

# Ensemble Attention Distillation for Privacy-Preserving Federated Learning

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## Abstract

We consider the problem of Federated Learning (FL) where numerous decentralized computational nodes collaborate with each other to train a centralized machine learning model without explicitly sharing their local data samples. Such decentralized training naturally leads to issues of imbalanced or differing data distributions among the local models and challenges in fusing them into a central model. Existing FL methods deal with these issues by either sharing local parameters or fusing models via online distillation. However, such a design leads to multiple rounds of inter-node communication resulting in substantial bandwidth consumption, while also increasing the risk of data leakage and consequent privacy issues. To address these problems, we propose a new distillation-based FL framework that can preserve privacy by design, while also consuming substantially less network communication resources when compared to the current methods. Our framework engages in inter-node communication using only publicly available and approved datasets, thereby giving explicit privacy control to the user. To distill knowledge among the various local models, our framework involves a novel ensemble distillation algorithm that uses both final prediction as well as model attention. This algorithm explicitly considers the diversity among various local nodes while also seeking consensus among them. This results in a comprehensive technique to distill knowledge from various decentralized nodes. We demonstrate the various aspects and the associated benefits of our FL framework through extensive experiments that produce state-of-the-art results on both classification and segmentation tasks on natural and medical images.

## 1. Introduction

Modern deep learning algorithms rely on massive annotated datasets for many practical applications [63, 21, 18]. In most cases, however, this data is physically located across multiple disparate locations and regulated by dif-

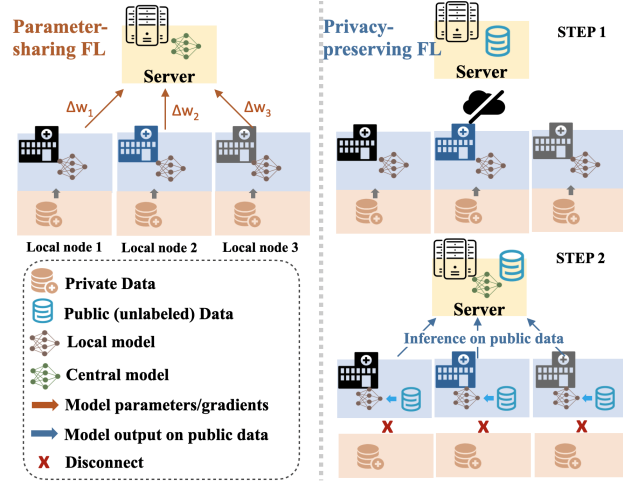


Figure 1. A schematic illustration of the proposed privacy-preserving federated learning framework compared to traditional update or parameter-sharing based federated learning frameworks. Traditional FL frameworks transfer gradients or parameter updates produced with private data from local nodes to a server, risking privacy leakage. Our framework only transfers products of unlabeled public data.

ferent entities. This results in the challenges of centralizing the physically dispersed data, with the primary concerns being privacy and network bandwidth issues. Consequently, federated learning (FL) [46, 54, 17] has emerged as an important topic where a single centralized model is trained in a distributed, decentralized fashion using model fusion/distillation techniques.

While some similarities exist, there are many more unique challenges that make FL substantially different from distributed learning. First, privacy is a critical consideration, and maintaining it is of utmost importance, particularly for applications such as healthcare [44, 3]. Second, one needs to be able to train centralized models efficiently and not get bogged down by network communication issues. Communication bandwidth may be quite pronounced depending on the task (e.g. image and video applications) and quantity of data (e.g., model parameters) being shared

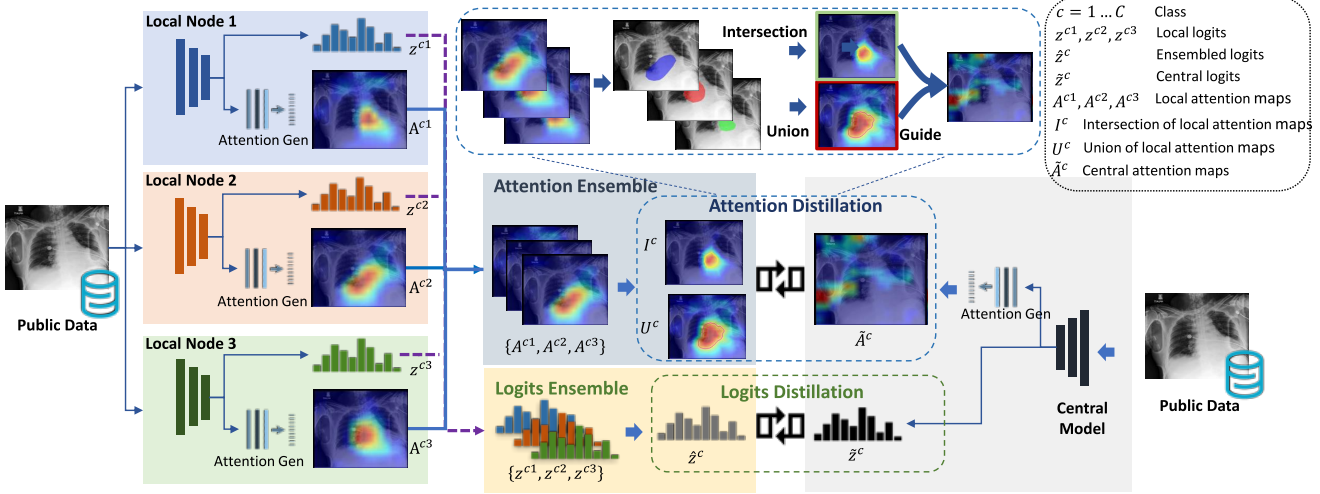


Figure 2. Overview of the proposed FedAD framework. Attention Avg + Logits Avg

among local nodes. Finally, given that raw data typically resides locally, the data collection and preprocessing mechanisms may vary substantially among the local nodes, leading to a situation where common assumptions such as independent and identically distributed sampling do not hold.

