Federated Graph Classification over Non-IID Graphs

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Abstract

Federated learning has emerged as an important paradigm for training machine learning models in different domains. For graph-level tasks such as graph classification, graphs can also be regarded as a special type of data samples, which can be collected and stored in separate local systems. Similar to other domains, multiple local systems, each holding a small set of graphs, may benefit from collaboratively training a powerful graph mining model, such as the popular graph neural networks (GNNs). To provide more motivation towards such endeavors, we analyze real-world graphs from different domains to confirm that they indeed share certain graph properties that are statistically significant compared with random graphs. However, we also find that different sets of graphs, even from the same domain or same dataset, are non-IID regarding both graph structures and node features. To handle this, we propose a graph clustered federated learning (GCFL) framework that dynamically finds clusters of local systems based on the gradients of GNNs, and theoretically justify that such clusters can reduce the structure and feature heterogeneity among graphs owned by the local systems. Moreover, we observe the gradients of GNNs to be rather fluctuating in GCFL which impedes high-quality clustering, and design a gradient sequence-based clustering mechanism based on dynamic time warping (GCFL+). Extensive experimental results and in-depth analysis demonstrate the effectiveness of our proposed frameworks.

1 Introduction

Federated learning (FL) as a distributed learning paradigm that trains centralized models on decentralized data has attracted much attention recently [28, 53, 25, 18, 17]. FL allows local systems to benefit from each other while keeping their own data private. Especially, for local systems with scarce training data or lack of diverse distributions, FL provides them with the potentiality to leverage the power of data from others, in order to facilitate the performance on their own local tasks. One important problem FL concerns is data distribution heterogeneity, since the decentralized data, collected by different institutes using different methods and aiming at different tasks, are highly likely to follow non-identical distributions. Prior works approach this problem from different aspects, including optimization process [25, 18], personalized FL [13, 6, 8], clustered FL [9, 15, 2], etc.

As more advanced techniques are developed for learning with graph data, using graphs to model and solve real-world problems becomes more popular. One important scenario of graph learning is graph classification, where models such as graph kernels [44, 34, 36, 45] and graph neural networks [21, 43, 49, 46, 47, 48] are used to predict graph-level labels based on the features and structures of graphs. One real scenario of graph classification is molecular property prediction, which is an important task in cheminformatics and AI medicine. In the area of bioinformatics, graph classification can be used to learn the representation of proteins and classify them into enzymes or non-enzymes. For collaboration networks, sub-networks can be classified regarding the information of research areas, topics, genre, etc. More applicable scenarios include geographic networks, temporal networks, etc.

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Since the key idea of FL is the sharing of underlying common information, as [23] discusses that real-world graphs preserve many common properties, we become curious about the question, whether real-world graphs from heterogeneous sources (e.g., different datasets or even divergent domains) can provide useful common information among each other? To understand this question, we first conduct preliminary data analysis to explore real-world graph properties, and try to find clues about common patterns shared among graphs across datasets. As shown in Table 1, we analyze four typical datasets from different domains, i.e., PTC_MR (molecular structures), ENZYMES (protein structures), IMDB-BINARY (social communities), and MSRC_21 (superpixel networks). We find them to indeed share certain properties that are statistically significant compared to random graphs with the same numbers of nodes and links (generated with the Erdős–Rényi model [7, 10]). Such observations confirm the claim about common patterns underlying real-world graphs, which can largely influence the graph mining models and motivates us to consider the FL of graph classification across datasets and even domains. More details and discussion about Table 1 can be found in Appendix A.

Table 1: Data analysis on important graph properties shared among real-world graphs across different domains. For example, large Kurtosis values [32] indicate long-tail distribution of node degrees, which is observed in ENZYMES, IMDB-BINARY, and MSRC_21; similar average shortest path lengths are observed in PTC_MR, ENZYMES, and MSRC_21, although their actual graph sizes are rather different; large CC are observed in ENZYMES, IMDB-BINARY, and MSRC_21 and large LC are observed in almost all graphs.

Property	kurtosis of degree distribution			avg. shortest path length			largest component size (LC, %)			clustering coefficient (CC)		
	real	random	p-value	real	random	p-value	real	random	p-value	real	random	p-value
PTC_MR (molecules)	2.1535	2.4424	0.9999	3.36	2.42	~ 0	100	82.68	~ 0	0.0095	0.1201	~ 0
ENZYMES (proteins)	3.0106	2.8243	0.0027	4.44	2.56	~ 0	98.24	97.69	0.2054	0.4516	0.1425	~ 0
IMDB-BINARY (social)	8.9262	2.2791	~ 0	1.48	1.54	~ 0	100	99.93	0.0023	0.9471	0.5187	~ 0
MSRC_21 (superpixel)	3.6959	2.9714	~ 0	4.09	2.81	~ 0	100	99.43	~ 0	0.5147	0.0655	~ 0

Although common patterns exist among graph datasets, we can still observe certain heterogeneity. In fact, the detailed graph structure distributions and node feature distributions can both diverge due to various reasons. To demonstrate this, we design and evaluate a structure heterogeneity measure and a feature heterogeneity measure in different scenarios (c.f. Section 4.1). We refer to the graphs possibly with significant heterogeneity in our cross-dataset FL setting as non-IID graphs, which concerns both structure non-IID and feature non-IID, where naïve FL algorithms like FedAvg [28] can fail and even backfire (c.f. Section 6.2). Moreover, as the heterogeneity varies from case to case, a dynamic FL algorithm is needed to keep track of such heterogeneity of non-IID graphs while conducting collaborative model training.

Due to the observations that the graphs in one client can be similar to those in some clients but not the others, we get motivated by [2] and find it intuitive to consider a clustered FL framework, which assigns local clients to multiple clusters with less data heterogeneity. To this end, we propose a novel graph-level clustered FL framework (termed GCFL) through integrating the powerful graph neural networks (GNNs) such as GIN [43] into clustered FL, where the server can dynamically cluster the clients based on the gradients of GNN without additional prior knowledge, while collaboratively training multiple GNNs as necessary for homogeneous clusters of clients. We theoretically analyze that the model parameters of GNN indeed reflect the structures and features of graphs, and thus using the gradients of GNN for clustering in principle can yield clusters with reduced heterogeneity of both structures and features. In addition, we conduct empirical analysis to support the motivation of clustered FL across heterogeneous graph datasets in Appendix A.

Although GCFL can theoretically achieve homogeneous clusters, during its training, we observe that the gradients transmitted at each communication round fluctuate a lot (c.f. Section 5.1), which could be caused by the complicated interactions among clients regarding both structure and feature heterogeneity, making the local gradients towards divergent directions. In the vanilla GCFL framework, the server calculates a matrix for clustering only based on the last transmitted gradients, which ignores the client's multi-round behaviors. Therefore, we further propose an improved version of GCFL with gradient-series-based clustering (termed GCFL+).

