#### Neural Networks compression

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1 / 78

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#### Context: deep neural networks (DNN) are growing fast



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# Scaling up

Why? : better performance, more complex tasks ...

- ... but
	- $\blacktriangleright$  The computational complexity, memory usage and energy consumption of deep networks are increasingly growing, both during training and inference
	- $\blacktriangleright$  Limitation for the deployment on resource-constrained devices (mobile, embedded systems) where energy is often a limited resource
	- $\blacktriangleright$  High environmental impact
	- $\blacktriangleright$  Limitation for real-time applications

## The computational bottlenecks



## The computational bottlenecks



## The computational bottlenecks



#### **Training a DNN model**

#### **Data movement**

- to compute units
- · send partial results back to memory

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#### Improve performance and power efficiency



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## **Outline**

#### [Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

#### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

## <span id="page-8-0"></span>**Outline**

#### [Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

#### **[Sparsification](#page-71-0)**

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

#### <span id="page-9-0"></span>[Quantization](#page-8-0) **[Generalities](#page-9-0)**

[Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

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#### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

# Quantization: definition

Quantization is the process of constraining an input from a continuous or large set of values (such as the real numbers) to a discrete set (such as the integers).

Example: convert floating-point numbers to lower precision (e.g., 8-bit integers).

Example: [https://www.qualcomm.com/news/onq/2019/03/](https://www.qualcomm.com/news/onq/2019/03/heres-why-quantization-matters-ai) [heres-why-quantization-matters-ai](https://www.qualcomm.com/news/onq/2019/03/heres-why-quantization-matters-ai)



24 bits per pixel

#### Low precision formats





#### Modern hardware

- $\blacktriangleright$  Nvidia tensor cores
- Google TPUs

 $\blacktriangleright$  FPGAs

## Benefits of quantization

#### **Quantization effects: the good**

#### Memory usage Power consumption Latency Silicon area storage needed for weights and enerav is significantly reduced for less memory access and simpler 8-bit arithmetic and below and activations is proportional to both computations and memory computations lead to faster runtimes requires less area than larger the bit width used accesses and reduced latency bit width FP compute units ADD energy (pJ) **Memory access** MULT area (µm<sup>2</sup>) **FP32** INT<sub>8</sub> energy (pJ) 111.125 **INT32 INT8** INT32 FP16 **FP32** INT8 **FP16 FP32** 111 Cache (64-bit) 0.03  $O.1$  $0.4$  $0.9$ 282 3495 1640 7700 **8KB** 10 91010101 0101010 32KB 20 **30x energy reduction** 27x area reduction 1MB  $100$ MULT energy (pJ) ADD area (um<sup>2</sup>) **DRAM** 1300-INT8 INT32 FP16 **FP32** 2600 INT8 INT32 **FP16 FP32**  $0.2$  $3.1$  $1.1$  $3.7$ Up to  $4x$ 36 137 1360 4184 energy 18.5x energy reduction 116x area reduction reduction

Sources: Mark Horowitz (Stanford), energy based on ASIC, area based on TSMC 45nm process Wikimedia Commons (co) (a)

11

# 8-bit Integer (INT8)

- $\triangleright$  INT8 quantization is widely used for model deployment in edge devices and mobile platforms.
- $\blacktriangleright$  Key benefits of INT8 quantization:
	- $\blacktriangleright$  Memory efficiency: 8-bit integers require one-quarter of the memory compared to FP32.
	- $\blacktriangleright$  Faster inference: Lower bit-width operations are less computationally expensive, reducing both memory access time and arithmetic operations.
	- $\triangleright$  Energy savings: Typically results in lower power consumption, especially in specialized hardware like TPUs, GPUs, and edge AI accelerators.
- ▶ Example: Intel's Neural Network Processor (NNP) and Google's Edge TPUs provide dedicated support for INT8 inference, reducing energy consumption by up to 4x compared to FP32.

# 4-bit Quantization

- $\triangleright$  4-bit quantization is an emerging trend, focused on ultra-low-power AI applications.
- $\blacktriangleright$  Benefits of 4-bit quantization:
	- $\blacktriangleright$  Extremely low memory and storage requirements.
	- $\triangleright$  Significant reduction in energy usage, particularly for inference.
- $\blacktriangleright$  Trade-offs:
	- $\triangleright$  **Accuracy loss:** Aggressive quantization can cause a notable decrease in model accuracy.
	- $\blacktriangleright$  Requires specialized hardware that supports low-precision operations.
- $\blacktriangleright$  4-bit quantization is still an area of active research, focusing on improving the trade-off between energy savings and model accuracy.

## Quantization: the bad



low precision  $=$  low accuracy

Challenge: reduce precision without harming accuracy

# Mixed Precision Quantization

- $\triangleright$  Mixed precision quantization involves applying different bitwidths to different parts of the model.
- $\triangleright$  Example: Weights can be quantized to 8 bits while activations remain at 16 bits.
- $\triangleright$  This helps to balance model accuracy with computational efficiency.
- $\blacktriangleright$  Particularly useful if not all parameters are equally important for model expressivity

## Quantization formats





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#### How to quantize?

# Limited (dynamic) range

⇒ value magnitude range

FP32:  $(10^{-38}, 10^{38})$ 

~76 orders of magnitude

FP8:  $(10^{-2}, 10^{2})$ 

~ 4 orders of magnitude

#### How to quantize?

Want to quantize  $X$  to int8

$$
X = \begin{bmatrix} 0.97 & 0.64 & 0.74 & 1.00 \\ 0.58 & 0.84 & 0.84 & 0.81 \\ 0.00 & 0.18 & 0.90 & 0.28 \\ 0.57 & 0.96 & 0.80 & 0.81 \end{bmatrix}
$$

We don't directly quantize it, but rather a scaled version to avoid overflows.

