Deep Hierarchical Super-Resolution for Scientific Data Reduction and Visualization

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Abstract—We present an approach for hierarchical super resolution (SR) using neural networks on an octree data representation. We train a hierarchy of neural networks, each capable of \(2^x \times \) upscaling in each spatial dimension between two levels of detail, and use these networks in tandem to facilitate large scale factor super resolution, scaling with the number of trained networks. We utilize these networks in a hierarchical super resolution algorithm that up scales multiresolution data to a uniform high resolution without introducing seam artifacts on octree node boundaries. We evaluate application of this algorithm in a data reduction framework by dynamically downscaling input data to an octree-based data structure to represent the multiresolution data before compressing for additional storage reduction. We demonstrate that our approach avoids seam artifacts common to multiresolution data formats, and show how neural network super resolution assisted data reduction can preserve global features better than compressors alone at the same compression ratios.

Index Terms—Octree, Super Resolution, Neural Networks.

1 INTRODUCTION

Data analysis and visualization are essential tasks for scientists to understand large-scale output from simulations. Typically, scientists run simulations which generate high resolution spatiotemporal data and write some portion of the generated data to storage for post-hoc analysis. However, with supercomputers reaching exascale computing capability (at least 1 exaFLOPS) this decade, scientists are able to generate more simulation data than on previous generation’s petascale supercomputers. For example, Nyx [3] is a astrophysics simulation for bayonic and cold dark matter that can output up to 4096\(^3\) size volumes per simulated timestep [13] which take up over 4 TB each, and the Johns Hopkins Turbulence Database (JHTDB) hosts computational fluid dynamics simulation results up to 8192\(^3\) [7] which take up to 16 TB per timestep. However, current storage technology cannot save the data at the rate they are being generated [11], making it infeasible to store this much raw data for post-hoc analysis and visualization.

One approach for avoiding this I/O limitation is to use error-bounded lossy compressors like ZFP [39] and SZ [12] to reduce the data’s footprint before saving it to persistent storage. Error-bounded lossy compressors ensure that a target error bound will be satisfied after decompression. While some compression methods aim to preserve features found in specific data types, such as the work by Liang et al. [17] to compress vector fields, error-bound compressors generally only consider the error between the raw floating point value and its (de)compressed value when compressing, which can distort spatial features present in the data when visualizing.

An alternate approach for avoiding the I/O bottleneck is to simply discard a portion of simulated data and recover it post-hoc via super resolution algorithms. Data is discarded by saving timesteps of the simulation infrequently, such as saving once every 100 timesteps, or by spatially subsampling the data, such as from \(N^3\) to \((\frac{N}{2})^3\). Interpolation techniques like linear or bicubic interpolation are used to fill in the discarded data post-hoc, but can lead to overly smooth results and missing high-frequency features important for analysis. Due to the recent advancements of machine learning (ML), super resolution (SR) methods applied to scientific data have seen better feature re-

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construction than interpolation methods [18][20][73]. Using neural networks (NNs) for non-error-bounded lossy compression has been studied extensively for image compression with results exceeding JPEG and JPEG2000 [15][55][57][59]. A major benefit of machine learning for data reconstruction is ML’s ability to preserve global features well due to the learned statistics across large spatial and temporal domains. These global features are often learned implicitly, but can be learned explicitly via loss functions. Although these SR models for scientific data reconstruct global features well, they are limited single scale factor SR. Scientific data is often modeled with hierarchical structures such as an octree, but current SR models cannot support this data format.

In this paper, we aim to give scientists a data reduction algorithm that uses neural network super resolution along with compressors in the data reduction-reconstruction pipeline to preserve global features better than compressors alone. Since a single network trained for \(N\times\) SR would only offer two compression options - either the data is downsampled or it is not - we give more flexibility to data reduction by training a hierarchy of networks \(G\) to allow \(2\times, 4\times, \ldots, 2^G\times\) super resolution in each spatial dimension, which allows larger compression ratios for the same error bounds. We further improve compression ratios for the same error-bound by dynamically downscaling the data into an octree, a multiresolution data format. To accommodate this data format, we design a multiresolution super resolution algorithm for our hierarchy of neural networks to upscale this octree data format to a uniform resolution with minimal seam artifacts, which are prevalent in multiresolution data [42][61][63]. Our process is as follows:

1. **Pre-training** We collect full resolution simulation output and train a hierarchy of SR neural networks, one model at a time, with each model responsible for \(2\times\) SR between downsampling levels (..., \(8\times \rightarrow 4\times \rightarrow 2\times \rightarrow 2\times \rightarrow \) full resolution).

2. **Error bounded data modelling and compression** Data is dynamically downsampled in a trial and error fashion into an octree-based data format conditional on a user set error bound. After downscaling, data is compressed with a compressor while still obeying the error bound.

3. **Post-hoc data decompression and reconstruction** In post-hoc analysis, data is decompressed to the octree format and finally upsampled using our multiresolution super resolution algorithm with the pre-trained hierarchy of networks.

In our results, we evaluate the trained neural network hierarchy and verify that they outperform baseline interpolation techniques. We demonstrate our data reduction framework’s effectiveness at global feature preservation by qualitatively and quantitatively comparing reconstructed data with and without our method at the same compression limit.
We review related works in three general data reduction techniques: hierarchical data representations, super resolution methods for scientific data, and compression techniques for scientific data. Our work is part of a larger category of research called AI4VIS or ML4VIS, which applies artificial intelligence algorithms to enhance or improve parts of the visualization and analysis process scientists across all fields use. We recommend the surveys by Wu et al. [69] and Wang et al. [65] for recent efforts in this area of research.

