SpreadGNN: Serverless Multi-task Federated Learning for Graph Neural Networks

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Abstract

Graph Neural Networks (GNNs) are the first choice methods for graph machine learning problems thanks to their ability to learn state-of-the-art level representations from graph-structured data. However, centralizing a massive amount of real-world graph data for GNN training is prohibitive due to user-side privacy concerns, regulation restrictions, and commercial competition. Federated Learning is the de-facto standard for collaborative training of machine learning models over many distributed edge devices without the need for centralization. Nevertheless, training graph neural networks in a federated setting is vaguely defined and brings statistical and systems challenges. This work proposes SpreadGNN, a novel multi-task federated training framework capable of operating in the presence of partial labels and absence of a central server for the first time in the literature. SpreadGNN extends federated multi-task learning to realistic serverless settings for GNNs, and utilizes a novel optimization algorithm with a convergence guarantee, Decentralized Periodic Averaging SGD (DPA-SGD), to solve decentralized multi-task learning problems. We empirically demonstrate the efficacy of our framework on a variety of non-I.I.D. distributed graph-level molecular property prediction datasets with partial labels. Our results show that SpreadGNN outperforms GNN models trained over a central server-dependent federated learning system, even in constrained topologies. The source code is publicly available at https://github.com/FedML-AI/SpreadGNN.

1 Introduction

Graph Neural Networks (GNNs) [17] are expressive models that can distill structural knowledge into highly representative embeddings. While graphs are the representation of choice in domains such as social networks[3,31], knowledge graphs for recommendation systems [6], in this work we focus on molecular graphs that are the core of drug discovery, molecular property prediction [13, 23] and virtual screening [56, 64]. Molecular graphs differ from their more well known counterparts such as social network graphs. First, each molecule is a graph representation of the basic atoms and bonds that constitute the molecule and hence the size of the graph is small. Second, even though each graph may be small there are numerous molecules that are being developed continuously for varied use cases. Hence, what they lack in size they make up for it in structural heterogeneity. Third, molecules can be labeled along multiple orthogonal dimensions. Since each graph has multiple labels the learning itself can be characterized as multi-task learning. For instance, whether a molecule has potentially harmful interaction with a diabetics drug, or whether that molecule can turn toxic under certain conditions are distinct labels. Molecular property analysis and labeling requires wet-lab experiments, which is time-consuming and resource-costly. As a consequence, many entities may

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only have partially labeled molecules even if they know the graph structure. Finally, molecules are coveted inventions and hence entities often possess proprietary graph representation that cannot be shared with other institutions for competitive and regulatory reasons. But training collectively over a private set of molecular graphs can have immense societal benefits such as accelerated drug discovery.

Federated Learning (FL) is a distributed learning paradigm that addresses this data isolation problem via collaborative training. In this paradigm, training is an act of collaboration between multiple clients (such as research institutions) without requiring centralized local data while providing a certain degree of user-level privacy [42, 26]. However there are still challenges and shortcomings to training GNNs in a federated setting. As shown in [20], federated GNNs perform poorly in a non-iid setting. This setting (Figure 1) is the typical case in molecular graphs since each owner may have different molecules and even when they have the same molecular graph each owner may have an incomplete set of labels for each molecule. The left half of Figure 1 shows a simpler case where all the clients can communicate through a central server. But in practice the presence of a central server is not feasible when multiple competing entities may want to collaboratively learn. The challenges are further compounded by the lack of a central server as shown in the right half of the Figure 1. Thus, it remains an open problem to design a federated learning framework for molecular GNNs, for a realistic setting, in which clients only have partial labels and one in which there is no reliance on a central server. This is the problem we seek to address in this work.

We propose a multitask federated learning framework called SpreadGNN that operates in the presence of partial labels and absence of a central server as shown in Figure 1. We use the word task and class interchangeably, with each label being composed of multiple tasks. First, we present a multitask learning (MTL) formulation to learn from partial labels. Second, in our MTL formulation, we utilize decentralized periodic averaging stochastic gradient descent (DPA-SGD) to solve the serverless MTL optimization problem, and also provide a theoretical guarantee on the convergence properties for DPA-SGD, which further verifies the rationality of our design.

We evaluate SpreadGNN on graph level molecular property prediction and regression tasks. We synthesize non-I.I.D. and partially labeled datasets by using curated data from the MoleculeNet [69] machine learning benchmark. With extensive experiments and analysis, we find that SpreadGNN can achieve even better performance than FedAvg [44], not only when all clients can communicate with each other, but also when clients are constrained to communicate with a subset of other clients. We plan on publishing the source code of SpreadGNN as well as related datasets for future exploration.

2 SpreadGNN Framework

2.1 Preliminary: Federated Graph Neural Networks for Graph Level Learning

We seek to learn graph level representations in a federated learning setting over decentralized graph datasets located in edge servers which cannot be centralized for training due to privacy and regulation restrictions [20]. For instance, compounds in molecular trials [55] may not be shared across entities because of intellectual property or regulatory concerns. Under this setting, we assume that there are $K$ clients in the FL network, and the $k^{th}$ client has its own dataset
\[ D^{(k)} := \left\{ (G^{(k)}_i, y^{(k)}_i) \right\}^{N^{(k)}}_{i=1}, \text{ where } G^{(k)}_i = (Y^{(k)}_i, \mathbf{e}^{(k)}_i) \text{ is the } i^{th} \text{ graph sample in } D^{(k)} \text{ with node} \]

\& edge feature sets \[ X^{(k)} = \left\{ x^{(k)}_m \right\}_{m \in V^{(k)}}^{N^{(k)}} \text{ and } Z^{(k)} = \left\{ e^{(k)}_{m,n} \right\}_{m,n \in V^{(k)}}^{N^{(k)}} , \text{ where } y^{(k)}_i \text{ is the corresponding multi-class label of } G^{(k)}_i. \]

\[ N^{(k)} \text{ is the sample number in dataset } D^{(k)}, \text{ and } N = \sum_{k=1}^{K} N^{(k)}. \]

Each client owns a GNN with a readout, to learn graph-level representations. We will call this model of a GNN followed by a readout function, a **graph classifier**. Multiple clients are interested in collaborating to improve their GNN models without necessarily revealing their graph datasets. In this work, we build our theory upon the Message Passing Neural Network (MPNN) framework [14, 56] as most spatial GNN models [29, 65, 17] can be unified into this framework. The forward pass of an MPNN has two phases: a message-passing phase (Eq (1)) and an update phase (Eq (2)). For each client, we define the graph classifier with an L-layer GNN followed by a readout function as follows:

\[ m^{(k,\ell+1)}_i = \text{AGG} \left( \left\{ M^{(k,\ell+1)}_\psi \left( h^{(k,\ell)}_i, h^{(k,\ell)}_j, e_{i,j} \right) \mid j \in N_i \right\} \right), \ell = 0, \ldots, L - 1 \]  

