GACT: Activation Compressed Training for Generic Network Architectures

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Abstract

Training large neural network (NN) models requires extensive memory resources, and Activation Compressed Training (ACT) is a promising approach to reduce training memory footprint. This paper presents GACT, an ACT framework to support a broad range of machine learning tasks for generic NN architectures with limited domain knowledge. By analyzing a linearized version of ACT’s approximate gradient, we prove the convergence of GACT without prior knowledge on operator type or model architecture. To make training stable, we propose an algorithm that decides the compression ratio for each tensor by estimating its impact on the gradient at run time. We implement GACT as a PyTorch library that readily applies to any NN architecture. GACT reduces the activation memory for convolutional NNs, transformers, and graph NNs by up to $8.1 \times$, enabling training with a $4.2 \times$ to $24.7 \times$ larger batch size, with negligible accuracy loss.

1. Introduction

In recent years, we have witnessed the trend of using larger and larger neural network (NN) models to deliver improved accuracy and generalization in various machine learning tasks (Devlin et al., 2018; Fedus et al., 2021). However, training these models requires a considerable amount of on-device GPU memory. Unfortunately, the increase of GPU memory capacity has been relatively slow, leading to a fundamental barrier to the development of large NN models.

Activation Compressed Training (ACT) is a promising approach to reduce the memory footprint of models during training. As all layers’ activations need to be kept in the memory for computing the gradients during training, ACT reduces memory consumption by compressing these saved activations. Prior work (Chakrabarti & Moseley, 2019; Fu et al., 2020; Chen et al., 2021; Evans & Aamodt, 2021) has shown the effectiveness of ACT by reducing activation footprint by up to $12 \times$ with 2-bit activations.

Although ACT has already demonstrated impressive compression capabilities, previous work on ACT is restricted to specific NN architectures. For example, ActNN (Chen et al., 2021) is a quantization framework for convolutional NNs only; Mesa (Pan et al., 2021) proposes a per head/layer quantization method for vision transformers; and AC-GC (Evans & Aamodt, 2021) derives convergence error bound for different types of operators separately.

Developing a generic ACT framework is challenging. Theoretically, convergence guarantees must be made without assumptions on the network architecture. Algorithmically, the framework should find effective compression strategies for all kinds of networks automatically. From the system perspective, the framework should support arbitrary NN operations, including user-defined ones.

In this work, we propose GACT, a general framework for ACT that is agnostic to the NN architecture. Neither specialized mathematical derivations nor customized implementation is needed to support different operators. To enable this, we develop a general convergence theory by analyzing the stochastic gradient (SG) introduced by ACT. We show that the SG can be well approximated by a linearized version, which is unbiased to stochastic compressors. The variance of the linearized gradient has a particularly simple structure that allows a numerical algorithm to predict the variance given a compression strategy. Then, we generate the strategy by approximately solving an integer program.

We implement our method as a library based on PyTorch that can be quickly integrated into real-world machine learning systems. The library also provides several optimization levels to explore the trade-off between memory and speed. We demonstrate the flexibility and efficiency of GACT on various tasks, including image classification, object detection, text, and graph node classification. Our evaluation shows that GACT can reduce activation memory by up to $8.1 \times$, enabling training with a $24.7 \times$ larger batch size on the...
same GPU. In sum, our main contributions are as follows:

- We propose a general convergence theory for ACT.
- We develop an algorithm that automatically estimates the sensitivity of each compressed tensor and selects the optimal compression strategy.
- We build efficient implementation of GACT in PyTorch with an easy-to-use API that can also be combined with other memory-saving techniques seamlessly.

2. Related Work

Activation Compressed Training. ACT has been applied to convolutional NNs using different compressors, such as quantizers (Chakrabarti & Moseley, 2019; Fu et al., 2020; Chen et al., 2021), JPEG (Evans et al., 2020), or scientific data compression algorithms (Jin et al., 2021; Evans & Aamodt, 2021). ACT is also applied to transformers (Pan et al., 2021) and graph NNs (Anonymous, 2022).

However, the existing theory for ACT (Chakrabarti & Moseley, 2019; Fu et al., 2020; Chen et al., 2021; Evans & Aamodt, 2021) relies on the case-by-case analysis of specific network operators, such as convolution, ReLU, and batch normalization. It also requires dedicated implementations for each operator. On the contrary, GACT focuses on the generality of activation compressed training, not a specific quantizer design, which is the main topic of previous work. Instead of assuming that the network is a stack of layers, GACT formulates the problem as a computational graph of operators. This is general enough to cover transformers (Vaswani et al., 2017), graph NNs (Kipf & Welling, 2016), second-order derivatives, and unknown future architectures.

Reduced Precision Training. Apart from ACT, reduced precision training (Micikevicius et al., 2018; Wu et al., 2018; Wang et al., 2018c; Banner et al., 2018; Chen et al., 2020a; Sun et al., 2020) performs calculations directly on low precision data, reducing the computation cost and memory footprint simultaneously. To achieve this, specialized kernels are used to calculate on low precision data in contrast, ACT only considers storage, and it can thus use more flexible compression strategies and achieve a much better compression ratio with the same accuracy loss.

Memory-Efficient Training. Gradient checkpointing (Chen et al., 2016; Jain et al., 2019) trades computation for memory by dropping some of the activations in the forward pass from memory and recomputing them in the backward pass. Swapping (Kirisame et al., 2020; Huang et al., 2020; Wang et al., 2018b; Peng et al., 2020) offloads activation or model parameters to an external memory (e.g., CPU memory). Recent work (Beaumont et al., 2021) explores the possibility of combining the gradient checkpointing and swapping. All these methods save memory by storing fewer tensors on the GPU. In contrast, GACT compresses the saved tensors and is complementary to these approaches. Moreover, the generality of GACT enables easy combination with these methods, which we explore in this paper.

3. Formulation

We first present the mathematical formulation of our activation compressed training (ACT) framework. As we would like to develop a general ACT algorithm, applicable to a wide range of NN architectures, we make minimal assumptions on our formulation. Throughout the paper, we define the variance of a vector $x$ as $\text{Var}[x] = \mathbb{E}[||x||^2] - \mathbb{E}[x]^2$.

3.1. Activation Compressed Training

In this work, we abstract the forward propagation as two functions $\ell(x; \theta)$ and $h(x; \theta)$. Both take a datum $x$ and the model parameter $\theta$ as the input. The loss function $\ell(x; \theta)$ outputs the loss of the network $\theta$ on datum $x$. The context function $h(x; \theta)$ outputs tensors to be stored in the memory for computing the gradients, which are referred as the context. Assume that the context consists of $L$ tensors, where each tensor $h^{(i)}(x; \theta)$ is represented by a flattened $D_i$-dimensional vector. Denote $h(x; \theta) = (h^{(i)}(x; \theta))_{i=1}^L$. Our notations are somewhat unconventional in the sense that we do not explicitly define each layer’s activation. We do not even assume that there is a NN. It could be any computational graph that saves context tensors.

