

Binary Graph Neural Networks

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Abstract

Graph Neural Networks (GNNs) have emerged as a powerful and flexible framework for representation learning on irregular data. As they generalize the operations of classical CNNs on grids to arbitrary topologies, GNNs also bring much of the implementation challenges of their Euclidean counterparts. Model size, memory footprint, and energy consumption are common concerns for many real-world applications. Network binarization allocates a single bit to parameters and activations, thus dramatically reducing the memory requirements (up to 32x compared to single-precision floating-point numbers) and maximizing the benefits of fast SIMD instructions on modern hardware for measurable speedups. However, in spite of the large body of work on binarization for classical CNNs, this area remains largely unexplored in geometric deep learning. In this paper, we present and evaluate different strategies for the binarization of graph neural networks. We show that through careful design of the models, and control of the training process, binary graph neural networks can be trained at only a moderate cost in accuracy on challenging benchmarks. In particular, we present the first dynamic graph neural network in Hamming space, able to leverage efficient k -NN search on binary vectors to speed-up the construction of the dynamic graph. We further verify that the binary models offer significant savings on embedded devices. Our code is publicly available on Github¹.

1. Introduction

Standard CNNs assume their input to have a regular grid structure, and are therefore suitable for data that can be well-represented in an Euclidean space, such as images, sound, or videos. However, many increasingly relevant types of data do not fit this framework [5]. Graph theory offers a broad mathematical formalism for modeling interactions, and is therefore commonly used in fields such as network sciences [12], bioinformatics [24, 40], and recommender systems

¹https://github.com/mbahri/binary_gnn

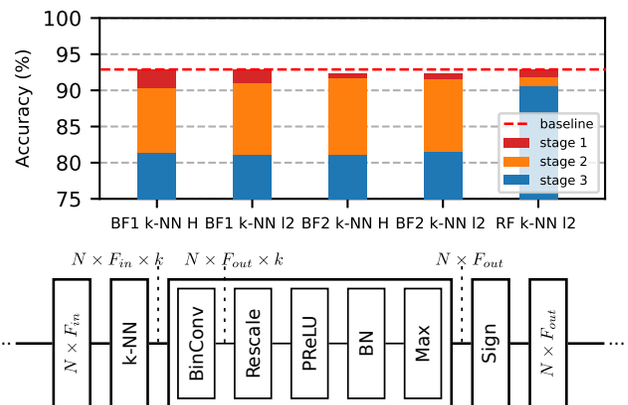


Figure 1. **Top:** Test accuracy of different binarization schemes at all stages of our cascaded distillation protocol (baseline: 92.89%). **Bottom:** The “BF2” variant of our XorEdgeConv operator.

[37], as well as for studying discretisations of continuous mathematical structures such as in computer graphics [4]. This motivates the development of machine learning methods able to deal with graph-supported data. Among them, Graph Neural Networks (GNNs) generalize the operations of CNNs to arbitrary topologies by extending the basic building blocks of CNNs such as convolutions and pooling to graphs. Similarly to CNNs, GNNs learn deep representations of graphs or graph elements, and have emerged as the best performing models for learning on graphs as well as on 3D meshes with the development of advanced and increasingly deep architectures [33, 18].

As the computational complexity of the networks and the scale of graph datasets increase, so does the need for faster and smaller models. The motivations for resource-efficient deep learning are numerous and also apply to deep learning on graphs and 3D shapes. Computer vision models are routinely deployed on embedded devices, such as mobile phones or satellites [2, 32], where energy and storage constraints are important. The development of smart devices and IoT may bring about the need for power-efficient graph learning models [27, 61, 9]. Finally, models that require GPUs for inference can be expensive to serve, whereas CPUs are typically more affordable. This latter point is especially relevant

to the applications of GNNs in large-scale data mining on relational datasets, such as those produced by popular social networks, or in bioinformatics [35].

While recent work has proposed algorithmic changes to make graph neural networks more scalable, such as the use of sampling [21, 60] or architectural improvements [15, 10] and simplifications [56], our approach is orthogonal to these advances and focuses on compressing existing architectures while preserving model performance. Model compression is a well-researched area for Euclidean neural networks, but has seen very little application in geometric deep learning. In this paper, we study different strategies for binarizing GNNs.

Our contributions are as follows:

- We present a binarization strategy inspired by the latest developments in binary neural networks for images [7, 36] and knowledge distillation for graph networks
- We develop an efficient dynamic graph neural network model that constructs the dynamic graph in Hamming space, thus paving the way for significant speedups at inference time, with negligible loss of accuracy when using real-valued weights
- We conduct a thorough ablation study of the hyperparameters and techniques used in our approach
- We demonstrate real-world acceleration of our models on a budget ARM device

Notations Matrices and vectors are denoted by upper and lowercase bold letters (*e.g.*, \mathbf{X} and \mathbf{x}), respectively. \mathbf{I} denotes the identity matrix of compatible dimensions. The i^{th} column of \mathbf{X} is denoted as \mathbf{x}_i . The set of real numbers is denoted by \mathbb{R} . A graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ consists of *vertices* $\mathcal{V} = \{1, \dots, n\}$ and *edges* $\mathcal{E} \subseteq \mathcal{V} \times \mathcal{V}$. The *neighborhood* of vertex i , denoted by $\mathcal{N}(i) = \{j : (i, j) \in \mathcal{E}\}$, is the set of vertices adjacent to i . Other mathematical notations are summarized in Table 5 of the Supplementary Material.