We conduct extensive experiments with various settings to demonstrate the effectiveness of our frameworks. Moreover, we provide in-depth analysis on the capability of them on reducing both structure and feature heterogeneity of clients through clustering. Lastly, we analyze the convergence of our frameworks. The experimental results show surprisingly positive results brought by our novel setting of cross-dataset/cross-domain FL for graph classification, where our GCFL+ framework can effectively and consistently outperform other straightforward baselines.

2 Related works

Federated Learning Federated learning (FL) has gained increasing attention as a training paradigm under the setting where data are distributed at remote devices and models are collaboratively trained under the coordination of a central server. FedAvg was first proposed by [27] which illustrates the general setting of an FL framework. Since the original FedAvg relies on the optimization by SGD, data of non-IID distribution will not guarantee the stochastic gradients to be an unbiased estimation of the full gradients, thus hurting the convergence of FL. In fact, multiple experiments [53, 25, 18] have shown that the convergence will be slow and unstable, and the accuracy will degrade with FedAvg when data at each client are statistically heterogeneous (non-IID). [53, 15, 12] proposed different data sharing strategies to tackle the data heterogeneity problem by sharing the local device data or serverside proxy data, which still requires certain public common data, whereas other studies explored the convergence guarantee under the non-IID setting by assuming bounded gradients [39, 51] or additional noise [19]. There are also works seeking to reduce the variance of the clients [26, 18, 25]. Furthermore, multiple works have been proposed to explore the connection between model-agnostic meta-learning (MAML) and personalized federated learning [8, 4]. They aim to learn a generalizable global model and then fine-tune it on local clients, which may still fail when data on local clients are from divergent domains with high heterogeneity. Some personalized FL works [6, 24] studied the bi-level problem of optimization which decouples the local and global optimization, but having each client maintain its own model can lead to high communication cost among the clients and the server. While personalization in the FL setting can address the client heterogeneity problem to some extent, the clustered FL framework [33] can incorporate personalization at the group level to keep the benefits of personalized FL and reduce the communication cost simultaneously.

Federated Learning on Graphs Although FL has been intensively studied with Euclidean data such as images, there exist few studies about FL for graph data. [22] first introduced FL on graph data, by regarding each client as a node in a graph. [3] studied the cross-domain heterogeneity problem in FL by leveraging Graph Convolutional Networks (GCNs) to model the interaction between domains. [29] studied spatio-temporal data modeling in the FL setting by leveraging a Graph Neural Network (GNN) based model to capture the spatial relationship among clients. [5] proposed a generalized federated knowledge graph embedding framework that can be applied for multiple knowledge graph embedding algorithms. Moreover, there are several works exploring the GNNs under the FL setting: [16, 54, 40] focused on the privacy issue of federated GNNs; [37] incorporated model-agnostic metalearning (MAML) into graph FL, which handled non-IID graph data while also preserving the model generalizability; [52] studied the missing neighbor generation problem in the subgraph FL setting; [38] proposed a computationally efficient way of GCN architecture search with FL; [11] implemented an open FL benchmark system for GNNs. Most existing works consider node classification and link prediction on graphs, which cannot be trivially applied to our graph classification setting.

3 Preliminaries

3.1 Graph Neural Networks (GNNs)

[42] provides a taxonomy that categorizes Graph Neural Networks (GNNs) into recurrent GNNs, convolutional GNNs, graph autoencoders, and spatial-temporal GNNs. In general, given the structure and feature information of a graph G=(V,E,X), where V,E,X denote nodes, links and node features, GNNs target to learn the representations of graphs, such as a node embedding $h_v \in \mathbb{R}^{d_v}$, or a graph embedding $h_G \in \mathbb{R}^{d_G}$. A GNN typically consists of message propagation and neighborhood aggregation, in which each node iteratively gathers the information propagated by its neighbors, and aggregates them with its own information to update its representation. Generally, an L-layer GNN can be formulated as

$$h_v^{(l+1)} = \sigma(h_v^{(l)}, agg(\{h_u^{(l)}; u \in \mathcal{N}_v\})), \forall l \in [L],$$
(1)

where $h_v^{(l)}$ is the representation of node v at the l^{th} layer, and $h_v^{(0)} = x_v$ is the node feature. \mathcal{N}_v is neighbors of node v, $agg(\cdot)$ is an aggregation function that can vary for different GNN variants, and σ represents an activation function.

For a graph-level representation h_G , it can be pooled from the representations of all nodes, as

$$h_G = readout(\{h_v; v \in V\}), \tag{2}$$

where $readout(\cdot)$ can be implemented as mean pooling, sum pooling, etc, which essentially aggregates the embeddings of all nodes on the graph into a single embedding vector to achieve tasks like graph classification and regression.

3.2 The FedAvg algorithm

McMahan et al. [28] proposed an SGD-based aggregating algorithm, FedAvg, based on the fact that SGD is widely used and powerful for optimization. FedAvg is the first basic FL algorithm and is commonly used as the starting point for more advance FL framework design.

The key idea of FedAvg is to aggregate the updated model parameters transmitted from local clients and then re-distribute the averaged parameters back to each client. Specifically, given m clients in total, at each communication round t, the server first samples a partition of clients $\{\mathbb{S}_i\}^{(t)}$. For each client \mathbb{S}_i in $\{\mathbb{S}_i\}^{(t)}$, it trains the model downloaded from the server locally with its own data distribution \mathcal{D}_i for E_{local} epochs. The client \mathbb{S}_i then transmits its updated parameters $w_i^{(t)}$ to the server, and the server will aggregate these updates by

$$w^{(t+1)} = \sum_{i=1}^{m} \frac{|D_i|}{|D|} w_i^{(t)}, \tag{3}$$

where $|D_i|$ is the size of data samples in client \mathbb{S}_i and |D| is the total size of samples over all clients. After generating the aggregated parameters (the global model updates), the server broadcasts the new parameters $w^{(t+1)}$ to remote clients, and at the (t+1) round clients use $w^{(t+1)}$ to start their local training for another E_{local} epochs.

4 The GCFL framework

4.1 Non-IID structures and features across clients

From Table 1 we notice that real-world graphs tend to share certain general properties across different graphs, datasets and even domains, which motivates the graph-level FL framework. However, there still exist differences when the detailed graph structures and node features are being considered. In Table 2, we present the average pair-wise structure heterogeneity and feature heterogeneity among graphs in a single dataset, a single domain, and across different domains. Specifically, for structure heterogeneity, we use the Anonymous Walk Embeddings (AWEs) [14] to generate a representation for each graph, and compute the Jensen-Shannon distance between the AWEs of each pair of graphs; for feature heterogeneity, we calculate the empirical distribution of feature similarity between all pairs of linked nodes in each graph, and compute the Jensen-Shannon divergence between the feature similarity distributions of each pair of graphs.