The current state-of-the-art FL techniques approach the aforementioned issues by repeatedly sharing local model parameters or their gradients during the training process [34, 47, 12, 24, 58, 49, 19, 30, 8]. However, for many applications, these parameter-based communication methods suffer from impractical network bandwidth overhead, are limited only to models with homogeneous architectures, and more importantly, have many known security weaknesses [4]. While some methods have taken a step towards data protection in medical imaging [26, 27], there are also counter arguments in the literature that show local private data can get exposed as a result of using publicly shared gradients in the FL technical pipeline [61].

Distillation-based techniques that aggregate locally-computed logits [16, 22, 4, 28, 62] form the basis for another line of work for building central models from multiple local models, helping eliminate the need for each local model to follow the same architecture. While some recent methods distill with public data to get around data privacy issues [22, 4], they assume both the public and private data are sampled from the same underlying distribution. This assumption invariably exposes private data to security risks and attacks. While the recently proposed FedDF [28] method provides some relaxations (e.g., public data can be unlabeled and domain robust, i.e., sampled from another domain), it still exchanges model parameters recursively, resulting in privacy vulnerabilities due to model memorization [61, 4]. Despite the known bottleneck of communication in FL, all the above methods jointly (online) optimize the central and local models by synchronizing lo-

cal inferred predictions. This approach requires a high degree of synchronization and communication bandwidth. It is clear that these co-distillation methods require recursive communication primarily because these methods ensemble with dark knowledge, such as averaging to soften labels, leaving structural knowledge unexplored. In other words, in addition to distilling the final *what* (i.e., logits), using more feature information depicting the *why* (e.g., attention maps) should lead to improved performance and efficiency, which is largely ignored by the current state-of-the-art.

In this work, we present a new distillation-based federated learning framework to address the aforementioned issues (Figure 1). First, our framework presents stronger privacy guarantees of local data by only using model outputs of unlabeled public data during distillation *without any exchange of local model parameters or gradients*. This, by design, eliminates the vulnerabilities identified by prior work. Second, in our framework, local models are fully trained and then distilled to the central server, in contrast to prior works [28, 4, 22] that synchronously update local models through online distillation.

Our key insight is that such well-trained local models, as opposed to incremental snapshots of “half-baked” models [28, 4, 22], provide more structural knowledge about their expertise. This design choice immediately enables top-down class-specific attention maps that capture the fully-trained local model’s reasoning process (note that for methods that do online distillation, this would not be possible since their attention maps would be incomplete due to incremental training). We ensemble local knowledge with both predicted logits and these attention maps, capturing each local model’s final output as well as the underlying reasoning process. We also use these attention maps to capture the knowledge diversity across models and reach a consensus to effectively coordinate local expertise in the FL

paradigm. This in-depth ensemble strategy enables our federated distillation to be completed in an offline fashion in a single round (which we call one-shot), helping keep local model training independent and asynchronous. This results in a FL framework that is substantially more efficient and flexible when compared to prior art.

To demonstrate efficacy, we conduct extensive experiments on CIFAR10/100 and large-scale chest x-ray datasets. We also show our framework is flexible to be used in other tasks by conducting preliminary proof-of-concept experiments on the segmentation tasks, where we demonstrate the state-of-the-art privacy/performance trade-offs compared to prior methods.

To summarize, our key contributions are below:

- We propose a one-shot federated learning framework with one-way distillation to explicitly preserve the privacy of local data by only distilling model outputs on unlabeled and domain robust public data.
- Our framework addresses the communication inefficiencies of prior work by communicating high-level logits and model-agnostic attention maps.
- We introduce a seminal distillation algorithm that aggregates structural knowledge with explicit balance between both local model diversity as well as consensus to deal with the inherent heterogeneity of decentralized federated learning.
- We show that the proposed framework can be extended to other applications such as semantic segmentation with evaluations on Cityscapes and BraTS dataset.

## 2. Related Work

Our proposed FL framework is related to areas such as knowledge transfer and the type of FL algorithms and how they deal with issues such as privacy. Here, we briefly review methods that are relevant to these topics.

**Knowledge Transfer.** Recent methods can be categorized based on their ensemble strategy and the kind of information distilled to the student. Following the work of Hinton et al. [11], there has been much progress in model ensemble, with a particular focus on the student-teacher learning paradigm [43]. These techniques aggregate the knowledge of multiple teachers before distilling it to the student model. This has led to a variety of aggregation schemes, with gate learning being quite popular in the supervised setting [43, 1, 53, 48, 59]. In semi-supervised and self-supervised scenarios, techniques based on relative sample similarity have been proposed [56, 52]. Going beyond the use of soft label distillation [11], recent approaches have explored the transfer of structural knowledge such as intermediate feature representations [41], Gram matrices [55],

maximum mean discrepancy [14], or mutual information [40]. Finally, there have also been some attempts at combining both structure as well as label ensemble strategies, with FEED [39] and knowledge flow [29] being notable examples. In contrast to these techniques, our feature-level ensemble method is label-free, model agnostic, and can also be used in heterogeneous knowledge distillation scenarios.