2. Related Work

Knowledge distillation uses a pretrained *teacher* network to supervise and inform the training of a smaller *student* network. In logit matching [22], a cross-entropy loss is used to regularize the output logits of the student by matching them with a blurred version of the teacher’s logits computed using a softmax with an additional temperature hyperparameter. More recent works also focus on matching internal activations of both networks, such as attention volumes in [59], or on preserving relational knowledge [44, 31, 49].

Quantized and Binary Neural Networks Network quantization [19, 62] refers to the practice of lowering the numerical precision of a model in a bid to reduce its size and

speed-up inference. Binary Neural Networks (BNNs) [25] push it to the extreme and use a single bit for weights and activations. The seminal work of XNOR-Net [47] showed that re-introducing a small number of floating point operations in BNNs can drastically improve the performance compared to using pure binary operations by reducing the quantization error. In XNOR-Net, a dot product \star between real vectors \mathbf{a} and \mathbf{b} of dimension n is approximated by $\mathbf{a} \star \mathbf{b} \approx (\text{sign}(\mathbf{a}) \otimes \text{sign}(\mathbf{b}))\alpha\beta$, where $\beta = \frac{1}{n}\|\mathbf{a}\|_1$ and $\alpha = \frac{1}{n}\|\mathbf{b}\|_1$ are rescaling constants. XNOR-Net++ [7] proposed to instead learn a rescaling tensor Γ , with shared factors to limit the number of trainable parameters and avoid overfitting. Finally, in Real-to-Binary networks [36], the authors compile state-of-the-art techniques and improve the performance of binary models with knowledge distillation.

Graph Neural Networks Graph Neural Networks were initially proposed in [20, 48] as a form of recursive neural networks. Later formulations relied on Fourier analysis on graphs using the eigendecomposition of the graph Laplacian [6] and approximations of such [11], but suffered from the connectivity-specific nature of the Laplacian. Attention-based models [38, 14, 51, 50] are purely spatial approaches that compute a vertex’s features as a dynamic weighting of its neighbours’. Spatial and spectral approaches have been unified [29] and shown to derive from the more general neural message passing [17] framework. We refer to recent reviews on GNNs, such as [57], for a comprehensive overview, and focus only on the operators we binarize in this paper.

The message-passing framework offers a general formulation of graph neural networks:

$$\mathbf{x}_i^{(l)} = \gamma^{(l)} \left(\mathbf{x}_i^{(l-1)}, \square_{j \in \mathcal{N}(i)} \phi^{(l)} \left(\mathbf{x}_i^{(l-1)}, \mathbf{x}_j^{(l-1)}, \mathbf{e}_{ij}^{(l-1)} \right) \right), \quad (1)$$

where \square denotes a differentiable symmetric (permutation-invariant) function, (*e.g.* \max or \sum), ϕ a differentiable kernel function, γ is an MLP, and \mathbf{x}_i and \mathbf{e}_{ij} are features associated with vertex i and edge (i, j) , respectively.

The EdgeConv operator is a special case introduced as part of the Dynamic Graph CNN (DGCNN) model [54] and defines an edge message as a function of $\mathbf{x}_j - \mathbf{x}_i$:

$$\begin{aligned} \mathbf{e}_{ij}^{(l)} &= \text{ReLU} \left(\boldsymbol{\theta}^{(l)} (\mathbf{x}_j^{(l-1)} - \mathbf{x}_i^{(l-1)}) + \phi^{(l)} \mathbf{x}_i^{(l-1)} \right) \quad (2) \\ &= \text{ReLU} \left(\boldsymbol{\Theta}^{(l)} \tilde{\mathbf{X}}^{(l-1)} \right) \quad (3) \end{aligned}$$

where $\tilde{\mathbf{X}}^{(l-1)} = \left[\mathbf{x}_i^{(l-1)} \parallel \mathbf{x}_j^{(l-1)} - \mathbf{x}_i^{(l-1)} \right]$, $\boldsymbol{\theta}$ and ϕ are trainable weights, and $\boldsymbol{\Theta}$ their concatenation.

The output of the convolution is the max aggregation ($\square = \max$) of the edge messages:

$$\mathbf{x}_i^{(l)} = \max_{j \in \mathcal{N}(i)} \mathbf{e}_{ij}^{(l)} \quad (4)$$

While the EdgeConv operator is applicable to graph inputs, the main use case presented in [54] is for point clouds, where the neighbours are found by k -Nearest Neighbours (k -NN) search in feature space before each convolutional layer. DGCNN is the first example of a dynamic graph architecture, with follow-up work in [26].

The GraphSAGE [21] operator introduced inductive learning on large graphs with sampling and can also be phrased as a message passing operator:

$$\mathbf{x}_i^{(l)} = \text{Norm} \left(\text{ReLU} \left(\mathbf{W}^{(l)} \left[\mathbf{x}_i^{(l-1)} \parallel \text{Aggr}_{j \in \mathcal{N}(i)} \mathbf{x}_j^{(l-1)} \right] \right) \right) \quad (5)$$

Where Aggr is a symmetric aggregation function such as max, sum or mean; Norm denotes the ℓ_2 normalization, and \mathbf{W} is a tensor of learnable weights.

