

FLEET: Online Federated Learning via Staleness Awareness and Performance Prediction

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Federated learning (FL) is very appealing for its privacy benefits: essentially, a global model is trained with updates computed on mobile devices while keeping the data of users local. Standard FL infrastructures are however designed to have no energy or performance impact on mobile devices, and are therefore not suitable for applications that require frequent (*online*) model updates, such as news recommenders.

This article presents FLEET, the first *Online FL* system, acting as a middleware between the Android operating system and the machine learning application. FLEET combines the privacy of Standard FL with the precision of online learning thanks to two core components: (1) I-PROF, a new lightweight profiler that predicts and controls the impact of learning tasks on mobile devices, and (2) ADASGD, a new adaptive learning algorithm that is resilient to delayed updates.

Our extensive evaluation shows that Online FL, as implemented by FLEET, can deliver a $2.3\times$ quality boost compared to Standard FL while only consuming 0.036% of the battery per day. I-PROF can accurately control the impact of learning tasks by improving the prediction accuracy by up to $3.6\times$ in terms of computation time, and by up to $19\times$ in terms of energy. ADASGD outperforms alternative FL approaches by 18.4% in terms of convergence speed on heterogeneous data.

CCS Concepts: • Security and privacy \rightarrow Privacy-preserving protocols; • Hardware \rightarrow Power estimation and optimization; • Theory of computation \rightarrow Distributed computing models;

Additional Key Words and Phrases: Federated learning, online learning, asynchronous gradient descent, profiling, mobile Android devices

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1 INTRODUCTION

The number of edge devices and the data produced by these devices have grown tremendously over the past 10 years. In 2009, mobile phones only generated 0.7% of the worldwide data traffic; however, in 2018, this number exceeded 50% [76]. This exponential growth is raising challenges both in terms of scalability and privacy. As the volume of data produced by mobile devices explodes, users expose increasingly detailed and sensitive information, which in turn becomes more costly to store, process, and protect. This dual challenge of privacy and scalability is pervasive in **machine learning (ML)** applications such as recommenders, image-recognition apps, and personal assistants. These ML-based application often operate on highly personal and possibly sensitive content, including conversations, geolocation, or physical traits (faces, fingerprints), and typically require tremendous volumes of data for training their underlying ML models. For example, people in the United States ages 18 to 24 type on average around 900 words per day (128 messages per day [70] with an average of 7 words per message [17]). The Android next-word prediction service is trained on average with sequences of 4.1 words [37], which means that each user generates around 220 training samples daily. With tens of millions or even billions of user devices [8], scalability issues arise.

Federated learning. To address this dual privacy and scalability challenge, large industrial players are now seeking to exploit the rising power of mobile devices to reduce the demand on their server infrastructures while, at the same time, protecting the privacy of their users. Federated learning (FL) is a new computing paradigm (spearheaded among others by Google [11, 42, 74]) where a central server iteratively trains a global model (used by an ML-based application) without the need to centralize the data. The iterative training orchestrated by the server consists of the following synchronous steps for each update. Initially, the server selects the contributing mobile devices and sends them the latest version of the model. Each device then performs a learning task based on its local data and sends the result back to the server. The server aggregates a predefined number of results (typically a few hundred [8]) and finally updates the model. The server drops any results received after the update. FL is "privacy-ready" and can provide formal privacy guarantees by using standard techniques such as secure aggregation and differential privacy [9].

The standard use of FL has so far been limited to a few lightweight and extremely privacy-sensitive services, such as next-word prediction [85], but its popularity is bound to grow. Privacy-related scandals continue to unfold [54, 55], and new data protection regulations come into force [31, 78]. The popularity of FL is clearly visible in two of the most popular ML frameworks (namely TensorFlow and PyTorch) [29, 71], and also in the rise of startups such as S20.ai [72] or SNIPS (now part of Sonos) [75], which are betting on private decentralized learning.

Limitation of Standard FL. These are encouraging signs, but we argue in this work that Standard FL [8] is unfortunately not effective for a large segment of ML-based applications, mainly due to its constraint for high device availability: the selected mobile devices need to be idle, charging, and connected to an unmetered network. This constraint removes any impact perceived by users but also limits the availability of devices for learning tasks. Google observed lower prediction accuracy during the day, as few devices fulfill this policy, and these generally represent a skewed population [85]. With most devices available at night, the model is generally updated every 24 hours.