Example: for int8 range= $[-128, 127] = [min_{int8}, max_{int8}]$ 

#### How to quantize?

$$
X=\begin{bmatrix} 0.97 & 0.64 & 0.74 & 1.00 \\ 0.58 & 0.84 & 0.84 & 0.81 \\ 0.00 & 0.18 & 0.90 & 0.28 \\ 0.57 & 0.96 & 0.80 & 0.81 \end{bmatrix}
$$

#### Compute scaling factor

$$
s = \frac{\max_{\text{int8}} - \min_{\text{int8}}}{x_{\max} - x_{\min}} = \frac{127 + 128}{x_{\max} - x_{\min}} = 255
$$

Scale

$$
X_s = s(X - x_{\min}) + \min_{int8}
$$

Quantize

round $(X_s)$ 

Clip the value

$$
X \approx \text{clip}(\text{round}(X_s))
$$

#### Outside the representable range: clip function

The basic idea of the clip function is to restrict values to a predefined range (min\_val, max\_val). The clip function is defined as:

$$
clip(x, min_val, max_val) = \begin{cases} min_val & \text{if } x < min_val \\ x & \text{if } min_val \le x \le max_val \\ max_val & \text{if } x > max_val \end{cases}
$$

For example:

 $clip(5, 0, 10) = 5$  (within the range)  $clip(-3, 0, 10) = 0$  (clipped to min\_val)  $clip(15, 0, 10) = 10$  (clipped to max\_val)

During inference (i.e., for a trained network):



#### During inference (i.e., for a trained network):

· store network parameters in low precision



#### During inference (i.e., for a trained network):

- store network parameters in low precision
- store/compute intermediate signals in low precision



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During training:

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#### During inference (i.e., for a trained network):

- · store network parameters in low precision
- · store/compute intermediate signals in low precision



**During training:** 

· store/compute back propagated gradients in low precision

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# Types of rounding

 $\triangleright$  Traditional Rounding: Typically rounds a number to the nearest integer or a nearby fixed point.

**P** Round to Nearest: Rounds 1.5 to 2 and 2.4 to 2.

- **P** Round Down: Always rounds down (e.g., 2.8 becomes 2).
- **Round Up:** Always rounds up (e.g., 2.1 becomes 3).
- $\triangleright$  Stochastic Rounding: Rounds based on probability, depending on the fractional part of the number.
	- $\triangleright$  For example, if a number is 1.7, it might round to 1 with probability 0.3 and to 2 with probability 0.7.
- $\triangleright$  Key Difference: Stochastic rounding introduces randomness, whereas traditional rounding methods are deterministic.

"Stochastic Rounding: Implementation, Error Analysis, and Applications" M. Croci, M. Fasi, N. Higham, T. Mary, M. Mikaitis, 2021

## Introduction to Stochastic Rounding

- $\triangleright$  Stochastic rounding is a method of rounding numbers where the rounding direction is chosen probabilistically, based on the fractional part of the number.
- $\triangleright$  Unlike traditional rounding methods (e.g., round to nearest, round down), stochastic rounding introduces randomness into the rounding process.
- $\triangleright$  Commonly used in contexts like:
	- $\blacktriangleright$  Neural network quantization in low precision.
	- $\triangleright$  Computationally intensive simulations with low precision.

# Why Stochastic Rounding?

 $E(\hat{x}) = x$ 

 $\blacktriangleright$  In standard rounding methods, bias can accumulate, leading to systematic errors in computations.

- $\triangleright$  Stochastic rounding introduces randomness, which helps to:
	- **Reduce bias:** Prevents systematic overestimation or underestimation of values. This randomness in rounding ensures that the rounding error has an expected value of zero, leading to less bias.
	- $\blacktriangleright$  Preserve statistical properties: Maintains the expected value over a series of rounding operations.
	- $\blacktriangleright$  Improve model accuracy: In neural networks, reduces the impact of rounding errors on training and inference, especially when working with low-precision formats, where small errors can propagate and magnify over multiple layers.

#### How Stochastic Rounding Works

In Let  $x = n + f$ , where *n* is the integer part and *f* is the fractional part.

 $\blacktriangleright$  In stochastic rounding:

Round(x) = 
$$
\begin{cases} n \text{ with probability } 1 - f \\ n + 1 \text{ with probability } f \end{cases}
$$



Figure 2.1. Stochastic rounding rounds the real number x to the next smaller number |x| in F or to the next larger number  $\lceil x \rceil$  in F. In this example, RN rounds x to  $\lceil x \rceil$ , whereas mode 1 SR can round to either  $\lceil x \rceil$  or  $\lceil x \rceil$  but is more likely to round to  $\lceil x \rceil$ .