Model Compression in Geometric Deep Learning In [52], the authors propose to binarize the Graph Attention (GAT) operator [50], and evaluate their method on small-scale datasets such as Cora [39] and Pubmed [29]. In [53], the authors apply the XNOR-Net approach to GCN [29] with success, but also on small-scale datasets. Finally, [46] propose to binarize PointNet with tailored aggregation and scaling functions. At the time of writing, the Local Structure Preserving (LSP) module of [58] is the only knowledge distillation method specifically designed for GNNs. LSP defines local structure vectors LS_i for each node in the graph:

$$LS_{ij} = \frac{\exp(\text{SIM}(\mathbf{x}_i, \mathbf{x}_j))}{\sum_{k \in \mathcal{N}(i)} \exp(\text{SIM}(\mathbf{x}_i, \mathbf{x}_k))} \quad (6)$$

where SIM denotes a similarity measure, *e.g.*, $\|\cdot\|_2^2$ or a kernel function such as a Gaussian RBF kernel. The total local structure preserving loss between a student network s and a teacher t is then defined as:

$$L_{LSP} = \frac{1}{|\mathcal{V}|} \sum_{i \in \mathcal{V}} \sum_{j \in \mathcal{N}^u(i)} LS_{ij}^s \log \frac{LS_{ij}^s}{LS_{ij}^t}. \quad (7)$$

$\mathcal{N}^u(i) = \mathcal{N}^s(i) \cup \mathcal{N}^t(i)$ to support dynamic graph models.

3. Method

Eq. 1 is more general than the vanilla Euclidean convolution, which boils down to a single matrix product to quantize. We must therefore choose which elements of Eq. 1 to binarize and how: the node features \mathbf{x}_i , the edge messages \mathbf{e}_{ij} , and the functions \square , γ and ϕ may all need to be adapted.

Quantization We follow the literature and adopt the sign operator as the binarization function:

$$\text{sign}(x) = \begin{cases} 1 & \text{if } x \geq 0 \\ -1 & \text{if } x < 0 \end{cases}. \quad (8)$$

As the gradient of sign is zero almost everywhere, we employ the straight-through estimator [3] to provide a valid gradient. We use this method for both network weights and activations. Furthermore, we mean-center and clip the real latent network weights after their update in the backpropagation step.

Learnable rescaling Assuming a dot product operation (*e.g.* a fully-connected or convolutional layer) $\mathbf{A} \star \mathbf{B} \in \mathbb{R}^{o \times h \times w}$, we approximate it as in [7]:

$$\mathbf{A} \star \mathbf{B} \approx (\text{sign}(\mathbf{A}) \otimes \text{sign}(\mathbf{B})) \odot \Gamma, \quad (9)$$

with Γ a learned rescaling tensor. We use two constructions of Γ depending on the model. Channel-wise:

$$\Gamma = \alpha \in \mathbb{R}^{o \times 1 \times 1} \quad (10)$$

and one rank-1 factor per mode:

$$\Gamma = \alpha \otimes \beta \otimes \gamma, \quad \alpha \in \mathbb{R}^o, \beta \in \mathbb{R}^h, \gamma \in \mathbb{R}^w \quad (11)$$

Activation functions Recent work [8] has shown using non-linear activations in XNOR-Net - type blocks can improve the performance of binary neural networks, with PReLU bringing the most improvement.

Knowledge Distillation Inspired by [36], we investigate the applicability of knowledge distillation for the binarization of graph neural networks. For classification tasks, we use a logit matching loss [22] as the base distillation method. We also implemented the LSP module of [58].

Multi-stage training We employ a cascaded distillation scheme [36], an overview of which is shown in Figure 2.

Stage 1: We first build a real-valued and real-weighted network with the same architecture as the desired binary network by replacing the quantization function with tanh. We distillate the original (base) network into this first student network. We employ weight decay with weight $\lambda = 1e - 5$, logit matching, and LSP. We use the same initial learning rate and learning rate schedule as for the base network.

Stage 2: The model of stage 1 becomes the teacher, the student is a binary network with real-valued weights but binary activations. We initialize the student with the weights of the teacher. We employ weight decay with $\lambda = 1e - 5$, logit matching, and LSP. We use a smaller learning rate (*e.g.* 25%) than for stage 1 and the same learning rate schedule.

Stage 3: The model of stage 2 becomes the teacher, the student is a binary network with binary weights and binary activations. We use logit matching and LSP but no weight decay. The hyperparameters we used are available in Section 5.2. We did not observe a significant difference in models initialized randomly or using the weights of the teacher.