This constraint may be acceptable for some ML-based services but is problematic to what we call *online learning* systems, which underlie many popular applications such as news recommenders or interactive social networks (e.g., Facebook, Twitter, LinkedIn). These systems involve large amounts of data with high *temporality*, which generally become obsolete in a matter of hours or even minutes [56]. To illustrate the limitation of Standard FL, consider two users, Alice and Bob,

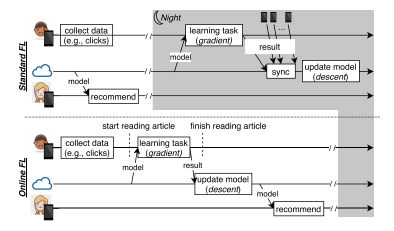


Fig. 1. Online FL enables frequent updates without requiring idle-charging-WiFi connected mobile devices.

who belong to a population that trains the ML model underlying a news recommendation system (Figure 1). Bob wakes up earlier than Alice and clicks on some news articles. To deliver fresh and relevant recommendations, these clicks should be used to compute recommendations for Alice when she uses the app, slightly after Bob. In Standard FL (upper half Figure 1), the device of Bob would wait until much later (when idle, charging, and connected to WiFi) to perform the learning task, thus negating the value of the task results for Alice. In an online learning setup (lower half of Figure 1), the activity of Bob is rapidly incorporated into the model, thereby improving the experience of Alice.

Challenges and contributions. In this article, we address the aforementioned limitation and enable Online FL. We introduce FLEET, the first FL system that specifically targets online learning, acting as a middleware between the operating system of the mobile device and the ML-based application. FLEET addresses two major problems that arise after forfeiting the high device availability constraint.

First, learning tasks may have an energy impact on mobile devices now powered on a battery. Given that learning tasks are generally compute intensive, they can quickly discharge the device battery and thereby degrade user experience. To this end, FLEET includes I-PROF (Section 2.2), our new profiling tool that predicts and controls the computation time and the energy consumption of each learning task on mobile devices. The goal of I-PROF is not trivial given the high heterogeneity of the devices and the performance variability even for the same device over time [62] (as we show in Section 3).

Second, as mentioned previously, synchronous training discards all late results arriving after the model is updated, thus wasting the battery of the corresponding devices and their potentially useful data. Frequent model updates call for small synchronization windows that, given the high performance variability, amplify this waste. We therefore replace the synchronous scheme of Standard FL with asynchronous updates. However, asynchronous updates introduce the challenge of *staleness*, as multiple users are now free to perform learning tasks at arbitrary times. A stale result occurs when the learning task was computed on an outdated model version; meanwhile, the global model has progressed to a new version. Stale results add noise to the training procedure, slow down, or even prevent its convergence [40, 88]. Therefore, FLEET includes ADASGD (Section 2.3), our new **stochastic gradient descent (SGD)** algorithm that tolerates staleness by dampening the impact of outdated results. However, some stale model updates may be computed on very rare data and

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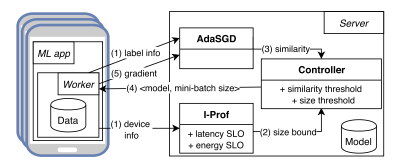


Fig. 2. The architecture of FLEET.

the staleness dampening may almost completely wipe out their contribution. Therefore, the final dampening factor also contains a similarity-based boosting that promotes model updates with a label distribution different from the label distribution of the past learning tasks.

We fully implemented the server side and the Android client of FLEET. We evaluate the potential of FLEET and show that it can increase the accuracy of a recommendation system (that employs Standard FL) by 2.3× on average, by performing the same number of updates but in a more timely (online) manner. Even though the learning tasks drain energy directly from the battery of the phone, they consume on average only 0.036% of the battery capacity of a modern smartphone per user per day. We also evaluate the components of FLEET on 40 commercial Android devices by using popular benchmarks for image classification. Regarding I-Prof, we show that 90% of the learning tasks deviate from a fixed **service-level objective (SLO)** of 3 seconds by at most 0.75 seconds in comparison to 2.7 seconds for the competitor (the profiler of MAUI [18]). The energy deviation from an SLO of 0.075% battery drop is 0.01% for I-Prof and 0.19% for the competitor. We also show that our staleness-aware learning algorithm (ADASGD) learns 18.4% faster than its competitor (DynSGD [40]) on heterogeneous data.

