initial matching is still an approximation of a linear assignment. QAAD_REASSIGNMENT (see Algorithm 2) conducts reassignments during gradient descent that ease the training of NeuralQAAD. At this, given a latent code, a sample of the input point cloud is predicted and the nearest neighbor from a sample of the target point cloud is found for each prediction. This can be efficiently done in parallel on a CPU using k-d trees or other suitable data structures. Newer GPU architectures that refrain from warp-based computation also allow for efficient GPU implementations. The k-d trees are build and queried with only a few samples per training step and, hence, add little overhead. Subsequently, the input points that are actually assigned to the nearest neighbors are also predicted. Reassignment is based on comparing the summed loss of both predictions under the actual matching with the summed loss of both predictions after a hypothetical pointwise swap. In case the latter loss is smaller, a reassignment is conducted. Hence, each reassignment is guaranteed to reduce the error defined by equation (15). More importantly, the optimizer is no longer biased towards minimizing the distance of an optimal linear assignment.

The distance measure in this algorithm can be arbitrarily chosen, but due to the targeted EM-kD, we rely on the mean squared error. As we only process point cloud subsets, we enforce a sampling procedure that outputs the same number of points for each patch. This allows us to efficiently implement NeuralQAAD as a grouped convolution. In contrast to previous implementations that sequentially run each MLP per patch, we can make use of all GPU resources and achieve a significant speedup that makes using hundreds of patches practical for the first time.

4 **Experiments**

In this section, we validate NeuralQAAD and its underlying assumptions. We start with an overview of the datasets used and some implementation details. Subsequently, we show that NeuralQAAD performs significantly better on high resolution points clouds than existing approaches. Specifically, we compare against the AtlasNetV2 deformation architecture (Deprelle et al., 2019), the only other state-of-the-art differentiable approach that is out-of-the-box applicable to point clouds with more than a few thousand points. Finally, we demonstrate the benefit of each NeuralQAAD design choice in an ablation study and the robustness of our approach in the number of sampled training points.

4.1 Datasets

Our experiments are based on the three different datasets COMA (Ranjan et al., 2018), D-Faust (Bogo et al., 2017) and Skulls. COMA, captures 20466 scans of extreme expressions from 12 different individuals captured by a multi-camera stereo setup. D-Faust, is a 4D full body dataset that includes 40000 multi-camera scans of humans in motion. We use about 6576 scans that contain more than 110000 points and about 13744 scans that contain more than 130000 points from COMA and D-Faust respectively. Skulls is a new artificially generated point cloud dataset constructed using a multilear model (Achenbach et al., 2018) that is based on a skull template fit-

ted to 42 CT scans of human skulls (Gietzen et al., 2019). The dataset consists of 4096 point clouds. Each point cloud is composed of 65536 points and includes fine-grained parts such as the nasal bones or teeth. In contrast to the other datasets, it consists not only of surfaces but also of volumetric structures. We will make this dataset public as a new high resolution point cloud test dataset.

4.2 Implementation Details

For all our experiments we implemented the NeuralQAAD folding operation with six fully connected layers with output dimensions 256-128-128-64-3 using PyTorch. In this implementation, the first four layers are shared across all patches (see Figure 1 for an architecture overview). To guarantee the same number of parameters for the AtlasNetV2 decoder while allowing it to be sufficiently deep, wide, and computationally tractable we construct the folding operation of AtlasNetV2 with four layers of size 128 (no sharing) and adapt the number of patches if needed. SELU (Klambauer et al., 2017) is applied as a nonlinearity to every but the output layer for both NeuralQAAD and AtlasNetV2. Adam (Kingma and Ba, 2015) is used as the optimizer and initialized with a learning rate of 0.001. We sample 4096 points for COMA as well as D-Faust and 2048 points for Skulls per instance and training step, resulting in low GPU memory consumption.

AtlasNetV2 is trained until we can no longer observe any significant improvement, which was the case for all data sets after 150 epochs at the latest. To show that our architecture is also superior without our novel QAAD training scheme we also train NeuralQAAD for 150 epochs with the AtlasNetV2 training scheme. Essentially, this means minimizing the augmented Chamfer loss on point cloud samples. For this purpose, we deploy a PointNet encoder, too, which we discard as soon as our QAAD training scheme is utilized. Nevertheless, the PointNet embeddings are reused to initialize the lookup table. For all experiments except the convergence NeuralQAAD experiment, we train for 450 additional epochs. All experiments are conducted on TITAN RTX GPUs and a AMD Ryzen[™] Threadripper[™] 3970X CPU. In comparison, the experiments of AtlasNetV2 have a noticeably longer runtime than those of NeuralQAAD since we use the released sequential implementation of AtlasNetV2. For instance, training AtlasNetV2 on D-Faust takes over a day whereas NeuralQAAD only requires two hours. For all experiments, we use the same hyperparameters for the auction algorithm i.e. 100 iterations with an epsilon of 1. The EM-kD is calculated with a subspace size of 1024.