# Stochastic Rounding (SR) in Neural Networks

#### Not a new idea:

Höhfeld M, Fahlman SE. 1992 "Probabilistic rounding in neural network learning with limited precision"

- $\triangleright$  Useful in NN, especially in low-precision formats like INT8 or FP16.
- $\blacktriangleright$  In low-precision arithmetic, rounding errors can significantly impact model performance due to the limited number of bits
- $\triangleright$  The bias introduced by quantization can be reduced, leading to:
	- Better accuracy: Helps to maintain the distribution of weights and activations, preserving model accuracy
	- $\blacktriangleright$  Improved robustness: Less sensitive to the propagation of rounding errors over multiple layers.
- $\triangleright$  SR can outperform traditional rounding methods in certain quantized neural networks:

Gupta S, Agrawal A, Gopalakrishnan K, Narayanan P. 2015 "Deep Learning with Limited Numerical Precision"

## Limitations of RN for low precision

"On Stochastic Roundoff Errors in Gradient Descent with Low- Precision Computation" Xia, L., Massei, S., Hochstenbach, M. E., Koren, B. (2024).



Fig. 2 Minimizing  $f(x) = (x - 1024)^2$  using GD with binary 8 ( $u = 2^{-3}$ ) and RN, where the red area indicates where stagnation occurs

#### SR for neural networks

Gupta S, Agrawal A, Gopalakrishnan K, Narayanan P. 2015 "Deep Learning with Limited Numerical Precision"



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#### Challenges and Limitations

- $\blacktriangleright$  Hardware support: Not all hardware accelerators natively support stochastic rounding, requiring custom implementations.
- $\blacktriangleright$  Higher complexity: The randomness involved in stochastic rounding can make it more difficult to analyze and debug models.

# **Quantization-Based Methods for Efficient DNN Inference**

14

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# **Quantization for efficient DNN inference**



- **+** no need for access to training pipeline
- **+** data-free or small calibration set used
- **+** usually fast, with simple API
- **-** lower accuracy at lower bit widths

### Post Training Quantization (PTQ) Quantization-Aware Training (QAT)



**-** access to training pipeline & labelled data

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- **-** longer training times
- **-** hyper-parameter tuning needed
- **+** higher accuracy in general

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### <span id="page-38-0"></span>[Quantization](#page-8-0)

[Generalities](#page-9-0)

### [Quantization aware training \(QAT\)](#page-38-0)

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[Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

# Training with Quantization

- $\blacktriangleright$  Quantization-Aware Training (QAT): A technique where quantization is simulated during training.
- $\blacktriangleright$  The model is trained with quantized weights and/or activations to maintain accuracy.
- $\triangleright$  During training, the gradients are computed as if the model were full precision.

# **QAT: backward path quantization simulation th quantization simulation**<br> **chappen:** How can we back propagate through quantization layers?<br>
—round-to-nearest does not have meaningful gradients<br>
(i.e., either zero or undefined everywhere)



### Problem: How can we back propagate through quantization layers?

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- ➡round-to-nearest does not have meaningful gradients
- ➡gradient-based training seems impossible

# **QAT: backward path quantization simulation**



[1] Estimating or Propagating Gradients Through Stochastic Neurons for Conditional Computation, *Bengio et al.*, arXiv:1308.3432, 2013

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# **QAT: backward path quantization simulation**



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# **QAT: backward path quantization simulation**



Schematic view of a QAT procedure with STE applied (adapted from [1])

[1] A Survey of Quantization Methods for Efficient Neural Network Inference, *Gholami et al.*, arXiv:2103.13630, 2021

 $A \cup B \rightarrow A \oplus B \rightarrow A \oplus B \rightarrow A \oplus B$ 

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# Early success of QAT: BinaryConnect

["BinaryConnect: Training Deep Neural Networks with binary weights during propagations", M. Courbariaux, Y. Bengio, JP David, 2016]

- ▶ BinaryConnect is a method for training neural networks with binary weights and activations.
- $\blacktriangleright$  During training, the weights are binarized (approximated to  $+1$  or -1).
- $\blacktriangleright$  The activations can also be binarized, reducing the computational cost of both forward and backward passes.

# How BinaryConnect Works

- $\triangleright$  Keeps double copy of the weights: binary weights  $W_{bin}$  and continuous weights W
- $\blacktriangleright$  Train the model using real-valued weights (for gradient computation), but at each step, constrain the weights to binary values after the weight update step.
- $\blacktriangleright$  Reduce memory and computation for inference, while still benefiting from continuous gradients during training (essential for SGD to work).

# How BinaryConnect Works

 $\blacktriangleright$  During forward pass:

 $\triangleright$  We compute the activations using binary weights

### $\blacktriangleright$  During backpropagation:

- $\blacktriangleright$  The gradients are computed as if the network were using real-valued weights.
- $\blacktriangleright$  The continous weights are updated based on these gradients.
- $\blacktriangleright$  The weights are binarized using the sign function:

$$
w_{\text{binary}} = \text{sign}(w)
$$

34 / 78

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# Algorithm: BinaryConnect

### Input:

- $\blacktriangleright$  Training data  $x_i, y_i$  for  $i = 1, \ldots, N$
- $\triangleright$  Neural network architecture (e.g., layers, activation functions)
- $\blacktriangleright$  Number of epochs  $T$
- **Learning rate**  $\eta$

**Output:** Trained binary weights  $W_{bin}$  and biases b (usually in full-precision)

# Algorithm: BinaryConnect

- 1. Initialize weights W and biases b with small random values
- 2. Initialize the binary weights  $W_{bin} = sign(W)$
- 3. For each epoch  $t = 1, \ldots, T$ :
	- 3.1 For each training sample  $(x_i, y_i)$ 
		- 3.1.1 Perform forward pass: calculate activations using the binary weights  $W_{bin}: a_i = \sigma(W_{bin}x_i + b)$
		- 3.1.2 Compute the loss  $L(x_i, y_i)$  (e.g., cross-entropy or mean squared error)
		- 3.1.3 Perform backward pass (backpropagation): compute the gradients  $\nabla L$  with respect to continuous weights W
		- 3.1.4 Update continuous weights using the continuous gradients  $\nabla L: W = \text{clip}(W - \eta \nabla_W L, -1, 1), b = b - \eta \nabla_b L$
		- 3.1.5 Update binary weights: binarize the continuous weights  $W_{bin}$  = sign(W)
- 4. Return the final binary weights  $W_{bin}$  and biases b