2 FLEET

FLEET incorporates two components we consider necessary in any system that has the ambition to provide both the (1) privacy of FL and (2) the precision of online learning systems. The first component is I-Prof, a *lightweight ML-based profiling* mechanism that controls the computation time and energy of the learning task by using ML-based estimators. The second component of FLEET is ADASGD, a new adaptive learning algorithm that tolerates stale updates by automatically adjusting their weight.

2.1 Architectural Overview

Similar to the implementation of Standard FL [8], FLEET follows a client-server architecture (Figure 2) where each user hosts a *worker* and the service provider hosts the *server* (typically in the cloud). In FLEET, the worker is a library that can be used by any mobile ML-based application (e.g., a news articles application). The model training protocol of FLEET is the following (the numbers refer to Figure 2):

(1) The worker requests a learning task and sends information regarding the labels of the local data along with information about the state of the mobile device. We introduce the purpose of this information in Steps 2 and 3.

¹https://github.com/gdamaskinos/fleet.

(2) I-Prof employs the device information to bound the workload size (hereinafter called the *mini-batch size bound*) that will be allocated to this worker such that the computation time and energy consumption approximate an SLO set by the service provider or negotiated with the user (details in Section 2.2). The *mini-batch size* is then set as the *min*(mini-batch size bound, local data size²).

- (3) ADASGD computes a *similarity* for the requested learning task with past learning tasks to adapt to updates with new data (details in Section 2.3).
- (4) To prevent the computation of learning tasks with low or no utility for the learning procedure, the controller checks if both the mini-batch size and the similarity values pass certain *thresholds* set by the service provider. If the check fails, the request of the worker is rejected; otherwise, the controller sends the model parameters and the mini-batch size to the worker and the learning task execution begins (details about setting these thresholds in Section 2.4).
- (5) Based on the mini-batch size returned by the server, the worker samples from its locally collected data, performs the learning task (i.e., one or multiple local model updates), and sends the result (i.e., the *gradient*) back to the server. On the server side, ADASGD updates the model after dynamically adapting this gradient based on its *staleness* and on its similarity value (details in Section 2.3).

The preceding protocol maintains the key "privacy-readiness" of Standard FL: the user data never leave the device.

2.2 Workload Bound via Profiling

In Online FL, a mobile device should be able to compute model updates at any time, not only during the night, when the mobile device is idle, charging, and connected to WiFi. Therefore, FLEET drops the constraint of Standard FL for high device availability. Hence, the learning task now drains energy directly from the battery of the device. Controlling the impact of a learning task on the user application in terms of energy consumption and computation time becomes crucial. To this end, FLEET incorporates a profiling mechanism that determines the workload size (i.e., the minibatch size) appropriate for each mobile device.

Best-effort solution. To highlight the need for a specific profiling tool, we first consider a naive solution in which users process data points until they reach the SLO either in terms of computation time or energy. At this point, a worker sends back the resulting "best-effort" gradient. The service provider cannot decide beforehand whether, for a given device, the cost (in terms of energy, time, and bandwidth) to download the model, compute, and upload the gradient is worth the benefit to the model. Updates computed on very small mini-batch sizes (by weak devices) will perturb the convergence of the overall model and might even negate the benefit of other workers.

To illustrate this point, consider the experiment of Figure 3. The figure charts the result of training a **convolutional neural network (CNN)** on CIFAR10 [44] under different combinations of "strong" and "weak" workers. The strong workers compute on a mini-batch size of 128, whereas the weak workers compute on a mini-batch size of 1. We observe that even 2 weak workers are enough to cancel the benefit of distributed learning—that is, the performance with 10 strong + 2 weak workers is the same as training with a single strong worker.

One way to avoid this issue could be to drop all of the gradients computed on a mini-batch size lower than a given bound or weigh them with a tiny factor according to the size of their underlying mini-batch. However, this way would waste the energy required to obtain these gradients. A profiler tool that can estimate the maximum mini-batch size (workload bound) that

²The size of the local data is available to the server via the label info.