**Federated Learning.** Most existing FL methods are either parameter-based or distillation-based. In parameter-based FL methods, each local model shares its parameters/gradients with the central server after every round of local training on its local data, following which the central server aggregates them, e.g., by averaging [34]. This result is then shared by the central server with the local nodes, which in turn update their corresponding local model and proceed with the next training round. This process is then repeated until the stopping criterion is met. A number of extensions to FedAVG [34, 49, 25, 13] have been proposed with new aggregation schemes such as momentum [12] or local weighting [25, 13], or new local training strategies such as the use of proximal term [24] or control variations[19]. However, such parameter/gradient sharing can certainly be a straightforward way of information exchange, it is highly susceptible to privacy leakage and stealth attacks, as also demonstrated elsewhere [61, 4].

On the other hand, distillation-based methods exchange model outputs (on local private data) [60, 45] rather than the parameters, leading to growing concerns on the privacy of local data. While some methods address this issue by distilling on public data [16, 22, 4], they often select public data based on some prior knowledge of private data, leading to similar security vulnerabilities as above. While the recently proposed FedDF [28] attempts to address this issue, it is quite inefficient (e.g., high network bandwidth) due to the iterative exchange of models over hundreds of rounds, which in turn also leads to more susceptibility to stealth attacks and hence privacy concerns. The framework most similar to ours is PATE [38], where all local models can be trained independently without inter-institutional communication, and the central model is trained with hard pseudo labels voted by local models. In contrast, our method preserves privacy by exchanging the soft prediction of domain robust public data, and further exploiting in-depth feature level information for high efficient communication.

**Privacy Concerns for FL.** Parameter-based FL methods have shown to be highly susceptible to privacy leakage [61, 4]. Furthermore, as noted above, some distillation-based FL methods are also at privacy risk when recursively model exchanges are involved [28]). Utilizing unlabeled public data during distillation, while also restricting the local nodes’ access to the server model, has been shown to be more resilient to attacks and guarantee privacy [38, 9]. Therefore, our framework explicitly protects local private

data by using one-shot communication, and public data from different domain or distributions.

### 3. Approach

We first begin with a formal introduction to the problem before discussing the design of our proposed framework, called Ensemble Attention Distillation federated learning (FedAD).

#### 3.1. Problem Definition

Let there be  $K$  local/private nodes with each node playing host to labeled (private) dataset  $\mathcal{D}_k = \{(x_k^i, y_k^i) | i = 1, \dots, |\mathcal{D}_k|\}$ , where  $y_k \in \mathcal{C}_k$ ,  $\mathcal{C}_k$  is the set of existing classes in dataset  $\mathcal{D}_k$ , and  $\mathcal{C}_k \subset \{1, \dots, C\}$  ( $C$  is the overall number of classes across all local nodes). The shared public dataset  $\mathcal{D}_0 = \{x_0^i | i = 1, \dots, |\mathcal{D}_0|\}$  can be accessed by any local node and can unlabeled or labeled.

As illustrated in Figure 2, in the first stage of FedAD, the model at each local node  $k$  is trained using the corresponding private data  $\mathcal{D}_k$  ( $\theta_k$  represents these local model parameters after local training). Since our proposed FedAD is agnostic to the type of network architecture, each local neural network can be customized to have its own architecture, helping adapt the model to its local data distribution.

In the second stage, we disconnect local datasets from the network to minimize security risks and protect privacy. Instead, the central public dataset  $\mathcal{D}_0$  is used at each local node as part of a one-way knowledge distillation framework from the local models to the central server. The set of local models and the central model form a teacher-student knowledge transfer setup where the teacher is a local-model ensemble (one local model at each local node).

#### 3.2. Logits Ensemble Distillation

During each distillation step  $t$ , we sample (randomly) a subset  $\mathcal{K}_t$  of models from all local nodes. This subset comprises a fraction  $\gamma$  of all local models. The standard distillation pipeline uses the Kullback-Leibler divergence to ensemble all teachers' soft labels:

$$\mathcal{L} = \sum_c p(y=c) \log \frac{p(y=c)}{q(y=c)}, \quad (1)$$

where  $p$  is the sample's probability being class  $c$  for the teacher model and  $q$  is the corresponding value for the student model. Given a sample  $x_0$  from the public dataset, let  $z^{ck} = f(x_0, \theta_k, c)$  be its logits (on class  $c$  from local model at node  $k$ ) and  $\tilde{z}^c = f(x_0, \theta_s, c)$ , where  $c \in \{1, \dots, C\}$  be the corresponding central model output. Given these notations, the standard ensemble strategy  $\tilde{z}^c = \frac{1}{|\mathcal{K}_t|} \sum_{k \in \mathcal{K}_t} z^{ck}$  uses the average operation on logits from all teachers' with

a softmax activation as:

$$p(y=c) = \frac{\exp(\tilde{z}^c / \tau)}{\sum_c \exp(\tilde{z}^c / \tau)}, q(y=c) = \frac{\exp(\tilde{z}^c / \tau)}{\sum_c \exp(\tilde{z}^c / \tau)} \quad (2)$$

It has been shown in prior work [11] that minimizing Eq. 1 with a high temperature parameter  $\tau$  is the same as to minimizing the  $\ell_2$  error between student and teacher logits, hence establishing a relationship between matching logits and the cross entropy objective. Without any loss of generality, let the activation be  $q^c = \sigma(\tilde{z}^c)$  and  $p^c = \sigma(\tilde{z}^c)$ .

Since the FL setting involves extreme heterogeneity, the standard distillation process discussed above is not directly suitable. This is because of the inability to tackle a general scenario where all local models do not share the same set of target classes. To explicitly consider this in our proposed framework, during our distillation procedure, we introduce a new variable, called the importance weight  $\omega$ , for each local model so as to capture the corresponding local data distribution used in the initial training step:

$$\tilde{z}^c = \sum_{k \in \mathcal{K}_t} \omega_k^c z^{ck}, \omega_k^c = \frac{N_k^c}{\sum_{k \in \mathcal{K}_t} N_k^c}, \quad (3)$$

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where  $N_k^c = \sum_{i=1}^{|\mathcal{D}_k|} (y_k^i = c)$  is the number of  $c$ -class samples used in training the local model at node  $k$ .