2.1 Hierarchical data representations for scientific data
Hierarchical data representations have been used in scientific applications to reduce computation and storage requirements by more efficiently allocating resources available. Rendering pipelines have utilized octrees [45] for efficient and scalable volume rendering by dynamically loading data from different octree depths depending on the current camera position and angle. Gobbetti et al. [16] organize large scalar fields into an octree structure and only feed the GPU information relevant to the current viewpoint and transfer function at each frame. Knoll et al. [31] create a technique for isosurface ray tracing for large octree volumes. Wilhelms and Van Gelder [68] use octrees for isosurface generation when the regions of interest are unknown. A key problem for rendering hierarchical data is the seam artifact that can occur between data blocks. Methods for reducing the effect of this issue are presented by Santos et al. [42], Wald et al. [61], and Wang et al. [63]. A survey of octree-based rendering methods is available by Knoll [30].

Aside from rendering, hierarchical data formats have also been used during data transfer to improve performance. Losasso et al. [33] create a method for simulating water and smoke on an unrestricted octree to capture small scale visual detail and allow for efficient solving of the Poisson equation. Popinet [48] uses quadtrees and octrees for a flexible and efficient approach to solving time-dependent incompressible Euler equations. Adaptive mesh refinement (AMR) schemes, a similar hierarchical data representation, have been utilized in numerous scientific software packages to improve performance by using a coarse mesh where there is little detail and fine meshes where precision is more important. Berger and Colella first design the AMR scheme for use with hyperbolic partial differential equations [8] and two-dimensional shock dynamics [7]. Different implementations of AMR are implemented in modern simulation packages such as Enzo [47], Chombo [44], LAVA [29], Nyx [3], and AMR-C [72].

These hierarchical representations can also be used for data reduction. Knoll et al. [31] use an octree representation that only uses a fraction of the original data’s memory footprint, similar to the approach of Velasco and Torres [60], to accelerate volume rendering. Bhatia et al. [9] introduce AMM, an adaptive multilinear mesh framework that reduces the memory footprint of raw data using tensor products of linear B-spline wavelets and allow a tradeoff between numerical precision and spatial resolution. Hoang et al. [24] use a precision-resolution tree, and an encoding of the tree, to perform data reduction on scalar field data. Ainsworth et al. [2] create a multilevel technique to compress univariate data that allows users to select a level that minimizes memory use while meeting some tolerance level. Another multilevel technique is presented in MGARD [38], which uses adaptive error based coefficient quantization to enable different tolerances at different levels of detail, and proposes to use the multilevel data decomposition as a preconditioner to terminate at an appropriate level.

There are only a few works combining ML with hierarchical data. Riegler et al. [49] create octree convolution, pooling, and upscaling layers for a neural network and then learn to classify octree data representing bathtubs, beds, dressers, etc. Tatarchenko et al. [56] create a deep convolutional decoder that learns to decode dense input to octree representations of 3D objects like a car.

2.2 Super resolution for scientific data
There are two categories of super resolution: spatial super resolution (SSR) and temporal super resolution (TSR). Spatial super resolution aims to increase the spatial resolution of input data. Zhou et al. [73] use a 3D convolutional neural network (CNN) to perform SSR on volumes with better feature reconstruction than trilinear interpolation or cubic-spline interpolation. Guo et al. [13] use three neural networks in parallel to perform SSR on 3D vector fields. Xie et al. create tempGAN [70], which upscale fluid flows for temporally consistent high resolution output. Fukami et al. compare two ML-based SSR methods for 2D fluid flow [14]. SSR-TVd by Han and Wang [19] uses a generative adversarial network (GAN) for upsampling volumes such that the upsampled output frame is temporally coherent with adjacent frames. A similar line of research to SSR in the computer vision community is called image super resolution (ISR), and we refer readers to a survey by Wang et al. [67] for recent techniques. With the recent advances in super resolution, Jakob et al. have created a public 2D flow dataset with varying Reynold’s number, citing that ML methods are a powerful method for interpolating flow maps and data reduction [25].

Temporal super resolution aims to increase the temporal resolution of input data. TSR-TVd by Han et al. [29] uses a convolutional LSTM to learn the recurrence between timesteps efficiently to recover interpolated timesteps with higher accuracy than linear interpolation. Other methods perform both spatial and temporal super resolution together with spatiotemporal super resolution (STSR). Fukami et al. [15] performs STSR on fluid flow data using two networks in a 3D+1D approach, where the spatial dimension is upsampled first, and then the temporal dimension. MeshfreeFlowNet [27], proposed by Jiang et al., trains a network to allow inference for arbitrarily sized spatiotemporal domains with PDEs impacting training through a loss function.

2.3 Compression for scientific data
Compressors for scientific data can be classified as lossless or lossy. Lossless compressors, like FPC [10], have perfect reconstruction of the original data. Lossy compressors do not perfectly reconstruct the original data, but usually deliver larger compression ratios than lossless compressors in exchange. Lossy compressors can be further classified as non-error-bounded or error-bounded. Error bounded lossy compressors allow a user to specify a local error which the compressor will guarantee is not exceeded. Our approach presented in this paper is a lossy error-bounded lossy compressor. For a comprehensive overview of state of the art scientific data compression, we refer to surveys by Le et al. [34], Balsa et al. [6], and Hoang et al. [23].

Error-bound lossy compressors can be classified as prediction- or transform-based. FPZIP [39] and SZ [12] are prediction-based error-bounded lossy compressors. FPZIP uses a Lorenzo predictor [25] followed by integer mapping on both the predicted and actual data. Then the residuals between the predicted and actual data are sectioned into entropy codes for transmission. SZ creates an improved Lorenzo prediction and dynamically encodes blocks of data with either the Lorenzo prediction or an optimized linear regression method based on which method performs best. ZFP [39] is a transformation-based compressor that analyzes each 4\textsuperscript{d} sized block and encodes them with a specified number of bits ber block through an orthogonal block transform and embedded coding.