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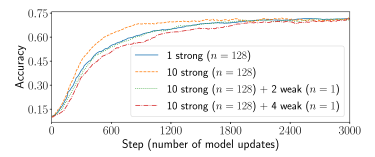


Fig. 3. Motivation for lower bounding the mini-batch size. The noise introduced by weak workers (i.e., with small mini-batch sizes) may be detrimental to learning.

a worker can compute is necessary for the controller to decide whether to reject the computation request of this worker, before the gradient computation. Unfortunately, existing profiling approaches [10, 13, 14, 18, 36, 45, 86] are not suitable because they are either relatively inaccurate (see Section 3.4) or they require privileged access (e.g., rooted Android devices) to low-level system performance counters.

I-Prof. Mobile devices have a significantly lower level of parallelism in comparison with cloud servers. For example, the graphical accelerators of mobile devices generally have 10 to 20 cores [6, 69], whereas the GPUs on a server have thousands of cores [63]. Given this low level of parallelism, even a relatively small mini-batch size can fill the processing pipelines. Hence, any additional workload will linearly increase the computation time and the energy consumption. Based on this observation, we built I-Prof., a lightweight profiler specifically designed for Online FL systems. We design I-Prof with three goals in mind: (1) operate effectively with data from a wide range of device types, (2) do so in a lightweight manner (i.e., introduce only a negligible latency to the learning task), and (3) rely only on the data available on a stock (non-rooted) Android device.

I-Prof employs an ML-based scheme to capture how the device features affect the computation time and energy consumption of the learning task. I-Prof predicts the largest mini-batch size a device can process while respecting both the time and the energy limits set by the SLO. To this aim, I-Prof uses two predictors: one for computation time and one for energy. Each predictor updates its state with data from the device information sent by the workers.

However, designing such predictors is tricky, as modern mobile phones exhibit a wide range of capabilities. For example, in a matrix multiplication benchmark, Galaxy S6 performs 7.11 Gflops, whereas Galaxy S10 performs 51.4 Gflops [47]. Figure 4 illustrates this heterogeneity on three different mobile devices by executing successive learning tasks of increasing mini-batch size ("up"). After reaching the maximum value, we let the devices cool down and execute subsequent learning tasks with decreasing mini-batch size ("down"). We present the results for the up-down part with the same color pattern, except for Honor 10 in Figure 4(b), which we split for highlighting the difference. Figure 4 illustrates that the linear relation changes for each device and for certain devices (Honor 10, Galaxy S7) also changes with the temperature. Note that Honor 10 shows an increased variance at the end of the "up" part (Figure 4(b)) that is attributed to the high temperature of the device. The variance is significantly smaller for the "down" part.

In the following, we describe how I-Prof predicts the mini-batch size (n) given a computation time SLO³ (t_{SLO}). The computation time linearly increases with the workload size—that is,

³The prediction method given an energy SLO is the same.

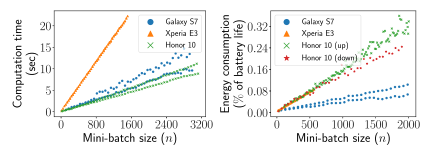


Fig. 4. The linear relation between the computation time and the mini-batch size depends on the specific device, and may even vary for the same device, depending on operation conditions such temperature.

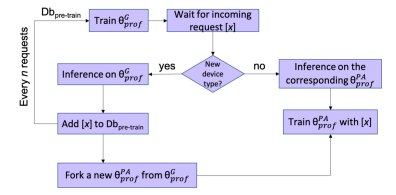


Fig. 5. The flowchart of I-Prof: the slope $\hat{\alpha}$ is predicted by a personalized per-device PA model $\theta_{\text{prof}}^{PA}$ except for the first request of each device type, which is directed to a cold-start linear regression model θ_{prof}^{G} .

 $t_{comp} = \alpha \cdot n$, where α depends on the device and its state. Considering the goal (i.e., $t_{comp} \rightarrow t_{SLO}$), the optimal mini-batch size is predicted as follows:

$$\hat{n} = \max\left(1, \frac{t_{SLO}}{\hat{\alpha}}\right). \tag{1}$$

I-Prof estimates the slope $\hat{\alpha}$ from the device characteristics and operational conditions using a method that combines linear regression and online **passive-aggressive** (PA) learning [16].