Following the observation above, we set  $\tau \rightarrow \infty$  and rewrite the logits learning objective as:

$$\mathcal{L}_{\text{logits}}(\tilde{\mathbf{z}}, \hat{\mathbf{z}}) = \frac{1}{C} \|\tilde{\mathbf{z}} - \hat{\mathbf{z}}\|, \quad (4)$$

MSE

where  $\tilde{\mathbf{z}} = [\tilde{z}^1, \dots, \tilde{z}^C]$  and  $\hat{\mathbf{z}} = [\hat{z}^1, \dots, \hat{z}^C]$ . Note that the aforementioned methods can be easily applied pixel-wise, making it readily applicable for segmentation tasks.

**Multi-label Classification:** We next show our proposed method can be easily extended to multi-class classification (to go with single class above). Here, we denote the private data as  $\mathcal{D}_k = \{(x_k^i, y_k^i) | i = 1, \dots, |\mathcal{D}_k|\}$  with  $y_k^i \in \{-1, 0, 1\}^C$  where -1, 0, and 1 represent unknown, negative and positive for class  $c$ . Finally, the adaptation requires a few more changes. Instead of softmax, we use the sigmoid  $p^c = \sigma(\tilde{z}^c)$  and  $q^c = \sigma(\tilde{z}^c)$  as the activation function. Next, we modify Eq. 3 by defining  $N_k^c = \sum_{i=1}^{|\mathcal{D}_k|} (y_k^i(c) = 1)$ .

#### 3.3. Attention Ensemble Distillation

Logits distillation discussed above essentially captures the divergence between the output vectors of teacher and student models. However, comparing only the output vectors in this fashion only ensures the outputs can match and does not necessarily mean the underlying structural knowledge, or the model's reasoning can be transferred. We posit that such knowledge, e.g., intermediate feature representations of models, can result in more accurate distillation of

knowledge, more so in cases such as FL with its high degree of heterogeneity in local data sources. While this can seem intuitive, there are practical challenges in implementing this method. Specifically, transferring mostly bulky matrices such as intermediate feature representations is quite resource intensive, e.g., from the perspective of network bandwidth burden, and furthermore, relies on restrictive requirements such as identical network architecture among the student and teacher models.

On the other hand, representations such as class-specific top-down attention maps generated with methods such as Grad-CAM [42] have been widely used in providing location cues for weakly supervised semantic segmentation, and has been shown to be effective in providing more precise and efficient guidance to the learning process [23]. Furthermore, recent studies such as [31] extend Grad-CAM to generic embedding models, making it applicable to any tasks that use or need a fully-connected feature extractor. We argue that such top-down model interpretations can transfer knowledge in a more efficient (instead of full feature tensors) and effective (instead of just the output vectors) way, without sacrificing communication efficiency or risking privacy leakage.

**Sampling by importance.** Let the attention map from local node  $k$  for class  $c$  be  $\mathbf{A}^{ck} \in \mathbb{R}^{HW}$ , where  $H, W$  are the size of attention maps, and the complete set of attention maps from all local nodes be  $\mathcal{A} = \{\mathbf{A}^{ck} | k \in \mathcal{K}_t, c = 1, \dots, C\}$ . Due to the high degree of heterogeneity among local nodes, we weight attention maps from each local node differently based on the sample distribution in its corresponding private dataset. Similar to FedVC [13] that resamples local nodes based on the probability proportional to local data size, we sample the attention maps based on the class-specific importance weight  $\omega_k^c$  in Eq. 3. That is, we independently decide on whether to use  $\mathbf{A}^{ck}$  with the probability  $\hat{\omega}_k^c$  in each batch.

To ensure at least  $\hat{K}$  samples are selected each time, we set  $\hat{\omega}_k^c = \max(1, \frac{\omega_k^c}{\text{Top}_{\hat{K}}(\omega_k^c)})$ , where  $\text{Top}_{\hat{K}}(\omega_k^c)$  denotes the  $\hat{K}$ -th maximum value among  $\{\omega_k^c | k \in \mathcal{K}_t\}$ . In our experiments, we set  $\hat{K} = 2$  by default. At each sample step  $t$ , we obtain a set of selected local indexes  $\hat{\mathcal{K}}_t^c$  for class  $c$  as:

$$\hat{\mathcal{K}}_t^c = \text{sample}(\{\hat{\omega}_k^c | k \in \mathcal{K}_t\}), \quad (5)$$

where  $\hat{\mathcal{K}}_t^c \subset \mathcal{K}_t$  and  $|\hat{\mathcal{K}}_t^c| \geq \hat{K}$ . Thus, the selected set of attention maps with respect to class  $c$  is  $\mathcal{A}_t^c = \{\mathbf{A}^{ck} | k \in \hat{\mathcal{K}}_t^c\}$ . To simplify notions, from this point on, we refer to  $\mathcal{A}_t^c$  as  $\mathcal{A}^c = \{\mathbf{A}^{ck} | k \in \hat{\mathcal{K}}_t^c\}$ .

**Attention bound constraint.** While there have been a few recent efforts in enforcing constraints directly on gradient-based attention maps [7, 50, 31], they cannot directly be applied to our scenario with an ensemble of attention maps produced by highly heterogeneous models.

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**Algorithm 1** FedAD on  $K$  local nodes with  $C$  classes.