4.3 Reconstruction of High Resolution Point Clouds

Skulls: The Skulls dataset contains point clouds that can be linearly reduced to 42 latent variables using principal component analysis. Additionally, in our experiments, we nonlinearly compress each skull to a latent vector of size 10. Although this seems to be an easy task at first sight, results of AtlasNetV2 demonstrate the contrary.

We process Skulls in a batch size of 16 instances and use 16 patches for both NeuralQAAD and AtlasNetV2. The visual results can be seen in Figure 2. NeuralQAAD is able to recover almost all low as well as high frequencies. In contrast, AtlasNetV2 fails in both. From the side view, it can be clearly recognized that detailed and non-manifold structures as teeth and the nasal area are lost by AtlasNetV2 to a huge extent but can be fully reconstructed by NeuralQAAD. In addition, rather coarse structures like the eye socket (front view) or the skullcap (front and side view) are badly recovered by AtlasNetV2, whereas NeuralQAAD does not suffer. We mainly attribute the better volumetric reconstruction capabilities to the improved training procedure. However, also the effects of weight sharing can be observed visually. AtlasNetV2 clearly struggles in gluing together individual patches. This does not seem to be an issue for NeuralQAAD that works on shared low level features. The visual impressions are strongly supported by the measured EM-kD losses stated in Table 1. Training on the augmented Chamfer loss alone leads to an improvement of 21% in terms of EM-kD. After training our approach to convergence, we even get an improvement of 78% and achieve a compression ratio of 3311:1 (# of dataset floats : # of network parameter and embedding floats).

D-Faust: The D-Faust dataset predominantly contains low-frequency structures. We compress each instance to a latent code of size 256. Training is conducted with batches of size 16 while using 128 patches for NeuralQAAD. To ensure comparability in the number of parameters AtlasNetV2 can only utilize 28 patches. Visual results are depicted in Figure 3a. Unlike the volumetric Skulls dataset, here we can recover a surface for each point cloud to facilitate visual comparison of reconstruction results. For surface reconstruction, we used the simple ball-pivoting algorithm (Bernardini et al., 1999) with identical hyperparameters for all experiments. From all perspectives, it becomes evident that AtlasNetV2 merges even coarse structured extremities and, as with Skulls, struggles to

Architecture	AtlasNetV2	NeuralQAAD & Encoder	NeuralQAAD			
Trained with	Aug. Chamfer	Aug. Chamfer	MSE	+ QAAD_GREEDY	+ QAAD_REASSIGN.	
Data Epochs	∞ (=150)	150	600	600	600	~
Skulls	30.681	24.004	419.230	18.312	10.481	6.708
D-Faust	0.076	0.023	0.187	0.015	0.011	0.010
COMA	200.083	139.230	2448.076	74.893	47.593	18.554

Table 1: Reconstruction EM-kD for the datasets Skull, D-Faust and COMA. Convergence is indicated with ∞ . On all datasets NeuralQAAD performances significantly better than the previous state-of-the-art AtlasNetV2 even if trained only with the augmented Chamfer loss. For the D-Faust dataset, NeuralQAAD achieves significant improvements with and without QAP training, indicating that the scalability of NeuralQAAD in the number of patches is the main performance factor. However, for the COMA dataset both the scalability in the number of patches as well as the QAP training procedure lead to huge performance jumps. For Skulls, mainly the novel QAP training procedure improves on the current state of the art.



Figure 2: Front and side views on NeuralQAAD reconstructions of Skulls (Achenbach et al., 2018). Each skull is compressed to a latent vector of size 10. Low as well as high frequencies can be mostly recovered by NeuralQAAD in contrast to AtlasNetV2 that fails on low frequencies like the skullcap or on high frequencies like the teeth.

stitch together individual patches. NeuralQAAD, by contrast, is able to almost fully recover all prevalent structures and forming a smooth surface across patch borders.

The EM-kD reconstruction losses for D-Faust stated in Tab. 1 show a significant improvement through NeuralQAAD. However, they tell a different story as for Skulls. Already after training NeuralQAAD with the augmented Chamfer distance, the EM-kD decreases by about 69%. Further, our training scheme leads to a total drop in EM-kD of 86% after convergence and achieves a compression ratio of 1144:1. Combined, both numbers indicate that the scalability of NeuralQAAD in the number of patches is the dominant factor of performance for D-Faust whereas the training scheme plays a minor role.