Given a dataset $X = \{x_n\}_{n=1}^N$, define the batch loss $L(\theta) := \frac{1}{N} \sum_{n=1}^N \ell(x; \theta)$. The dataset can be equivalently represented as an empirical data distribution $p_X(x) := \frac{1}{N} \sum_{n=1}^N \delta(x - x_n)$, where $\delta$ is the Dirac delta function. The batch loss can be written as $L(\theta) = \mathbb{E}_X[\ell(x; \theta)]$, where $\mathbb{E}_X$ denotes for taking expectation over $p_X$.

The network is trained with stochastic gradient descent (SGD) (Bottou, 2010). Starting from an initial model $\theta_0$, at the $t$-th iteration, SGD updates the model with:

$$
\theta_{t+1} \leftarrow \theta_t - \eta \nabla_\theta \ell(x; \theta_t),
$$

where $\eta$ is a learning rate, and the SG $\nabla_\theta \ell(x; \theta)$ is computed on a random datum $x \sim p_X$. Notice that $\mathbb{E}_X[\nabla_\theta \ell(x; \theta)] = \nabla_\theta L(\theta)$, i.e., the SG is an unbiased estimator of the batch gradient $\nabla_\theta L(\theta)$.

Crucially, the SG can be written in the form $\nabla_\theta \ell(x; \theta_t) = g(h(x; \theta_t); \theta_t)$. In other words, the back propagation only depends on the forward propagation through the context $h(x; \theta_t)$. The entire context must be kept in memory for computing the gradients. The context dominates the memory consumption in many applications.

ACT reduces the training memory footprint by compressing the context. Let $Q(h)$ be a compressor, which converts $h$
We refer to the key technique in this work is to construct an unbiased function \( g \) itself is unbiased.

**3.2. Convergence of ACT**

ACT is a lossy approximation of SGD, as it uses an approximate gradient \( g(Q(h); \theta) \). Therefore, some kind of theoretical guarantee is required for ACT to be useful. Fortunately, analyzing ACT is made significantly simpler by introducing an unbiased stochastic compressor \( Q(\cdot) \), such that \( \mathbb{E}_Q [Q(x)] = x \) for any \( x \). \( \mathbb{E}_Q [\cdot] \) means taking expectation over the compressor. In this way, \( g(Q(h); \theta) \) can be viewed as a stochastic estimator of the batch gradient \( \nabla L(\theta) \), but the randomness comes not only from the datum \( x \) but also the compressor \( Q(\cdot) \). Therefore, ACT is still an SGD algorithm. Standard analytical tools for SGD (Bottou et al., 2018) are applicable for studying ACT.

SGD algorithms have particular good properties when the SG is unbiased. In our case, this means \( \mathbb{E}_Q [g(Q(h); \theta)] = g(h; \theta) \). However, the SG is biased general, even when the stochastic compressor itself is unbiased.\(^1\)

The key technique in this work is to construct an unbiased approximation of the AC gradient by linearizing the gradient function \( g(\cdot; \theta) \). Consider the first-order Taylor expansion of \( g(\cdot; \theta) \) at \( h \):

\[
\hat{g}(Q(h); h, \theta) := g(h; \theta) + J(h, \theta) \Delta h, \tag{3}
\]

where \( J(h, \theta) := \frac{\partial g(h, \theta)}{\partial h} \) is a Jacobian matrix, \( \Delta h := Q(h) - h \) is the compression error. We further denote \( \hat{g}_{x\theta}(Q(h); h) := \hat{g}(Q(h); h, \theta) \mid_{h=h(x; \theta)} \) and \( J_{x\theta}(h) := J(h, \theta) \mid_{h=h(x; \theta)} \) for short. Since \( \mathbb{E} [\Delta h(x; \theta)] = 0 \), \( \hat{g}_{x\theta}(Q(h); h) \) is an unbiased SG. Furthermore, the approximation error is small:

**Proposition 1.** Assuming that \( g(h; \theta) \) is twice differentiable w.r.t. \( h \), and the second order derivative is bounded, then

\[
\mathbb{E} [\|g(Q(h); \theta) - \hat{g}_{x\theta}(Q(h); h)\|_2] = O(\text{Var}_Q [\Delta h]).
\]

Since \( \Delta h \) itself is unbiased, \( \text{Var}_Q [\Delta h] = \mathbb{E}_Q [\|\Delta h\|^2] \) is simply the expected compression error. Prop. 1 implies that the linearization error is bounded by the compression error. The linearized gradient \( \hat{g} \) is accurate if the compression is accurate. Using \( \hat{g} \) as a bridge, we arrive in the following convergence theorem:

**Theorem 1.** Assume that:

A1. \( L(\theta) \) is a continuous differentiable, \( \nabla L(\theta) \) is \( \beta \)-Lipschitz continuous.

A2. \( L(\theta) \) is bounded below by \( L_* \).

A3. \( g(h; \theta) \) is differentiable w.r.t. \( h \) and \( \exists b > 0 \), s.t. \( \forall \theta, \mathbb{E} [\|g(Q(h(x; \theta)); \theta) - \hat{g}_{x\theta}(Q(h); h)\|] \leq b \).

A4. \( \exists \sigma^2 > 0 \), s.t. \( \forall \theta, \text{Var} [\hat{g}_{x\theta}(Q(h); h)] \leq \sigma^2 \).

Then, for all \( \eta < \frac{1}{\beta^2} \), if we run ACT defined as Eq. (2) for \( T \) iterations, then we have

\[
\min_{t=0,..,T-1} \mathbb{E} [\|\nabla L(\theta_t)\|^2] \leq \frac{4(\mathcal{L}(\theta_0) - \mathcal{L}_*)}{\eta T} + 3b^2 + \eta \beta^2 \sigma^2
\]

**Remark:** The analytical technique used in Thm. 1 is rather standard, see Thm. 4.8 in Bottou et al. (2018). However, we consider the variance term \( \sigma^2 \) of the linearized gradient, rather than the SG itself. This formulation brings better analytical properties and an adaptive algorithm for determining the compression scheme, as we shall see soon in Sec. 4.

The convergence of ACT is affected by both the linearization error (A3) and the variance of the unbiased gradient \( \hat{g}(\cdot; \theta) \) (A4). The latter is characterized as:

**Proposition 2.** \( \text{Var} [\hat{g}_{x\theta}(Q(h); h)] = \mathbb{E}_x [\text{Var}_Q [\hat{g}_{x\theta}(Q(h); h)]] \), where the second term on the RHS equals to \( \mathbb{E}_x [\text{Var}_Q [J_{x\theta}(h)\Delta h]] = O(\text{Var}_Q [\Delta h]) \).

Prop. 2 separates the variance from different noise sources. \( \mathbb{E}_x [g(h(x; \theta); \theta)] \) is the variance raised by random sampling of data (“sampling variance”). \( \mathbb{E}_x [\text{Var}_Q [J_{x\theta}(h)\Delta h(x; \theta)]] \) is the variance raised by compression. Now, the convergence in Thm. 1 is depicted by \( 3b^2 + \eta \beta^2 \sigma^2 \). By Prop. 1, \( b^2 = O(\text{Var}_Q [\Delta h]^2) \). By Prop. 2, \( \sigma^2 = O(1) + O(\text{Var}_Q [\Delta h]) \), since the sampling variance is not affected by compression. Therefore, when the compres-
sion is accurate ($\Delta h \to 0$), the impact of the linearization error is negligible, and the variance of the unbiased gradient dominates. ACT behaves as if the AC gradient is unbiased.