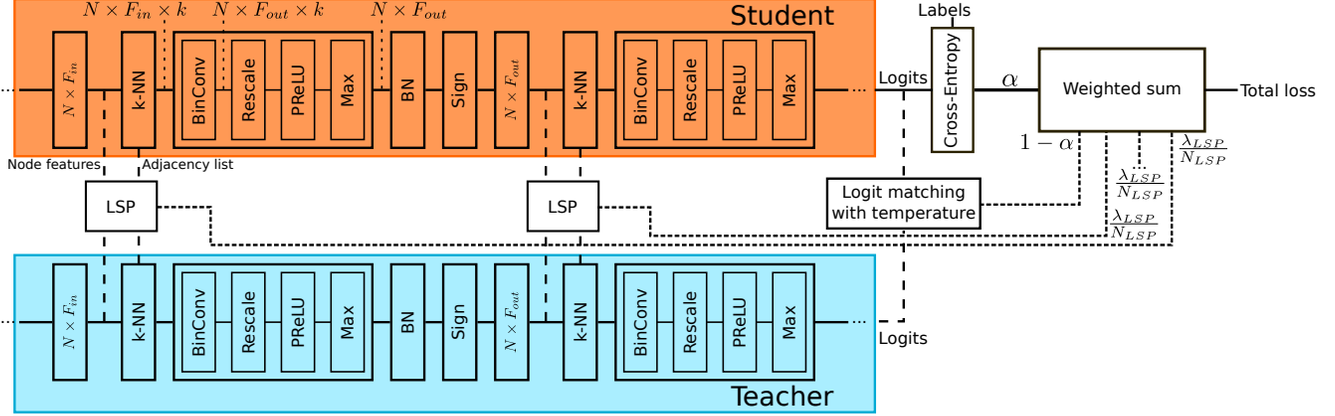


Figure 2. **Distillation with the "BF1" variant of XorEdgeConv**: the student model is more heavily quantized than the teacher. Knowledge transfer points equipped with LSP modules encourage similar dynamic graph feature distributions after each k -NN graph computation (except for the first, performed on the fixed 3D coordinates). Logit matching is used to further inform the training of the student.

Batch Normalization We investigate the importance of the order of the dot product and batch normalization operations for discretizing dot product operations within graph convolution operators. However, our base approach is to follow the XNOR-Net block structure [47] with learnable rescaling (*i.e.* XNOR-Net++ block). In particular, all fully-connected layers of MLPs that follow graph feature extraction layers are binarized using the XNOR-Net++ block.

4. Models

We choose the Dynamic Graph CNN model, built around the EdgeConv operator of Eq. 3 as our main case study. DGCNN has several characteristics that make it an interesting candidate for binarization. First, the EdgeConv operator is widely applicable to graphs and point clouds. Second, the operator relies on both node features and edge messages, contrary to other operators previously studied in GNN binarization such as GCN. Third, the time complexity of DGCNN is strongly impacted by the k -NN search in feature space. k -NN search can be made extremely efficient in Hamming space, and fast algorithms could theoretically be implemented for the construction of the dynamic graph at inference, provided that the graph features used in the search are binary, which requires a different binarization strategy than merely approximating the dense layer in EdgeConv.

For completeness, we also derive a binary SAGE operator.

4.1. Direct binarization

Our first approach binarizes the network weights and the graph features at the input of the graph convolution layers, but keeps the output real-valued. The network, therefore, produces real-valued node features. We replace the EdgeConv operator by a block similar to XNOR-Net++, using learnable rescaling and batch normalization pre-quantization.

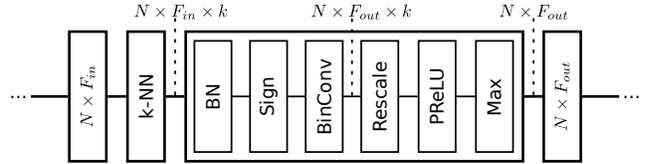


Figure 3. **The BinEdgeConv operator** ("RF" model in the experiments) can be deployed as a drop-in replacement for EdgeConv and follows the XNOR-Net++ approach to binarization.

We define the BinEdgeConv operator as:

$$\mathbf{e}_{ij}^{(l)} = \sigma \left(\text{sign}(\Theta^{(l)}) \otimes \text{sign} \left(\text{BN} \left(\tilde{\mathbf{X}}^{(l-1)} \right) \right) \odot \Gamma^{(l)} \right) \quad (12)$$

$$\mathbf{x}_i^{(l)} = \max_{j \in \mathcal{N}(i)} \mathbf{e}_{ij}^{(l)} \quad (13)$$

with σ the PReLU activation, and $\Gamma^{(l)}$ a real rescaling tensor. BinEdgeConv is visualized in Figure 3.

We use the same structure to approximate the MLP classifier. Similarly, we binarize Eq. 5 to get:

$$\mathbf{h}^{(l)} = \text{sign} \left(\text{BN} \left(\left[\mathbf{x}_i^{(l-1)} \parallel \text{Aggr}_{j \in \mathcal{N}(i)} \mathbf{x}_j^{(l-1)} \right] \right) \right) \quad (14)$$

$$\mathbf{x}_i^{(l)} = \text{Norm} \left(\sigma \left(\left(\text{sign}(\mathbf{W}^{(l)}) \otimes \mathbf{h}^{(l)} \right) \odot \Gamma^{(l)} \right) \right). \quad (15)$$

with σ the PReLU activation and $\Gamma^{(l)}$ following Eq. 10.

4.2. Dynamic Graph in Hamming Space

As mentioned, one advantage of binary node features is to enable fast computation of the k -Nearest Neighbours graph at inference time by replacing the ℓ_2 norm with the Hamming distance. We detail our approach to enable quantization-aware training with k -NN search on binary vectors.