Most lossy compressors focus on absolute and relative errors per floating point number, which cannot guarantee global metrics such as structural similarity index measure (SSIM) or power spectra. Our approach uses neural network super resolution to combined with state-of-the-art compression techniques as a preprocessing step.
of the art lossy compressors to preserve global features better than lossy compressors alone, detailed in Sect. 4.2.3.

3 Overview

An overview of our data reduction and reconstruction pipeline is shown in Fig. 1. Our primary goal is data reduction while retaining global properties of the input data, such as peak signal-to-noise ratio (PSNR) and SSIM, and avoiding artifacts during visualization. Trained neural networks have shown strong performance when recovering these global metrics while generating realistic looking output [14, 18, 19, 33, 66, 67]. However, a single trained SR neural network will only be capable of a fixed $N \times SR$ in each dimension, which limits data reduction and reconstruction because the network works optimally when data is downscaled by $N \times$.

To improve data reduction for target error-bounds, we use a hierarchy of neural networks $G$ where each network in the SR hierarchy is trained to perform $2 \times$ super resolution between two downsampling levels. By using these networks in tandem, the SR hierarchy is capable of $2 \times, 4 \times, \ldots, 2^{|G|} \times SR$, where $|G|$ is the number of networks in the hierarchy. For data reduction purposes, this gives more flexibility because now data can be downscaled to any of those target scale factors. We detail the SR hierarchy structure and architecture in Sect. 4.1.

Though the SR hierarchy is an improvement on supporting only a single scale factor, there is no way to target error-bounds between what each SR factor offers with uniform downsampling and upsampling. For example, if $4 \times$ SR gives 40 dB PSNR and $2 \times$ SR gives 50 dB PSNR, there is no way to achieve 45 dB PSNR with a static downsampling ratio.

To give more flexibility to meet error-bounds between downsampling levels, we introduce a error-bounded dynamic downsampling algorithm, outlined in Sect. 4.2 that will downslope spatial regions of a single volume to a multiresolution data format. Scientific data do not always have complex features uniformly distributed throughout an entire volume; the dynamic downsampling algorithm will keep detail in complex regions and reduce resolution more in regions that are simple to upscale. To model the multiresolution data, we use an octree that is optimized for our purposes of dynamic downsampling, which we call an SR-octree. After the dynamic downsampling, data is stitched together into a representative volume and compressed using a compressor.

Though the multiresolution format may reduce storage costs, standard neural networks are not designed to perform multiresolution upsampling, and upsampling each octree node individually would introduce seam artifacts between nodes. To upscale multiresolution data and avoid seam artifacts, we design an algorithm that does not upscale octree nodes individually, but instead begins at the coarsest level of detail and uniformly upsamples an entire volume one level at a time until reaching the original full resolution. This algorithm is detailed in Sect. 4.3.

4 Our Method

Our neural network hierarchy for super resolution is outlined in Sect. 4.1. Next, we discuss the multiresolution data format we use and the dynamic downsampling algorithm together in Sect. 4.2. Finally, we discuss our solution for upsampling the multiresolution data with minimal seam artifacts in Sect. 4.3.

4.1 Spatial Super Resolution Hierarchy

We use a hierarchy of SR networks to improve data reduction by allowing multiple $SR$ factors. Most SR models support a static upsampling ratio of $N \times$ in each spatial dimension, which limits data reduction because the network only supports data downscaled by exactly $N \times$. Therefore, the compression ratio is limited to either $1 \times$ (no data reduction) and $N^P \times$, where $N$ is the upsampling ratio the model supports, and $D$ is the number of dimensions.

In the computer vision field, researchers have created models that support multiple scale factor SR such as LapSRN [32] and SinGAN [52]. The PSNR degrades as the scale factor increases, but multiple SR factors gives the user a choice for the reconstruction quality/data reduction trade-off. In this paper, we use a hierarchy of identical generative adversarial networks (GANs), with each one trained to perform $2 \times$ super resolution in each spatial dimension between downsampling levels. We define downsampling level $0$ as full resolution and downsampling level $i$ data that is downscaled by a factor of $2^i$ in each dimension. Together, they form a hierarchy of networks $G$ that supports up to $2^{|G|} \times SR$. An example of a SR hierarchy is shown in Fig. 2(e).

4.1.1 Spatial super resolution generator

We use a super resolution generator architecture based on the enhanced super-resolution generative adversarial network (ESRGAN) [66], a state-of-the-art GAN for upsampling single images from a low resolution input to a high resolution output. The full architecture is depicted in Fig. 2. We use the ESRGAN architecture due to three of the main features of the network: (1) Pre-upsample learning, (2) pixel/voxel shuffling, and (3) residual learning.

Pre-upsample learning allows us to reduce our memory requirement during training. Pre-upsample learning means that the input remains in the low-resolution input space when convolution kernel weights are learned before being upsampled to the final output resolution. This method is the opposite of post-upsampling, which first upsamples the data and then learns weights for convolution kernels in the high-resolution space. Training and inferring in this high dimensional space with post-upsampling learning is much more computationally expensive than in the low dimensional space because there is twice the number of pixels/voxels in each dimension which leads to $4 \times$ (in 2D) or $8 \times$ (in 3D) as many convolution operations to happen per layer compared to the low-resolution input. Given the large volumes to train with and limited memory budget, pre-upsample learning is a better choice.

Voxel shuffling is an upsampling method that allows the neural network to directly learn the upsampling function while being free from biases that other upsampling methods have. Recent work for volumetric super resolution has found success with a method called voxel shuffling, first presented in TSR-TVD [20], which is the 3D counterpart of pixel shuffling [53]. Voxel shuffling permutes a volume of shape $[u^C, L, W, H]$ to $[C, uL, uW, uH]$, where $u$ is the upsampling ratio, $C$ is the number of channels in the data, and $L, W,$ and $H$ are the 3 spatial dimension sizes. In our case, $u = 2$, and each spatial dimension is...
which allow deeper layers to use feature maps from earlier in the network. RRDB have proven to be powerful for SR tasks due to the skip connections in the dense blocks between shallow and deeper features, giving them their name, shown in Fig. 2(b). First, each dense block creates a residual which is added back to the input. Second, the entire RRDB creates a residual which is added back to the previous generator. If the previous generators have learned to accurately reconstruct features at their scale, then there would be no reason to learn those features again in this discriminator.