As we can observe in Table 2, both graph structures and features demonstrate different levels of heterogeneity within a single dataset, a single domain, and across different domains. We refer to graphs with such structure and feature heterogeneity as non-IID graphs. Intuitively, directly applying naïve FL algorithms like FedAvg on clients with non-IID graphs can be ineffective and even backfiring. To be specific, structure heterogeneity makes it difficult for a model to capture the universally important graph structure patterns across different clients, whereas feature heterogeneity makes it hard for a model to learn the universally appropriate message propagation functions across different clients. How can we leverage the shared graph properties among clients while addressing the non-IID structures and features across clients?

4.2 Problem formulation

Motivated by our real graph data analysis in Tables 1 and 2, we propose a novel framework of Graph Clustered Federated Learning (GCFL). The main idea of GCFL is to jointly find clusters of clients with graphs of similar structures and features, and train the graph mining models with FedAvg among clients in the same clusters.

Specifically, we are inspired by the Clustered Federated Learning (CFL) framework on Euclidean data [33] and consider a clustered FL setting with one central server and a set of n local clients $\{S_1, S_2, \ldots, S_n\}$. Different from the traditional FL setting, the server can dynamically cluster the

Table 2: Summary of the average heterogeneity of features and structures for some datasets. In general, the structure heterogeneity increases from the settings of one dataset to across-dataset, and to across-domain. However, the feature heterogeneity is more case-by-case, and the high variances indicate that graphs could have large feature divergence even within the same dataset. Additionally, it is not necessarily true that one dataset itself should be more homogeneous (e.g., IMDB-BINARY).

dataset	IMDB-BINARY (social)	COX2 (molecules)	COX2 (molecules) PTC_MR (molecules)	COX2 (molecules) ENZYMES (proteins)	COX2 (molecules) IMDB-BINARY (social)
avg. struc. hetero. avg. feat. hetero.	$0.4406 (\pm 0.0397) 0.1785 (\pm 0.1226)$	$0.3246 (\pm 0.0145) 0.0427 (\pm 0.0314)$	$0.3689 (\pm 0.0540)$ $0.1837 (\pm 0.1065)$	$0.5082~(\pm 0.0399) 0.1912~(\pm 0.1000)$	$0.6079 (\pm 0.0331) 0.1642 (\pm 0.1006)$

clients into a set of clusters $\{\mathbb{C}_1,\mathbb{C}_2,\ldots\}$ and maintain m cluster-wise models. In our GCFL setting, each local client \mathbb{S}_i owns a set of graphs $\mathcal{G}_i=\{G_1,G_2,\ldots\}$, where each $G_j=(V_j,E_j,X_j,y_j)\in\mathcal{G}_i$ is a graph data sample with a set of nodes V_j , a set of edges E_j , node features X_j , and a graph class label y_j . The task on each local client \mathbb{S}_i is graph classification that predicts the class label $\hat{y}_j=h_k^*(G_j)$ for each graph $G_j\in\mathcal{G}_i$, where h_k^* is the collaboratively learned optimal graph mining model for cluster \mathbb{C}_k to which \mathbb{S}_i belongs. Our goal is to minimize the loss function $F(\Theta_k):=\mathbb{E}_{\mathbb{S}_i\in\mathbb{C}_k}[f(\theta_{k,i};\mathcal{G}_i)]$, for all clusters $\{\mathbb{C}_k\}$. The function $f(\theta_{k,i};\mathcal{G}_i)$ is a local loss function for client \mathbb{S}_i which belongs to cluster \mathbb{C}_k . In the meantime, we also aim to maintain a dynamic cluster assignment $\Gamma(\mathbb{S}_i)\to\{\mathbb{C}_k\}$ based on the FL process.

4.3 Technical design

GNNs are demonstrated to be powerful for learning graph representations and have been wildly used in graph mining. More importantly, the model parameters and their gradients of GNNs can reflect the graph structure and feature information (more details in Section 4.4). Thus, we use GNNs as the graph mining model in our GCFL framework.

Specifically, our GCFL framework can dynamically cluster clients by leveraging their transmitted gradients $\{\Delta\theta_i\}_{i=1}^n$, in order to maximize the collaboration among more homogeneous clients and eliminate the harm from heterogeneous clients. According to [33], if the data distribution of clients are highly heterogeneous, the general FL that trains the clients together cannot jointly optimize all their local loss functions. In this case, after some rounds of communication, the general FL will be close to the stationary point, and the norm of clients' transmitted gradients will not all tend towards zero. Therefore, clustering clients is needed as the general FL approaches to the stationary point. Here, we first introduce a hyper-parameter ε_1 as a criterion to decide whether to stop the general FL based on whether a stationary point is approached, that is,

$$\delta_{mean} = \| \sum_{i \in [n]} \frac{|\mathcal{G}_i|}{|\mathcal{G}|} \Delta \theta_i \| < \varepsilon_1.$$
 (4)

In the meantime, if there exist some clients still with large norms of transmitted gradients, it means that clients in the group are highly heterogeneous, and thus clustering is needed to eliminate the negative influence among them. We then introduce the second criterion with a hyper-parameter ε_2 to split the clusters when

$$\delta_{max} = \max(\|\Delta\theta_i\|) > \varepsilon_2 > 0. \tag{5}$$

The GCFL framework follows a top-down bi-partitioning mechanism. At each communication round t, the server receives m sets of gradients $\{\{\Delta\theta_{i_1}\}, \{\Delta\theta_{i_2}\}, \dots, \{\Delta\theta_{i_m}\}\}$ from clients in clusters $\{\mathbb{C}_1, \mathbb{C}_2, \dots, \mathbb{C}_m\}$. For a cluster \mathbb{C}_k , if δ^k_{mean} and δ^k_{max} satisfy the Eqs. 4 and 5, the server will calculate a cluster-wise cosine similarity matrix α_k , and its entries are used as weights for building a full-connected graph with nodes being all clients within the cluster. The Stoer–Wagner minimum cut algorithm [35] is then applied to the constructed graph, which bi-partitions the graph and divides the cluster $\mathbb{C}_k \to \{\mathbb{C}_{k1}, \mathbb{C}_{k2}\}$. The clustering mechanism based on Eqs. 4 and 5 can automatically and dynamically determine the number of clusters along the FL, while the two hyper-parameters ε_1 and ε_2 can be easily set through some simple experiments on the validation sets following [33].

