Evaluating the Feasibility of Sampling-Based Techniques for Training Multilayer Perceptrons

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Motivation

A scalability issue

- DNNs are often very large in scale:
 - Computationally expensive to train
 - Requiring powerful hardware, including expensive GPUs
 - ▶ CPUs are widely available! why not using them instead?

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Extended Database Technologies for Machine Learning

- Feasibility of the extension of sampling-based techniques for efficient training of deep neural networks (DNNs)
- On CPU machines with limited resources

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Two scalability directions

- Sampling-based approaches based on locality-sensitive hashing (LSH) [GIM99]
- 2 Monte-carlo (MC) estimations [Rob16]

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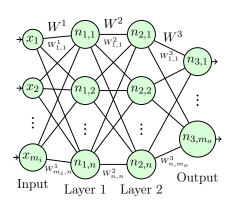
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Review: multi-layer perceptron (MLP)

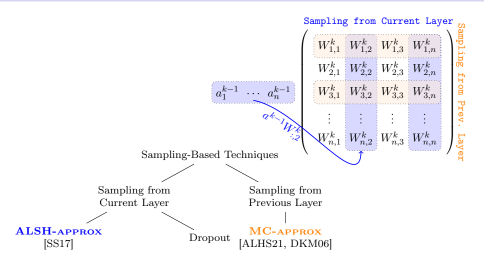
$$z^k = a^{k-1}W^k + b^k$$
$$a^k = f(z^k)$$



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Taxonomy of Sampling-Based Techniques



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Asymmetric Locality-Sensitive Hashing

A family \mathcal{H} is $(\tau, c, p_1 > p_2)$ -sensitive for c-NNS, if for all $h \in \mathcal{H}$:

$$sim(w, a) \ge \tau \implies \Pr[h(Q(a)) = h(P(w))] \ge p_1$$

 $sim(w, a) \le c\tau \implies \Pr[h(Q(a)) = h(P(w))] \le p_2$

For $w, a \in \mathbb{R}^n$ with $||w|| \leq C$, where C is a constant less than 1, and ||a|| = 1, they define the transformations P and Q for the inner product as follows.

$$P: \mathbb{R}^{n} \to \mathbb{R}^{n+m}, \quad w \mapsto \left[w; \|w\|^{2^{1}}, \dots, \|w\|^{2^{m}} \right]$$

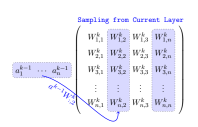
$$Q: \mathbb{R}^{n} \to \mathbb{R}^{n+m}, \quad a \mapsto \left[a; 1/2, \dots, 1/2 \right]$$
(1)

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ALSH-APPROX

• Only computes the high-value elements of $a^{k-1}W^k$ (called active nodes), skipping the computation for the small values.



Uses ALSH to find active nodes: the columns $W_{:,j}^k$ that collide with a^{k-1} into the same bin.

$$\operatorname{argmax}_{j} \left\langle W_{:,j}^{k}, a^{k-1} \right\rangle \approx \operatorname{argmin}_{j} \left\| Q\left(a^{k-1}\right) - P\left(W_{:,j}^{k}\right) \right\|$$

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Approximate Matrix Multiplication $A \times B$

- $\bullet AB_{i,j} = \sum_{t=1}^{n} A_{i,t}B_{t,j}$
- ullet Goal: A Monte-carlo method to estimate the sum over a subset of values of t
- Standard approach: uniformly select a sample of cells $t \in [n]$ \Rightarrow high estimation error
- Idea (weighted sampling): compute the values that are expected to be larger

$$p_t = \frac{\|A_{:,t}\| \cdot \|B_{t,:}\|}{\sum_{k=1}^n \|A_{:,k}\| \cdot \|B_{k,:}\|}$$

For a set C of c samples:

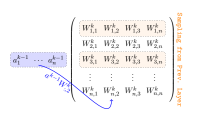
$$AB_{i,j} \approx \sum_{t \in C} \frac{1}{cp_t} A_{i,t} B_{t,j}$$

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MC-APPROX

- Computes all values of $a^{k-1}W^k$, but instead of exact computation, it estimates each value.
- It is more appropriate for mini-batch settings, where a matrix of activation vectors is multiplied to W^k .
- For single-row settings, it may not accurately estimate p_t values \Rightarrow its estimations are not accurate.



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Negative Result: ALSH-APPROX does not scale

Theorem

Let f be a linear activation function such that a = f(z) = z. Suppose for any node n_p^l , $\sum_{i \in \uparrow_p^l} a_i^{l-1} W_{i,p} = c \sum_{i \notin \uparrow_p^l} a_i^{l-1} W_{i,p}$. Then, $a_j^k = \bar{a}_j^k \left(\frac{c+1}{c}\right)^k$. That is,

$$e_j^k = \bar{a}_j^k \left(\left(\frac{c+1}{c} \right)^k - 1 \right)$$

Example: suppose c=5 (i.e., the weighted sum for the active nodes is five times that of the inactive nodes). Then,

1 (1 (1)	-1	0	0	4	_	0
k (number of layers)	1	2	3	4	5	6
$ m e_i^k/ar{a}_i^k$	0.2	0.44	0.72	1.07	1.48	1.98

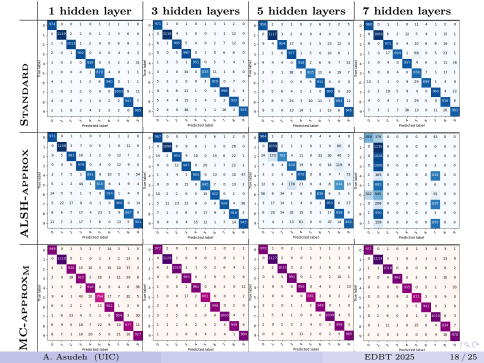
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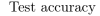
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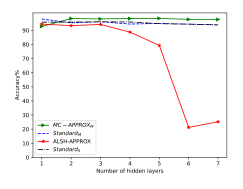


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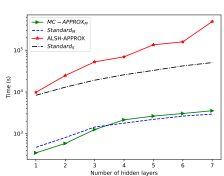


Time & Accuracy Comparison





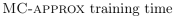
Training time

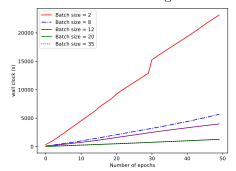


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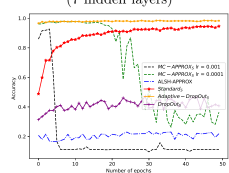
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Time & Accuracy Comparison: MC-APPROX_s failed





Validation accuracies (7 hidden layers)



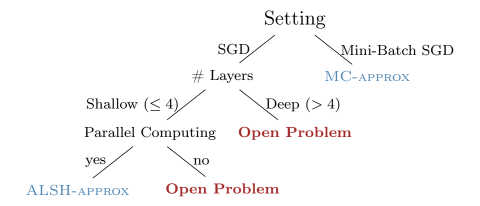
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Summary

Sampling-based Techniques for Training DNNs on CPU machines:



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Next Step Advertisement!

Efficient Matrix Multiplication for Accelerating Inference in Binary and Ternary Neural Networks

- Complexity: $O(\frac{n^2}{\log n})$ time and memory
- Experiments: up to 6X less memory
- On CPU. C++: up to 30X faster; NumPy: 24X faster
- LLM Inference (Llama3.0, Falcon3, etc.), without system-level optimizations:
 - ▶ on CPU: up to 5.24X faster; on GPU: up to 2.5X faster







Thank you!

• InDeX Lab: cs.uic.edu/~indexlab/



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References



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