We use a GAN for two reasons. First, they have shown to be powerful for reconstructing important high-frequency features for both images as well as scientific data. Second, we expect that output from a GAN, specifically Wasserstein GANs (WGANs) [4], will more accurately match the distribution of ground truth data, which is important when using multiple super resolution generators from the SR hierarchy in tandem. When training, only ground truth data is used as both input and target output. Consequently, each network is not conditioned to super-resolve output from previous generators as in works like LapSRN [32] and SinGAN [52]. Therefore, the SR hierarchy is only learning to upscale data from within the ground truth distribution. By constraining the output of each generator to a specific distribution, that of the real data, we should see less compounding errors as networks are used sequentially. This distribution difference minimization is precisely the goal of WGANs.

4.1.3 Loss functions

We use two components in our loss function: a reconstruction component and a adversarial component, with a weight factor for each. The loss function is computed as:

\[ \mathcal{L} = \lambda_0 \mathcal{L}_{\text{rec}} + \lambda_1 \mathcal{L}_{\text{adv}} \]  

where \( \lambda_0 \) is the weight factor for \( \mathcal{L}_{\text{rec}} \), our reconstruction loss, and \( \lambda_1 \) is the weight factor for \( \mathcal{L}_{\text{adv}} \), our adversarial loss.

For the following loss functions, we denote the output from our network as \( \hat{F} \) and the ground truth as \( F \). Our generator \( G \) performs \( G(F^{LR}) = \hat{F} \), and we denote our discriminator as \( D \).

Reconstruction loss. The reconstruction loss is defined as

\[ \mathcal{L}_{\text{rec}} = \frac{1}{L \times W \times H} \sum_{i \in \text{vols}} ||F_i - \hat{F}_i||_1 \]  

where \( || \cdot ||_1 \) is the \( l^1 \) norm. We use the \( l^1 \) norm instead of the \( l^2 \) norm because the \( l^2 \) norm tends to have overly-smoothed output [67]. \( \mathcal{L}_{\text{rec}} \) is subject to minimization for the generator during optimization.

Adversarial loss. We use the WGAN-gradient penalty (WGAN-GP) loss [17] for two reasons. First, we expect that while super resolving multiple times through a hierarchy, a compounding errors may occur. Since WGAN [4] aims to minimize the Earth-Mover’s distance (Wasserstein-1 distance) between the output of the neural network and the ground truth distributions, we can expect that neural network output should have an output distribution that better matches the ground truth data. This may help reduce compounding errors when performing high scale factor super resolution. Additionally, the gradient penalty increases training stability. The loss is defined as

\[ \mathcal{L}_{\text{adv}} = \mathbb{E}_{F \in \mathbb{P}_F} [D(F)] - \mathbb{E}_{\hat{F} \in \mathbb{P}_{\hat{F}}} [D(\hat{F})] \]  

where \( \mathbb{E}[] \) is the expectation operator, \( \mathbb{P}_F \) is the model distribution of upscaled volumes, and \( \mathbb{P}_{\hat{F}} \) is the data distribution of the full resolution ground truth volumes. \( \mathcal{L}_{\text{adv}} \) is subject to minimization for the generator through, which helps work against the vanishing gradient problem.

4.1.2 Spatial super resolution discriminator

Each generator is coupled with a patch discriminator, shown in Fig. 2(d), which classifies overlapping patches as real or fake. This network is designed to be relatively shallow with a small receptive field. Each Conv block (convolution, batch normalization, leaky rectified linear unit (LeakyReLU)) has a kernel size of 3 and a stride of 1. With only 5 layers there is a corresponding receptive field of 11 for each discriminator. This small receptive field is to prevent redundant learning across different scales. By keeping the receptive field small, the patch discriminator will only be judging the realness of local features in a small window, which were the responsibility of the paired generator to synthesize. Global features present in the data will either be ground truth low resolution data (which are already real), or are synthesized features from a previous generator. We use a GAN for two reasons. First, they have shown to be powerful for reconstructing important high-frequency features for both images as well as scientific data. Second, we expect that output from a GAN, specifically Wasserstein GANs (WGANs) [4], will more accurately match the distribution of ground truth data, which is important when using multiple super resolution generators from the SR hierarchy in tandem. When training, only ground truth data is used as both input and target output. Consequently, each network is not conditioned to super-resolve output from previous generators as in works like LapSRN [32] and SinGAN [52]. Therefore, the SR hierarchy is only learning to upscale data from within the ground truth distribution. By constraining the output of each generator to a specific distribution, that of the real data, we should see less compounding errors as networks are used sequentially. This distribution difference minimization is precisely the goal of WGANs.

The ESRGAN architecture makes use of residual learning, which has been shown to improve results and stability of deep convolutional neural networks [21]. Residual learning means that instead of learning an output directly, the network instead learns the difference between the input and the expected output. ESRGAN employs residual learning across many layers with \( d \) sequential residual-in-residual dense blocks (RRDB). RRDB have proven to be powerful for SR tasks due to the skip connections in the dense blocks between shallow and deeper features, which allow deeper layers to use feature maps from earlier in the network to influence weight learning. RRDBs also add two levels of residuals within them, giving them their name, shown in Fig. 2(b). First, each dense block creates a residual which is added back to the input to the block. Second, the entire RRDB creates a residual which is added back to its input. Each residual is multiplied by a constant \( \beta \) between 0 and 1 (chosen empirically) that controls the influence of each residual. This follows a technique called residual scaling which prevents instability in deep networks. After input has gone through the \( d \) RRDBs, that output is finally used as another residual and added back to a shallow feature map from before the first RRDB. All of the residuals used offer many pathways for gradients to be computed increased by a factor of 2, while the channel dimension reduces by a factor of 8. We use pixel shuffling in 2D, and voxel shuffling in 3D.