For a client \mathbb{S}_i in cluster \mathbb{C}_k , it tries to find $\theta_{k,i}$ that is close to the real solution $\theta_{k,i}^* = \arg\min_{\theta_i \in \Theta_k} f(\theta_{k,i}; \mathcal{G}_i)$. At a communication round t, the client \mathbb{S}_k transmits its gradient to the server

$$\Delta \theta_{k,i}^t = \hat{\theta}_{k,i}^t - \theta_{k,i}^{t-1}. \tag{6}$$

Since the server maintains the cluster assignments, it can aggregate the gradients cluster-wise by

$$\theta_k^{t+1} = \theta_k^t + \sum_{i \in [n_k]} \Delta \theta_{k,i}^t. \tag{7}$$

4.4 Theoretical analysis

We investigate the problem of graph FL with multi-domain data distribution, and use the gradient-based FL paradigm [27] to facilitate the model training. We theoretically analyze that the gradient-based FL algorithm on GNNs can in principle reduce the structure and feature heterogeneity in clusters, along with the task difference between data from different domains. We study two general problems in order to prove that the gradients can reflect the feature, structure, and task information.

Definition 4.1 Let a function $f: \mathcal{X} \to \mathcal{Y}$ which maps from the metric space (\mathcal{X}, d) to (\mathcal{Y}, d') , the function f is considered to have δ distortion if $\forall u, v \in \mathcal{X}$, $\frac{1}{\delta}d(u, v) \leq d'(f(u), f(v)) \leq d(u, v)$.

Theorem 4.1 (Bourgain theorem [1]) Given an n-point metric space (\mathcal{X}, d) and an embedding function f as defined above, $\forall u, v \in \mathcal{X}$, there exist an embedding mapped from (\mathcal{X}, d) to \mathbb{R}^k with the distortion of the embedding being $O(\log n)$.

Problem 1. In GCFL which involves the communication of the gradients between graphs with heterogeneous structures distributed among different clients, the structure and feature difference can be captured by the GNN gradients.

For simplicity, we solve Problem 1 with the GNN of Simple Graph Convolutions (SGC) [41], through the following two propositions.

Proposition 4.1 Given a graph G with fixed structure represented by the normalized graph Laplacian $\mathcal{L} = \widetilde{D}^{-\frac{1}{2}}\widetilde{A}\widetilde{D}^{-\frac{1}{2}}$, feature represented with X, and an SGC $f(\mathcal{L},X) = softmax(\mathcal{L}^KX\Theta)$ with weights Θ trained on graph G. If we have another graph G' with different structure \mathcal{L}' , the weight difference $||\Theta' - \Theta||_2$ is bounded with the structure difference.

Proposition 4.2 Given a graph G with fixed structure represented by the normalized graph Laplacian $\mathcal{L} = \widetilde{D}^{-\frac{1}{2}}\widetilde{A}\widetilde{D}^{-\frac{1}{2}}$, feature represented with X, and an SGC $f(\mathcal{L},X) = softmax(\mathcal{L}^KX\Theta)$ with weights Θ trained on graph G. If we have another graph G' with different feature \mathcal{X}' , the weight difference $||\Theta' - \Theta||_2$ is bounded with the feature difference.

We prove proposition 4.1 and 4.2 in Appendix B. We use the *Bourgain theorem* to bound the difference between embeddings generated with different graph structures/features, and prove that the feature and structure information of a graph is incorporated into the model weights (gradients). By proving that the model weights (gradients) are bounded with the structure/feature difference, we show that the gradients will change with the structure and feature. This further justifies that our proposed gradient based clustering framework GCFL is able to capture the structure and feature information. In addition, we also study the following problem, which allows our GCFL framework to be further extended to cross-task graph-level federated learning in the future.

Problem 2. The communicated gradients in GCFL can also capture the task heterogeneity.

Proposition 4.3 Given a graph G with structure represented by the normalized graph Laplacian $\mathcal{L} = \widetilde{D}^{-\frac{1}{2}}\widetilde{A}\widetilde{D}^{-\frac{1}{2}}$, and feature represented with X, if trained with different tasks, we will get the Simple SGC with bounded weights.

The proof of proposition 4.3 can be found in Appendix B.

5 GCFL+: improved GCFL based on observation sequences of gradients

5.1 Fluctuation of gradient norms

When observing the norm of gradients for each communication round in GCFL, as shown in Figure 1, we notice that: 1) the norm of gradients continuously fluctuates; 2) different clients can have divergent

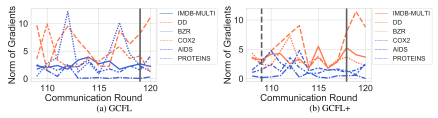


Figure 1: Norm of gradients versus communication round with six clients across datasets. Clients with datasets colored the same are split to the same cluster.

scales of gradient norms. The fluctuation of gradient norms and different scales indicate that the updating directions and distances of gradients for clients are diverse, which manifests the structure and feature heterogeneity in our setting again. In our vanilla GCFL framework, the server calculates a cosine similarity matrix based on the last transmitted gradients once the clustering criteria are satisfied. However, with the observation that the norm of gradients fluctuates along the communication round, albeit with the constraints of clustering criteria, GCFL clustering based on gradient-point could omit important client behaviors and be misled by noises. For example, in Figure 1 (a), GCFL performs clustering at round 119 based on the gradients at that round, which does not effectively find graphs with lower heterogeneity.

5.2 Technical design

Motivated by these observations, we propose an improved version of GCFL, named GCFL+, which conducts clustering by taking series of gradient norms into consideration. In the GCFL+ framework, the server maintains a multi-variant time-series matrix $Q \in \mathbb{R}^{\{n,d\}}$, where n is the number of clients and d is the length of a gradient series being tracked. At each communication round t, the server updates Q by adding in the norm of gradients $\|\Delta\theta_i^t\|$ to $Q(i,:) \in \mathbb{R}^d$ and remove the out-of-date one. GCFL+ uses the same clustering criteria as GCFL (Eqs. 4 and 5). If the clustering criteria are satisfied, the server will calculate a distance matrix β in which each cell is the pair-wise distance of two series of gradients. Here, we use a technique called dynamic time warping (DTW) [31] to measure the similarity between two data sequences. For a cluster \mathbb{C}_k , the server calculates its distance matrix as

$$\beta_k(p,q) = dist(Q(p,:), Q(q,:)), p, q \in idx(\{\mathbb{S}_i\}), \tag{8}$$

where $idx(\{S_i\})$ is the indices of all clients $\{S_i\}$ in cluster \mathbb{C}_k . With the distance matrix β , the server can perform bi-partitioning for clusters who meet the clustering criteria. As a result, in Figure 1 (b), GCFL+ performs clustering at round 118 based on the gradient sequence of length 10, which captures the longer-range behaviors of clients and effectively more homogeneous clusters.

6 Experiments

6.1 Experimental settings

Datasets We use a total of 13 graph classification datasets [30] from three domains including seven molecule datasets (MUTAG, BZR, COX2, DHFR, PTC_MR, AIDS, NCI1), three protein datasets (ENZYMES, DD, PROTEINS), and three social network datasets (COLLAB, IMDB-BINARY, IMDB-MULTI), each with a set of graphs. Node features are available in some datasets, and graph labels are either binary or multi-class. Details of the datasets are presented in Appendix C.