Key points:

- $\triangleright$  compute gradients as if the weights were continuous, even though they are binary:  $\nabla L(W)$  by using the straight-through estimator (STE) for the sign function.
- $\triangleright$  Update the weights in full precision and then binarize:

 $W = W - \eta \nabla L(W)$ 

# Binarizing the weights

Often it is better to binarize stochastically

$$
w_{bin} = \begin{cases} +1 & \text{with } p = \text{clip}(w, 0, 1) \\ -1 & \text{with } 1 - p \end{cases}
$$

instead that

 $W_{bin}$  = sign(W)

$$
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$$

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# Advantages of BinaryConnect

- $\blacktriangleright$  Extreme reduction of memory requirements: binary weights take only 1 bit per weight.
- $\blacktriangleright$  Faster computations: using binary values accelerates inference and training
- $\triangleright$  Suitable for embedded systems and mobile devices.
- $\triangleright$  Often retains near state-of-the-art performance despite the simplifications.

# Numerical results



# Challenges and Limitations

- ▶ Binarizing weights can lead to reduced model expressiveness and lower performance for complex tasks.
- $\triangleright$  While STE works well in practice for many cases, it is still an approximation: the learning process may be less efficient or may converge to suboptimal solutions in some scenarios.
- $\triangleright$  Not all types of neural network architectures are suitable for binary weights (CNN, GANs, transformers..)
- ▶ BinaryConnect may require more epochs or larger learning rates to achieve comparable performance to networks with continuous weights.
- $\triangleright$  The optimization landscape may also be more noisy or less smooth due to the discretization of the weights

### <span id="page-54-0"></span>[Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

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### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

# Post-Training Quantization

- $\triangleright$  Post-Training Quantization (PTQ) refers to the process of quantizing a pre-trained model without retraining.
- $\blacktriangleright$  Typically applied after training a high-precision model (e.g., 32-bit floating point) for deployment on resource-constrained devices.
- It involves converting the model's weights and/or activations to lower bitwidth (e.g., 8-bit integers).

# Methods for Post-Training Quantization

- $\triangleright$  Quantizing Weights: Weights can be quantized after training to reduce the model's size.
- $\triangleright$  Quantizing Activations: Activations are also quantized during inference to further reduce computational costs.
- $\triangleright$  **Calibration:** Calibration is used to select optimal scaling factors for quantization. It typically involves running a small dataset through the model to estimate the range of activations.

# Calibration Process

- $\triangleright$  Calibration helps to determine the scaling factors that best preserve the model's accuracy after quantization.
- $\triangleright$  Common calibration techniques include:
	- $\triangleright$  Min-Max Calibration: Finds the minimum and maximum values of activations and weights to determine the quantization range.
	- $\blacktriangleright$  Histogram-based Calibration: Uses a histogram of activation values to set more precise scaling factors.
- $\triangleright$  Calibration is especially important when quantizing activations to prevent a significant accuracy drop.

# <span id="page-58-0"></span>Challenges in Post-Training Quantization

- $\triangleright$  Calibration Sensitivity: The accuracy of the calibration process is critical, and improper calibration can lead to significant performance drops.
- $\blacktriangleright$  Automated Calibration Methods: Advances in machine learning-based calibration techniques that do not require manual tuning.
- $\triangleright$  Non-Uniform Distributions: Some models have highly non-uniform weight distributions, making it harder to quantize efficiently.

### <span id="page-59-0"></span>[Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

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### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)

# Limitations of classical quantization schemes

- $\blacktriangleright$  Traditional quantization methods usually focus on reducing bit-width uniformly across layers or channels.
- $\triangleright$  Limitations: Such methods can lead to significant accuracy loss because they ignore how sensitive to quantization the parameters in the different layers are.
- $\triangleright$  Solution: Use second-order information (i.e., the Hessian matrix) to guide the quantization process.
- $\blacktriangleright$  Hessian Matrix: Captures the sensitivity of the model's loss function with respect to the parameters, providing richer information for quantization decisions.

## Origins of Hessian aware quantization - 1994



Idea: Taylor development

$$
f(x) - f(\tilde{x}) \approx \nabla f(\tilde{x})^{\mathsf{T}}(x - \tilde{x}) + \frac{1}{2}(x - \tilde{x})^{\mathsf{T}} H(\tilde{x})(x - \tilde{x})
$$

We want to find parameters  $x_i$  that make  $f(x) - f(\tilde{x})$  small to suppress them.