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**Input:** Labeled private data  $\{\mathcal{D}_k\}$ , unlabeled public data  $\mathcal{D}_0$ , central model  $\theta_s$ , local models  $\{\theta_k\}$ ,  $T$  distillation steps, batch-size  $S$ , sample fraction  $\gamma$ .

**Local Training:** Train each local model  $\theta_k$  with  $\mathcal{D}_k$  Client train

**for** each distillation step  $t = 1, \dots, T$  **do**

- $\mathcal{K}_t \leftarrow$  random subset ( $\gamma$  fraction) from  $K$  locals
- $\mathbf{x}_0 \leftarrow$  a batch of public data from  $\mathcal{D}_0$  with size  $S$
- for** each local  $k \in \mathcal{K}_t$  **do** KD based on public data
- $\mathbf{z}^k, \mathbf{A}^k \leftarrow f(\mathbf{x}_0; \theta_k)$
- end for**
- $\widehat{\mathbf{z}} \leftarrow$  ensemble  $\{\mathbf{z}^k | k \in \mathcal{K}_t\}$  Weighted Logits Avg
- for** each class  $c = 1, \dots, C$  **do**
- $\widehat{\mathcal{K}}_t^c \leftarrow$  sample a subset from  $\mathcal{K}_t$  Eq. 5
- $\mathbf{I}^c, \mathbf{U}^c \leftarrow$  ensemble  $\mathcal{A}^c = \{\mathbf{A}^{ck} | k \in \widehat{\mathcal{K}}_t^c\}$  Eq. 6
- end for** min and max Attention
- $\widetilde{\mathbf{z}}, \widetilde{\mathbf{A}} \leftarrow f(\mathbf{x}_0, \theta_s)$
- Update:  $\theta_s \leftarrow \theta_s - \frac{1}{S} \nabla_{\theta_s} \mathcal{L}(\widetilde{\mathbf{z}}, \widehat{\mathbf{z}}, \widetilde{\mathbf{A}}, \mathbf{I}, \mathbf{U})$  Eq. 10

**end for**

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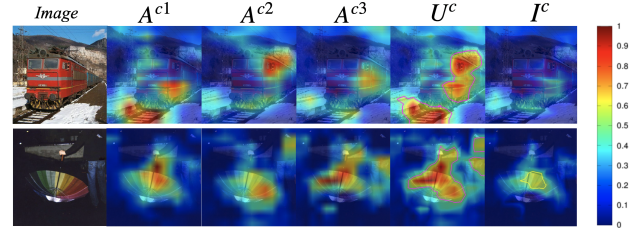


Figure 3. Illustration of Eq. 6. Suppose  $k \in \widehat{\mathcal{K}}_t^c = \{1, 2, 3\}$  with respect to class  $c$ ,  $\mathbf{A}^{ck}$  indicate local attention maps,  $\mathbf{U}^c$  and  $\mathbf{I}^c$  indicate union attention maps and intersection attention maps respectively.

Specifically, the training objective based on the  $\ell_1$  or  $\ell_2$  functions in these methods seek to enforce exactly the same activation strength in each location of the attention map, which in our case may likely introduce and amplify noise from local models. To achieve consensus while also maintaining the diversity that is inherent among the local nodes, we design a novel attention bound constraint based on the intersection and union of local attention maps with respect to the same input data and class output. Given the set of local attention maps  $\mathcal{A}^c = \{\mathbf{A}^{ck} | k \in \widehat{\mathcal{K}}_t^c\}$ , we denote  $\mathbf{I}^c, \mathbf{U}^c \in \mathbb{R}^{HW}$  as the intersection and union among all the local attention maps  $\mathcal{A}^c$  with respect to class  $c$ , respectively. Let  $h, w$  be the pixel index, we have

$$I_{hw}^c = \min_{k \in \widehat{\mathcal{K}}_t^c} A_{hw}^{ck}, U_{hw}^c = \max_{k \in \widehat{\mathcal{K}}_t^c} A_{hw}^{ck}, \quad (6)$$

where  $\mathbf{I}^c$  denotes a consensus on the high-response region among all the local attention maps, that has a high probability to comprise the object of interest. While  $\mathbf{U}^c$  considers all of the high-response regions among the local attention maps, it also preserves diversity of ‘expertise’ among the local models by means of the union operation. Figure 3 shows an example of such attention maps. Given the at-

tention map  $\mathbf{A}$ , let  $T(\cdot)$  be a soft-masking operation (with sigmoid) to have all values lie between 0 and 1 [23]:

$$T(\mathbf{A}) = \frac{1}{1 + \exp(-\rho(\mathbf{A} - b))}. \quad (7)$$

For simplicity, we denote  $\tilde{\mathbf{A}}^c$  as the attention map generated by central model with respect to class  $c$ . Considering the attention intersection  $\mathbf{I}^c$  as the consensus achieved by all locals, we enforce the high-response region in  $\tilde{\mathbf{A}}^c$  to explicitly include that of  $\mathbf{I}^c$  using our proposed attention intersection loss, defined as:

$$\mathcal{L}_{\text{inter}}(\tilde{\mathbf{A}}, \mathbf{I}) = -\frac{1}{C} \sum_c \frac{\sum_{hw} I_{hw}^c \cdot T(\tilde{A}_{hw}^c; \rho_1, b_1)}{\sum_{hw} I_{hw}^c}, \quad (8)$$