We design two settings that follow different data partitioning mechanisms, and the example real scenarios of the two settings can be found in Appendix A. The first setting (i.e., single-dataset) is to randomly distribute graphs from a single dataset to a number of clients, with each client holding a distinct set of about 100 graphs, among which 10% are held out for testing. In the second setting (i.e., multi-dataset), we use multiple datasets either from a single domain or multiple domains. Each client holds a graph dataset, among which 10% are held out for testing. In the first setting, we use NCI1, PROTEINS, and IMDB-BINARY from three domains and distribute them to 30, 10, 10 clients, respectively. In the second setting, we create three data groups including MOLECULES which consists of seven datasets from the molecule domain distributed into seven clients, BIOCHEM where we add three datasets from the protein domain into MOLECULES and distribute them into 10 clients, MIX where we add three datasets from the social domain into BIOCHEM and distribute them into 13 clients.

Baselines We use self-train² as the first baseline to test whether FL can bring improvements to each client through collaborative training. In self-train, each client firstly downloads the same randomly initialized model from the server and then trains locally without any communications. Then we implement two widely used FL baselines FedAvg [27] and FedProx [25], the latter of which can deal with data and system heterogeneity in non-graph FL. For the graph classification model, we use the same GIN [43] design, which represents the state-of-the-art GNN for graph-level tasks. We fix the GIN architecture and hyper-parameters through all baselines in order to control the experiments across different settings.

Parameter settings We use the three-layer GINs with hidden size of 64. We use a batch size of 128, and an Adam [20] optimizer with learning rate 0.001 and weight decay $5e^{-4}$. The μ for FedProx is set to 0.01. For all FL methods, the local epoch E is set to 1. The two important hyper-parameters ε_1 and ε_2 as clustering criteria vary in different groups of data, which are set through offline training for about 50 rounds following [33]. We run all experiments for five random repetitions on a server with 8 24GB NVIDIA TITAN RTX GPUs.

6.2 Experimental results

Federated graph classification within single datasets Conceptually, clients in this setting are more homogeneous. As can be seen from the results in Table 3, our framework can obviously improve the performance of graph classification over local clients. For the NCI1 dataset distributed on 30 clients, GCFL and GCFL+ achieve 13.27% and 14.75% performance gains over self-train on average, and GCFL and GCFL+ help 10-14 more clients than FedAvg and FedProx who fail to improve about half of clients. For the PROTEINS dataset on the total 10 clients, GCFL and GCFL+ achieve 7.29% and 7.81% average performance gains compared to self-train. For IMDB-BINARY on 10 clients, FedAvg and FedProx fail to help 5/10 and 4/10 clients respectively, while both GCFL and GCFL+ are able to improve all 10 clients. Overall, FedAvg can only help around half of the clients, which demonstrates that FedAvg can be ineffective even for decentralized graphs from a single dataset, because of the graph non-IIDness as shown in Table 2. In addition, in all three datasets, the minimum performance gain of clients over self-train using GCFL or GCFL+ is obviously larger than the minimum performance gain using FedAvg and FedProx. It indicates that even when some clients do not improve from self-train by using GCFL or GCFL+, they can achieve more comparable performance as self-train than using FedAvg and FedProx. These experimental results demonstrate that our frameworks are effective on the single-dataset multi-client FL setting.

Federated graph classification across multiple datasets — According to our data analysis in Tables 1 and 2, clients in such a setting are more heterogeneous. We conduct experiments with multiple datasets in two settings: single domain (using the data group MOLECULES), and across domains (using the data groups BIOCHEM and MIX). As can be seen from the results in Table 4, our frameworks GCFL and GCFL+ can effectively improve the performance of clients with distinct datasets. The results show 1.7% — 2.7% improvements of our frameworks compared to self-train. In all three data groups, our GCFL+ framework can improve twice as many as clients than FedAvg, and it achieves a ratio of 100% in MOLECULES to improve all clients' performance. The FedAvg failed to improve around 60% clients, which further demonstrates its ineffectiveness facing graph non-IIDness. Additionally, the GCFL+ framework also outperforms GCFL. In MIX, although GCFL can achieve the same ratio of improved clients as GCFL+, the GCFL+ framework has a much larger minimum gain of clients than GCFL. It indicates that by GCFL+ few clients that cannot benefit from others will not be degraded through the collaborating. These results indicate that graphs across datasets or even across domains are able to help each other through proper FL, which is a surprising and interesting start point for further study.

Effects of hyper-parameters ε_1 and ε_2 The hyper-parameter ε_1 is a stopping criterion for checking whether a general FL on the current set of clients is near the stationary point. Theoretically, ε_1 should be set as small as possible. The hyper-parameter ε_2 is more dependent on the number of clients and the heterogeneity among them. A smaller ε_2 will make the clients more likely to be clustered. When ε_1 and ε_2 are in the feasible ranges, their small variation can have little effect on the performance because the clustering results would largely remain the same. When ε_2 is set too large,

²We do not have a global model as baseline as datasets are from various domains and their tasks are divergent.

Table 3: Performance on the single-dataset-multi-client setting. We present the average accuracy and minimum gain over self-train on all clients, as well as the ratio of clients which get improved.

Dataset (# clients)	NCI1 (30)			PROTE	INS (10)		IMDB-BINARY (10)		
Accuracy	average	min gain	ratio	average	min gain	ratio	average	min gain	ratio
self-train	$0.6468(\pm0.053)$	_	_	$0.7213(\pm0.058)$	_	_	$0.7654(\pm0.057)$	_	_
FedAvg FedProx	$0.6474(\pm 0.076)$ $0.6437(\pm 0.072)$	-0.1333 -0.2400	14/30 16/30	$0.7490(\pm 0.034)$ $0.7556(\pm 0.036)$	-0.0615 -0.0923	6/10 7/10	$0.7596(\pm 0.049)$ $0.7746(\pm 0.048)$	-0.0800 -0.0600	5/10 6/10
GCFL GCFL+	0.7326(±0.052) 0.7422 (±0.053)	-0.0462 -0.1143	26/30 28 /30	0.7739(±0.043) 0.7776 (±0.037)	-0.0545 -0.0154	8/10 9 /10	0.8256(±0.059) 0.8299 (±0.052)	0.0182 0.0167	10 /10 10 /10

Table 4: Performance on the multi-dataset-multi-client setting. Metrics are the same as Table 3.