(1)

\[ h^{(k,\ell+1)}_i = U^{(k,\ell+1)}_\psi \left( h^{(k,\ell)}_i, m^{(k,\ell+1)}_i \right), \ell = 0, \ldots, L - 1 \]  

(2)

\[ \hat{y}^{(k)}_i = R_{\Phi_{\text{pool}}, \Phi_{\text{task}}} \left( \left\{ h^{(k,L)}_j \mid j \in V^{(k)}_i \right\} \right) \]  

(3)

where \[ h^{(k,0)}_i = x^{(k)}_i \] is the \( i^{th} \) vertex’s hidden state with the edge and vertex data from its neighbors to generate a new message. \[ M^{(k,\ell+1)}_\psi \] is the message generation function which takes the hidden state of current node \( h_i \), the hidden state of the neighbor node \( h_j \) and the edge features \( e_{i,j} \) as inputs to gather and transform neighbors’ messages. In other words, \[ M_\psi \] combines a vertex’s hidden state with the edge and vertex data from its neighbors to generate a new message. \[ U^{(k,\ell+1)}_\psi \] is the state update function that updates the model using the aggregated feature \( m^{(k,\ell+1)}_i \) as in Eq. (2). After propagating through \( L \) GNN layers, the final module of the graph classifier is a readout function \[ R_{\Phi_{\text{pool}}, \Phi_{\text{task}}} \] which allows clients to predict a label for the graph, given node embeddings that are learned from Eq. (2). In general the readout is composed of two neural networks: the pooling function parameterized by \( \Phi_{\text{pool}} \) and a task classifier parameterized by \( \Phi_{\text{task}} \). The role of the pooling function is to learn a single graph embedding given node embedding from Eq (2). The task classifier then uses the graph level embedding to predict a label for the graph.

To formulate GNN-based FL, using the model definition above, we define \( W = \{ \theta, \Psi, \Phi_{\text{pool}}, \Phi_{\text{task}} \} \) as the overall learnable weights. Note that \( W \) is independent of graph structure as both the GNN and Readout parameters make no assumptions about the input graph. Thus, one can learn \( W \) using a FL based approach. The the overall FL task can be formulated as a distributed optimization problem as:

\[ \min_W F(W) \overset{\text{def}}{=} \min_{W} \sum_{k=1}^{K} \frac{N^{(k)}}{N} \cdot f^{(k)}(W) \]  

(4)

where \[ f^{(k)}(W) = \frac{1}{N^{(k)}} \sum_{i=1}^{N^{(k)}} \mathcal{L}(\hat{y}^{(k)}_i, y^{(k)}_i) \] is the \( k^{th} \) client’s local objective function that measures the local empirical risk over the heterogeneous graph dataset \( D^{(k)} \). \( \mathcal{L} \) is the loss function of the global graph classifier.

With such a formulation, it might seem like an optimization problem tailored for a FedAvg based optimizers [44, 18, 51]. Unfortunately, in molecular graph settings this is not the case for the following reasons. (a) In our setting, clients belong to a decentralized, serverless topology. There is no central server that can average model parameters from the clients. (b) Clients in our setting possess incomplete labels, hence the dimensions of \( \Phi_{\text{task}} \) can be different on different clients. For instance, a client may only have a partial toxicity labels for a molecule, while another client may have a molecule’s interaction property with another drug compound. Even with such incomplete information from each client, our learning task is interested in classifying each molecule across multiple label categories. Federated Multitask Learning (FMTL) [51, 5] is a popular framework designed to deal with such issues. However, current approaches do not generalize to non-convex deep models such as graph neural networks. To combat these issues, we first introduce a centralized FMTL framework for graph neural networks in Section 2.2. But reliance on a central server may not be feasible in molecular graphs owned by multiple competing entities. Hence, we enhance this centralized FMTL to a serverless scenario in section 2.3, which we call SpreadGNN.
2.2 Federated Multi-Task Learning with Graph Neural Networks

Under the regularized MTL paradigm [13], we define the centralized federated graph MTL problem (FedGMTL) as follows:

\[
\min_{\theta, \Psi, \Phi_{\text{pool}}, \Phi_{\text{task}}} \sum_{k=1}^{K} \frac{1}{N_k} \sum_{i=1}^{N_k} L(\hat{y}_i^{(k)}, y_i^{(k)}) + R(\mathbf{W}, \Omega),
\]

\[
\text{s.t. } \Omega \geq 0 \text{ and } \text{Tr}(\Omega) = 1.
\]

where

\[
R(\mathbf{W}, \Omega) = \frac{1}{2} \lambda_1 \text{Tr}(\Phi_{\text{task}} \Omega^{-1} \Phi_{\text{task}}^T) + \frac{1}{2} \sum_{\chi \in \{\theta, \Psi, \Phi_{\text{pool}}, \Phi_{\text{task}}\}} \lambda_{\chi} ||\chi||_F^2
\]

is the bi-convex regularizer introduced in [74]. The first term of the Eq. (5) models the summation of different empirical loss of each client, which is what Eq. (4) exactly tries to address. The second term serves as a task-relationship regularizer with \(\Omega \in \mathbb{R}^{S \times S}\) being the covariance matrix for \(S\) different tasks constraining the task weights \(\Phi_{\text{task}} = [\Phi_{\text{task},1}, \ldots, \Phi_{\text{task},S}] \in \mathbb{R}^{d \times S}\) through matrix trace \(\text{Tr}(\Phi_{\text{task}} \Omega^{-1} \Phi_{\text{task}}^T)\). Recall that each client in our setting may only have a partial set of tasks in the labels of its training dataset, but still needs to make predictions for tasks it does not have in its labels during test time. This regularizer helps clients relate its own tasks to tasks in other clients. Intuitively, it determines how closely two tasks \(i\) and \(j\) are related. The closer \(\Phi_{\text{task},i}\) and \(\Phi_{\text{task},j}\) are, the larger \(\Omega_{i,j}\) will be. If \(\Omega\) is an identity matrix, then each node is independent to each other. But as our results show, there is often a strong correlation between different molecular properties. This compels us to use a federated learning model.

Figure 2-a depicts the FedGMTL framework where clients’ graph classifier weights are using an FL server. While the above formulation enhances the FMTL with a constrained regularizer that can be used for GNN learning, we still need to solve the final challenge, which is to remove the reliance on a central server to perform the computations in Eq. (5). Therefore, we propose a Decentralized Graph Multi-Task Learning framework, SpreadGNN, and utilize our novel optimizer, Decentralized Periodic Averaging SGD (DPA-SGD) to extend the FMTL framework to our setting. The decentralized framework is shown in the Figure 2-c. Note that each client’s learning task remains the same but the aggregation process differs in SpreadGNN.