where  $\tilde{\mathbf{A}} = [\tilde{\mathbf{A}}^1, \dots, \tilde{\mathbf{A}}^C]$ ,  $\mathbf{I} = [\mathbf{I}^1, \dots, \mathbf{I}^C]$ ,  $\rho_1 = 10$ , and  $b_1 = 0.6$ . With the attention union  $\mathbf{U}^c$  including all the high-response regions among all locals, we enforce the high-response region in  $\tilde{\mathbf{A}}^c$  to be explicitly inside that of  $\mathbf{U}^c$  using our attention union loss  $\mathcal{L}_{\text{union}}$ . The intuition here we seek each high-response pixel in the  $\tilde{\mathbf{A}}^c$  to have support from at least one local model to take model diversity into account. The  $\mathcal{L}_{\text{union}}$  loss is defined as:

$$\mathcal{L}_{\text{union}}(\tilde{\mathbf{A}}, \mathbf{U}) = -\frac{1}{C} \sum_c \frac{\sum_{hw} \tilde{A}_{hw}^c \cdot T(U_{hw}^c; \rho_2, b_2)}{\sum_{hw} \tilde{A}_{hw}^c}, \quad (9)$$

where  $\mathbf{U} = [\mathbf{U}^1, \dots, \mathbf{U}^C]$ ,  $\rho_2 = 10$ , and  $b_2 = 0.3$ .

Our method optimizes the learning of the central model so that  $\tilde{\mathbf{A}}^c$  is encouraged to activate in the pixels which are activated in  $\mathbf{I}^c$ , and penalized for activating in the pixels not activated in  $\mathbf{U}^c$ . The attention bound constraint combines the two constraints balancing the local consensus and diversity. This is a relaxed constraint as it is capable of tolerating up to  $|\hat{\mathcal{K}}_t^c| - 1$  incorrect/biased attention maps, and maintaining high robustness to outliers, which is particularly important to tackle with heterogeneity in the FL setting. The overall loss for optimization is

$$\mathcal{L} = \mathcal{L}_{\text{logits}}(\tilde{\mathbf{z}}, \hat{\mathbf{z}}) + \mathcal{L}_{\text{inter}}(\tilde{\mathbf{A}}, \mathbf{I}) + \mathcal{L}_{\text{union}}(\tilde{\mathbf{A}}, \mathbf{U}). \quad (10)$$

**Segmentation.** Crucially, our framework by no means is limited to just classification problems. The distillation algorithm of our proposed FedAD framework can be used to aggregate knowledge for other kinds of tasks as well, e.g., segmentation. In this case, we aggregate predicted masks  $\mathbf{z}^{ck}$  in Eq. 5 and  $N_k^c$  is counted pixel-wisely. Thus the logits in Eq. 4 can be written as  $\tilde{\mathbf{z}} = [\tilde{z}^1, \dots, \tilde{z}^C]$  and  $\hat{\mathbf{z}} = [\hat{z}^1, \dots, \hat{z}^C]$ . The other modification is we obtain the class-specific attention map of local model through activation:  $\mathbf{A}^{ck} = \sigma(\mathbf{z}^{ck})$ , where  $\sigma$  can be softmax or sigmoid. More details are in supplementary materials.

The overall process is explained in Algorithm 1.

Accuracy(%)	Shared Param.	CIFAR-10		CIFAR-100	
		$\alpha = 1$	$\alpha = 0.1$	$\alpha = 1$	$\alpha = 0.1$
FedAvg [34]	Y	78.57 ±0.22	68.37 ±0.50	42.54 ±0.51	36.72 ±1.50
FedProx [24]	Y	76.32 ±1.95	68.65 ±0.77	42.94 ±1.23	35.74 ±1.00
FedAvgM [12]	Y	77.79 ±1.22	68.63 ±0.79	42.83 ±0.36	36.29 ±1.98
FedDF [28]	Y	80.69 ±0.43	71.36 ±1.07	47.43 ±0.45	39.33 ±0.03
FedMD [22]	N	80.37 ±0.37	69.23 ±1.31	45.83 ±0.58	39.86 ±0.78
<i>Standalone</i>	N	61.11 ±24.90	28.99 ±27.24	27.49 ±14.76	16.31 ±15.75
FedAD	N	<b>82.48</b> ±0.21	<b>73.11</b> ±1.25	<b>50.34</b> ±0.33	<b>48.43</b> ±1.01

Table 1. Results on CIFAR-10 and CIFAR-100 with ResNet-8 when  $\gamma=0.4$  and  $K=20$ , comparing our FedAD with several existing parameter-based [34, 24, 12, 28] and distillation-based [22] FL methods. *Standalone*: mean/std accuracy of all local models.

Accuracy (%) ↑	$\alpha = 1$	FedDF [28]	FedAD (Ours)			
		$\gamma = 0.4$	$\gamma = 0.4$	$\gamma = 0.8$	$\gamma = 1$	
		70.69	82.48	83.16	<b>83.68</b>	
Bandwidth (GB) ↓			29.4	<b>10.1</b>	22.1	
					27.6	

Table 2. FL communication efficiency on CIFAR-10. In FedDF, both logits and parameters are of type float64, whereas attention maps are of type float16.

aggregation scheme	[28, 13]	Eq. 3	Eq. 3	Eq. 3
logits distillation	$\tau = \infty$	$\tau = 3$	$\tau = \infty$	$\tau = \infty$
attention distillation	✗	✗	✗	✓
Accuracy(%)↑	65.73	67.81	72.67	<b>73.11</b>

Table 3. Ablation study on CIFAR-10 with ResNet-8,  $\gamma=0.4$ ,  $K=20$ , and  $\alpha=0.1$ . We compare our aggregation scheme (Eq. 3) with the existing aggregation schemes [28, 13], and different temperatures for logits distillation, we only list the result with typical value  $\tau=3$  [11].

## 4. Experiments

We conduct a number of experiments on natural images/medical images for both classification/segmentation tasks. Additionally, we experiment on text classification tasks and provide the results in the supplementary.