The ESRGAN architecture makes use of residual learning, which has been shown to improve results and stability.
network, to make the generator capture the data distribution of ground truth data well, and is subject to maximization for the discriminator, so that the discriminator can accurately tell real high-resolution volumes apart from synthesized volumes from the generator.

4.2 Dynamic downscaling to a multiresolution data format

Now that we have an NN hierarchy $G$ that can perform $2^i \times$ SR, for $2 \leq i \leq |G|$, we give more flexibility to the data reduction process by dynamically downsampling data according to the presence of features instead of uniformly downsampling. Scientific data do not always have complex features present throughout the entire volume, and large regions of a volume may be constant. Therefore, instead of uniformly downsampling a full volume by a single scale factor, we use an octree data representation and dynamic downsampling algorithm to reduce a volume dynamically while still meeting an error bound after SR.

4.2.1 Dynamic downscaling

Our dynamic downsampling algorithm is described with the following pseudocode, beginning with the full resolution volume $F$.

1. Downscale the input data $D$ by $2 \times$ in each dimension to $D \downarrow$.
2. Reconstruct $\hat{F}$, the full resolution approximation of $F$, from the octree data including $D \downarrow$ by using our multiresolution SR algorithm (covered in Sect. 4.3).
3. Check to see if the error between $F$ and $\hat{F}$ is acceptable.
   a) If the error is less than the error-bound, see if $D \downarrow$ can be downscaled further (return to step 1 with $D \downarrow$).
   b) Otherwise, reset $D \downarrow$ to $D$. Split $D$ into its octants, and recurse on each octant (return to step 1 for each octant).

Note that step 2 reconstructs $\hat{F}$, the full resolution approximated volume of $F$, the ground truth. Though this is more expensive than checking octree nodes individually, we upscale all nodes together using our multiresolution super resolution algorithm because the CNN will create artifacts along boundaries due to zero padding. In neural networks, convolution layers use zero padding to guarantee that the output feature maps have the same spatial dimensions as the input to the layer. Guaranteed output size is desirable for specific operations like pixel/voxel shuffling. As a drawback, when convolutional kernels convolve with the edges and corners, the zero padding will influence the convolved feature map and increase error. This effect is magnified in deeper networks since multiple convolution layers with zero padding will feed in to each other. Zero padding already influences boundary voxels of output from an SR neural network, but performing SR on individual octants would increase the number of voxels affected by zero padding leading to an increased error along the boundaries between upscaled octants. Therefore, we develop the multiresolution super resolution algorithm to eliminate zero padding influence which gives us an approximation $\hat{F}$ that has no seams and reports and accurate error for our error-bound.

We have three stopping criteria for any octant to not be subdivided further (3b in the above mentioned algorithm). First, if the size of an octant is only $2^D$ regardless of the downsampling level, where $D$ is the number of dimensions, then it is not useful to split any more, since those octants of size $1^D$ cannot be downscaled further. Second, octants cannot be downscaled past $\text{MAXDSL} = |G|$ because the neuronal network hierarchy cannot perform SR past that scale factor. Our final stopping condition is a $\text{minChunk}$ hyperparameter that limits the algorithm from spatially subdividing if a data chunk only represents $\text{minChunk}$ voxels in each dimension. Smaller $\text{minChunk}$ will trade off compression rate for time.

The worst case time complexity of our dynamic downsampling algorithm is $O(2 \times \left\lceil \log_2 N \right\rceil \cdot \text{MAXDSL} \cdot \text{SR})$, where $O(\cdot)$ is the time complexity, $N$ is the spatial dimension with the fewest number of voxels in it, and $O(\text{SR})$ is the time complexity for the multiresolution super resolution algorithm. The worse case happens when region can be downscaled at all. The best case time complexity is $O(\text{MAXDSL} \cdot O(\text{SR}))$, and happens when the entire volume can be downscaled to the lowest resolution while still meeting the error criteria. Since neural network inference can be relatively slow compared to other upsampling algorithms (i.e. $O(\text{SR})$ is large, so step 2 in the pseudocode for dynamic downsampling at the beginning of Sect. 4.2 is computationally expensive), we use linear interpolation as an efficient conservative heuristic/proxy for the neural network’s reconstruction quality during dynamic downsampling. This reduces $O(\text{SR})$, but limits data reduction since the reconstructed data will likely have higher quality than the bound given. Since we use the interpolation heuristic, we perform a check after the dynamic downsampling finishes that uses the NN hierarchy instead of the interpolation proxy to ensure that the SR reconstructed data does meet the target error bound. If it is not met (which empirically has never happened), the dynamic downsampling process is repeated with a target error bound that is progressively more strict until the error bound is met.

![Fig. 3. An example of the progression of a volume (square) and SR-octree (tree structure beneath each square) during the dynamic downsampling process using a quadtree for a simple visualization. Data starts at full resolution with no downsampling in (1). A trial and error step failed, so it was subdivided the input into its quadrants in (2). The same is repeated on the top left quadrant in (3). During (4, 5), the trial and error for the light orange (2 ×) octants were successful, so that data is kept at downsampling level 1. The final example SR-octree is shown in (10).](image)

4.2.2 SR-octree

To model the multiresolution data in our dynamic downsampling algorithm, we create a modified octree data structure called an SR-octree. Our SR-octree is designed to optimize data reduction time for our dynamic downsampling algorithm by minimizing the number of nodes in the tree and keeping data contiguous until it is spatially divided.