2.3 SpreadGNN: Serverless Federated MTL for GNNs

To transition to the serverless setting, we introduce a novel optimization method called Decentralized Periodic Averaging SGD (DPA-SGD). The main idea of DPA-SGD is that each client applies SGD locally, and synchronizes all parameters with only its neighbors during a communication round that occurs every \(\tau\) iterations. A decentralized system in which all clients are not necessarily connected to all other clients also makes it impossible to maintain one single task covariance matrix \(\Omega\). Thus we propose using distinct covariance matrices \(\Omega_k\) in each client that are efficiently updated using the exchange mechanism illustrated in Figure 3. We formalize this idea as follows. Consider one particular client \(m\) having task weights \(\Phi_{\text{task},m} \in \mathcal{R}^{d \times S_m}\) where \(S_m\) is the number of tasks that client \(m\) has. Recall that the regularizer in Eq. (6) is how various clients learn to predict tasks that are not within its \(S_m\).
In the decentralized setting, we emphasize that clients can collectively learn the exhaustive set of tasks, even when clients may not have access to some of the classes in each multi-class label. That is and \( \bigcup_i S_i = S \). Then, the new non-convex objective function can be defined as:

\[
\min_{\theta, \psi, \Phi_{pool}, \Phi_{\text{task}}} \sum_{k=1}^{K} \frac{1}{N_k} \sum_{i=1}^{N_k} \left[ l(y_i^{(k)}(x_i^k, z_k^k; W_k), y_i^{(k)}) + \frac{1}{2} \lambda_1 \text{Tr}(\Phi_{\text{task},i} \Omega_k^{-1} \Phi_{\text{task},i}^T) \right]
+ \frac{1}{2} \sum_{\chi \in \{\theta, \psi, \Phi_{pool}, \Phi_{\text{task}}\}} \lambda_\chi \|\chi\|_F^2,
\]

\( \quad \text{s.t.} \ \Omega_k \geq 0 \quad \text{and} \quad \text{Tr}(\Omega_k) = 1, \quad k = 1, 2, \ldots, K. \)

where \( W_k = \{\theta, \psi, \Phi_{pool}, \Phi_{\text{task},k}\} \) is the set of all learnable weights for client \( k \); \( \mathcal{M}_k = k \cup N_k \) is the neighbor set for client \( k \) including itself. This gives rise to: \( \Phi_{\text{task},i} = [\Phi_{\text{task},i}^1 | \Phi_{\text{task},i}^2 | \ldots | \Phi_{\text{task},i}^{\mathcal{M}_k}] \in \mathbb{R}^{d \times |\mathcal{M}_k|} \) which is the task weight matrix for client \( k \) and its neighbors and \( \| \) represents the row-wise union operation. The matrix \( \Omega_k \in \mathbb{R}^{(|\mathcal{M}_k| \times |\mathcal{M}_k|)} \) represents the correlation amongst all the available tasks for the set \( \mathcal{M}_k \).

To solve this non-convex problem, we apply the alternating optimization method presented in [74], where \( W_k \) and \( \Omega_k \) are updated in an alternative fashion.

**Optimizing \( W_k \)**: For simplicity, we define \( \Omega = \{\Omega_1, \Omega_2, \ldots, \Omega_K\} \) to represent the set of correlation matrices for all clients. Fixing \( \Omega \), we can use SGD to update \( W_k \) jointly. Let \( L = \sum_{k=1}^{K} \frac{1}{N_k} \sum_{i=1}^{N_k} l(y_i^{(k)}(x_i^k, z_i^k; W_k), y_i^{(k)}) \). Then, our problem can then be reformulated as:

\[
G(W_k|\Omega) = L + \frac{1}{N_k} \sum_{i=1}^{N_k} \frac{1}{2} \lambda_1 \text{Tr}(\Phi_{\text{task},i} \Omega_k^{-1} \Phi_{\text{task},i}^T) + \frac{1}{2} \sum_{\chi \in \{\theta, \psi, \Phi_{pool}, \Phi_{\text{task}}\}} \lambda_\chi \|\chi\|_F^2. \quad (8)
\]

where the summation in (8) is amongst all the nodes connected to node \( k \). Then the gradient formualations for each node are:

\[
\frac{\partial G(W_k|\Omega)}{\partial \Phi_{\text{task},i}} = \frac{\partial L}{\partial \Phi_{\text{task},i}} + \lambda_1 \frac{1}{N_k} \Phi_{\text{task},i} \Omega_k^{-1} + \lambda_2 \Phi_{\text{task},i}
\quad (9)
\]

\[
\frac{\partial G(W_k|\Omega)}{\partial \chi} = \frac{\partial L}{\partial \chi} + \lambda_\chi \chi, \quad \forall \chi \in \{\theta, \psi, \Phi_{pool}\}
\quad (10)
\]

**Optimizing \( \Omega_k \in \Omega \)**: In [74], an analytical solution for \( \Omega \) is equal to \( (\Phi_{\text{task}}^T \Phi_{\text{task}})^{\frac{1}{2}} \). However, this solution is only applicable for the centralized case. This is because The missing central node forbids averaging parameters globally. So here we propose a novel way to update each \( \Omega_k \in \Omega \):

\[
f_{\text{align}}(\Omega_k^{(t+1)}) \leftarrow \eta \frac{1}{|\mathcal{M}_k|} \sum_{i=1}^{|\mathcal{M}_k|} f_{\text{align}}(\Omega_k^{(t)}) + f_{\text{align}}((\Phi_{\text{task},i}^T \Phi_{\text{task},i})^{\frac{1}{2}}) \quad (11)
\]

\[
es_{\text{align}}(\Omega_k^{(t+1)}) \leftarrow \eta \frac{1}{|\mathcal{M}_k|} \sum_{i=1}^{|\mathcal{M}_k|} e_{\text{align}}(\Omega_k^{(t)}) + e_{\text{align}}((\Phi_{\text{task},i}^T \Phi_{\text{task},i})^{\frac{1}{2}}) \quad (12)
\]

\[
es_{\text{align}}(\Omega_k^{(t+1)}) \leftarrow \eta \frac{1}{|\mathcal{M}_k|} \sum_{i=1}^{|\mathcal{M}_k|} e_{\text{align}}(\Omega_k^{(t)}) + e_{\text{align}}((\Phi_{\text{task},i}^T \Phi_{\text{task},i})^{\frac{1}{2}}) \quad (13)
\]

\[
es_{\text{align}}(\Omega_k^{(t+1)}) \leftarrow \eta \frac{1}{|\mathcal{M}_k|} \sum_{i=1}^{|\mathcal{M}_k|} e_{\text{align}}(\Omega_k^{(t)}) + e_{\text{align}}((\Phi_{\text{task},i}^T \Phi_{\text{task},i})^{\frac{1}{2}}) \quad (14)
\]

\[
es_{\text{align}}(\Omega_k^{(t+1)}) \leftarrow \eta \frac{1}{|\mathcal{M}_k|} \sum_{i=1}^{|\mathcal{M}_k|} e_{\text{align}}(\Omega_k^{(t)}) + e_{\text{align}}((\Phi_{\text{task},i}^T \Phi_{\text{task},i})^{\frac{1}{2}}) \quad (15)
\]
The first averaging term can incorporate the nearby nodes correlation into its own. It should be noticed that each $\Omega_i$ may have a different dimension (different number of neighbors), so this averaging algorithm is based on the node-wised alignment in the global $\Omega$. The $f_{\text{align}}$ function is illustrated in Figure 3. Here, $f_{\text{align}}$ operates on each client and The second term captures the new correlation between its neighbors as shown in Figure 3. We refer readers to to Algorithm 1 in Appendix 10.