Edge feature The central learnable operation of EdgeConv is $\Theta [\mathbf{x}_i \parallel \mathbf{x}_j - \mathbf{x}_i]$ as per Eq. 3, where the edge feature is $\mathbf{x}_j - \mathbf{x}_i$. Assuming binary node features, the standard subtraction operation becomes meaningless. Formally, for $\mathbf{x}_1, \mathbf{x}_2 \in \mathbb{R}^n$ with \mathbb{R}^n the n -dimensional Euclidean vector space over the field of real numbers,

$$\mathbf{x}_1 - \mathbf{x}_2 := \mathbf{x}_1 + (-\mathbf{x}_2) \quad (16)$$

by definition, with $(-\mathbf{x}_2)$ the additive inverse of \mathbf{x}_2 . Seeing binary vectors as elements of vector spaces over the finite field \mathbb{F}_2 , we can adapt Eq. 16 with the operations of Boolean algebra. The addition therefore becomes the Boolean exclusive or (XOR) \oplus , and the additive inverse of $(-x)_{\mathbb{F}_2}$ is x itself ($x \oplus x = 0$). With our choice of quantizer (Eq. 8), $\mathbf{x}_i, \mathbf{x}_j \in \{-1, 1\}^n$ and we observe that $\mathbf{x}_i \oplus \mathbf{x}_j = -\mathbf{x}_i \odot \mathbf{x}_j$. We therefore base our binary EdgeConv operator for binary node features, XorEdgeConv, on the following steps:

$$\mathbf{e}_{ij}^{(l)} = \sigma \left(\text{sign}(\Theta^{(l)}) \otimes \tilde{\mathbf{X}}_b^{(l-1)} \odot \Gamma^{(l)} \right) \quad (17)$$

$$\mathbf{x}_i^{(l)} = \text{sign} \left(\max_{j \in \mathcal{N}(i)} \mathbf{e}_{ij}^{(l)} \right) \quad (18)$$

with $\tilde{\mathbf{X}}_b^{(l-1)} = [\mathbf{x}_i^{(l-1)} \parallel -\mathbf{x}_j^{(l-1)} \odot \mathbf{x}_i^{(l-1)}]$, $\Theta^{(l)}$ a set of learnable real parameters and $\Gamma^{(l)}$ a real rescaling tensor. We further investigate the practical importance of the placement of the batch normalization operation, either before or after the aggregation function, by proposing two variants:

$$\mathbf{x}_i^{(l)} = \text{sign} \left(\text{BN} \left(\max_{j \in \mathcal{N}(i)} \mathbf{e}_{ij}^{(l)} \right) \right) \quad (19)$$

shown as part of Figure 2 and

$$\mathbf{x}_i^{(l)} = \text{sign} \left(\max_{j \in \mathcal{N}(i)} \text{BN} \left(\mathbf{e}_{ij}^{(l)} \right) \right) \quad (20)$$

drawn in Figure 1. Here, the main difference lies in the distribution of the features pre-quantization.

Nearest Neighbours Search The Hamming distance between two binary vectors \mathbf{x}, \mathbf{y} is $d_H(\mathbf{x}, \mathbf{y}) = \|\mathbf{x} \oplus \mathbf{y}\|_H$ where $\|\cdot\|_H$ is the number of non-zero bits, and can be efficiently implemented as `popcount(x xor y)`. We note that this relates our approach to previous work on efficient hashing [41, 43, 30] and metric learning [42], especially given the dynamic nature of the graph. Unfortunately, like the sign function, the Hamming distance has an ill-defined gradient, which hinders its use as-is for training. We therefore investigate two continuous relaxations. (1) we use the standard ℓ_2 norm for training, since all norms are equivalent in finite dimensions. (2) we observe that the matrix of pairwise Hamming distances between d -dimensional vectors \mathbf{x}_i

valued in $\{-1, 1\}$ can be computed in a single matrix-matrix product up to a factor 2 as (see Eq. 5 of [34]):

$$\mathbf{D} = -(\mathbf{X}\mathbf{X}^T - d\mathbf{I}_d) \quad (21)$$

with \mathbf{X} the matrix of the \mathbf{x}_i stacked row-wise, and \mathbf{I}_d the identity matrix. We investigate both options.

Local structure With binary node features, we now have to choose how to define the local structure similarity measure of Eq. 6. One option is to use the standard Gaussian RBF as in the real-valued case. Another option is to define the similarity in Hamming space, like for the k -NN search. We therefore investigate the following similarity metric:

$$\text{SIM}(\mathbf{x}_i, \mathbf{x}_j) = e^{-\|\mathbf{x}_i \oplus \mathbf{x}_j\|_H} \quad (22)$$

For vectors $\mathbf{x}, \mathbf{y} \in \{-1, 1\}^n$, we note that $\|\mathbf{x} \oplus \mathbf{y}\|_H = \frac{1}{2} \sum_{k=1}^n (-x_k y_k + 1)$.

5. Experimental Evaluation

We perform a thorough ablation study of our method on Dynamic Graph CNN. The model binarized according to the method of Section 4.1 and using the BinEdgeConv operator of Eq. 12 is referred to as **RF** for "Real graph Features". The model binarized according to Section 4.2 and using the XorEdgeConv operator is referred to as **BF1**, if following Eq. 19, or **BF2**, if following Eq. 20. We evaluate DGCNN on the ModelNet40 classification benchmark, as in [54]. We implement Γ as per Eq. 11 for the **RF** model and Eq. 10 for the **BF** models. In the next paragraphs, the numbers in parentheses refer to the corresponding lines of Table 1.