The D-Faust experiments reveal another interesting observation. In Figure 4 the trainable input point clouds are shown. Although initialized with a random example of the respective dataset, only for D-Faust a full deformation can be noticed. The *egg-like* structure seems to be favorable if strong low frequency changes occur which is not the case for Skulls and COMA (see next Section).

COMA: COMA is the most challenging dataset

as each point cloud contains almost as many points as those of D-Faust but is restricted to the facial area and, hence, captures an enormous amount of details. As for D-Faust, we compress each instance into a latent code of size 256, conduct training with a batch size of 16, and make use of 128 patches for NeuralQAAD as well as 28 patches for AtlasNetV2. Surface reconstructions are shown in Figure 3b. For both instances shown, NeuralQAAD is found to capture way more fine-grained structures than AtlasNetV2 while being less noisy. Even coarse structures like the ear in the first instance can not be recovered by AtlasNetV2.

The EM-kD outcomes stated in Table 1 suggest that for COMA the number of patches as well as the QAP training scheme are essential and neither alone suffices. Training NeuralQAAD on the augmented Chamfer distance achieves a 30% better performance than AtlasNetV2. Applying the QAP training procedure further improves NeuralQAAD by 90% overall after convergence and results in a compression ratio of 748:1.



Figure 3: a) Front and side views on NeuralQAAD reconstructions of D-Faust (Bogo et al., 2017). The differences between NeuralQAAD and AtlasNetV2 are clearly visible as AtlasNetV2 is not able to recover even coarse structures like extremities. b) Front and side views on NeuralQAAD reconstructions of COMA (Ranjan et al., 2018). As for D-Faust, most detailed structures are reconstructed by NeuralQAAD whereas AtlasNetV2 even looses coarse structures.



Figure 4: Combined trainable input patches after QAP training for the evaluated dataset. Colors reflect assignment to patches. For Skulls and COMA, datasets mostly defined by changes in high frequencies, the input point clouds looks like noisy instances. D-Faust, mostly defined by changes in low frequencies, exhibits a fully deformed input point cloud.

4.4 Ablation Study & Scalability

In this section we ground the major and novel design choices made for NeuralQAAD. In the preceding section, we already discussed the number of patches as well as shared low-level features as the key success factors for D-Faust and COMA proving the concept of weight sharing. However, we did not discuss the isolated impact of QAAD_GREEDY and QAAD_REASSIGNMENT, yet.



Figure 5: EM-kD for AtlasNetV2, NeuralQAAD architecture trained on the augmented Chamfer distance, and NeuralQAAD trained with our training scheme related to the number of sampled points per training step. The Skulls dataset is used.

In a baseline experiment (see *MSE & Decoder* in Table 1) we trained NeuralQAAD to minimize the MSE of a random assignment. As expected, this culminates in extraordinary bad results as the criterion does not even follow the properties of a LAP. In the subsequent experiment (see QAAD_GREEDY in Table 1), we observe that QAAD_GREEDY solves

this issues but can clearly not be made responsible for the overall performance of NeuralQAAD. Additionally applying QAAD_REASSIGNMENT (see QAAD_REASSIGNMENT in Table 1) and, hence, following the smoothing bias of neural networks, boosts the results again. In summary, the conducted experiments strongly support the capabilities and the necessity of the proposed contributions.

The behavior of NeuralQAAD, in the case where the number of sampled points is reduced, is very robust. Figure 5 shows that NeuralQAAD, with and without our training scheme, is always superior to AtlasNetV2, regardless of the number of sampled points. More importantly, when the number of sampled points is reduced, the performance loss is significantly lower than with AtlasNetV2. The observed robustness to sampling demonstrates the efficiency and scalability of NeuralQAAD with respect to the size of the point clouds to be processed. This also equips NeuralQAAD for the future, in which the resolution of point clouds will most probably continue to increase in most areas.

5 Conclusion

We introduced a new scaleable and robust point cloud autodecoder architecture called NeuralQAAD together with a novel training scheme. Its scalability comes from low level feature sharing across multiple foldable patches. In addition, refraining from classical encoders makes NeuralQAAD robust to sampling. Our novel training scheme is based on two newly developed algorithms to efficiently determine an approximate solution that follows the smoothing bias of neural networks. We showed that NeuralQAAD provides better results than the previous state-of-the-art applicable to high resolution point clouds. Our comparisons are based on the EM-kD, a novel scalable and fast upper bound for the EMD. In our experiments, the EM-kD has proven to reasonably reflect visual differences between point clouds.

The next steps will be to make our approach applicable to generative models and to bridge the gap to correspondence problems. Although at first glance generative tasks seem to be a straightforward extension, preliminary results have shown an unstable training process. Recent advancements in continuous learning might be adaptable to diminish the effects of sampling.

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