> $\mathbf{A} \oplus \mathbf{B} \rightarrow \mathbf{A} \oplus \mathbf{B} \rightarrow \mathbf{A} \oplus \mathbf{B} \rightarrow \mathbf{A} \oplus \mathbf{B} \rightarrow \mathbf{A}$ 49 / 78

# OBD pruning

### Simplifying assumptions:

At convergence:  $\nabla f(\tilde{x}) \approx 0$ 

$$
\triangleright \frac{1}{2}(x-\tilde{x})^{\mathsf{T}}H(\tilde{x})(x-\tilde{x})=\frac{1}{2}\sum_{i}H_{i,i}x_{i}^{2}+\frac{1}{2}\sum_{i\neq j}H_{i,j}x_{i}x_{j}
$$

- $\triangleright$  We neglect the last term to reduce cost
- $\blacktriangleright$   $\frac{1}{2}\sum_i H_{i,i}x_i^2$  can be efficiently computed by backpropagation with cost similar to that of the gradient

50 / 78

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**Saliency** of a parameter:  $s_i = H_{i,i} x_i^2/2$ 

# OBD procedure

### Procedure:

- 1. Choose a reasonable network architecture
- 2. Train the network until a reasonable solution is obtained
- 3. Compute the second derivatives for each parameter
- 4. Compute the saliencies for each parameter
- 5. Sort the parameters by saliency and delete some low-saliency parameters
- 6. Iterate to step 2

# Numerical results



Figure 1: (a) Objective function (in dB) versus number of parameters for OBD (lower curve) and magnitude-based parameter deletion (upper curve). (b) Predicted and actual objective function versus number of parameters. The predicted value (lower curve) is the sum of the saliencies of the deleted parameters.

# HAWQ: Hessian AWare Quantization

HAWQ: Hessian AWare Quantization of Neural Networks with Mixed-Precision, Z. Dong, Z. Yao, A. Gholami, M. Mahoney, K. Keutzer, 2019

- $\blacktriangleright$  HAWQ leverages the Hessian matrix of the loss function to guide the quantization of weights.
- $\blacktriangleright$  The Hessian matrix provides second-order information about the importance of each weight for the model's performance.
- $\triangleright$  The idea behind HAWQ is to reduce the precision (bit-width) of the weights that have low sensitivity to the loss, while preserving precision in important weights.
- $\triangleright$  This method helps minimize accuracy degradation during quantization, while achieving substantial compression.

# How to measure sensitivity?

- $\triangleright$  Compute the eigenvalues of the Hessian of each block in the network.
- $\triangleright$  Important: it is not possible to explicitly form the Hessian since the size of a block can be quite large.
- $\triangleright$  Solution: compute the Hessian eigenvalues without explicitly forming it, using a matrix-free power iteration algorithm

## Why the eigenvalues of the Hessian?



Fig. 1: Top eigenvalue of each individual block of pre-trained ResNet20 on Cifar-10 (Left), and Inception-V3 on ImageNet (Right). Note that the magnitudes of eigenvalues of different blocks varies by orders of magnitude. See Figure  $\delta$  and  $\bar{\beta}$  in appendix for the 3D loss landscape of other blocks.



Fig. 2: 1-D loss landscape for different blocks of ResNet20 on Cifar-10. The landscape is plotted by perturbing model weights along the top Hessian eigenvector of each block, with a magnitude of  $\epsilon$  (i.e.,  $\epsilon = 0$  corresponds to no perturbation).

# <span id="page-68-0"></span>Hessian-Aware Quantization (HAWQ) Overview

- 1 Train the model with full precision (e.g., FP32).
- 2 Compute the sensitivity measure based on the Hessian eigenvalues
- 3 Apply adaptive quantization based on the sensitivity of the weights.
- 4 Fine-tune the model with quantized weights to minimize accuracy loss (quantization-aware multi-stage re-training)

<span id="page-69-0"></span>**Algorithm 2: Hessian AWare Quantization** 

**Input:** Block-wise Hessian eigenvalues  $\lambda_i$  (computed from Algorithm  $\overline{1}$ , and block size  $n_i$  for  $i=1,\cdots,b.$ for  $i = 1, 2, ..., b$  do // Compute Quantization Precision  $S_i = \lambda_i/n_i$ // See Eq.  $|5|$ Order  $S_i$  in descending order and to determine relative quantization precision for each block. Compute  $\Delta W_i$  based on Eq. 2. **for**  $i = 1, 2, ..., b$  **do** // Fine-Tuning Order  $\Omega_i = \lambda_i ||\Delta W_i||^2$ // See Eq.  $6$ Order  $\Omega_i$  in descending order and perform block-wise fine-tuning

$$
\Delta W_i = Q(W_i) - W_i
$$

Fine tuning intuition : first fine-tune layers that have high curvature, which cause more perturbations a[fte](#page-68-0)[r q](#page-70-0)[u](#page-68-0)[an](#page-69-0)[t](#page-70-0)[i](#page-58-0)[z](#page-59-0)[a](#page-70-0)[ti](#page-71-0)[o](#page-7-0)[n](#page-8-0)[.](#page-70-0)

 $\Omega$ 57 / 78

Block	Laver(s)	Layer Type	Parameter Size	Weight bit	Activation bit
Block 0	Laver <sub>0</sub>	Conv	4.32e2	8	8
Block 1	Layer 1-2	Conv	4.61e3	6	$\overline{4}$
Block 2	Laver 3-4	Conv	4.61e3	6	$\overline{4}$
Block 3	Laver 5-6	Conv	4.61e <sub>3</sub>	8	$\overline{4}$
Block 4	Layer 7-8	Conv	1.38e4	3	$\overline{4}$
Block 5	Layer 9-10	Conv	1.84e4	3	$\overline{4}$
Block 6	Laver 11-12	Conv	1.84e4	3	$\overline{4}$
Block 7	Layer 13-14	Conv	5.53e4	$\overline{2}$	$\overline{4}$
Block 8	Layer 15-16	Conv	7.37e4	$\overline{c}$	$\overline{4}$
Block 9	Layer 17-18	Conv	7.37e4	$\overline{2}$	$\overline{4}$
Block 10	Laver 19	FC	6.40e2	3	8

<span id="page-70-0"></span>Table VI: Block seperation and final block precision of ResNet20 on Cifar-10. Here we abbreviate convolutional laver as "Conv," fully connected layer as "FC."