Balance functions (16-18, 23-24) Recent work [46] has uncovered possible limitations in binary graph and point cloud learning models when quantizing the output of the max-pooling aggregation of batch-normalized high-dimensional features. Similarly, the authors of [52] claim that a balance function is necessary to avoid large values in the outputs of the dot product operations when most pre-quantization inputs are positive.

We evaluate two strategies for re-centering the input of `sign`, namely mean-centering, and median-centering (thus ensuring a perfectly balanced distribution of positive and negative values pre-quantization). We evaluate these techniques for the max aggregation of edge messages ("edge balance", e.g. between the framed block and the `sign` operation in Figure 1) and for the max and average pooling operations before the MLP classifier ("global balance").

We can see in Table 1 that in all cases, the addition of balance functions actually lowered the performance of the models. This suggests that using batch normalization prior to quantization, as is common in the binary CNN literature, is sufficient at the message aggregation level and for producing graph embedding vectors.

#	Model	Distillation	Stage	k -NN	LSP	λ_{LSP}	Activation	Global balance	Edge balance	Acc (%)
1	BF1	None	-	H	-	-	PReLU	-	-	61.26
2	BF1	Direct	-	H	-	-	PReLU	-	-	62.84
3	BF1	Cascade	3	H	-	-	PReLU	-	-	81.04
4	BF1	Cascade	3	H	-	-	None	-	-	74.80
5	BF1	Cascade	2	H	ℓ_2	100	PReLU	-	-	90.32
6	BF1	Cascade	3	H	ℓ_2	100	PReLU	-	-	81.32
7	BF1	Cascade	3	H	ℓ_2	100	ReLU	-	-	76.46
8	BF1	None	-	ℓ_2	-	-	PReLU	-	-	62.97
9	BF1	Direct	-	ℓ_2	-	-	PReLU	-	-	61.95
10	BF1	Cascade	3	ℓ_2	-	-	PReLU	-	-	81.00
11	BF1	Cascade	2	ℓ_2	ℓ_2	100	PReLU	-	-	91.00
12	BF1	Cascade	3	ℓ_2	ℓ_2	100	PReLU	-	-	81.00
13	BF2	None	-	H	-	-	PReLU	-	-	59.72
14	BF2	Direct	-	H	-	-	PReLU	-	-	59.00
15	BF2	Cascade	3	H	-	-	PReLU	-	-	79.58
16	BF2	Cascade	3	H	-	-	PReLU	-	Mean	51.05
17	BF2	Cascade	3	H	-	-	PReLU	-	Median	75.93
18	BF2	Cascade	3	H	-	-	None	-	Median	71.96
19	BF2	Cascade	2	H	ℓ_2	100	PReLU	-	-	91.57
20	BF2	Cascade	3	H	ℓ_2	100	PReLU	-	-	81.08
21	BF2	Cascade	3	H	ℓ_2	100	None	-	-	76.09
22	BF2	Cascade	3	H	ℓ_2	100	ReLU	-	-	76.22
23	BF2	Cascade	3	H	ℓ_2	100	PReLU	Mean	-	67.87
24	BF2	Cascade	3	H	ℓ_2	100	PReLU	Median	-	60.82
25	BF2	None	-	ℓ_2	-	-	PReLU	-	-	57.90
26	BF2	Direct	-	ℓ_2	-	-	PReLU	-	-	59.12
27	BF2	Cascade	3	ℓ_2	-	-	PReLU	-	-	80.11
28	BF2	Cascade	2	ℓ_2	ℓ_2	100	PReLU	-	-	91.53
29	BF2	Cascade	3	ℓ_2	ℓ_2	100	PReLU	-	-	81.52
30	RF	None	-	ℓ_2	-	-	PReLU	-	-	79.30
31	RF	Direct	-	ℓ_2	-	-	PReLU	-	-	72.69
32	RF	Cascade	3	ℓ_2	-	-	PReLU	-	-	91.05
33	RF	Cascade	3	ℓ_2	ℓ_2	100	PReLU	-	-	90.52
34	RF	Cascade	3	ℓ_2	ℓ_2	100	None	-	-	89.71
35	RF	Cascade	3	ℓ_2	ℓ_2	100	ReLU	-	-	89.59
36	Baseline	-	-	-	-	-	ReLU	-	-	92.89

Table 1. Different variants and ablations of our binarized DGCNN models on the ModelNet40 dataset.

Non-linear activation (3-4,6-7,17-18,20-22,33-35) Since the sign operation can be seen as acting as an activation applied on the output and to the weights of the XorEdgeConv operator, we compare the models with binary node features with PReLU, ReLU, or no additional activation in Table 1. We can see the PReLU non-linearity offers significant improvements over the models trained with ReLU or without non-linearity in the edge messages, at the cost of a single additional fp32 parameter - the largest improvement being observed for the models that apply either median-centering or batch normalization before the quantization operation.