4. Adapting the Compression Rate

In a network, some context tensors (such as those stored for computing the cross entropy loss) are extremely sensitive, a small amount of compression would result in diverged training, while other tensors are quite robust to compression. Therefore, we must apply different amounts of compression for each context tensor. As a general framework, we have no prior knowledge of the users’ model architecture, so we designed an algorithm to infer the sensitivity for each context tensor and determine their compression rate automatically.

There is a tradeoff between the compression error and the storage requirement. We represent the storage requirement of the compressed context in \textit{bits per dimension}. We assume that $b_l$ bits/dim. are used for compression $h_l^{(i)}$, and $Q_{b_l}(h_l^{(i)})$ be the compression result. Let $b = (b_l)_{l=1}^L$ be a compression scheme, $Q_{b_l}(h) := \{Q_{b_l}(h_l^{(i)})\}_{i=1}^L$, and $\Delta_l h = Q_{b_l}(h) - h$.

4.1. Structure of Variance

As discussed in Sec. 3.2, when the compression is relatively accurate, the variance plays the main role in determining the convergence. Therefore, we would like to investigate how the compression scheme would impact the variance. Formally, we are interested in:

$$V(b; h; \theta) := \text{Var}_Q[\hat{g}(Q_{b_l}(h); h, \theta)].$$

Once $V(b; h; \theta)$ is known, we can find the minimum variance compression scheme under a given total bits budget $B$, by solving the integer programming problem:

$$\min_b V(b; h(x; \theta), \theta), \quad \text{s.t.} \quad \sum_{l=1}^L b_l D_l \leq B, \quad (4)$$

where $D_l$ is the dimensionality of $h_l^{(i)}$. To proceed, we need the following assumptions on the compressor $Q_{b_l}(\cdot)$:

**Assumption B1:** The compressed result is element-wise uncorrelated. That is, for any $i \neq j$, $\text{Cov}[Q_{b_l}(h_i), Q_{b_l}(h_j)] = 0$.

**Assumption B2:** For compressing $h_l^{(i)}(x; \theta)$ to $b_l$ bits/dim., the compression error can be written in the form

$$\text{Var} [\Delta_l h_l^{(i)}(x; \theta)] \leq R_{l_j}(x; \theta) S(b_l), \quad \text{where} \quad S(b_l) \text{is a known function. This isolates the effect of $b_l$ through the unary factor $S(b_l)$.}$$

Both assumptions can be achieved by a stochastic rounding quantizer (Courbariaux et al., 2015), where $R_{l_j}(x; \theta) = \frac{1}{4} \left( \max_{k} h_k^{(l)} - \min_{k} h_k^{(l)} \right)^2$ and $S(b) = (2^b - 1)^{-2}$. See Appendix A.4 for the derivations.

The following theorem reveals the structure of the variance:

**Theorem 2.** Under assumptions B1, B2, there exists a family of functions $\{c_l(h, \theta)\}_{l=1}^L$, such that the compression variance can be written in the form

$$V(b; h, \theta) \leq \sum_{l=1}^L c_l(h, \theta) S(b_l). \quad (5)$$

4.2. Computing Sensitivity

Thm. 2 reveals two good properties of the variance: (1) the impact of compressing different context tensors simply sums up, without affecting each other; and (2) the compression scheme only impacts the variance through $S(b_l)$.

Both properties are brought about by linearization. Since $S(\cdot)$ is a known function, we only need to know $c_l(h, \theta)$ to solve problem Eq. (4). $c_l(h, \theta)$ can be understood as the sensitivity of the AC gradient to the compression of the $l$-th tensor. We can compute $c_l(h, \theta)$ numerically by leveraging the idempotence of compressing a tensor:

**Assumption B3:** If $h = Q(h')$ for some $h'$ with non-zero probability, then $Q(h) = h$ and $\text{Var}_Q[Q(h)] = 0$.

Let $Q_b^{-1}(h) = \{Q_{b_1}(h_1^{(L)}), \ldots, h^{(i)}, \ldots, Q_{b_L}(h_L^{(1)})\}$ be some tensors, where every tensor except $h^{(i)}$ is compressed. Plug $h = Q_b^{-1}(h)$ into Eq. (5), and use B3, we have

$$V(b; Q_b^{-1}(h), \theta) \leq c_l(Q_b^{-1}(h), \theta) S(b_l).$$

The left hand side can be approximated by taking $\hat{g}(Q_{b_l}(h); h, \theta) \approx g(Q_{b_l}(h); h, \theta)$. Assume that $c_l(\cdot, \theta)$ is reasonably continuous, we have

$$c_l(h, \theta) \approx \text{Var}_Q[g(Q_{b_l}(h); \theta)]_{h=Q_b^{-1}(h)} / S(b_l).$$

The variance can be replaced by empirical variance.

Alg. 1 illustrates this idea. To compute $\text{Var}_Q[\hat{g}(Q_{b_l}(h); \theta)]$ at $h = Q_b^{-1}(h)$, we keep the random seeds fixed for all the compressors except the $l$-th one. We compute the empirical variance by two evaluations of $g(Q_{b_l}(h); \theta)$, which are two NN iterations (forward + backward propagation).

Finally, we assume that $c(h, \theta)$ remains stable for different mini-batches $h$, and along the training trajectory ($\theta_t$). Therefore, we maintain a $c_l$ for each tensor $l$, which is updated by periodically running Alg. 1. Eq. (4) is approximately solved by the $O(L \log_2 L)$ greedy algorithm (Chen et al., 2021).

Another useful feature of this approach is predicting failure (in an \textit{a posteriori} manner). If the compression vari-
import torch
import gact
model = ... # user defined model
controller = gact.controller(model, opt_level='L2')
controller.install_hook()

# training loop
for epoch in ...
for iter in ...
    ......
    # instruct gact how to perform forward and backward
    def fwdbwdprop()
    output = model(data)
    loss = loss_func(output, target)
    optimizer.zero_grad()
    loss.backward()
    controller.iterate(fwdbwdprop)

Figure 2. Usage example of GACT

5. System Implementation

We implemented GACT as a lightweight library in PyTorch. Users can use GACT for any NN architecture with several lines of code change. GACT uses low-level PyTorch hooks to capture context tensors, so it supports arbitrary operators, including custom operators defined by users. We implemented efficient CUDA kernels to infer tensor sensitivity and to perform compression during run time. GACT uses the same per-group quantizer in ActNN (Chen et al., 2021) as the compressor. However, GACT differs from ActNN in several aspects. ActNN relies on manual analytical deduction to compute the sensitivity for different operators, while GACT infers tensor sensitivity automatically, as described in Sec. 4.2. Moreover, ActNN performs layer-level quantization. It has to implement an activation compressed version for each operator and substitute operators during the training (e.g., replace torch.nn.Conv2d with actnn.Conv2d). In contrast, GACT runs at tensor level and uses a single hook interface to compress saved tensors for all operators.