# <span id="page-71-0"></span>**Outline**

### [Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

### [Sparsification](#page-71-0)

[Pruning](#page-72-0) [Structured sparsification](#page-101-0)
#### <span id="page-72-0"></span>[Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

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#### [Sparsification](#page-71-0) [Pruning](#page-72-0) [Structured sparsification](#page-101-0)

# Pruning

Definition: Select some neurons and/or weights and suppress them (set to zero)



How to choose?

# Magnitude pruning

- $\blacktriangleright$  Many of the learned weights have small magnitudes and contribute little to the network's performance.
- $\triangleright$  Magnitude pruning removes weights that have small magnitudes, reducing the complexity of the model.
- $\blacktriangleright$  Pruning criterion: given a weight  $w_i$ , prune if

 $|w_i| < \tau$ 

62 / 78

with  $\tau$  chosen pruning threshold.

# Pruning Process

- 1. Train the full Model
- 2. Compute Magnitudes of the weights
- 3. Sort weights based on their absolute magnitudes.
- 4. Set threshold  $\tau$  (Top-k pruning, Percentage pruning, Global threshold)
- 5. Prune the weights that are below the threshold (set them to zero)
- 6. Retrain the model (fine-tune) for a few more epochs with the found mask. This allows the model to adjust to the new sparsity pattern and recover any lost performance.

# Magnitude pruning

Advantage: pruning small weights can also act as a form of regularization, helping the model generalize better by reducing overfitting.

#### Variants:

- $\blacktriangleright$  Layer-wise pruning
- ▶ Structured Pruning (removes entire neurons, filters (in CNNs), or channels in the network). This leads to a more structured sparsity pattern and can take advantage of hardware optimizations for matrix or tensor operations.
- Iterative Pruning: prune the network iteratively, pruning a small fraction of weights at each step and retraining the model after each pruning phase. This helps in minimizing performance degradation.

## Other pruning criteria

- $\triangleright$  Gradient pruning: reducing the number of parameters that are updated in each iteration by setting to zero small gradients  $\rightarrow$ limitation: saturation (vanishing gradients), better Hessian pruning
- $\blacktriangleright$  L<sub>1</sub> regularization  $\rightarrow$  limitation: tuning of  $\lambda$

$$
regloss = loss + \lambda ||w||_1
$$

### Lottery tickets

The Lottery Ticket Hypothesis: Finding Sparse, Trainable Neural Networks, J.Frankle, M. Carbin, 2019

#### Objective: Find a subnetwork of a large network, such that, if trained starting from the same  $w_0$  maintains the same performance as the large network



### Lottery ticket algorithm

- $\blacktriangleright$  Choose  $w_0$  random
- ▶ Train the full network starting from  $w_0$  and get  $w^*$
- ▶ Prune the network based on the magnitude of  $w^*$ : select a mask m (binary matrix  $0/1$ ) and set  $w_p = w \cdot m$
- Reset  $w_p = w_0 \cdot m$
- $\blacktriangleright$  Train sub network with just weights  $w_p$

#### What do we expect from the subnetwork?



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#### Training time  $\# \text{Iter}(M, I)$  of model M with initialization /



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#### Training time  $\#\text{Iter}(M, I)$  of model M with initialization / given  $(D, A, H, L)$  (datasets, learning algorithm, hyperparameters, loss)

First iteration at which it reaches minimum validation loss



 $(M, I)$  is said to learn faster than  $(M', I')$  on  $(D, A, H, L)$  if

 $\#\text{Iter}(M,I) \leq \#\text{Iter}(M',I')$ 

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**Remark 1:** If the cost of one iteration of  $(D, A, H, L)$  for  $(M', I')$  is much cheaper than for  $(M, I)$ , then the actual training time on a machine for  $M'$ could be smaller than the one for M.

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**Remark 1:** If the cost of one iteration of  $(D, A, H, L)$  for  $(M', I')$  is much cheaper than for  $(M, I)$ , then the actual training time on a machine for M' could be smaller than the one for  $M$ . For instance if  $M'$  is a smaller model than  $M$ , then an iteration for  $M'$  is likely to be cheaper than for  $M$ : be cautious if  $\#\text{Iter}(M, I) \leq \#\text{Iter}(M', I').$ 

**Remark 2:** Doing gradient-descent, if  $M'$  is a subnet of M and if it is trained by computing all the gradients of  $M$  and then zeroing the ones not in  $M'$ , then an iteration for  $M'$  should have the same cost as for M. Remark 1 does not apply in this case.

#### **Notations**

#### **Notations:**  $s \in [0, 1]$  = level of sparsity,  $M_s$  = subnet of M of sparsity s

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Figure: Dashed lines show the average training time for random subnets with random initializations of given sparsity of a fixed original model.

#Iter(original model)  $\leq$  #Iter(random  $M_s$ )

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Figure: Dashed lines show the average top 1 accuracy for random subnets with random initializations of given sparsity of a fixed original model.