### 2.3.1 Convergence Properties of DPA-SGD

In this section, we present our convergence analysis for DPA-SGD. Our analysis explains why DPA-SGD works well in the federated setting. In the previous section, we present DPA-SGD algorithm. Before any formalism, we first introduce a binary symmetric node connection matrix $M \in \mathbb{R}^{K \times K}$ to record the connections in the network, where $M_{i,j}$ is a binary value that indicates whether node $i$ and $j$ are connected or not. We assume that $M$ satisfies the following: (a) $M1_K = 1_K$ (b) $M^T = M$ (c) $\max\{|\lambda_2(M)|, \ldots, |\lambda_K(M)|\} < \lambda_1(M) = 1$ where $\lambda$ is the eigenvalues of $M$. Note that, if $M = I$ (Identity matrix), then every nodes are independent and update the parameters respectively. If $M = J$ (one for each element), the model is degenerated into centralized model. Our convergence analysis satisfies the assumptions [4] listed in Appendix 10. Here, we present mathematically simple and convenient update rule of DPA-SGD:

$$X_{t+1} = (X_t - \eta G_t) \cdot M_t,$$

where $X_t = [x_t^{(1)}, \ldots, x_t^{(K)}]$ is the matrix of our interest (e.g. $W^{(t)}_k$), $G_t$ is the gradient and $M_t$ is the connection matrix at time $t$. When $t \mod \tau = 0$, $M_t = M$, otherwise, $M_t$ equals to $I_K$. In equation (12), multiplying $I_K/K$ on both side, defining the averaged model $u_t = X_t I_{K/K}$, we obtain the following update:

$$u_{t+1} = u_t - \eta g_t = u_t - \eta \left[ \frac{1}{K} \sum_{i=1}^{K} g(x_t^{(i)}) \right]$$

Next, we will present our analysis on the convergence of the above-averaged model $u_t$. For non-convex optimization, previous works on SGD convergence analysis use the average squared gradient norm as an indicator of convergence to a stationary point [4].

**Theorem 1 (Convergence of DPA-SGD)** Assume that Assumption 1 and Definition (13) hold. Then, if the learning rate $\eta$ satisfies the following condition:

$$\eta L + \frac{\eta^2 L^2 \tau^2}{1 - \zeta} \left( \frac{2\zeta^2}{1+\zeta} + \frac{2\zeta}{1-\zeta} + \frac{\tau - 1}{\tau} \right) \leq 1$$

where $\tau$ is the averaging period(one synchronization per $\tau$ local updates), and $\zeta = \max\{|\lambda_2(M)|, \ldots, |\lambda_m(M)|\}$, and all local models are initialized at a same point $x_0$, then after $K$ iterations the average squared gradient norm is bounded as

$$\mathbb{E} \left[ \frac{1}{T} \sum_{t=1}^{T} \|\nabla F(u_t)\|^2 \right] \leq \frac{2\|F(x_t) - F_{\text{inf}}\|}{\eta T} + \frac{\eta L \sigma^2}{K} + \eta^2 L^2 \sigma^2 \left( \frac{1+\zeta^2}{1-\zeta^2} \tau - 1 \right)$$

Theorem 1 shows that DPA-SGD algorithm is a trade-off between convergence rate and system optimization including communication-efficiency(communication rounds) and convergence speed. find a recent work [68] has got the same result in an unified framework. Our work differs from this work in that it does not provide adequate theoretical analysis and empirical evaluation for federated learning. The proof of Theorem 1 is given in the Appendix 10.

### 3 Experiments

#### 3.1 Setup

**Implementation.** All experiments are conducted on a single GPU server equipped with 2 Nvidia Geforce GTX 1080Ti GPUs and an AMD Ryzen 7 3700X 8-Core Processor. Our models are built on top of the FedML framework [19] and PyTorch [48].
Table 1: Dataset summary used in our experiments.

| Dataset | # Molecules | Avg # Nodes | Avg # Edges | # Tasks | Task Type | Evaluation Metric |
|---------|-------------|-------------|-------------|---------|-----------|-------------------|
| SIDER   | 1427        | 33.64       | 35.36       | 27      | Classification | ROC-AUC           |
| Tox21   | 7831        | 18.51       | 25.94       | 12      | Classification | ROC-AUC           |
| MUV     | 93087       | 24.23       | 76.80       | 17      | Classification | ROC-AUC           |
| QM8     | 21786       | 7.77        | 23.95       | 12      | Regression    | MAE               |

Table 2: Results on the molecular property prediction task. SpreadGNN uses a complete topology: all clients communicated with each other. Communication period $\tau = 1$

|                | GraphSAGE | GAT       |
|----------------|-----------|-----------|
|                | FedAvg    | FedGMTL  | SpreadGNN | FedAvg    | FedGMTL  | SpreadGNN |
| SIDER          | 0.582     | 0.629    | **0.5873**| 0.5857    | 0.61     | **0.6034**|
| Tox21          | 0.5548    | 0.6664   | **0.585** | 0.6035    | 0.6594   | **0.6056**|
| MUV            | 0.6578    | 0.6856   | **0.703** | 0.709     | 0.6899   | **0.713**  |
| QM8            | 0.02982   | 0.03624  | **0.02824**| 0.0392    | 0.0488   | **0.0315**|

Multi-task Dataset. We use molecular datasets from the MoleculeNet [69] machine learning benchmark in our evaluation. In particular, we evaluate our approach on molecular property prediction datasets described in Table 1. The label for each molecule graph is a vector in which each element denotes a property of the molecule. Properties are binary in the case of classification or continuous values in the case of regression. As such, each multitask dataset can adequately evaluate our learning framework.

Non-I.I.D. Partition for Quantity and Label Skew. We introduce non-I.I.D.ness in two additive ways. The first is a non-I.I.D. split of the training data based on quantity. Here we use a Dirichlet distribution parameterized by $\alpha$ to split the training data between clients. Specifically, the number of training samples present in each client is non-I.I.D. The second source of non-I.I.D.ness is a label masking scheme designed to represent the scenario in which different clients may possess partial labels as shown in Figure 1. More specifically, we mask out a subset of classes in each label on every client. In our experiments, the sets of unmasked classes across all clients are mutually exclusive and collectively exhaustive. This setting simulates a worst case scenario where no two clients share the same task. However our framework is just as applicable when there is label overlap. Such masking, introduces a class imbalance between the clients making the label distribution non-I.I.D. as well.

Models. In order to demonstrate that our framework is agnostic to the choice of GNN model, we run experiments on two different GNN architectures: GraphSAGE [17] and GAT [65].