5.1. General API

As shown in Fig. 2, the interface of GACT is straightforward and intuitive, requiring the user to (i) initialize the GACT controller and specify an optimization level (Line 5); (ii) install hooks (Line 6); and (iii) instruct GACT how to perform forward and backward propagation (Lines 13-17) and pass it as a function (fwdbwdprop) to the controller (Line 19). We require users to specify (iii) because GACT needs to numerically run the forward and backward pass to infer tensor sensitivity. Although fwdbwdprop is passed to the controller every iteration, it is only called internally every adapt_interval iterations when tensor sensitivity changes. As shown in Sec. 6.1, tensor sensitivity stabilizes quickly after the first several epochs, adapt_interval can thus be set to a large number, introducing negligible impact on training speed.

5.2. System Architecture

Fig. 1 shows an overview of GACT. The GACT controller has three modules: Adaptative Algorithm; Compressor; and Decompressor. In the forward pass, the controller uses PyTorch pack_hook to capture all context tensors. Then Adaptive Algorithm infers tensor sensitivity based on gradients and assigns higher bits to more sensitive tensors, as described in Sec. 4.2. The bits information is used to instruct Compressor to perform quantization. In the backward pass, Decompressor dequantizes context tensors and uses unpack_hook to send the dequantized results back to the PyTorch’s auto differentiation engine. The controller is also responsible for swapping quantized tensors to the CPU and prefetching them back during the backward propagation if swapping is enabled.

5.3. Identifying Tensors to Quantize

The pack_hook and unpack_hook process all types of context tensors, including activation, parameters trained by the optimizer, and training states such as running mean/variance used by batch normalization. To guarantee that only the activations are quantized, we filter out saved parameters by recording the data pointers of all the model parameters before training, and we skip quantization if the input tensor pointer exists in the parameter pointer set. Similarly, GACT does not quantize training states by checking if the input tensor requires gradients.

However, using hooks blindly disables some memory-saving optimization. For example, in a transformer’s self-attention layer, the keys, query, value tensors are all calculated from the same input tensor. The saved objects of the three operations thus all refer to the same tensor. In this case, PyTorch triggers the pack_hook three times. If we perform quantization blindly, we waste computation resources and introduce extra memory consumption because the same underlying tensor is quantized and saved more than once. GACT avoids duplication by generating footprints for each input context tensor. We use the CUDA data pointer, sampled data points, and the tensor statistics (e.g., sum) as the footprint. GACT manages all quantized context tensors and uses the footprint to differentiate them. If a tensor is already quantized, GACT will skip quantization and return previous results directly.
5.4. Parallel Swap and Prefetch

To further reduce activation memory, we combine GACT with swapping. All compressed tensors are offloaded to the CPU during the forward pass and swapped back in the backward pass. Here, we replace the original tensor with quantized activation, as data movement is more expensive than computation. Swapping the original tensor saves the quantization overhead but adds more data movement cost between CPU and GPU. As shown in Sec. 6.4, quantization overhead is much smaller than copying full-precision data to CPU in modern GPU architecture.

Furthermore, we create two new streams (swap in/out) to parallelize the computation and swapping operation to reduce the swap overhead. The forward computation and swap-out process happen in parallel during the forward pass. During the backward pass, in each layer the swap-in stream is responsible for prefetching the compressed activation of the previous layer to avoid synchronization overhead. We leverage the CUDA event to ensure tasks in different streams are executed in the correct order.

5.5. Other Memory Optimizations

Gradient checkpointing. Gradient checkpointing (Chen et al., 2016) works by dividing the NN into segments. The algorithm only stores the inputs of each segment and recomputes the dropped activations segment by segment during backpropagation. The memory consumption is thus the cost of storing the inputs of all segments plus the maximum memory cost to backpropagate each segment. When combined with gradient checkpointing, GACT can reduce the memory consumption of both parts. GACT reduces the memory consumption of the first part by quantizing the segment inputs. Moreover, the activations saved during the recompute phase are also quantized, reducing the memory cost of the second part. Combining GACT with gradient checkpointing might introduce more training noise because the recompute starts from quantized segment inputs, making the forward pass of recompute phase not exact. However, in Sec. 6.4, we show the noise introduced by forwarding from the quantized tensors is negligible.

Memory efficient self-attention. When the batch size is very large, the single layer after dequantization occupies a large amount of memory and prevents the batch size from increasing further. We observe this problem in transformer-based models where self-attention has quadratic space complexity in terms of sequence length. To reduce the memory footprint of the self-attention layer, we implement the algorithm introduced in (Rabe & Staats, 2021) that achieves linear space complexity, and combines it with GACT.

5.6. Optimization level

To exploit the trade-off between memory saving and training speed, GACT provides several optimization levels. Higher levels can save more memory but with more overhead. Tab. 1 lists these optimization levels. L1 uses per-group quantization with the adaptive algorithm. L2 combines per-group quantization with swapping and prefetching. For transformer-based models, CB1 combines GACT with gradient checkpointing. CB2 further reduces the peak memory by adding efficient self-attention to CB1.

6. Experiments

We first demonstrate the effectiveness of the GACT adaptive algorithm. We further apply GACT to a wide range of machine learning tasks, including image classification, object detection, text, and graph node classification. We compare the training accuracy and activation compression rate for full precision, adaptive 4/3/2 (using GACT to adaptively decide quantization bits with an average of 4/3/2 bit) and fix-4 bit (quantizing all tensors uniformly with 4 bits). Next, we study the trade-off between compression rate and training throughput and compare GACT with other state-of-the-art memory-saving methods. Lastly, we demonstrate the flexibility of GACT by exploring the possibility of combining it with other memory optimization methods (CB1, CB2 as listed in Table 1). We use open-source model implementations for all tasks.

6.1. Compression Strategy

We first test the effectiveness of our adaptive compression rate algorithm for training VGG-11 (Simonyan & Zisserman, 2015) on ImageNet. Fig. 3(a) plots the inferred per-tensor sensitivity \( c_l \) and the corresponding optimal bits/dim. GACT assigns more bits to more sensitive layers. The context tensor saved by the cross-entropy loss operator is most sensitive. A small amount of compression leads to a huge gradient variance. This makes sense since the loss is the first operator to back-propagate through, where the error accumulates. Therefore, GACT assigns 32 bits/dim. for the tensors in the classification head. With the adaptive algorithm, GACT with an average of 4 bits/dim. achieves smaller gradient variance than uniformly assigning 8 bits/dim. for all the tensors, as shown in Fig. 3(b). Finally, Fig. 3(c) shows that the sensitivity \( c_l(h; \theta_l) \) remains stable during training. Therefore, periodically updating \( c_l \) at a large
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(a) Inferred per-tensor $c_l$ (line) and bits/dim. (bar) for VGG-11. Layers with * have a preceding ReLU layer with shared context. drop=dropout, loss=cross entropy loss.

(b) Gradient variance. (c) Evolution of the per-tensor sensitivity. Each line is $c_l$ for a tensor.

Figure 3. Effectiveness of the adaptive algorithm.