Top1(trained random  $M_s$ )  $\leq$  Top1(trained original model)

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Average sparse subnets learn slower than the average trained full **model:** for *M* and *s* considered in the experiments and empirical means:

 $\mathbb{E}_l\# \text{Iter}(M, I) \leq \mathbb{E}_{M_s,l} \# \text{Iter}(M_s, I)$  $\max_{l} \# \text{Iter}(M, l) \leq \min_{M \leq l} \# \text{Iter}(M_s, l)$  for s not too close from 100%

Average sparse subnets learn slower than the average trained full **model:** for *M* and *s* considered in the experiments and empirical means:

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Average sparse subnets are less accurate than the average trained full model: for M and s considered in the experiments and empirical means<sup>.</sup>

 $\mathbb{E}_I$ Top1 $(M, I) \geq \mathbb{E}_{M_s, I}$ Top1 $(M_s, I)$  $\min_{l} \text{Top1}(M, l) \ge \max_{M_s, l} \text{Top1}(M_s, l)$  for s not too close from 100%

지하다 지수는 어려운 사람들이 그를 지수야 없다.



Is it possible to find early on during training a sparse subnet that trains faster than the original model without accuracy degradation?



### Definition of lottery tickets:  $(M_s, I)$  versus  $(M, I)$

Lottery ticket: Fix  $(D, A, H, L)$ . Consider a model and an initialization  $(M, I)$ . A lottery ticket is a submodel  $(M_s, I)$  of sparsity s of  $(M, I)$ .

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Definition of lottery tickets:  $(M_s, I)$  versus  $(M, I)$ 

**Lottery ticket:** Fix  $(D, A, H, L)$ . Consider a model and an initialization  $(M, I)$ . A lottery ticket is a submodel  $(M_s, I)$  of sparsity s of  $(M, I)$ .

• Average lottery ticket learns slower than the average full model:

 $\mathbb{E}_{l} \# \text{Iter}(M, I) \leq \mathbb{E}_{M_{\rm c}} \# \text{Iter}(M_{\rm s}, I),$  $\forall M. \forall s.$ 

Definition of lottery tickets:  $(M_s, I)$  versus  $(M, I)$ 

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 $\mathbb{E}_{l} \# \text{Iter}(M, I) \leq \mathbb{E}_{M_{\rm c}} \# \text{Iter}(M_{\rm s}, I),$  $\forall M. \forall s.$ 

• Average lottery ticket is less accurate than the average trained full model:

 $\mathbb{E}_I$ Top1 $(M, I) \geq \mathbb{E}_{M_{\rm c}}$  Top1 $(M_{\rm s}, I)$ ,  $\forall M, \forall s$ .

## Definition: Winning tickets

**Winning lottery ticket:** Fix  $(D, A, H, L, Top1)$  with Top1 a measure of accuracy. Consider a model and an initialization  $(M, I)$ . A lottery ticket is a submodel  $(M_s, I)$  of sparsity s is winning if:

• it learns faster than the original model:  $\#\text{Iter}(M_{s}, I) \leq \#\text{Iter}(M, I)$ 

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• it is more accurate than the original model:  $\text{Top1}(M, I) \leq \text{Top1}(M_s, I)$ 

# <span id="page-98-0"></span>Algorithm to find winning tickets

Algorithm of Iterative Pruning to find winning tickets:

- $\blacktriangleright$  Train original model
- **If** Layer-wise, prune  $p = 20\%$  of the weights with the smallest magnitude ( $p/2\%$  for the output layer)
- $\blacktriangleright$  Iterate until desire sparsity is achieved

Novelty: IMP find subnets that can be trained efficiently from the start for unprecedented small level of sparsity without degradation of accuracy

### Empirical performance of tickets found by Iterative Pruning

**Learns faster**<sup>1</sup> than the original model for  $3.6\% \le s \le 100\%$ :  $\mathbb{E}_l \# \text{Iter}(\text{IMP}(M, s), l) \leq \mathbb{E}_l \# \text{Iter}(M, l)$ 



Figure: Dashed lines: random  $M_s$ , I. Solid lines: random IMP $(M, s)$ , I.

<sup>1</sup>The training time decreases from  $s = 100\%$  to  $s = 21\%$  at which point early-stopping occurs 38% earlier than for the original model, then it increases

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#### Empirical performance of tickets found by Iterative Pruning

More accurate<sup>2</sup> than the original model for  $3.6\% \leqslant s \leqslant 100\%$ :  $\mathbb{E}_I$ Top1(IMP $(M, s), I) \geq \mathbb{E}_I$ Top1 $(M, I)$ 



<sup>2</sup>The accuracy increases from  $s = 100\%$  to  $s = 13.5\%$  where it gained 0.3%, then it decreases  $QQ$ 

#### <span id="page-101-0"></span>[Quantization](#page-8-0)

[Generalities](#page-9-0) [Quantization aware training \(QAT\)](#page-38-0) [Post training quantization \(PTQ\)](#page-54-0) [Mixed precision quantization](#page-59-0)

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[Sparsification](#page-71-0) [Pruning](#page-72-0) [Structured sparsification](#page-101-0) Aim: replace dense weight matrices with structured ones (e.g., sparse, low-rank, Fourier transform).

These methods have not seen widespread adoption:

- $\triangleright$  in end-to-end training due to unfavorable efficiency-quality tradeoffs,
- $\triangleright$  in dense-to-sparse fine-tuning of pretrained models due to lack of tractable algorithms to approximate a given dense weight matrix

### Monarch networks

["Monarch: Expressive Structured Matrices for Efficient and Accurate Training", T. Dao et all, 2022]

#### Monarch matrices:

- $\blacktriangleright$  hardware-efficient (they are parameterized as products of two block-diagonal matrices for better hardware utilization)
- $\triangleright$  expressive (they can represent many commonly used transforms).
- $\blacktriangleright$  The problem of approximating a dense weight matrix with a Monarch matrix, though nonconvex, has an analytical optimal solution.