Network Topology. We first evaluate our framework in a complete topology in which all clients are connected to all other clients to measure the efficacy of our proposed regularizer. We then perform ablation studies on the number of neighbors of each client to stress test our framework in the more constrained setting.

Hyperparameters. We use Adam [28] as the client optimizer in all of our experiments. For our Tox21, MUV and QM8 we use an 8 client topology. For SIDER we use a 4 client topology. A more comprehensive hyperparameter list for network topology and models can be found in the Appendix 8.

3.2 Results

We use a central, server dependent FedAvg system as the baseline of comparison. More specifically, all clients are involved in the averaging of model parameters in every averaging round.

Our results summarized in Table 2 demonstrate that SpreadGNN (third column) outperforms a centralized federated learning system that uses FedAvg (first column) when all clients can communicate with all other clients. This shows that by using the combination of the task regularizer in equation 6 and the DPA-SGD algorithm, we can eliminate the dependence on a central server and enable clients to learn more effectively in the presence of missing molecular properties in their respective labels. Additionally, the results also show that our framework is agnostic to the type of GNN model.
being used in the molecular property prediction task since both GraphSAGE and GAT benefit from our framework. Our framework also works in the case a trusted central server is available (second column). The presence of a trusted central server improves accuracy in a few scenarios. However, SpreadGNN provides competing performance in a more realistic setting where central server is not feasible.

3.3 Sensitivity Analysis

Task Regularizer. Figure 4 illustrates the effect of $\lambda_1$ on both regression as well as classification tasks. Interestingly, regression is much more robust to variation in $\lambda_1$ while classification demands more careful tuning of $\lambda_1$ to achieve optimal model performance. This implies that the different properties in the regression task are more independent than the properties in the classification task.

Network Topology. The network topology dictates how many neighbors each client can communicate with in a communication round. While Table 2 shows that SpreadGNN outperforms FedAvg in a complete topology, Figure 5 shows that our framework performs outperforms FedAvg even when clients are constrained to communicate with fewer neighbors. We can also see that it’s not just $n_{neighbors}$ that matters, the topology in which clients are connected does too. When $n_{neighbors} = 2$, a ring topology outperforms a random topology, as a ring guarantees a path from any client to any other client. Thus, learning is shared indirectly between all clients. The same is not true in a random topology. More ablation studies on the the network topology are included in the Appendix 9.

Period. The communication period $\tau$ is another important hyperparameter in our framework. In general, our experiments suggest that a lower period is better, but this is not always the case. We include an ablation study on $\tau$ to support this claim in the Appendix 9.
4 Related Works

**Molecular Representation Learning.** [52] encode the neighbors of atoms in the molecule into a fix-length vector to obtain vectors-space representations. To improve the expressive power of chemical fingerprints, [12, 7] use CNNs to learn rich molecule embeddings for downstream tasks like property prediction. [27, 57, 58] explore the graph convolutional network to encode molecular graphs into neural fingerprints. To better capture the interactions among atoms, [14] proposes to use a message passing framework and [70, 30] extend this framework to model bond interactions.

**FL.** Early examples of research into federated learning include [31, 32, 43]. To address both statistical and system challenges, [61] and [5] propose a multi-task learning framework for federated learning and its related optimization algorithm, which extends early works in distributed machine learning [59, 71, 72, 21, 40, 60]. The main limitation of [61] and [5], however, is that strong duality is only guaranteed when the objective function is convex, which can not be generalized to the non-convex setting.[2, 25, 41, 67, 3, 39] extends federated multi-task learning to the distributed multi-task learnings setting, but not only limitation remains same, but also nodes performing same amount of work is prohibitive in FL.

**Federated Graph Neural Networks.** [63] and [45] use computed graph statistics for information exchange and aggregation to avoid node information leakage. [22, 75] utilize cryptographic approaches to incorporate into GNN learning. [66] propose a hybrid of federated and meta learning to solve the semi-supervised graph node classification problem in decentralized social network datasets. [46] uses an edge-cloud partitioned GNN model for spatio-temporal traffic forecasting tasks over sensor networks. The previous works do not consider graph learning in a decentralized setting.

**Stochastic Gradient Descent Optimization.** In large scale distributed machine learning problems learning, synchronized mini-batch SGD is a well-known method to address the communication bottleneck by increasing the computation to communication ratio [10, 35, 8]. It is shown that FedAvg [32] is a special case of local SGD which allow nodes to perform local updates and infrequent synchronization between them to communicate less while converging fast [62, 68, 73, 38]. Decentralized SGD, another approach to reducing communication, was successfully applied to deep learning [24, 23, 37]. Asynchronous SGD is a potential method that can alleviate synchronization delays in distributed learning [50, 8, 15, 47, 11], but existing asynchronous SGD doesn’t fit for federated learning because the staleness problem is particularly severe due to the reason of heterogeneity in the federated setting [9].

5 Conclusion

In this work, we propose SpreadGNN, a framework to train graph neural networks in a serverless federated system. We motivate our framework through a realistic setting, in which clients involved in molecular machine learning research cannot share data with each other due to privacy and competition. Moreover, we further recognize that clients may only possess partial labels of their data. Through experimental analysis, we show for the first time, that training graph neural networks in a federated manner does not require a centralized topology and that our framework can address the non-iidness in dataset size and label distribution across clients. SpreadGNN can outperform a central server depend baseline even when clients can only communicate with a few neighbors. To support our empirical results, we also provide a convergence analysis of the DPA-SGD optimization algorithm used by SpreadGNN.

**Broader impact**

Our method can be used to train graph neural networks for molecular property prediction tasks in a serverless manner. Our method can have immense societal benefits such as accelerated drug discovery. One limitation of our method is that although federated learning can protect privacy to a certain extent, recent studies have shown that it cannot guarantee that data or models are not leaked. Therefore, if researchers want to deploy our method to the actual production environment as a public service, we recommend analyzing inherent security and privacy risks comprehensively and adequately. It is necessary to add security and privacy-related components, such as Differential Privacy and Secure Aggregation.
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Appendix

This section includes additional information that a reader might find useful. Apart from the proof of Theorem 1, we include the algorithm sketch for SpreadGNN using DPA-SGD, a more detailed description of the datasets we used, hyperparameter configurations used in our experiments and additional ablation studies on communication period and network topology.

6 Algorithm Sketch

Algorithm 1 SpreadGNN: Serverless Multi-task Federated Learning for Graph Neural Networks

Require: initial parameters for each node $W_k^{(t=0)} = \{\theta^{(t=0)}, \Psi^{(t=0)}, \Phi^{(t=0)}_\text{pool}, \Phi^{(t=0)}_{\text{task},k}\}$ and $\Omega^{(t=0)} = (\Omega_1^{(t=0)}, \Omega_2^{(t=0)}, \ldots, \Omega_K^{(t=0)})$; learning rate $\eta$; maximum number of global iterations $T$, maximum number of client epochs $E$; communication period $\tau$.