Table 2. For classification, we train VGG11 (Simonyan & Zisserman, 2015), ResNet-50 (He et al., 2016), and Swin-Tiny (Liu et al., 2021) on ImageNet (Deng et al., 2009). For object detection, we train RetinaNet (Lin et al., 2017), Faster R-CNN (Ren et al., 2015) on Coco (Lin et al., 2014). We report accuracy on validation sets (Div. indicates diverge) and the compression rate of context tensors (numbers in brackets) for both tasks.

| Task | Model     | FP32 | GACT Adapt 4bit (L1) | GACT Adapt 2bit |
|------|-----------|------|----------------------|-----------------|
| Cls. | VGG11     | 68.75| 68.77 (2.84×)        | 68.49 (3.34×)   |
|      | ResNet-50 | 77.29| 76.96 (6.69×)        | 76.13 (11.39×)  |
|      | Swin-tiny | 81.18| 80.92 (7.44×)        | 77.91 (13.73×)  |
| Det. | Faster R-CNN | 37.4 | 37.0 (4.86×) | 36.1 (6.81×) |
|      | RetinaNet | 36.5 | 36.3 (3.11×) | Div.          |

interval is reasonable, and this introduces negligible impact on training speed.

6.2. Optimization level

We apply GACT on various computer vision tasks, including image classification and object detection, as shown in Fig. 2. We also vary the average bits used by the adaptive algorithm to explore the memory accuracy trade-off. On both tasks, GACT L1 achieves comparable (< 0.5% accuracy drop) or even better results than the full precision training, while reducing activation memory by up to 7.44×. Here, we list the accuracy of FP32 as the strongest accuracy baseline. For other lossy methods we consider in Sec. 6.3, the accuracy is no better than FP32, and we list their training accuracy in Appendix C. Notice that here GACT Adapt 2bit diverges on the detection task. This is because, as shown in Sec. 3.2, although ACT has unbiased gradients, the compression error and learning rate affect the convergence. When using 2 bit, the compression error is large and the learning rate has to be reduced accordingly to guarantee convergence. However, we do not want to slow training by decreasing the learning rate. All experiments are run with the same learning rate as the full precision. Therefore when compression error is large, the training diverges. Furthermore, we observe that the memory reduction varies among networks because GACT does not quantize intermediate states, and the size of intermediate states differs between networks. For example, in VGG11, when the batch size is 128, GACT reduces the saved tensor size from 5889MB to 2080MB, among which 78% (1494MB) is used to store the intermediate index for the max-pooling layer that is not quantized by GACT.

Next, we demonstrate the flexibility of GACT by applying it to a wider variety of natural language processing (NLP) and graph machine learning (Graph) tasks. We run multiple seeds for each task, and we report the mean±std of accuracy/F1 across runs as shown in Tab. 3. We include the detailed experimental setup in Appendix B. For both NLP and Graph tasks, GACT L1 achieves comparable training results with FP32, introducing less than 0.3% accuracy/F1-score drop, while reducing activation memory by 4.18× to 7.93×. Moreover, the results are stable across runs, introducing similar accuracy variance as FP32. We also show the training results of fix-4bit quantization, where all tensors are uniformly quantized with 4 bits. As shown in Tab. 3, fix-4 bit quantization causes significant accuracy/F1-score loss on various graph models. For Bert-large, fixed-4 bit quantization works fine because all the context tensors have similar sensitivity. On the other hand, GACT L1, using a similar amount of memory as always quantizing each layer to 4 bits, still performs on par with full precision training on all the models. This shows the necessity of using adaptive algorithms to assign bits based on tensor sensitivity for stabilized training. Moreover, for Bert-large and three graph models (GCN/GAT/GCNII), GACT converges and gives lossless results with 3 bits. Remarkably, across all the graph models, training with 2-bit GACT causes little accuracy loss (< 1%). This shows the robustness of our adaptive algorithm.

6.3. Memory Saving and Computational Overhead

Settings and baselines. We implement the benchmark with PyTorch 1.10 and measure the memory saving and overhead of GACT on an AWS g4dn.4xlarge instance, which has a 16GB NVIDIA T4 GPU and 64GB CPU memory. On ResNet-50, we compare with ActNN (Chen et al., 2021), a dedicated quantization framework for convolutional NNs, and DTR (Kirisame et al., 2020), a state-of-the-art rematerialization method for dynamic graphs. “swap” is a simple
Table 3. Accuracy and activation compression rate for NLP and Graph tasks. Accuracy that drops > 1% is in italic font.

| Model | Dataset | FP32 | Fix 4bit | GACT Adapt 4bit (L1) | GACT Adapt 3bit | GACT Adapt 2bit |
|-------|---------|------|---------|---------------------|----------------|----------------|
| GCN   | Flickr  | 51.17 ± 0.19 | 50.93 ± 0.16 (7.56×) | 51.08 ± 0.18 (7.93×) | 51.14 ± 0.18 (11.34×) | 51.20 ± 0.18 (17.56×) |
|       | Reddit  | 95.33 ± 0.07 | 94.42 ± 0.11 (7.55×) | 95.32 ± 0.07 (7.90×) | 95.31 ± 0.07 (9.79×) | 95.34 ± 0.06 (13.68×) |
|       | Yelp    | 39.86 ± 0.94 | 39.85 ± 1.22 (5.94×) | 40.06 ± 0.74 (6.42×) | 40.21 ± 0.82 (7.46×) | 39.89 ± 1.45 (9.00×) |
|       | ogbn-axiv | 71.51 ± 0.65 | 68.61 ± 0.77 (7.54×) | 71.35 ± 0.36 (8.09×) | 70.82 ± 0.95 (10.45×) | 70.87 ± 0.66 (13.75×) |
| GAT   | Flickr  | 52.40 ± 0.28 | 35.34 ± 11.90 (4.23×) | 52.26 ± 0.31 (4.34×) | 51.68 ± 1.13 (5.04×) | 51.62 ± 1.19 (5.46×) |
|       | Reddit  | 95.95 ± 0.06 | 59.37 ± 11.48 (4.12×) | 96.02 ± 0.09 (4.29×) | 95.96 ± 0.06 (4.64×) | 95.82 ± 0.06 (5.24×) |
|       | Yelp    | 52.41 ± 0.69 | 36.09 ± 13.70 (4.04×) | 52.18 ± 0.38 (4.18×) | 51.63 ± 0.83 (4.53×) | 51.15 ± 0.53 (5.24×) |
|       | ogbn-axiv | 71.68 ± 0.54 | 54.64 ± 5.62 (3.04×) | 71.80 ± 0.47 (5.09×) | 71.47 ± 0.50 (6.14×) | 71.21 ± 0.68 (6.98×) |
| GCNII | Flickr  | 52.37 ± 0.16 | 52.28 ± 0.16 (4.84×) | 52.31 ± 0.16 (4.91×) | 52.36 ± 0.16 (5.54×) | 52.23 ± 0.15 (6.44×) |
|       | Reddit  | 96.32 ± 0.24 | 86.50 ± 1.08 (4.51×) | 96.11 ± 0.22 (4.52×) | 96.01 ± 0.33 (5.16×) | 95.94 ± 0.29 (5.92×) |
|       | Yelp    | 62.33 ± 0.20 | 62.21 ± 0.22 (5.26×) | 62.28 ± 0.26 (5.34×) | 62.53 ± 0.36 (6.29×) | 62.33 ± 0.37 (7.28×) |
|       | ogbn-axiv | 72.52 ± 0.12 | 44.57 ± 5.01 (6.54×) | 72.28 ± 0.35 (6.74×) | 72.22 ± 0.28 (7.98×) | 71.74 ± 0.26 (10.24×) |
| Bert-large | MNLI   | 68.74 ± 0.24 | 85.98 ± 0.16 (7.55×) | 86.61 ± 0.11 (7.38×) | 86.68 ± 0.08 (9.13×) | 84.24 ± 0.74 (12.87×) |
|       | SST-2   | 93.69 ± 0.30 | 93.46 ± 0.23 (7.55×) | 93.54 ± 0.52 (7.30×) | 93.20 ± 0.37 (9.05×) | 91.90 ± 1.04 (12.94×) |
|       | MRPC    | 88.20 ± 0.02 | 87.36 ± 0.19 (7.55×) | 87.90 ± 0.10 (7.40×) | 87.69 ± 0.07 (9.19×) | 82.54 ± 0.38 (12.91×) |
|       | QNLI    | 92.29 ± 0.14 | 92.34 ± 0.07 (7.55×) | 92.44 ± 0.07 (7.42×) | 92.43 ± 0.31 (9.19×) | 90.74 ± 0.13 (12.95×) |