Dataset (# domains)	MOLECULES (1)			віосн	IEM (2)		MIX (3)		
Accuracy	average	min gain	ratio	average	min gain	ratio	average	min gain	ratio
self-train	$0.7543(\pm0.017)$	_	_	$0.7129(\pm0.016)$	_	_	$0.7001(\pm0.034)$	_	_
FedAvg FedProx	$0.7524(\pm 0.026) \\ 0.7668(\pm 0.032)$	-0.0132 -0.0054	3/7 5/7	$0.6944(\pm 0.027) \\ 0.7053(\pm 0.26)$	-0.1467 -0.1000	4/10 5/10	$0.6886(\pm 0.023) \\ 0.6897(\pm 0.026)$	-0.1233 -0.1367	5/13 5/13
GCFL GCFL+	$0.7661(\pm 0.016)$ $0.7745(\pm 0.030)$	0.0010 0.0010	7 /7 7 /7	0.7172(±0.019) 0.7312 (±0.031)	-0.0700 -0.0300	7/10 8 /10	0.7056(±0.019) 0.7121 (±0.021)	-0.1400 -0.0233	10 /13 10 /13

the performance will be similar as applying a basic FL algorithm directly (i.e. with a single cluster). When ε_2 is set too small, more clusters with smaller sizes or even single clients will be generated. We provide additional experimental results regarding the performance of GCFL and GCFL+ w.r.t. varying ε_1 and ε_2 in Figure 2. As shown in the subfigures, some points that represent varying ε_2 overlap for each ε_1 , which indicates that varying ε_2 in a certain range w.r.t. the fixed ε_1 leads to similar performance. Looking at a ε_2 , within a certain range of ε_1 , we can find the performance often fluctuating within a 0.01 variance. The results show that the performance of GCFL and GCFL+ are not very sensitive to the changes of ε_1 and ε_2 in reasonable ranges.

6.3 Structure and feature analysis in clusters

We conduct an in-depth analysis to explore the clustering results of GCFL and GCFL+. As can be seen in Figure 3, after being clustered by GCFL and GCFL+, the overall structure and feature heterogeneity of clients' graphs within clusters are reduced significantly compared to the original values, especially for the multiple dataset setting (Figure 3c and 3d). For the one dataset setting (Figure 3a and 3b), since features all fall in the same space, pairs of clients tend to have more homogeneous features. Therefore, the feature heterogeneity only gets reduced slightly after clustering. Unlike feature heterogeneity, the structure heterogeneity within clusters decreases significantly. In the setting of multiple datasets, as shown in Figure 3c and 3d, both structure and feature heterogeneity decrease significantly, which is intuitive since datasets across domains usually tend to have higher heterogeneity, as discussed in 4.1. We also look into the clusters and find that datasets from the same domains are more likely to be clustered together, while datasets from different domains also constantly get clustered together and benefit each other. For example, the clustering of GCFL+ corresponding to Figure 3d groups two social networks COLLAB and IMDB-BINARY together with PROTEINS and also several molecules datasets, and there is also a cluster of NCI1, DD, and IMDB-MULTI which are molecules, proteins and social networks, respectively. These analysis manifests that domains of datasets can verify the sanity of clusters to some extent, but one cannot solely rely on such prior knowledge to determine the optimal clusters, which demonstrates the necessity of our frameworks with the ability of performance-driven dynamic clustering along the process of FL.

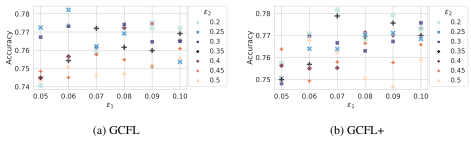


Figure 2: Performance of GCFL and GCFL+ on MOLECULES w.r.t varying ε_1 and ε_2 .

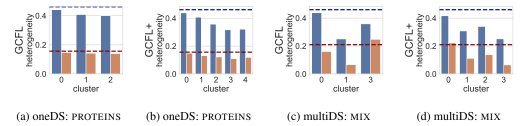


Figure 3: Structure (blue) and feature (red) heterogeneity within clusters found by GCFL and GCFL+. Dashed lines denote the heterogeneity over all clients before clustering.

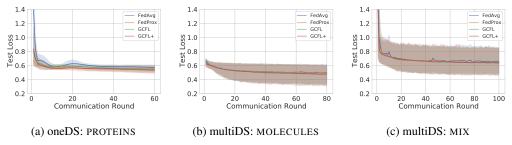


Figure 4: Average with standard deviation of the training curves of all clients.

6.4 Convergence analysis

We visualize the testing loss with respect to the communication round to show the convergence of GCFL and GCFL+ compared with the standard federated learning baselines. Figure 4 shows the training curves on two settings, which illustrates that GCFL and GCFL+ achieves similar convergence rate as FedProx, which is the state-of-the-art FL framework dealing with non-IID Euclidean data. We also notice that both GCFL, GCFL+ and FedProx can converge to a lower loss compared with FedAvg, which corroborates our consideration of the non-IID problem in our setting.

6.5 More results in Appendix

In Table 3 and 4, we averaged the accuracy across all clients for presentation simplicity. To understand the detailed performance by clients and clusters, we present different Violin plots in Appendix D. Besides, we also show more results regarding various settings (overlapping clients, real vs. synthetic node features, standardized gradient-sequence matrix in GCFL+, etc) in Appendix D.

7 Conclusion

In this work, we propose a novel setting of cross-dataset and cross-domain federated graph classification. The techniques (GCFL and GCFL+) we develop allow multiple data owners holding structure and feature non-IID graphs to collaboratively train powerful graph classification neural networks without the need of direct data sharing. As the first trial, we focus on the effectiveness of FL in this setting and have not carefully studied other issues such as data privacy, although it is intuitive to preserve the privacy of clients by introducing an encryption mechanism (e.g. applying orthonormal transformations), and to prevent from adversarial scenarios by clustering out the malicious clients. Due to its evident motivations and proofs on the effective FL in a new setting, we believe this work can serve as a stepping stone for many interesting future studies.

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A More Motivating Examples and Analysis

Additional explanations for Table 1 We provide two real examples of how graph properties can decide important graph patterns as the signals for graph classifications. The first example is about social networks. Among the graph properties, social networks (such as IMDB-BINARY shown in Table 1) have a significant high clustering coefficient (CC), which means they contain many triangles as a graph pattern. This may be an important signal for effective graph classification, which is consistent with commonly used social network classification methods such as motif-counting. On the other hand, if a set of random graphs does not possess high CC, the prominent graph patterns might be some subtrees or long edges, where the training of a graph classification model is not likely to facilitate the capturing of triangles, and thus not beneficial for the classification of social networks. The second example is about protein networks and superpixel networks (such as ENZYMES and MSRC_21 shown in Table 1). ENZYMES is a protein dataset, in which a graph is an enzyme with secondary structure elements as nodes and the links connect nodes if they are neighbors in space. MSRC 21 is a semantic image processing dataset, in which nodes represent superpixels and links are constructed if two superpixels are adjacent. Although the two datasets are from totally different domains, they are both formed by spatial structures, which is reflected in Table 1 where the two datasets have very close values regarding multiple properties (degree distribution, shortest path length, CC). For ENZYME, it is known that the tertiary structure of proteins is essential and necessary for their biological activities, and the tightly knit groups are important signals for classifying an enzyme into different catalyzed levels. For MSRC 21, the environment information is also essential for identifying the superpixels or describing objects in the images. Thus, there exists the case that two cross-domain datasets contain certain significant graph patterns that correspond to different dataset-specific meanings and tasks, but can be shared across datasets towards the training of more powerful graph models.