#### Monarch matrices

**Definition 3.1.** Let  $n = m^2$ . An  $n \times n$  *Monarch matrix* has the form:

$$
\mathbf{M} = \mathbf{PLP}^{\top} \mathbf{R},
$$

where **L** and **R** are block-diagonal matrices, each with m blocks of size  $m \times m$ , and **P** is the permutation that maps  $[x_1, \ldots, x_n]$  to  $[x_1, x_{1+m}, \ldots, x_{1+(m-1)m}, x_2, x_{2+m}, \ldots,$  $x_{2+(m-1)m},\ldots,x_m,x_{2m},\ldots,x_n].$ 



Figure 2: Monarch matrices are parametrized as products of two block-diagonal matrices up to permutation, allowing efficient multiplication algorithm that leverages batch matrix multiply.

We can interpret  $P=P^{\mathcal{T}}$  as follows: it reshapes a vector  $x$  of size n as a matrix of size  $m \times m$ , transposes the matrix, then converts back into a vector of size n.

### Link with butterfly matrices

In Let B butterfly matrix of size n where n is a power of 4.

$$
B = B_1 \dots B_{\log_2(n)/2} B_{\log_2(n)/2} \dots B_{\log_2(n)}
$$

- ▶ R is block-diagonal with  $m = \sqrt{n}$  dense blocks, each block of size  $m \times m$
- If L' is composed of  $m \times m$  blocks of size  $m \times m$ , where each block is a diagonal matrix:

$$
L' = \begin{bmatrix} D_{11} & \dots & D_{1m} \\ \dots & \dots & \dots \\ D_{m1} & \dots & D_{mm} \end{bmatrix}
$$

- $\blacktriangleright$  L' can be written as block-diagonal with the same structure as R after permuting the rows and columns.
- $\blacktriangleright$   $L = PL'P^{T}$ : up to permuting rows and columns,  $L'$  is also a block-diagonal matrix of m dense blocks, each of size  $m \times m$ .
- $\triangleright$  B butterly implies B monarch

#### Special case of matrix factorization algorithm - cf. Cours 10 Sparse matrix factorization

 $\rightarrow$  Solve a series of block SVDs

### Compression - end-to-end (E2E)

Replacing dense matrices with Monarch matrices in Vision Transformer ViT, MLP-Mixer (ImageNet), and GPT-2 (WikiText-103) can speed up training by up to  $2\times$  without sacrificing model quality

Table 1: The performance of Monarch matrices and ViT / MLP-Mixer on ImageNet, including the number of parameters and FLOPs. We measure the Top-1 accuracy and the training time speedup compared to the corresponding dense model.

Model	ImageNet acc. Speedup Params FLOPs			
$Mixer-S/16$	74.0	٠	18.5M	3.8G
Monarch-Mixer-S/16	73.7	$1.7\times$	7.0M	1.5G
$Mixer-B/16$	77.7	۰	59.9M	12.6G
Monarch-Mixer-B/16	77.8	$1.9\times$	20.9M	5.0G
$ViT-S/16$	79.4	٠	48.8M	9.9G
Monarch-ViT-S/16	79.1	$1.9\times$	19.6M	3.9G
$ViT-B/16$	78.5	۰	86.6M	17.6G
Monarch-ViT-B/16	78.9	$2.0\times$	33.0M	5.9G

Table 2: Performance of Monarch matrices and GPT-2-Small/Medium on WikiText-103, including the # of parameters and FLOPs. Monarch achieves similar perplexity (ppl) but 2.0× faster.


## Compression - Denso-to-sparse

**Procedure**: BERT pretrained weights, approximate them with Monarch matrices, and finetune the resulting model on the 9 GLUE tasks (collection of nine natural language understanding tasks).

**Result:** Monarch finetuned model with similar quality to the dense BERT model, but with  $1.7\times$  faster finetuning speed.<br>Table 8: The performance of Monarch matrices in finetuning BERT on GLUE.





## Pixelated butterfly

Pixelated Butterfly: Simple and Efficient Sparse Training for Neural Network Models, Dao et all, 2022

- $\triangleright$  An approach similar to the previous one, but that uses butterfly factorizations
- $\triangleright$  As classical butterfly matrices are not hardware efficient, they propose variants of butterfly (block and flat) to take advantage of modern hardware.



Recent development : "Fast inference with Kronecker-sparse matrices" A. Gonon, L. Zheng, P. Carrivain, Q. Le, 2024 (GPU matrix multiplication algorithms specialized for Kronecker-sparse matrices) 

## Exercise: quantization of neural networks

Objective: Understand and apply quantization techniques to neural networks. Experiment with a post-training quantization technique. Evaluate the trade-offs between model size, inference speed, and accuracy.

## Exercise Overview:

- 1. Train a basic neural network on a dataset (e.g., MNIST).
- 2. Apply post-training quantization to the trained model.
- 3. Compare the performance of the quantized models with the original floating-point model:
	- $\triangleright$  Model Performance: check how accuracy changes with quantization and quantization-aware training.
	- Inference Speed: measure the impact of quantization on the inference time. Does it decrease? Why?
	- $\triangleright$  Model Size: check how much the model size is reduced by quantization.