Binary node features and k -NN We now study the final performance of our models depending on whether we use BinEdgeConv (real node features) or XorEdgeConv. Looking at the final models (stage 3) in Table 1, the model with

real-valued node features that performs k -NN search with the ℓ_2 norm (32) performs comparably with the full floating-point model (36). On the other hand, we saw a greater reduction in accuracy with the binary node features for the full binary models (6,12,20,29), and comparable accuracy whether we use the ℓ_2 norm (12,29) or the relaxed Hamming distance (6,20). However, as reported in Table 1, using real weights (stage 2) with binary node features and k -NN search performed in Hamming space (5,11,19,28) matched the performance of the original floating point model (36).

We found stage 3 to be crucial to the final model’s performance, and sensitive to the choice of learning rate and learning rate schedule. This suggests that, although more research and parameter tuning is necessary to maximize the performance of the full binary networks in Hamming space, dynamic graph networks that learn binary codes and con-

struct the dynamic graph in Hamming space can be trained with minimal reduction in performance.

Impact of LSP The node features of the teacher and of the students are always real-valued at stage 1. Stage 2 was carried out using either the Gaussian RBF similarity or Eq. 22 for the student (which may have binary node features) and the Gaussian RBF for the teacher. Stage 3 uses either similarity measure for both the teacher and student. We also report the results of distilling the baseline DGCNN (full floating-point) model into a BF1 or BF2 full-binary model using the similarity in Hamming space for the student.

We saw inconsistent improvements when using LSP with the Gaussian RBF (ℓ_2), as seen in Table 1 (3,6,10,12,15,20,27,29,32,33). This suggest the usefulness of the additional structure preserving knowledge is situational, as it can both increase (3,4,15,20,27,29) or decrease model performance (32,33). Contrary to the models trained using k -NN search performed in Hamming space, the models trained with distillation using Eq. 22 did not match the performance of their Gaussian ℓ_2 counterparts, as shown in Table 2, which we conjecture to be due to poor gradients.

Model	Stage	KNN	LSP	λ_{LSP}	Acc. (%)
BF1	2	H	H	100	38.21
BF1	2	ℓ_2	H	100	38.94
BF2	2	H	H	100	63.25
BF2	2	ℓ_2	H	100	64.71
BF1	3	H	H	100	16.29
BF1	3	ℓ_2	H	100	20.34
BF2	3	ℓ_2	H	100	9.40
BF2	3	H	H	100	11.47
BF1	Direct	ℓ_2	H	100	23.34
BF2	Direct	H	H	100	30.23
BF2	Direct	ℓ_2	H	100	32.17
BF1	Direct	H	H	100	36.47

Table 2. Performance of models trained with LSP using the Hamming-based similarity of Eq. 22 (H) at different stages and for direct distillation. Compared to the models trained using the Gaussian RBF (ℓ_2) similarity, low performance was observed.

Cascaded distillation (1-3,13-15,25-27,30-32) Table 1 compares distilling the baseline network directly into a binary network, training from scratch, and the three-stage distillation. We observed consistently higher performance with the progressive distillation, confirming its effectiveness.

Large-scale inductive learning with GraphSAGE We benchmark our binarized GraphSAGE on the OGB-Products and OGB-Proteins node property prediction datasets [23], which are recent and challenging (2,449,029 nodes, 61,859,140 edges for OGB-Product) benchmarks with standardized evaluation procedures, compared to the more com-

monly used ones, such as Cora [39] (2708 nodes, 5429 edges) used in [52] or Reddit [21] (232,965 nodes, 114,615,892 edges) used in [53]. Notably, the Proteins dataset is challenging due to the high average node degree and small graph diameter, which may exacerbate limitations of GNNs [1].

Model	OGBN-Products		OGBN-Proteins	
	Mean acc.	Std. acc.	Mean acc.	Std. acc.
SAGE fp32	0.7862	0.0043	0.7734	0.0041
SAGE bin f.s.	0.7300	0.0156	0.7497	0.0047
SAGE bin l.m.	0.7260	0.0153	-	-
GCN fp32	0.7564	0.0021	0.7254	0.0044

Table 3. Final test accuracy on the OGB-Products and OGB-Proteins node property prediction benchmarks, averaged over 10 runs. "f.s.": from scratch. "l.m.": logit matching.

We implemented BinSAGE according to Eq. 15 - details of the architecture can be found in Section 5.2. For OGB-Products, we use logit matching only for distillation and no PReLU activation. For OGB-Proteins, we use PReLU activations and no distillation, as the task is multi-label classification, and the very large number of edges made using LSP impractical. We use channel-wise rescaling only for both to maximize scalability. On OGB-Products, we did not observe a statistically significant difference between training the model from scratch and three-stage distillation with logit matching: in both cases, the full binary model came within 5-6% of the full-precision model. On OGB-Proteins, the simple binary network trained from scratch is within 3% of the accuracy of the full-precision network and outperforms the full-precision GCN. This suggests other strategies to improve model scalability, in this case sampling, can be successfully combined with our binarisation method.

5.1. Speed on embedded hardware

In order to measure the speed improvements yielded by our binary conversion scheme, we benchmark it on a Raspberry Pi 4B board with 4GB of RAM and a Broadcom BCM2711 Quad core Cortex-A72 (ARM v8) 64-bit SoC clocked at 1.5GHz, running Manjaro 64-bit. The Pi is a popular, cheap, and readily available ARM-based platform, and is thus a good fit for our experiments.