Additional analysis for clustered FL With analysis from Table 2, it shows significant statistical heterogeneity regarding features and structures in graphs. To motivate clustered FL, we conduct additional experiments regarding the connections between such statistical heterogeneity and the power of GNNs. Specifically, we use PROTEINS as one example dataset, and pair it with other datasets from the same and different domains. We focus on structure heterogeneity here and use one-hot degree features in order to avoid the effects of feature heterogeneity. We then train a GIN model with basic FedAvg on all pairs, and analyze the GIN performance versus the structural heterogeneity as defined in our paper on PROTEINS in Figure 5. As observed from Figure 5: (1) at first, the GNN on one graph dataset (PROTEINS) can benefit from the federated learning on another graph dataset, and that benefit becomes larger as the

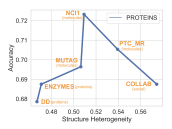


Figure 5: GIN Performance on PROTEINS (proteins) federated averagely trained with datasets from the same or different domains.

structure heterogeneity between two graphs becomes larger (DD, ENZYMES, MUTAG, NCI1), which clearly supports the benefit of cross-dataset/cross-domain graph federated learning. However, (2) as the heterogeneity becomes too large, the performance of GNN starts to degenerate (PTC_MR, COLLAB), which clearly supports our design of clustered federated learning. Our whole framework is built to achieve a good trade-off between (1) and (2), which we achieved to some extent, but can further improve on it in future studies.

Example real scenario for the single-dataset setting In this work, we use the public IMDB datasets to mimic the real scenario for social networks. For example, TikTok as an emerging video sharing platform has branches in different countries nowadays, among which direct data sharing is illegal. However, the users in different countries naturally reside in social networks with similar properties. It is then very viable to perform graph federated learning across branches in different countries towards the training of more powerful graph learning models for tasks such as group/community profiling.

Example real scenario for multi-dataset setting In this work, we synthesized the extreme setting where each client holds data from totally different domains for experimental purposes, to establish the benefit of cross-domain collaboration. A realistic scenario in a less extreme setting can be about various types of apps on mobile phones. For example, different types of iPhone users may leverage

healthcare apps and social media apps and opt in their data collection. As a consequence, the two apps will both hold some user data that may complement each other but cannot be directly shared across the departments. Since Apple owns both departments, it is then easier for them to collaborate through a federated learning framework.

B Missing Proofs in Section 4.4

B.1 Proof of Proposition 4.1

Assume the structure difference between graph G and G' is bounded with

$$||\mathcal{L}' - \mathcal{L}||_2^2 = ||E_{\mathcal{L}}||_2^2 \le \epsilon_{\mathcal{L}},\tag{9}$$

The difference between the original weights Θ and the weights trained on the new graph structure Θ' is represented as

$$||\Theta' - \Theta||_2^2 = ||(\mathcal{L}'X)^{-1}Y' - (\mathcal{L}X)^{-1}Y||_2^2$$

= $||X^{-1}(\mathcal{L}'^{-1}Y' - \mathcal{L}^{-1}Y)||_2^2$. (10)

Given that $||\mathcal{L}\cdot\mathcal{L}'||_2^2 = ||\mathcal{L}\cdot(\mathcal{L}+E_{\mathcal{L}})||_2^2 \geq ||\mathcal{L}E_{\mathcal{L}}||_2^2$. Let $||\mathcal{L}E_{\mathcal{L}}||_2^2 = \delta_{\mathcal{L}}$, then we can get $||\mathcal{L}'^{-1} - \mathcal{L}^{-1}||_2^2 = ||E_{\mathcal{L}^{-1}}|| \leq \frac{\epsilon_{\mathcal{L}}}{\delta_{\mathcal{L}}}$.

Given the Bourgain theorem [50], the difference between the embedding Y and Y' is bounded with

$$||\mathcal{Y}' - \mathcal{Y}||_2^2 = ||E_Y||_2^2 \le \epsilon_Y,$$
 (11)

The weight difference can then be bounded with

$$||\Theta' - \Theta||_{2}^{2} \leq ||X^{-1}||_{2}^{2}||(\mathcal{L}^{-1} + E_{\mathcal{L}^{-1}})(Y + E_{Y}) - \mathcal{L}^{-1}Y||_{2}^{2}$$

$$= ||X^{-1}||_{2}^{2}||\mathcal{L}^{-1}E_{Y} + E_{\mathcal{L}^{-1}}Y + E_{\mathcal{L}^{-1}}E_{Y}||_{2}^{2}$$

$$\leq ||X^{-1}||_{2}^{2} \left[\epsilon_{Y}||\mathcal{L}^{-1}||_{2}^{2} + \frac{\epsilon_{\mathcal{L}}}{\delta_{\mathcal{L}}}||Y||_{2}^{2} + \frac{\epsilon_{\mathcal{L}}\epsilon_{Y}}{\delta_{\mathcal{L}}}\right].$$
(12)

With the trained SGC, the feature and graph structure is fixed as X and \mathcal{L} . Thus the weight difference is bounded with X and \mathcal{L} .

B.2 Proof of Proposition 4.2

Assume the feature difference between graph G and G' is bounded with

$$||X' - X||_2^2 = ||E_X||_2^2 < \epsilon_X, \tag{13}$$

The difference between the original weights Θ and the weights trained on the new graph structure Θ' is represented as

$$||\Theta' - \Theta||_2^2 = ||(\mathcal{L}X')^{-1}Y' - (\mathcal{L}X)^{-1}Y||_2^2$$
(14)

Given that $||X \cdot X'||_2^2 = ||X \cdot (X + E_X)||_2^2 \ge ||XE_X||_2^2$. Let $||XE_X||_2^2 = \delta_X$, then we can get $||X'^{-1} - X^{-1}||_2^2 = ||E_{X^{-1}}|| \le \frac{\epsilon_X}{\delta_X}$.