Results. We compare the training throughput of GACT against other memory saving systems in Fig. 4. On ResNet-50, GACT achieves similar throughput as ActNN (ActNN optimization L5 is not listed because it optimizes PyTorch memory allocation, which is unrelated to quantization and can also be applied to GACT), but ActNN enables training with a larger batch size. This is expected because ActNN implements efficient, customized layers for different operators in convolutional NNs. For Bert-large, Zero-offload fails quickly because it only offloads optimizer states that occupy a small portion of total memory to CPU. GACT L1 outperforms Mesa because Mesa only compresses tensors to 8 bit. When the batch is bigger, the activation size of each segment becomes the memory bottleneck and prevents gradient checkpointing from increasing the batch size. Moreover, combining GACT with gradient checkpointing and efficient self-attention further reduces the peak memory, increasing the batch size by up to 24.7×. Meanwhile, it introduces a small throughput overhead compared with the original gradient checkpointing. Across all the network architectures, GACT enables training with a 4.2× to 24.9× larger batch size under the same memory budget.

Network scaling. With GACT, we can construct larger models or train with a higher image resolution. Tab. 4 compares the largest model we can train against full precision. With the same batch size and memory budget, GACT can scale a ResNet-152 to 7.0× deeper, 3.6× wider or 3.0× higher resolution. Similarly, Bert-large can be scaled to 2.0× deeper or 1.6× wider. In GCN, GACT enables training 10.0× deeper and 1.7× wider network. Overall, GACT maintains 75% - 136% original training throughput.

6.4. Other Optimizations

We evaluate the idea of combining GACT with swapping on Bert-large-cased. As shown in Tab. 5, swapping compressed tensors is faster than swapping the original ones because communication between CPU and GPU is more time-consuming than computation. Combining GACT with
Table 4. Largest models GACT can train with 16G GPU memory. In ResNet (batch size=64), D (depth): number of layers, W (width): base width of the bottleneck block, R (resolution): width and height of input images. In Bert-large (batch size=16) and GCN, D (depth): number of transformer/gcn blocks, W (width): hidden size.

| Dim   | Maximum Value | Throughput (TFLOPS) |
|-------|--------------|----------------------|
|       | FP | L1 | L2 | FP | L1 | L2 |
| ResNet-152 | D | 160 | 640 | 1124 | 0.43 | 0.47 | 0.41 |
|         | R | 232 | 548 | 716 | 0.41 | 0.39 | 0.44 |
| Bert-large | D | 32 | 56 | 64 | 0.67 | 0.56 | 0.53 |
|         | W | 1280 | 1488 | 2032 | 0.68 | 0.61 | 0.60 |
| GCN | D | 24 | 152 | 240 | 0.20 | 0.14 | 0.15 |
|       | W | 2464 | 3948 | 4244 | 0.36 | 0.38 | 0.40 |

Table 5. Swap and prefetch speed/memory on Bert-large.

| Algorithm | Speed (sequence/s) | Peak Mem. (MB) | Total Mem. (MB) |
|-----------|---------------------|----------------|-----------------|
| FP32 | 16.41 | 9573 | 9527 |
| FP32 + swap | 6.02 | 5215 | 5093 |
| GACT swap | 12.95 | 5426 | 5325 |
| GACT swap + prefetch | 14.02 | 5426 | 5324 |

swapping increases training speed by up to 2.3×. Notice here that the peak memory use of “GACT swap” is slightly higher than “FP32 + swap” because GACT does not quantize and swap intermediate states such as running mean/var of BatchNorm layer. Moreover, prefetch increases the speed by about 7% with negligible memory overhead.

We next demonstrate combining GACT with gradient checkpointing (CB1). Gradient checkpointing is performed at the beginning of each transformer block, thus avoiding saving tensors generated within the block. We then apply GACT with gradient checkpointing, where the saved tensors are quantized with 4 bits. As shown in Tab. 6, the accuracy is unaffected. We also compare the activation memory and peak memory of CB1 and CB2 in Tab. 7. AM2 denotes the peak activation memory, which is the size of saved tensors after re-forwarding the first transformer block. When batch size = 288, compared with gradient checkpointing on full precision (FP32), CB1 and CB2 reduce the peak activation size by 4.7× and 5.4× respectively.

7. Conclusion

This paper presents GACT, an ACT framework for generic NN architectures. We prove the convergence of GACT without prior knowledge about operator type or network architecture by analyzing a linearized approximation of ATC’s gradients. With the adaptive algorithm, GACT achieves negligible accuracy loss on various tasks, reducing activation memory by up to 8.1× and enabling training with up to 24.7× batch size compared with full precision training.

Table 6. Accuracy of Bert-large-cased on SST-2 and QNLI datasets

| Algorithm | SST-2 | QNLI | Algorithm | SST-2 | QNLI |
|-----------|-------|------|-----------|-------|------|
| FP32 | 93.58 | 92.42 | CB1 | 93.81 | 92.26 |

Table 7. Memory use of different algorithms on Bert-large. AM1: Activation size before backward, AM2: Activation size after re-forwarding the first transformer block. When batch size = 288, L0 runs out of memory, and therefore it is not listed below.

| Batch Size | Algorithm | AM1 (MB) | AM2 (MB) | Peak Mem. (MB) |
|------------|-----------|----------|----------|----------------|
| 16         | L0        | 4434     | -        | 9573           |
|            | FP32 + CKPT | 210     | 394     | 5541           |
|            | CB1       | 37      | 99      | 5286           |
|            | CB2       | 31      | 79      | 5269           |
| 288        | FP32 + CKPT | 3783   | 7092     | 12885          |
|            | CB1       | 515     | 1497    | 8251           |
|            | CB2       | 486     | 1307    | 8102           |

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A. Proof of Theorems

A.1. Theorem 1: Convergence of ACT

Assume that:

A1. \( L(\theta) \) is a continuous differentiable, \( \nabla L(\theta) \) is \( \beta \)-Lipschitz continuous.