We benchmark four DGCNN models, in order to measure the speedup for each subsequent optimization. The specifics of each model are given in Table 4. The input size is set to 1024 points with a batch size of 8, 40 output classes, and 20 nearest neighbors. We convert our models to Tensorflow Lite using LARQ [16], an open-source library for binarized neural networks, and benchmark them using the LARQ Compute Engine (LCE) tool. Once converted, the smallest model's file size is only 341KB down from 7.2MB, for a 20x reduction.

Figure 4 shows the benchmark results. Our optimized binary model halves the run-time, thus achieving a substantial speedup. Peak memory usage is also significantly reduced,

Model	Binary Weights	Binary Features	Hamming Distance
DGCNN			
BDGCNN RF	✓		
BDGCNN BF	✓	✓	
BDGCNN BF H	✓	✓	✓

Table 4. Features of benchmarked models. Hamm Dist = Pairwise Hamming distance instead of ℓ_2 , implemented in ARM NEON operations on bit-packed features (simulated).

from 575MB to 346MB. It is to be noted that DGCNN is a complex model with costly operations, such as concatenation and top- k , that are not made faster by binarization (denoted as "incompressible ops" in Figure 4). We provide a profiling of the models in Appendix B of the Supplementary Material.

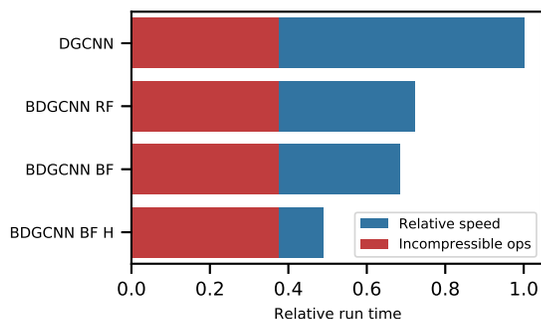


Figure 4. Relative run time on a Raspberry Pi 4B compared to the base DGCNN model. A 2x speedup is achieved by our final optimized model. Run times computed with the LCE benchmark tool over 50 runs.

Unfortunately, we did not have an optimized version of the Hamming distance in LCE at the time of writing. Thus, the final result is simulated by profiling the run-time of an un-optimized implementation, and estimating the savings we would get with ARM NEON instructions. It is theoretically possible to treat 64 features at a time, and achieve a 64x speedup (or higher by grouping loads and writes with `vlld4`). We use 32x as a *conservative estimate* since we couldn't account for LCE's bit-packed conversion.

5.2. Implementation details

For DGCNN, we follow the architecture of [55]. For GraphSAGE, we use the baseline architecture of the OGB benchmarks [23]; that is, three layers with 256 hidden features and mean aggregation. We use three knowledge transfer points for LSP on DGCNN, one after each EdgeConv layer except for the first layer (the k -NN and graph features are computed on the 3D coordinates of the point clouds and do not change). All binary models use binary inputs for the convolution and dense layers. For DGCNN, the final layer of the MLP classifier is kept real-weighted, as is customary in the binary neural network literature due to the small number of parameters, but the input features are binarized. For GraphSAGE, all three layers have binary weights.

Our models are implemented in Pytorch [45]. We use the implementation of DGCNN by the authors as a starting

point, and Pytorch Geometric [13] for GraphSAGE and OGB [23]. We use the Adam optimizer [28]. We train the DGCNN models for 250 epochs for stage 1 and 350 epochs for stages 2 and 3, on 4 Nvidia 2080 Ti GPUs. The initial learning rate of stage 1 is set to $1e-3$ and for stage 2 to $2.5e-4$, with learning rate decay of 0.5 at 50% and 75% of the total number of epochs. For stage 3, we set the learning rate to $1e-3$ and decay by a factor of 0.5 every 50 epochs. We trained GraphSAGE according to the OGB benchmark methodology, using the provided training, validation, and test sets. We trained all models for 20 epochs and averaged the performance over 10 runs. For GraphSAGE, we used ℓ_2 regularization on the learnable scaling factors only, with a weight $\lambda = 1e-4$. For logit matching, we set $T = 3$ and $\alpha = 1e-1$. For LSP, we set $\lambda_{LSP} = 1e2$.

6. Conclusion

In this work, we introduce a binarization scheme for GNNs based on the XNOR-Net++ methodology and knowledge distillation. We study the impact of various strategies and design decisions on the final performance of binarized graph neural networks, and show that our approach allows us to closely match or equal the performance of floating-point models on difficult benchmarks, with significant reductions in memory consumption and inference time. We further demonstrate that dynamic graph neural networks can be trained to high accuracy with binary node features, enabling fast construction of the dynamic graph at inference time through efficient Hamming-based algorithms, and further relating dynamic graph models to metric learning and compact hashing. Our DGCNN in Hamming space nearly equals the performance of the full floating point model when trained with floating point weights, and offers competitive accuracy with large speed and memory savings when trained with binary weights. We believe higher performance can already be obtained with this model by adjusting the learning rate schedule in the final distillation stage. Future work will investigate further improving the accuracy of the models, theoretical properties on binary graph convolutions, and inference with fast k -NN search in Hamming space.

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