Like most octrees, children of a parent node represent the 8 octants that compose the parent. Unique to the SR-octree is a $\text{downsampling level}$ attribute on each node that represents the factor that data has been downscaled in each spatial dimension. This attribute allows us to represent contiguous large regions with a uniform downsampling ratio with one node, because our dynamic downsampling algorithm will only subdivide nodes when it is necessary. In Fig. 3, we show what the SR-octree looks like during the dynamic downsampling process. White nodes are “dummy nodes” that hold no data and only point to their child octants. Nodes are colored according to their downsampling level shown by the key on the left.

4.2.3 Compression stage

After the SR-octree $T$ is output from the dynamic downsampling algorithm, we use SZ, a state of the art error-bounded lossy compressor, to compress the multiresolution data. Though we use SZ, any compressor supporting grid-structured floating point data may be used.

Instead of compressing each node’s data separately, which would have a large overhead for each node, we combine all data into a single $\text{representative volume}$, shown in Fig. 1b)-left, to compress. To fill
the representative volume with the SR-octree data, we stitch together all nodes into an empty full resolution volume. Since some data is downscaled, we have empty space in the volume, which we choose to fill the the average data value so that run-length encoding can efficiently compress these areas.

Starting with a full resolution empty volume might be excessive if all data has been downscaled by a minimum scale factor. For example, if all nodes in the SR-octree have a downsampling level of at least 1, then we can instead stitch together the SR-octree nodes into an empty volume that is half of the original volume size in each spatial dimension. This extends for any minimum downsampling level.

After creating the representative volume, we compress with SZ. We want to compress without distorting the data such that the reconstructed volume’s error after SR is still acceptable. Since CNNs learn spatial statistics, we use a pointwise error bound. We use a bound of $\epsilon = \sqrt{\max - \min}$ for SZ’s pointwise error-bound compression, where $P$ is the reconstructed PSNR in decibels and $F$ is the original volume. This $\epsilon$ guarantees that the PSNR of the (de)compressed representative volume will be at least $P - \epsilon$ dB, and serves as a starting point for our target error bound that scales with the reconstructed PSNR so that compression scales with the error-bound. We perform a sanity check here that performs decompression and multiresolution super resolution with the NN hierarchy to ensure that error-bound is still met, which empirically has always be the case. When the error-bound is not met, we use a binary search approach to find the optimal $\epsilon$.

Fig. 4. An example of upscaling multiresolution data with neural networks by (a) upscaling nodes individually (b) our proposed multiresolution super resolution algorithm.

4.3 Multiresolution super resolution for SR-octrees

Lastly, we present our algorithm for upscaling our SR-octree multiresolution data format while avoiding seam artifacts. A naive solution to upscale the multiresolution data is to use the pre-trained neural networks to perform super resolution on each data chunk individually based on the downsampling factor each node is saved with. The upsampled individual chunks would finally be stitched together into a single full resolution volume of data. However, this approach introduces seam artifacts, or boundary artifacts, between data chunks, illustrated in Fig. 4. Seams are a problem common to hierarchical rendering techniques [42,61,63], but is even more pronounced when using neural networks due to zero padding influence outlined in Sect. 4.2.

Therefore, we require an approach that upscales all data together instead of separately to avoid seam artifacts. Though there is data distributed across many different downsampling levels, the only level that can uniformly represent the data without any separate upsampling is at the largest downsampling level, since smaller downsampling levels can be reduced further to the largest downsampling level. This gives us a uniform low resolution version of the raw data that we can begin the upsampling process at that will not create seam artifacts when upscaled.

4.3.1 Upscaling process

Our upsampling process begins at the largest downsampling level $\text{MAXDSL} = |G|$, where $G$ is our SR hierarchy. Since all data is saved with at most $\text{MAXDSL}$, this means that all data nodes can be individually downscaled to the maximum downsampling level and stitched together into a single volume. From here, we upscale this volume to $\text{MAXDSL} - 1$ which will not introduce seam artifacts since the full volume was upscaled at once. The upscaled volume is now an approximation of the ground truth raw data at $\text{MAXDSL} - 1$ with no seam artifacts. However, we do not need to use approximated voxels when downsampling data exists in the SR-octree. If a node in the octree has $\text{downsampling level} \leq \text{MAXDSL} - 1$, then we opt to replace the approximated voxel values in our volume with the more accurate voxel values from the node representing them. We call these approximated values that should be overwritten stale voxels. We can repeat our upscaling to stale voxel overwriting until we reach $\text{downsampling level} = 0$. Since we are only ever upsampling the full volume instead of many small volumes, we are not introducing seam artifacts on node boundaries.

4.3.2 Downscaling process

In our upsampling process (Sect. 4.3.1), we claim that all data can be downscaled uniformly to $\text{MAXDSL} = |G|$, and begin upsampling one level at a time starting there. However, downsampling all nodes to $\text{MAXDSL}$ and putting them together into a volume is not always possible because very small nodes (limited by $\text{minChunk}$ in Sect. 4.2) cannot always be downscaled to $\text{MAXDSL}$. As an example, a single voxel node cannot be downscaled by a factor of 64.

We resolve this issue with a downsampling process which mirrors our upsampling process. The primary issue is that some small nodes at low downsampling levels cannot be downscaled to $\text{MAXDSL}$ in one large downsampling step. Instead, we can take many smaller downsampling steps which mirrors how we take many small upsampling steps. Our SR-octree construction guarantees that single voxel nodes at $\text{MINDSL}$, where $\text{MINDSL}$ is the minimum downsampling level present in any node of an SR-octree $T$, must have siblings with the exact same size (a single voxel) and downsampling level. Therefore, these siblings can be joined to make a larger node that is $2^D$ that can be downscaled by $2 \times$.