Given the Bourgain theorem [50], the difference between the embedding Y and Y' is bounded with

$$||\mathcal{Y}' - \mathcal{Y}||_2^2 = ||E_Y||_2^2 \le \epsilon_Y,$$
 (15)

The weight difference can then be bounded with

$$||\Theta' - \Theta||_{2}^{2} = ||(X'^{-1}\mathcal{L}^{-1}Y' - X^{-1}\mathcal{L}^{-1}Y||_{2}^{2}$$

$$= ||(X'^{-1}\mathcal{L}^{-1}(Y + E_{Y}) - X^{-1}\mathcal{L}^{-1}Y||_{2}^{2}$$

$$= ||(X'^{-1}\mathcal{L}^{-1}Y + X'^{-1}\mathcal{L}^{-1}E_{Y} - X^{-1}\mathcal{L}^{-1}Y||_{2}^{2}$$

$$= ||(X'^{-1}\mathcal{L}^{-1}Y - X^{-1}\mathcal{L}^{-1}Y + X'^{-1}\mathcal{L}^{-1}E_{Y}||_{2}^{2}$$

$$= ||(X'^{-1} - X^{-1})\mathcal{L}^{-1}Y + X'^{-1}\mathcal{L}^{-1}E_{Y}||_{2}^{2}$$

$$= ||E_{X^{-1}}\mathcal{L}^{-1}Y + (X + E_{X})^{-1}\mathcal{L}^{-1}E_{Y}||_{2}^{2}$$

$$= ||E_{X^{-1}}\mathcal{L}^{-1}Y + (\mathcal{L}X + \mathcal{L}E_{X})^{-1}E_{Y}||_{2}^{2}$$

$$\leq \frac{\epsilon_{X}}{\delta_{Y}}||\mathcal{L}^{-1}Y||_{2}^{2} + \frac{\epsilon_{X}^{2}\epsilon_{Y}}{\delta_{Y}}||(\mathcal{L}X)^{-1}||_{2}^{2} + \epsilon_{X}\epsilon_{Y}||(\mathcal{L}X)^{-1}||_{2}^{4}$$

With the trained SGC, the feature and graph structure is fixed as X and \mathcal{L} . Thus the weight difference is bounded.

B.3 Proof of Proposition 4.3

The difference between the weights trained with different tasks can be written as

$$||\Theta_i - \Theta_j||_2^2 = ||(\mathcal{L}X)^{-1}(Y_i - Y_j)||_2^2.$$
(17)

With the Bourgain theorem, we have the transformed embedding bounded with

$$||Y_i - Y_j||_2^2 = ||E_Y||_2^2 \le \epsilon_Y. \tag{18}$$

By substituting $||Y_i - Y_j||_2^2 = ||E_Y||_2^2$, we have

$$||\Theta_i - \Theta_j||_2^2 = ||(\mathcal{L}X)^{-1}E_Y||_2^2$$

$$\leq \epsilon_Y ||(\mathcal{L}X)^{-1}||_2^2$$
(19)

C Dataset Details

We provide details of the datasets we use in Table 5.

Table 5: The statistics of datasets.

dataset			statistics			dataset	statistics					
	#graphs	avg. #nodes	avg. #edges	#classes	node features		#graphs	avg. #nodes	avg. #edges	#classes	node features	
MUTAG	188	17.93	19.79	2	original	ENZYMES	600	32.63	62.14	6	original	
BZR	405	35.75	38.36	2	original	DD	1178	284.32	715.66	2	original	
cox2	467	41.22	43.45	2	original	PROTEINS	1113	39.06	72.82	2	original	
DHFR	467	42.43	44.54	2	original	COLLAB	5000	74.49	2457.78	3	degree	
PTC_MR	344	14.29	14.69	2	original	IMDB-BINARY	1000	19.77	96.53	2	degree	
AIDS	2000	15.69	16.20	2	original	IMDB-MULTI	1500	13.00	65.94	3	degree	
NCI1	4110	29.87	32.30	2	original						-	

D More Detailed Experiment Results

Violin plots instead of tables Figures 6, 7, and 8 show the detailed experiment results regarding more various client settings including overlapped vs. non-overlapped data partitioning, original vs. synthetic node features, standardized vs. non-standardized multi-variant time-series matrix in GCFL+. Since we want to provide more detailed results by clients, we use violin plots to display the distributions of performance gains of all clients compared to self-train, instead of the average numbers in Tables 3 and 4. In Figure 6 and 7, each violin represents a distribution of all clients' performance gain using one algorithm, and in Figure 8 each violin represents a distribution of all clients' performance gain on one dataset or data group. In Figures 6, 7, and 8, the blue left sides of violins are corresponding to the results in the main tables 3 and 4.

Overlapping versus non-overlapping For distributing one dataset to multiple clients, we compare the two settings of allowing overlapping (same graphs appearing multiple clients) and not. As can be seen in Figure 6, our frameworks can also improve on overlapped clients.

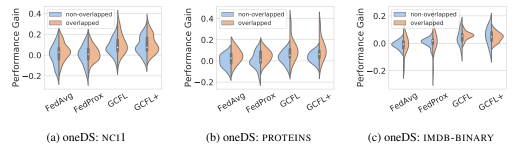


Figure 6: Distributions of performance gains of all clients with overlapped versus non-overlapped data partitioning.

Original node features versus one-hot degree features Apart from the original node features, we also use one-hot node degree features, in order to study the influence of node features. Figure (7a, 7b) and (7c, 7d) show the comparisons between original features and one-hot degree features on the single-dataset (oneDS) setting and the multi-dataset (multiDS) setting, respectively. Overall, our frameworks can consistently improve when using one-hot degree features. However, as in Figure 7c, the performance gains decreased when using only one-hot node degrees, which can be because of the decease of feature heterogeneity.

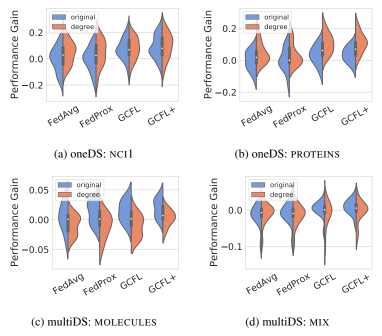


Figure 7: Distributions of performance gains of all clients using original node features versus one-hot degree features on the oneDS (top) and multiDS (bottom) settings.

Standardized versus non-standardized multi-variant time-series matrix For the GCFL+ framework, we compared the performance gains of clients using the standardized or non-standardized multi-variant time-series matrix $Q \in \mathbb{R}^{\{n,d\}}$. By standardization, only the trends of gradients' fluctuation are considered and the scales are ignored. The standardization step is performed before calculating the distance matrix β as

$$Q'(i,:) = Q(i,:)/std(Q(i,:)), i = 0, 1, \dots, n.$$
(20)

As shown in Figure 8, the average performance gains of standardization and non-standardization are similar.

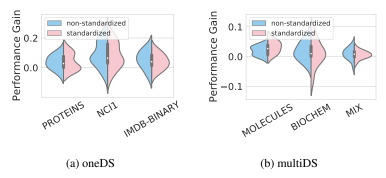


Figure 8: Distributions of performance gains of all clients using standardized gradient-sequence matrix versus non-standardized gradient-sequence matrix in GCFL+ on the oneDS and multiDS settings.