1: for all nodes: $k = 1, 2, \ldots, K$ in parallel do
2: for $t = 1$ to $T$ do
3: for $k = 1$ to $E$ (epoch loop) do
4: for $m \in MB$ (mini-batch loop) do
5: Read a minibatch $m$
6: Calculate gradient: $g(W_k^{(t,m)}) = \frac{\partial G(W_k^{(t,m)}|\Omega^{(t,m)})}{\partial W_k^{(t,m)}}$
7: Update the local $k$th optimization variables:
   $W_k^{(t+1,m)} \leftarrow W_k^{(t,m)} - \eta g(W_k^{(t,m)})$
   $\Omega_k^{(t+1,m)} \leftarrow (\Phi_{\text{task},M_k}^T \Phi_{\text{task},M_k})^{1/2} \text{Tr}((\Phi_{\text{task},M_k}^T \Phi_{\text{task},M_k})^{1/2}) / |M_k|$
8: end for
9: end for
10: if $t$ mod $\tau = 0$ then
11: Perform aggregation and alignment over neighbors for node $k$:
   $W_k^{(t+1)} \leftarrow \frac{1}{|M_k|} \sum_{j=1}^{|M_k|} W_k^{(t)}$
   $f_{\text{align}}(\Omega_k^{t+1}) \leftarrow \eta (\sum_{j=M_k \setminus k} \frac{1}{N_j} f_{\text{align}}(\Omega_j^{(t)}) + f_{\text{align}}(\frac{(\Phi_{\text{task},M_k}^T \Phi_{\text{task},M_k})^{1/2}}{\text{Tr}((\Phi_{\text{task},M_k}^T \Phi_{\text{task},M_k})^{1/2})}))/|M_k|$
12: end if
13: end for
14: end for

7 Dataset Details

Table 1 summarizes the necessary information of benchmark datasets [69]. The details of each dataset are listed below:

- SIDER [33], or Side Effect Resource, the dataset consists of marketed drugs with their adverse drug reactions.
- Tox21 [1] is a dataset which records the toxicity of compounds.
- MUV [53] is a subset of PubChem BioAssay processed via refined nearest neighbor analysis. Contains 17 tasks for around 90 thousand compounds and is specifically designed for validation of virtual screening techniques.
- QM8 [49] is composed from a recent study on modeling quantum mechanical calculations of electronic spectra and excited state energy of small molecules.

7.1 Feature Extraction Procedure for Molecules

The feature extraction is in two steps:

1. Atom-level feature extraction and Molecule object construction using RDKit [34].
2. Constructing graphs from molecule objects using NetworkX [16].

Atom features, shown in Table 3, are the atom features we used. It’s exactly the same set of features as used in [54].

Table 3: Atom features

| Features         | Size | Description                                                                 |
|------------------|------|-----------------------------------------------------------------------------|
| atom type        | 100  | Representation of atom (e.g., C, N, O), by its atomic number                |
| formal charge    | 5    | An integer electronic charge assigned to atom                              |
| number of bonds  | 6    | Number of bonds the atom is involved in                                    |
| chirality        | 5    | Number of bonded hydrogen atoms                                            |
| number of H      | 5    | Number of bonded hydrogen atoms                                            |
| atomic mass      | 1    | Mass of the atom, divided by 100                                           |
| aromaticity      | 1    | Whether this atom is part of an aromatic system                            |
| hybridization    | 5    | SP, SP2, SP3, SP3D, or SP3D2                                              |

8 Model Hyperparameters

8.1 Model Architecture

As explained in section 2.1 our model is made up of a GNN and Readout. The GNNs we use are GAT [65] and GraphSAGE [17]. Each accepts input node features $X_v \in \mathbb{R}^{|V_M| \times d_{input}}$ and outputs node embeddings $h_v \in \mathbb{R}^{|V_M| \times d_{node}}, v \in V_M$. Where $V_M$ is the set of atoms in molecule $M$. Given the output node embeddings from the GNN the Readout function we use is defined as follows:

$$R_{\Phi_{pool}, \Phi_{task}}(h_v, X_v) = \text{MEAN}(\text{ReLU}(\Phi_{task}(\text{ReLU}(\Phi_{pool}(X_v \parallel h_v))))))$$

where $\parallel$ represents the row wise concatenation operation. $\Phi_{pool} \in \mathbb{R}^{(d_{node}+d_{input}) \times d_{pool}}$ and $\Phi_{task} \in \mathbb{R}^{d_{pool} \times d_{out}}$ are learnable transformation matrices. $d_{out}$ represents the number of classes/tasks present in the classification label. The MEAN operation here is a column wise mean. Note that while our general description of the readout in section 2.1 does not include the input features as part of the input, we find that including the input features leads to better generalization.

8.2 Hyperparameter Configurations

For each task, we utilize grid search to find the best results. Table 4 lists all the hyper-parameters ranges used in our experiments. All hyper-parameter tuning is run on a single GPU. The best hyperparameters for each dataset and model are listed in Table 5. The batch-size is kept 1. This pertains to processing a single molecule at a time. The number of GNN layers were fixed to 2 because having too many GNN layers result in over-smoothing phenomenon as shown in [36]. For all experiments, we used Adam optimizer.

9 Detailed Ablation Studies

9.1 Effect of Communication Period $\tau$

Figures 6 & 7 illustrate the effect of communication period SIDER and Tox21 datasets. As we increase the communication period $\tau$ more, model performance decreases. However, selecting $\tau = 5$ can sometimes be better than averageing & exchanging each round. This indicates that, tuning $\tau$ is important for while controlling the tradeoff between the performance and the running time.
Table 4: Hyperparameter Range for Experiments

| hyper-parameter | Description | Range |
|-----------------|-------------|-------|
| Learning rate   | Rate of speed at which the model learns. | [0.00015, 0.001, 0.0015, 0.0025, 0.015, 0.15] |
| Dropout rate    | Dropout ratio | [0.3, 0.5, 0.6] |
| Node embedding dimension (d_{node}) | Dimensionality of the node embedding | 64 |
| Hidden layer dimension | GNN hidden layer dimensionality | 64 |
| Readout embedding dimension (d_{pool}) | Readout Hidden Layer Dimensionality | 64 |
| Graph embedding dimension (d_{out}) | Dimensionality of the final graph embedding | 64 |
| Attention heads | Number of attention heads required for GAT | 1-7 |
| Alpha | LeakyRELU parameter used in GAT model | 0.2 |
| Rounds | Number of federating learning rounds | 150 |
| Epoch | Epoch of clients | 1 |
| Number of clients | Number of users in a federated learning round | 4-10 |
| Communication Period | Exchanging Period between clients | 1 |

Table 5: Hyperparameters used in our experiments. For SpreadGNN we use a communication period $\tau = 1$ and a complete topology (all clients connected to all other clients) in all experiments.