### 4.1. Classification

In constructing local training sets, we use heterogeneous data splits using a Dirichlet distribution as in prior works [12]. The value of  $\alpha$  controls the degree of non-IID-ness: a plus infinite  $\alpha$  indicates identical local data distribution, and a smaller  $\alpha$  indicates higher non-IID-ness. While distilling knowledge with task-relevant data from the same domain is ideal, FedAD is compatible with using public data from different domains. Here, we thus consider a more general heterogeneous setting where the public data come from different domains. Note we save augmentation seeds locally and transmit the predictions of each seed during distillation

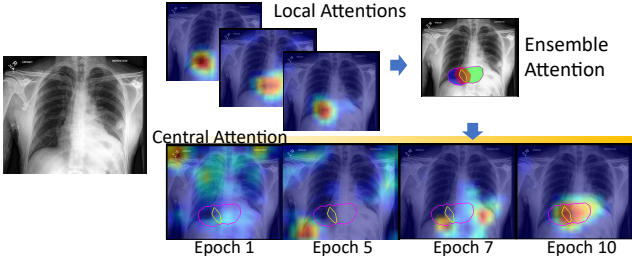


Figure 4. One example showing how ensemble attention can effectively guide the central model focus on the right region.

Methods	Distill	Test	mAUC(%)↑	Distill	Test	mAUC(%)↑
FedMD[22]	C <sup>3</sup>	C	71.23	X <sup>3</sup>	X	77.66
	C <sup>3</sup> +X <sup>3</sup>	C	70.97	C <sup>3</sup> +X <sup>3</sup>	X	77.02
FedAD	C <sup>3</sup> alone	C	75.08 ± 4.44	X <sup>3</sup> alone	X	73.67 ± 7.31
	C <sup>3</sup>	C	75.17	X <sup>3</sup>	X	82.62
	C <sup>3</sup> +X <sup>3</sup>	C	75.55	C <sup>3</sup> +X <sup>3</sup>	X	<b>82.65</b>
Centralized	C <sup>c</sup>	C	79.73	X <sup>c</sup>	X	82.58
	C <sup>c</sup> +X <sup>c</sup>	C	<b>80.47</b>	C <sup>c</sup> +X <sup>c</sup>	X	80.91

Table 4. Results on CXR14 and CheXpert when private data is inside/cross domain. We split each dataset with  $K_d = 3$ ,  $\alpha = 1, \gamma = 1$ . ‘C’ refers to CXR14, ‘X’ refers to CheXpert, and  $\{C, X\}^n$  denotes  $n$  local nodes with data from the dataset, where an additional *alone* denotes mean/std of all local models.  $\{C, X\}^c$  denotes centralized training all local data.

for classification tasks.

#### 4.1.1 CIFAR-10/100 Classification

We first consider CIFAR-10/100 [20]. For a fair comparison, we follow the same setup as in FedDF [28], using CIFAR-10 and CIFAR-100 as private datasets, and CIFAR-100 and a downsampled version of ImageNet ( $32 \times 32$ ) as the corresponding public datasets for CIFAR-10 and CIFAR-100 respectively. We use test datasets on the central server and report the average accuracy over three different random seeds. Implementation details are provided in the supplementary material.

The comparison in Table 1 shows that our method outperforms current state-of-the-art methods by a large margin, particularly for the higher non-IID-ness scenario. Next, we compare the accuracy and communication efficiency at different sample fractions  $\gamma = \{0.4, 0.8, 1.0\}$  with the current state-of-the-art method [28] in Table 2, where our method achieves higher accuracy on CIFAR-10 while consuming significantly lower communication bandwidth. We observed that the distillation result is not sensitive to data precision, which has also been shown in prior work [43], and hence used float16 for attention maps. In Table 3, we show the results of ablation studies to validate the efficacy of our proposed ensemble and distillation strategy.

#### 4.1.2 Chest X-Ray Images Classification

With privacy being a particularly important topic for real-world medical applications, we believe our proposed FL framework will be a good fit to facilitate privacy-preserving learning across various hospital sites. To this end, we evaluate our method on cross-domain, cross-site learning with private data, which is relatively under-explored in the contemporary FL methods.

We use NIH chestX-ray14 (NIH CXR14) [51] and CheXpert [15] as two domains where private data come from. To ensemble their labels, we disregard ambiguous categories such as Effusion, Pleural Effusion, and Pleural Other, and other samples labeled “Support Device”, leaving 86,524 images in NIH CXR14 and 64,346 images in CheXpert for training across 14 classes. Of these, NIH CXR14 has annotations for 12 classes and CheXpert for 8 classes, with 6 of these classes overlapping across both datasets. For public data, we use 26,684 x-ray images in the RSNA Pneumonia Detection Challenge public data <sup>1</sup> without using their labels. For  $K$  local nodes, each private dataset is distributed to  $K_d = K/2$  local nodes. Implementation details and ablation studies are in supplementary material.

Figure 4 illustrates the effectiveness of bound attention constraint during ensemble distillation. In Table 4, we compare FedAD to FedMD[22] on multi-label chest-x-ray image classification under the same settings, where one can note our FedAD outperforms FedMD by a significant margin on both test datasets. While distilling and testing on the data from the same, cross domain with FedAD obtains the best result for CheXpert dataset, distilling from one domain gives comparable performance. Note that our proposed FedAD uses additional unlabeled public data during distillation when compared to the centralized training setting. Under this cross-domain setting, the trained model is capable of classifying all 14 classes, whereas training with a single domain can only classify 12 and 8 classes, respectively due to the annotation limitations as noted above.