This motivates the following downsampling process:

1. Set $\text{MINDSL} = 0$.
2. Combine all single voxel data nodes that have $\text{downsampling level} = \text{MINDSL}$ to a single node of size $2^D$.
3. Downscale all nodes with $\text{downsampling level} = \text{MINDSL}$ by $2 \times$ each in each spatial dimension.
4. Set $\text{MINDSL} = \text{MINDSL} + 1$.
5. If $\text{MINDSL} = \text{MAXDSL}$, finish. Else, return to the step 2.

After finishing, all data is downscaled to a single volume at $\text{MAXDSL}$.

Since we downscale regions of data by a factor of 2 multiple times, we require the following property in the downsampling algorithm used. Given a full resolution frame $F$, scale factor $S$, and downsampling algorithm $\text{down}(F, S) : \mathbb{R}^{|F|} \rightarrow \mathbb{R}^{|F|/2}$, it must hold that $\text{down}(F, 2 \times S) = \text{down}(\text{down}(F, S), S)$. When this property holds, there is no effect from downsampling a region multiple times compared to downsampling it once at a large downsampling factor. For instance, downsampling a region by a factor of $2 \times$ three times would be the same as downsampling it by $8 \times$ once. We see this property in all pooling algorithms (min, max, mean pooling) as well as subsampling.

4.3.3 Multiresolution super resolution end-to-end, optimization, and time complexity

To summarize the multiresolution super resolution algorithm, the downsampling process will first combine and downscale nodes, starting from the smallest downsampling level to the largest downsampling level, by a factor of $2 \times$ at a time. Then, the uniform low resolution data is upscaled by the neural network one downsampling level at a time, with stale voxels overwritten with non-approximated data when available. Since all data is upscaled uniformly step by step, there are no seams introduced, and the neural network does not suffer from boundary artifacts at each node’s edges because zero padding is minimized.

The time complexity of this SR-octree super resolution algorithm with a caching optimization is $\mathcal{O}(|N| + \text{MAXDSL}(|\mathcal{O}(DS) + \mathcal{O}(SR)|))$. }
5 Evaluation

We evaluate our framework through a series of experiments that compare our approach to baseline approaches across three datasets. Information about each dataset is listed in Sect. 5.1. In Sect. 5.2, we describe hyperparameters and training procedure used for training the NN hierarchies that perform SR on low-resolution data as described in Sect. 4.1 and illustrated with Fig. 2(e). In Sect. 5.3, we evaluate the neural network SR results against bilinear interpolation and bicubic interpolation baselines for 2D datasets, and trilinear interpolation baseline for 3D datasets. Sect. 5.4 compares our method with and without dynamic downscaling. In Sect. 5.5, we evaluate our data reduction/reconstruction pipeline by comparing PSNR/SSIM metrics over varying compression ratios, power spectra, and renderings for our method against our baseline state of the art lossy compressor SZ [36].

5.1 Datasets

We experiment with three scalar field datasets from Johns Hopkins Turbulence Databases (JHTDB) [35] that are time-varying results from a direct numerical simulation (DNS) for turbulent fluid flow. Information about each dataset and corresponding trained hierarchies are shown in Table 1.

Table 1. Iso3D magnitude is a 3D velocity magnitude dataset from a DNS of turbulent fluid flow at Taylor-scale Reynolds number $R_{t} \approx 433$, hosted in the “isotropic1024coarse” dataset in the JHTDB. We take the middle z-axis slice ($z=512$) of the Iso3D magnitude dataset to create an Iso2D magnitude dataset for testing our approach with 2D data. Lastly, we use the pressure field from the “mixing” dataset in JHTDB (Mixing3D pressure).

5.2 Network training and hyperparameters

One neural network hierarchy is trained per dataset, with settings for each shown in Table 1. We perform a training/test set split by training on the first $k$ timesteps and testing on the final $N - k$ timesteps, where $N$ is the total number of timesteps for the dataset (# train + # test in Table 1), and $k$ is the number of testing timesteps. We train this way to mimic a use case where a network is trained on data already available and then applied on newly generated simulation results.

During training, networks are optimized with the Adam optimizer [28] with default decay rates $\beta_1 = 0.9$, $\beta_2 = 0.999$ for the first and second moment estimation. We use $\lambda_0 = 1.0$ and $\lambda_1 = 0.1$ for our loss function defined in Sect. 4.1.3. All networks use 96 kernels in every convolution layer (see Fig. 2) and 3 RRDBs to achieve a good balance of computational efficiency/memory use and reconstruction quality. Following Wang et al. [60], the RRDBs use $\beta = 0.2$ for residual scaling, explained in Sect. 4.1.1. Table 1 shows how many models are in each trained hierarchy. We use PyTorch's DistributedDataParallel to train in parallel across 8 NVidia A100 40GB Tensor Core graphics cards on a single NVidia DGX A100 system. Since the full resolution 3D data cannot fit in the GPU’s memory during training, we crop training volumes to $128^3$ and use a random starting position for the crop to increase training example diversity. Each model is trained for 50 epochs, at which point each had converged. Training time takes between 3–5 hours, but is only necessary once per dataset. Storage cost for each model is listed in Table 1.

5.3 Evaluating neural network hierarchy SR

We provide PSNR/SSIM charts for each scale factor trained in each network in Fig. 5 plotted against interpolation baselines. Our results show that the NNs are properly trained and outperform the baseline SR methods for each scale factor for both PSNR and SSIM. This results also verifies that using multiple trained neural networks in tandem for SR factors of $4 \times$ or higher does not cause unforeseen reconstruction artifacts which is important because the networks were not trained conditional on the output of previous networks.