| Parameters | SIDER | Tox21 | MUV | QMS |
|------------|-------|-------|-----|-----|
| ROC-AUC Score | 0.582 | 0.58 | 0.6584 | 0.705 |
| Partition alpha | 0.2 | 0.2 | 0.2 | 0.2 |
| Learning rate | 0.0015 | 0.0015 | 0.0015 | 0.0015 |
| Dropout rate | 0.3 | 0.3 | 0.3 | 0.3 |
| Node embedding dimension | 64 | 64 | 64 | 64 |
| Hidden layer dimension | 64 | 64 | 64 | 64 |
| Readout embedding dimension | 64 | 64 | 64 | 64 |
| Graph embedding dimension | 64 | 64 | 64 | 64 |
| Attention Heads | NA | NA | NA | NA |
| Leaky ReLU alpha | 0.2 | 0.2 | 0.2 | 0.2 |
| Number of Clients | 4 | 4 | 8 | 8 |
| Task Regularizer | NA | NA | NA | NA |
| ROC-AUC Score | 0.585 | 0.6584 | 0.695 | 0.705 |
| Partition alpha | 0.1 | 0.1 | 0.1 | 0.1 |
| Learning rate | 0.0015 | 0.0015 | 0.0015 | 0.0015 |
| Dropout rate | 0.3 | 0.3 | 0.3 | 0.3 |
| Node embedding dimension | 64 | 64 | 64 | 64 |
| Hidden layer dimension | 64 | 64 | 64 | 64 |
| Readout embedding dimension | 64 | 64 | 64 | 64 |
| Graph embedding dimension | 64 | 64 | 64 | 64 |
| Attention Heads | NA | NA | NA | NA |
| Leaky ReLU alpha | 0.2 | 0.2 | 0.2 | 0.2 |
| Number of Clients | 8 | 8 | 8 | 8 |
| Task Regularizer | NA | NA | NA | NA |
| ROC-AUC Score | 0.585 | 0.6584 | 0.705 | 0.705 |
| Partition alpha | 0.3 | 0.3 | 0.3 | 0.3 |
| Learning rate | 0.002 | 0.0025 | 0.0025 | 0.0025 |
| Dropout rate | 0.3 | 0.3 | 0.3 | 0.3 |
| Node embedding dimension | 64 | 64 | 64 | 64 |
| Hidden layer dimension | 64 | 64 | 64 | 64 |
| Readout embedding dimension | 64 | 64 | 64 | 64 |
| Graph embedding dimension | 64 | 64 | 64 | 64 |
| Attention Heads | NA | NA | NA | NA |
| Leaky ReLU alpha | 0.2 | 0.2 | 0.2 | 0.2 |
| Number of Clients | 8 | 8 | 8 | 8 |
| Task Regularizer | NA | NA | NA | NA |
| ROC-AUC Score | 0.585 | 0.6584 | 0.705 | 0.705 |
| Partition alpha | 0.3 | 0.3 | 0.3 | 0.3 |
| Learning rate | 0.0015 | 0.0015 | 0.0015 | 0.0015 |
| Dropout rate | 0.3 | 0.3 | 0.3 | 0.3 |
| Node embedding dimension | 64 | 64 | 64 | 64 |
| Hidden layer dimension | 64 | 64 | 64 | 64 |
| Readout embedding dimension | 64 | 64 | 64 | 64 |
| Graph embedding dimension | 64 | 64 | 64 | 64 |
| Attention Heads | NA | NA | NA | NA |
| Leaky ReLU alpha | 0.2 | 0.2 | 0.2 | 0.2 |
| Number of Clients | 8 | 8 | 8 | 8 |
| Task Regularizer | NA | NA | NA | NA |

Table 5: Hyperparameters used in our experiments. For SpreadGNN we use a communication period $\tau = 1$ and a complete topology (all clients connected to all other clients) in all experiments.
9.2 Effect of Serverless Network Topology

Figure 8 illustrates the effect of varying topologies on SpreadGNN on the Sider dataset when using GraphSage as the GNN. The qualitative behavior is similar to Figure 5, in that when each client is connected to more neighbors, the local model on each client is more robust. However, when the total number of clients involved in the network is smaller, the effect of topology is understated and the total number of neighbors matters more. Recall that in an 8 client network, when each client was restricted to being connected to only 2 neighbors, random connections performed worse than a ring topology, meaning that the topology mattered as much as mere number of neighbors. However, in the case of the 4 client network in Figure 8 there is a minimal difference between a 2 neighbor random configuration and a 2 neighbor ring configuration.

10 Proof for Convergence of DPA-SGD

In our analysis, we assume that following properties hold [4]:

- The objective function $F(\cdot)$ is L-Lipschitz.
- $F(\cdot)$ is lower bounded by $F_{inf}$ such that $F(\cdot) \geq F_{inf}$.
- The full gradient of the objective function $F(\cdot)$ is approximated by stochastic gradient $g$ on a mini-batch $\epsilon_k$ with an unbiased estimate: $E_{\epsilon_k}[g(\hat{x})] = \nabla F(\hat{x})$.
- The variance of stochastic gradient $g(\hat{x})$ given a mini-batch $\epsilon_k$ is upper bounded and $\text{Var}_{\epsilon_k}(g(\hat{x})) \leq \beta \|
abla F(\hat{x})\|^2 + \sigma^2$, $\exists \beta, \sigma^2 \geq 0$, $\forall k$. 


In order to get a more clear view of our algorithm, we reformulate the loss function on each worker as follows:

\[
f_k \left( \Gamma, \Phi_{\text{task},s,t}, \Omega^{(k)}, \xi^{(k)} \right) := L \left( y^{(k)}_t (X_{i,k}, Z_{i,k}; W_k), y^{(k)}_t \right) + \frac{1}{2} \lambda_1 \text{Tr} (\Phi_{\text{task},s,t}^{\top} \Omega^{-1}_k \Phi_{\text{task},s,t}) \\
+ \frac{1}{2} \sum_{\chi \in \{\theta, \Phi_{\text{pool}}, \Phi_{\text{task}}\}} \lambda_\chi \| \chi \|_F^2,
\]

where we include all the parameters that is different on each worker to be \( \delta^{(k)} := \{\Phi_{\text{task},s,t}\} \), \( \Gamma \) denotes the shared parameters, and \( \xi^{(k)}_i \) is the random variable that denotes the data samples \( (X_{i,k}, Z_{i,k}, y^{(k)}_t) \).