A2. \( L(\theta) \) is bounded below by \( L_* \).

A3. \( g(h; \theta) \) is differentiable w.r.t. \( b \) and \( 3b > 0 \), s.t. \( \forall \theta, \mathbb{E} \| g(Q(h(x, \theta)); \theta) - \hat{g}(h(x, \theta); \theta) \| \leq b. \)

A4. \( \exists \sigma^2 > 0 \), s.t., \( \forall \theta, \text{Var} [\hat{g}(h(x, \theta))] \leq \sigma^2. \)

Then, for all \( \eta < \frac{1}{2\beta} \), if we run ACT defined as Eq. (2) for \( T \) iterations, then we have

\[
\min_{t=0, \ldots, T-1} \mathbb{E} \left[ \|\nabla L(\theta_t)\|^2 \right] \leq \frac{4(L(\theta_0) - L_*)}{\eta T} + 3b^2 + \eta \beta \sigma^2
\]

**Proof.** Denote \( m := \nabla_b L(\theta_t), \epsilon := \hat{\epsilon}(h(x, \theta_t); \theta_t) - m, d := g(Q(h(x, \theta_t)); \theta_t) - \hat{g}(h(x, \theta_t); \theta_t) \). Then, by A3 and A4, we have

\[
\mathbb{E} [\epsilon] = \mathbb{E} [g(h(x, \theta_t); \theta_t) - \nabla_b L(\theta_t)] + \mathbb{E} [\langle J(x, \theta_t), \Delta Q(h(x, \theta_t)) \rangle] = 0.
\]

(6)

By Eq.s (6,8)

\[
\mathbb{E} \left[ (m, m + d + \epsilon) \right] \geq \|m\|^2 - \|m\| \|d\| + \mathbb{E} [\epsilon] \geq \|m\|^2 - \|m\| \|b\|.
\]

(10)

By Eq.s (6,7,8), and \( \|x + y\|^2 \leq 2 \|x\|^2 + 2 \|y\|^2 \),

\[
\mathbb{E} \left[ \|m + d + \epsilon\|^2 \right] = \mathbb{E} \left[ \|m + d\|^2 \right] + \mathbb{E} \left[ \|\epsilon\|^2 \right] \leq 2 \mathbb{E} [\|m\|^2] + 2 \mathbb{E} [\|d\|^2] + 2 \mathbb{E} [\|\epsilon\|^2] = 2 \mathbb{E} [\|m\|^2] + 2b^2 + \sigma^2.
\]

(11)

Taking expectation on both sides of Eq. (9), plug in Eq.s (10, 11), and use \( \eta < \frac{1}{2\beta} \), we have

\[
\mathbb{E} [L(\theta_{t+1})] \leq L(\theta_t) - \eta (\|m\|^2 - \|m\| \|b\|) + \frac{\beta \eta^2}{2} (2 \mathbb{E} [\|m\|^2] + 2b^2 + \sigma^2).
\]

(9)

Completing the squares,

\[
\mathbb{E} [L(\theta_{t+1})] \leq L(\theta_t) - \frac{\eta}{2} (\|m\| - b)^2 + \frac{3\beta \eta^2}{2} (2b^2 + \sigma^2).
\]

Taking expectation on both sides and sum up for \( t = 0, \ldots, T - 1 \),

\[
\mathbb{E} [L(\theta_T) - L(\theta_0)] \leq -\frac{\eta}{2} \sum_{t=0}^{T-1} \mathbb{E} [\|\nabla L(\theta_t)\| - b]^2 + \frac{3\beta \eta^2 T}{2} (2b^2 + \sigma^2).
\]

Reorganize the terms,

\[
\mathbb{E} [\|\nabla L(\theta_t)\| - b]^2 \leq \frac{2(L(\theta_0) - L(\theta_T))}{\eta T} + \eta \beta (2b^2 + \sigma^2).
\]

Let \( \eta T = \text{argmin}_{\eta} \mathbb{E} [\|\nabla L(\theta_t)\|]. \), and use A1, we have

\[
\mathbb{E} (\|\nabla L(\theta_{t+1})\| - b)^2 \leq \frac{2(L(\theta_0) - L_*)}{\eta T} + \eta \beta (2b^2 + \sigma^2).
\]

Use \( (a + b)^2 \leq 2a^2 + 2b^2 \), we have

\[
\mathbb{E} [\|\nabla L(\theta_t)\|^2] \leq \frac{4(L(\theta_0) - L_*)}{\eta T} + (2\beta \eta + 2)b^2 + \eta \beta \sigma^2 \leq \frac{4(L(\theta_0) - L_*)}{\eta T} + 3b^2 + \eta \beta \sigma^2.
\]
A.2. Proposition 1: The Linearization Error

Proof. Consider the gradient function \( g(Q(h(x; \theta); \theta)) \), whose output is a \( P \)-dimensional vector. Since it is twice differentiable, we construct the Taylor’s expansion at \( h(x; \theta) \) with Lagrange remainder:
\[
\exists H_1, \ldots, H_P, \text{s.t.} \forall i, \; g_i(Q(h(x; \theta); \theta)) = g_i(h(x; \theta); \theta) + J_i(x, \theta) \Delta h(x, \theta) + \Delta h(x, \theta)^\top H_i \Delta h(x, \theta),
\]
where \( J_i(h(x; \theta), \theta) := \frac{\partial g_i(h(x; \theta); \theta)}{\partial h} \). By the assumption, there exists \( P > 0 \), such that the linearization error is
\[
\|g(Q(h(x; \theta)); \theta) - \hat{g}(h(x; \theta); h(x; \theta), \theta)\|_1 = \sum_{i=1}^{P} \Delta h(x, \theta)^\top H_i \Delta h(x, \theta) \leq \gamma P \|\Delta h(x, \theta)\|^2.
\]
Taking expectation,
\[
\mathbb{E}[\|g(Q(h(x; \theta)); h(x; \theta), \theta) - \hat{g}(h(x; \theta); h(x; \theta), \theta)\|_1] \leq \mathbb{E}[\|g(Q(h(x; \theta)); \theta) - \hat{g}(h(x; \theta); h(x; \theta), \theta)\|_1] \leq \gamma P \text{Var}[\Delta h(x, \theta)] = O(\text{Var}[\Delta h(x, \theta)]).
\]

A.3. Proposition 2: The Order of the Variance

The following proposition is convenient for isolating the different noise sources.