5.4 Dynamic downscaling and SR-octree evaluation

In Fig. 6, we can see that using our approach (in blue) that dynamically downscales data to the SR-octree allows for more flexibility in terms of target PSNR/SSIM error-bounds than our approach with dynamic downsampling disabled (in orange). To disable dynamic downsampling in our approach, we use the same algorithm presented in Sect. 4.2 but do not split data into its octants when uniform downscaling cannot occur (step 3b). The result of this augmented process is a uniformly downscaled volume instead of a multiresolution volume. Since data cannot be downscaled at arbitrary scale factors, only $2^i$ for $1 \leq i \leq |G|$, where $G$ is the NN hierarchy, the compression rate will increase as $i$ increases, but the only data points will be on those few samples between $i = 1$ and $i = |G|$. Since error-bounds between those points will default to the lower downscaling level to have an acceptable error, our approach without dynamic downsampling and an SR-octree follows a step function. Introducing our SR-octree and dynamic downsampling allows for PSNR/SSIM targets to bridge the gap between downsampling...
levels more smoothly, and enables higher compression ratios for given error-bounds as a result.

5.5 Data reduction and reconstruction evaluation
Quantitative results with PSNR/SSIM over compression ratios.
The charts in Fig. 6 shows PSNR and SSIM over compression ratios for our approach against SZ alone. Since our neural networks can begin downscaling around the 45-55dB PSNR mark (shown in Fig. 5), our framework’s best performance is under that PSNR error-bound. In Iso2D and Iso3D, our approach can offer up to 4× to the compression of SZ alone with the same or larger error bound in some cases between 30-40 dB PSNR, with results more comparable to SZ alone above 50 PSNR. The 3D Mixnet test has the worse results relative to the other two datasets tested which may be due to the training and test set division of the dataset. From the “mixing” dataset information page from JHTDB [41], we notice that the turbulent Reynolds number is much larger on average in the first part (training timesteps) of the dataset compared to the later part (testing timesteps). The network may have learned features from the more turbulent timesteps, which gives worse results on the testing timesteps with less turbulence. Our results could be improved for the Mixing3D dataset by adjusting the training data, or with a fine-tuned error-bound value for post-processing compression in our data reduction pipeline to allow SZ to do more compression.

Quantitative results with power spectra. In Fig. 7 we observe comparable performance between our method and SZ alone in the preservation of power spectra for reconstructed data at similar compression ratios. The 1D power spectrum is created by taking the average power of a 2D/3D FFT (depending on dataset dimensionality) at varying wavenumbers up to the Nyquist frequency for each reconstructed volume at a fixed compression ratio. In the Iso2D data, we can see that our output almost aligns with the ground truth up to wavenumber k=100. We also show the power spectrum for the SR-octree, which is the multiresolution saved data format that is input to our multiresolution super resolution algorithm. We notice that the SR-octree has larger power in higher frequencies compared to our final output due to how the multiresolution super resolution algorithm for each trial and error step.

One limitation of our approach is that the spikes associated with block artifacts are completely removed, demonstrating that our approach removes the seam artifacts. Preservation of strict pointwise error bounds. The last limitation we address is neural networks weakness during our data reduction framework for pointwise error bounds. Empirically, we find that compression ratios at a fixed pointwise error bound for our approach are worse than SZ alone. Neural networks learn from loss values that average error across the entire spatial domain which can give a good result on average, but does not punish the network for having high pointwise error outliers, resulting in poor pointwise error-bound performance.

6 LIMITATIONS
This section comments on the limitations of our approach and how future work may address those limitations. We address the computational efficiency of our dynamic downscaling, the use of 2× scale factor SR between the neural network hierarchy levels, and neural network’s weakness at pointwise error bounds.

Data reduction computational efficiency. One limitation of our approach is the time it takes for data reduction because we use our multiresolution super resolution algorithm for each trial and error step. Recall that during dynamic downscaling we use linear interpolation as a conservative approximation for the quality the neural network will produce, though we continue use the neural networks to perform SR during data reconstruction. Though the interpolation proxy is faster than the NNs during data reduction, our data reduction process can still take up to 4 minutes depending on volume size, shown in Table 1. To improve data reduction speed at the cost of compression rate, nodes could be processed in parallel individually instead of upscaled together. Time complexity can also be improved if the trial and error steps in our pipeline can be eliminated. Alternatively, the error-bound could be removed and the data reduction algorithm could operate as a non-error bounded lossy compressor instead.

Handling of non-power-of-two sized volumes. Another limitation is our use of a 2× scale factor upsampling within each neural network in the SR hierarchies. This limits our SR capabilities to only 2i, for 1 ≤ i ≤ |G|. Therefore, data that have spatial dimensions that are not a power of 2 are not compatible, and will need to be resampled or cropped. To accommodate data with more spatial dimension sizes, our hierarchy could be extended to have 3×, 5×, and other prime number scale ratios. This would increase what spatial dimensions are supported, but is not exhaustive, and also requires many more neural networks to be trained. For support of arbitrary spatial dimension sizes, another kind of NN architecture is needed.

7 CONCLUSION AND FUTURE WORK
In this paper, we present a multiresolution super resolution algorithm for upsampling hierarchical data using neural networks without introducing seam artifacts on node boundaries. The multiresolution super resolution algorithm is comprised of a downscaling process followed by an upsampling process to avoid introducing seam artifacts between nodes. We demonstrate one use of this algorithm within a data reduction and
reconstruction pipeline for scientific data. Our quantitative results show that our method can enhance SZ’s compression capability with better PSNR/SSIM at similar compression ratios, especially as compression ratios grow. Our qualitative results show that our method avoids seam artifacts between node boundaries while also retaining visual features better than SZ alone. In future work, the compressor and neural network architecture can be swapped out for other compatible compressors or architectures. We experiment with SZ and ESRGAN, but any compressor capable of compressing grid-structured or multiresolution data could be used instead, and any neural network architecture capable of 2× SR can be used as the generator in the NN hierarchy. Specialized network architectures, such as a network created for vector field SR,
may perform better in their specialized domain. Outside of data reduction, the multiresolution super resolution algorithm we propose may be useful in other scientific visualization algorithms, such as realtime multiresolution volume rendering.

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