Therefore, the original objective function an be cast into the following form:

\[
F \left( \Gamma, \delta^{(1:K)}, \Omega^{(1:K)} \right) := \frac{1}{n} \sum_{i=1}^{K} E_{\xi^{(k)}} f_k \left( \Gamma, \delta^{(k)}, \Omega^{(k)}, \xi^{(k)} \right).
\]

Notice that the updating rule for \( \Gamma, \delta^{(1:K)} \), and \( \Omega^{(1:K)} \) are different in that:

\[
\Gamma_{t+1} = \Gamma_t - \eta \frac{\partial F}{\partial \Gamma_t} M, \\
\delta^{(k)}_{t+1} = \delta^{(k)}_t - \eta \frac{\partial F}{\partial \delta^{(k)}_t}, \\
\Omega^{(k)}_{t+1} = \text{argmin}_{\Omega^{(k)}} F \left( \Gamma, \delta^{(1:K)}_t, \Omega^{(1:K)}_t \right).
\]

From the update rule of \( \Omega^{(1:K)}_t \), we know that

\[
\leq \frac{L}{2} \text{Tr} \left( \left( \delta^{(1:K)}_{t+1} - \delta^{(1:K)}_t \right)^2 \right) + L \eta \text{Tr} \left( \frac{\partial F}{\partial \delta^{(1:K)}_t} \right) + \frac{\eta}{2} \text{Tr} \left( \frac{\partial F}{\partial \Gamma_t} \right) \leq \frac{\eta}{2} \text{Tr} \left( \frac{\partial F}{\partial \Gamma_t} \right)
\]

where the overline \( \overline{\text{tr}} \) denotes the expectation operation \( \overline{\text{tr}} \), \( \sigma^2_\Gamma \) and \( \sigma^2_\delta \) are the variance bounds for the stochastic gradients of \( \Gamma \) and \( \delta_t \) respectively. To estimate the upper bound for \( \overline{\text{tr}} \left( \frac{\partial F}{\partial \Gamma_t} \right) \),
we utilize convexity and Lipschitzness as follows:

$$
E \left\| \frac{\partial F}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 = \frac{1}{n^2} \sum_{i=1}^n \left( \frac{\partial F_i}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t(i)} \right)^2 \leq \frac{1}{n} \sum_{i=1}^n \left\| \frac{\partial F_i}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t(i)} \right\|_F^2 \\
\leq \frac{L^2}{n} \sum_{i=1}^n \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2.
$$

Therefore, we re-write the lower bound (16) as

$$
-\eta \left(1 - \frac{L \eta}{2} \right) E_t \left\| \frac{\partial F}{\partial \delta_t(i \cap k)} \right\|_F^2 - \frac{\eta}{2} E_t \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 \eta \left(1 - \frac{L \eta}{2} \right) E_t \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 + \frac{\eta L^2}{2n} E_t \sum_{i=1}^n \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2 + \frac{L \eta^2 (\sigma^2_f + \sigma^2 \gamma)}{2K}.
$$

Summing the inequality above for all time-steps \( t = 0, \ldots, T \), we get

$$
-\eta \left(1 - \frac{L \eta}{2} \right) \sum_{t=0}^T E_t \left\| \frac{\partial F}{\partial \delta_t(i \cap k)} \right\|_F^2 - \frac{\eta}{2} \sum_{t=0}^T E_t \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 + \frac{\eta L^2}{2n} \sum_{t=0}^T E_t \sum_{i=1}^n \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2 + \frac{L \eta^2 (\sigma^2_f + \sigma^2 \gamma) T}{2K}.
$$

(17)

The main challenge however now becomes bounding the term with \( \sum_{t=0}^T E_t \sum_{i=1}^n \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2 \). Bounding it requires to derive another lower bound and using an available result. First re-write the

$$
E \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2
$$

by utilizing Frobenius-norm properties:

$$
E \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 = E \left\| \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 + \frac{\partial f}{\partial \Gamma_t} \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2
$$

$$
= E \left\| \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 + E \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 + 2E \left\langle \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t}, \frac{\partial f}{\partial \Gamma_t} \right\rangle
$$

$$
= E \left\| \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 + E \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2
$$

Then bound each term as follows:

$$
\leq n \sigma^2 + E \sum_{i=1}^n \left( \frac{\partial F_i}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t} \right)^2 + \left( \frac{\partial F}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t} \right)^2 + \left( \frac{\partial F}{\partial \Gamma_t} - \frac{\partial F}{\partial \Gamma_t} \right)^2 \leq n \zeta^2 + n \zeta^2 + n \zeta^2,
$$

where first term is bounded by its stochastic gradient variance. To bound the second term \( E \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 \) above, first bound the term with sum of the individual components’ norm and then use add-subtract trick used with \( \frac{\partial F}{\partial \Gamma_t} \) and \( \frac{\partial F}{\partial \Gamma_t} \). Then, use the facts \( \sum_{i=1}^n E \left\| \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 \leq L^2 E \sum_{i=1}^n \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2 \) (an intermediate result derived above) and \( \sum_{i=1}^n E \left\| \frac{\partial f}{\partial \Gamma_t} - \frac{\partial f}{\partial \Gamma_t} \right\|_F^2 \leq n \zeta^2 \), while \( \zeta \) being the spectral gap of matrix \( M \). Then, from [64], we have

$$
\sum_{t=1}^T \frac{1}{n} E \left\| \Gamma_t - \Gamma_t(i) \right\|_F^2 \leq \frac{2}{(1 - \zeta^2)} \sum_{t=1}^T \frac{1}{n} \left\| \frac{\partial f}{\partial \Gamma_t} \right\|_F^2.
$$
with the previously derived bound over \( \sum_{t=1}^{T} \eta T \| \frac{\partial f}{\partial \Gamma_t} \|_F^2 \) bound leading to another intermediate bound

\[
\left(1 - \frac{2\eta^2 L^2}{(1 - \zeta)^2}\right) \sum_{t=0}^{T} \sum_{i=1}^{n} \mathbb{E} \left\| \Gamma_t - \Gamma_t^{(i)} \right\|_F^2 \leq \frac{2\eta^2}{(1 - \zeta)^2} \left(n\sigma_t^2 + n\zeta^2\right) + n \sum_{t=1}^{T} \mathbb{E} \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 \\

\sum_{t=0}^{T} \sum_{i=1}^{n} \mathbb{E} \left\| \Gamma_t - \Gamma_t^{(i)} \right\|_F^2 \leq C_1 \eta^2 n \left(\sigma_t^2 + \zeta^2\right) T + \sum_{t=1}^{T} \mathbb{E} \left\| \frac{\partial F}{\partial \Gamma_t} \right\|_F^2 \tag{18}
\]

where \( C_1 \) being a function of the spectral gap.

Finally, combining (17) with the intermediate bound (18) together first, bounding the Frobenius norms, and dividing both sides by \( T \) (to average in the end), we get the desired lower bound

\[
\frac{2[F(x_0) - F_{\text{inf}}]}{\eta T} + \frac{\eta L\sigma^2}{K} + \eta^2 L^2 \sigma^2 \left(\frac{1 + e^2}{1 - \zeta^2} - 1\right)
\]

where \( \eta \) is the learning rate that satisfies the given conditions. \( x_0 = \{ \Gamma_0, \delta_0^{(1:K)}, \Omega_0^{(1:K)} \} \) is the initial starting point and \( \sigma^2 = \sigma_\Gamma^2 + \sigma_\delta^2 \) is the total variance bound over stochastic gradients of \( \Gamma \) & \( \delta \).

Our work differs from [68] in that it does not provide adequate theoretical analysis and empirical evaluation for federated learning.