Proposition A. (Law of Total Variance)
\[
\text{Var}[X] = \mathbb{E}[\text{Var}[X \mid Y]] + \text{Var}[\mathbb{E}[X \mid Y]].
\]

Proof. By definition
\[
\text{Var}[\hat{g}(h(x; \theta); h(x; \theta), \theta_i)] = \text{Var}[g(h(x; \theta); \theta)] + \text{Var}[J(h(x; \theta), \theta) \Delta h(x, \theta)],
\]
where \( \text{Var}[g(h(x; \theta); \theta)] \) is the noise introduced by subsampling the data \( x \). By law of total variance,
\[
\text{Var}[J(h(x; \theta), \theta) \Delta h(x, \theta)] = \mathbb{E}_X [\text{Var}_Q [J(h(x; \theta); \theta_i) \Delta h(x, \theta)]] + \mathbb{E}_X [\mathbb{E}_Q [J(h(x; \theta); \theta_i) \Delta h(x, \theta)]]
\]
where
\[
\text{Var}_Q [J(h(x; \theta); \theta_i) \Delta h(x, \theta)] = \mathbb{E}_Q [\|J(h(x; \theta); \theta_i) \Delta h(x, \theta)\|^2] \leq \mathbb{E}_Q [\|J(h(x; \theta); \theta_i)\|^2 \|\Delta h(x, \theta)\|^2] = \|J(h(x; \theta); \theta_i)\|^2 \mathbb{E}_Q [\|\Delta h(x, \theta)\|^2] = O(\text{Var}[\Delta h(x, \theta)]).
\]

A.4. Proposition 3: The Structure of the Variance

Before investigating the structure of \( \text{Var}_Q [J(h(x; \theta_i) \Delta h(x, \theta))] \), let’s do some recap: the parameter \( \theta_i \) is a \( P \)-dimensional vector; the context difference \( \Delta h(x, \theta) \) is a \( D \)-dimensional vector, and \( J(x; \theta_i) \) is a \( P \times D \) matrix. Recall that \( \Delta h(x, \theta) \) is the concatenation of \( L \)-vectors, \( \Delta h^{(l)}(x, \theta) \), and let \( J^{(l)}(x, \theta) := \frac{\partial}{\partial h} g \left( \left( (h^{(l)}(x; \theta))_{l=1}^L \right)_{l=1}^L \right), \) which is a \( P \times D_l \) matrix. Furthermore, let \( h^{(l)}_j(x, \theta) \) be the \( j \)-th dimension, and \( J^{(l)}_j(x, \theta) \) be its \( j \)-th column.

To proceed, we need to make the following assumptions to the compressor \( Q(\cdot) : \mathbb{R}^D \rightarrow \mathbb{R}^D \):

B1: The compressed result is element-wise uncorrelated. That is, for any \( i \neq j \), \( \text{Cov}[Q(h)_i, Q(h)_j] = 0 \).

B2: For compressing a vector \( h \) to \( b \) bits, the compression variance of each dimension can be written in the form \( \text{Var}[Q(h)_j] \leq R_j(h) S(b) \), where \( S(\cdot) \) is a known function.

Both assumptions can be achieved by a stochastic rounding (Courbariaux et al., 2015) quantizer, where
\[
Q(h)_j = \begin{cases} T_{h,b}^{-1} ([T_{h,b}(h)_j]) & \text{w.p. } T_{h,b}(h)_j - [T_{h,b}(h)_j] \\ T_{h,b}^{-1} (\lfloor T_{h,b}(h)_j \rfloor) & \text{otherwise} \end{cases},
\]
where \( T_{h,b}(h)_j = (2^b - 1) \frac{h_j - \min_j h}{\max_j h - \min_j h} \). Since each dimension is quantized independently, B1 is met. Moreover,
\[
\text{Var}[Q(h)_j] \leq \frac{1}{4} \left( \frac{\max_j h - \min_j h}{\max_j h - \min_j h} \right)^2 (2^b - 1)^2 = R_j(h) S(b),
\]
where

\[ R_j(h) = \frac{1}{4} \left( \frac{\max_j h - \min_j h}{h_j - \min_j h} \right)^2, \quad S(b) = (2^b - 1)^{-2}. \]

**Proof.** By definition,

\[ J(h; \theta) \Delta h = \sum_{l=1}^{L} \sum_{j=1}^{D_l} J_j^{(l)}(h; \theta_t) \Delta h_j^{(l)}. \]

Using Assumption B1, we have

\[
\text{Var}_Q [J(h; \theta) \Delta h] = \mathbb{E}_Q \left[ \left\| \sum_{l=1}^{L} \sum_{j=1}^{D_l} J_j^{(l)}(h; \theta_t) \Delta h_j^{(l)} \right\|^2 \right] \\
= \sum_{l=1}^{L} \sum_{j=1}^{D_l} \mathbb{E}_Q \left[ \left\| J_j^{(l)}(h; \theta_t) \Delta h_j^{(l)} \right\|^2 \right] \\
= \sum_{l=1}^{L} \sum_{j=1}^{D_l} \left\| J_j^{(l)}(h; \theta_t) \right\|^2 \text{Var}_Q \left[ \Delta h_j^{(l)} \right]
\]

Using Assumption B2, we have

\[
\text{Var}_Q [J(h; \theta) \Delta h] \leq \sum_{l=1}^{L} \sum_{j=1}^{D_l} \left\| J_j^{(l)}(h; \theta_t) \right\|^2 R_l(h) S(b_l) = \sum_{l=1}^{L} c_l(h, \theta) S(b_l),
\]

where \( c_l(\theta, h) := R_l(h) \left\| J_j^{(l)}(h; \theta_t) \right\|^2 \).

**B. Experiment Setup**

**B.1. Node classification task on graphs**

We conduct experiments on four node classification datasets with standard splits, including Flickr, Reddit, Yelp from GraphSAINT (Zeng et al., 2019), and ogbn-arxiv from Open Graph Benchmark (OGB) (Hu et al., 2020). The four datasets cover extensive downstream applications with different scales. We use accuracy as the evaluation metric for multi-class classification and micro-F1 for multi-label classification. We run ten seeds (0 to 9) and report the average accuracy across runs.

We evaluate GACT on three representative GNN models, including GCN (Kipf & Welling, 2016), GAT (Veličković et al., 2017), and GCNII (Chen et al., 2020b) under the full-batch training setting. All three models are implemented by CogDL (Cen et al., 2021), a toolkit for graph neural networks.

**B.2. Text classification task**

We select four largest datasets, MNLI, QQP, SST-2, and QNLI, from the GLUE benchmark (Wang et al., 2018a). The four datasets cover different aspects of natural language understanding, including sentiment classification, natural language inference and paraphrase detection. We use the mainstream transformer implementation (Wolf et al., 2020) to train Bert-large (Devlin et al., 2019). We run three seeds (42, 43, 44) and report F1 for QQP, accuracy for the others.

**C. Training Accuracy of Baselines**

For all the baselines we compared in Sec. 6.3, only ActNN, Mesa, and ZeRO-Offload are lossy methods. All other methods are lossless and have the same training accuracy as FP32. For ResNet-50 on ImageNet, the training accuracy for FP32, GACT, ActNN L2, and ActNN L3 are 77.3, 77.0, 77.4, and 76.9. For Bert-Large on SST-2, the accuracy for FP32, GACT, Mesa, and ZeRO-Offload are 93.7, 93.5, 93.8, and 93.3. For Swin-tiny on ImageNet, the training accuracy for FP32, GACT, and Mesa are 81.2, 81.0, and 81.3 respectively.