#### 4.2 Extension to Segmentation

We evaluate our method on segmentation on the Cityscapes dataset and 3D Brain Tumor Segmentation (BraTS) dataset. Note each local model predicts one time on the public data, thus the communication cost is independent of the distillation iterations.

##### 4.2.1 Cityscapes Segmentation

For natural images, we use Cityscapes dataset [5] for semantic segmentation of urban street scenes with 50 cities. In constructing local training sets, we split 2975 training images into three subsets based on the countries the images

<sup>1</sup><https://www.kaggle.com/c/rsna-pneumonia-detection-challenge>

Method	mIoU (%)↑	Pixel Accuracy(%)↑
<i>Standalone</i> (mean ± std)	$64.11 \pm 10.58$	$93.74 \pm 1.79$
<i>Centralized</i>	<b>75.65</b>	<b>95.87</b>
PATE [38]	64.48	93.99
FedAD	$\tau=1$	58.56
	$\tau=\infty$	71.86
	w/ att.	<b>72.97</b>
		<b>95.07</b>

Table 5. Segmentation results on Cityscapes. We calculate IoU between the ground truth mask and the predicted segmentation mask for each class, and report mean IoU of all classes.

Method	$73.38 \pm 3.44$	<i>Centralized</i>	82.13
	Li <i>et al.</i> [26]	FedMD [22]	FedAD
Average Dice (%)↑	84.33	75.71	77.85
Bandwidth (GB)↓	64.37	2154.84	<b>13.36</b>
Privacy (NO Shared Param.)	✗	✓	✓
Flexibility	online	online	on/offline
One-shot	✗	✗	✓

Table 6. Segmentation results on BraTS. We compare average Dice score over three types of tumor regions, communication bandwidth, privacy guarantee, and other attributes such as flexibility.

are collected from, e.g., Germany, France and Switzerland. We use 1,525 unlabeled test images as public data. Note the cities where the public images are collected from are different from with either of that of the private data. We test 19 classes on the validation dataset of cityscapes which contains 500 images.

We adopt PSPNet [57] with ResNet101 [10], with the backbone pretrained on ImageNet [6]. We use mini-batch SGD with the learning rate initialized as 0.01 and decreased to 0 in 100,000 iterations, and the same data augmentation as [32]. In Table 5, we compare with the other offline decentralized learning method PATE [38], which is also privacy-guaranteed (no shared weights/gradients). With the same locally trained models, our FedAD demonstrates superior performance on both mIoU and pixel accuracy. For ablation study, Table 5 also compares with the commonly used [32, 33]  $\tau = 1$  case, showing the efficacy of our distillation method for knowledge aggregation.

#### 4.2.2 Brain Tumor Segmentation

For medical images, we use the BraTS 2018 dataset [35, 2] that contains multi-parametric preoperative 3D MRI scans of 285 subjects with brain tumors. Each subject was scanned under the T1-weighted, T1-weighted with contrast enhancement, T2-weighted, and T2 fluid-attenuated inversion recovery (T2-FLAIR) modalities. Following the experimental protocol in prior work [26], we use 242 subjects for the training set and 43 subjects for held-out test set. The training set is stratified into three subsets according to the institution the data is originated (“2013”, “CBICA”, “TCIA”) and assigned each to federated local client. We

use half of the unlabeled validation set of the BraTS 2020 dataset [35, 2] as the public data comprising 62 subjects independent of either private dataset.

While the aforementioned distillation method is designed for general tasks which use cross entropy objective, we optimize with dice loss [36] for brain tumor segmentation. We thus modify Eq. 4 with soft dice to constrain the predicted mask  $\hat{z}^c$  to match the aggregated masks  $\tilde{z}^c$ . Let  $p^c = \sigma(\hat{z}^c)$  be the predicted probabilities of 3D voxel being class  $c$ ,  $q^c = \sigma(\tilde{z}^c)$  be the 3D soft pseudo labels, we have:

$$\mathcal{L}_{\text{logits}} = -\frac{1}{C} \sum_c \frac{2 \cdot \sum p^c \cdot q^c}{\sum (p^c)^2 + \sum (q^c)^2 + \epsilon}, \quad (11)$$

where the summation on  $p^c$ ,  $q^c$  are voxel-wise, and  $\epsilon$  is a small constant to avoid numerical instability. Note in Eq. 8 and Eq. 9 the summation in numerator and denominator are modified to be voxel-wise as well for this 3D case.

We use the same network backbone as [26]. The training strategy is the same as [37]. We train each local model individually with 20,000 iterations, with local-to-central distillation taking 5,000 iterations. The weight decay is 5e-4 and 0 for local training and distillation, respectively. Table 6 compares the segmentation performance and other utilities with parameter-based [26] and distillation-based [22] federated methods. We can note that our method achieves the state-of-the-art privacy/performance trade-offs compared with other existing methods.

## 5. Conclusions

In this work, we propose a **one-shot** federated learning framework, called FedAD, that can in principle preserve local data privacy using only unlabeled and domain robust public data and be efficient in the utilization of available network bandwidth resources when compared to competing prior art. Another key challenge in federated learning is its inherent heterogeneity, which can manifest itself in various ways: different differing domain distributions, different local model architectures, or simply the diversity in knowledge across all local models. To comprehensively address these issues, our framework also includes a novel knowledge distillation algorithm that is based on ensemble distillation of prediction logits as well as structural knowledge by means of model attention. Extensive experiments on classification and segmentation tasks with both natural image and medical image datasets demonstrated the efficacy of FedAD while also not risking privacy leakage. Furthermore, with experiments and analyses using cross-domain and heterogeneous data distributions, we also demonstrated FedAD’s applicability in real-world cross-institutional learning with medical imaging data.

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