The input to this method is a set of device features based on measurements available through the Android API, namely available memory, total memory, temperature, and sum of the maximum frequency over all CPU cores. However, these features only encode the computing power of a device. For the prediction based on the energy SLO, I-Prof also needs a feature that encodes the energy efficiency of each device. We choose this additional feature as the energy consumption per non-idle CPU time. We show in our evaluation (Section 3.4) that these features achieve our three design goals. Given a vector of device features (\mathbf{x}), and a vector of model parameters (θ_{prof}), the slope $\hat{\alpha}$ is estimated as $\hat{\alpha} = \mathbf{x}^T \theta_{\text{prof}}$.

I-Prof uses a cold-start linear regression model θ_{prof}^G for the first request of each user device. We pretrain the cold-start model using ordinary least squares with an offline dataset (Figure 5). This dataset consists of data collected by executing requests from a set of training devices with a

 $^{^4\}mathrm{CPU}$ time spent by processes executing in user or kernel mode.

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mini-batch size increasing from 1 until a value such that the computation time reaches twice the SLO. I-Prof periodically re-trains the cold-start model after appending new data (device features). The training cost is negligible given the small number of features.

Furthermore, I-Prof creates a personalized model $\theta_{\text{prof}}^{PA}$ for every new device model (e.g., Galaxy S7) and employs it for every following request coming from this particular model. I-Prof bootstraps the new model with the first request (for which the cold-start model is used to estimate the computation time). For all of the following learning tasks that result in pairs of $(\mathbf{x}^{(k)}, \alpha^{(k)})$, I-Prof incrementally updates a PA model [16] as $\theta_{\text{prof}}^{(k+1)} = \theta_{\text{prof}}^{(k)} + \frac{f^{(k)}}{\|\mathbf{x}^{(k)}\|^2} \mathbf{v}^{(k)}$, where $\mathbf{v}^{(k)} = sign(\alpha^{(k)} - \mathbf{x}^{(k)T}\theta_{\text{prof}}^{(k)})\mathbf{x}^{(k)}$ denotes the update direction and f the loss function:

$$f(\boldsymbol{\theta}_{\text{prof}}, \boldsymbol{x}, \alpha) = \begin{cases} 0 & \text{if } |\boldsymbol{x}^T \boldsymbol{\theta}_{\text{prof}} - \alpha| \le \epsilon \\ |\boldsymbol{x}^T \boldsymbol{\theta}_{\text{prof}} - \alpha| - \epsilon & \text{otherwise.} \end{cases}$$
 (2)

The parameter ϵ controls the sensitivity to prediction error and thereby the aggressiveness of the regression algorithm—that is, the smaller the value of ϵ , the larger the update for each new data instance (more aggressive).

I-Prof focuses solely on the time and energy spent during an SGD computation. Despite network costs (particularly when transferring models) having also an important impact, they fall outside the scope of this work, as one can rely on prior work [4, 52, 67] to estimate the time and energy of network transfers within FLEET.

2.3 Adaptive SGD

The server-driven synchronous training of Standard FL is not suitable for Online FL, as the latter requires frequent updates and needs to exploit contributions from all workers, including slow ones (Section 1). Therefore, we introduce ADASGD, an asynchronous learning algorithm that is robust to stale updates. ADASGD is responsible for aggregating the gradients sent by the workers and updating the application model ($\theta_{\rm app}$). Each update takes place after ADASGD receives K gradients. The aggregation parameter K can be either fixed or based on a time window (e.g., update the model every 1 hour). The model update is as follows:

$$\boldsymbol{\theta}_{\text{app}}^{(t+1)} = \boldsymbol{\theta}_{\text{app}}^{(t)} - \gamma_t \sum_{i=1}^K \min\left(1, \Lambda(\tau_i) \cdot \frac{1}{sim(\boldsymbol{x}_i)}\right) \cdot G(\boldsymbol{\theta}_{\text{app}}^{(t_i)}, \xi_i), \tag{3}$$

where γ_t is the learning rate, $t \in \mathbb{N}$ denotes the global logical clock (or step) of the model at the server (i.e., the number of past model updates), and $t_i \leq t$ denotes the logical clock of the model that the worker receives. $G(\theta_{\text{app}}^{(t_i)}, \xi_i)$ is the gradient computed by the client w.r.t. the model $\theta_{\text{app}}^{(t_i)}$ on the mini-batch ξ_i drawn uniformly from the local dataset x_i .

The workers send gradients asynchronously that can result in *stale* updates. The *staleness* of the gradient ($\tau_i := t - t_i$) shows the number of model updates between the model pull and gradient push of worker i. One option is to directly apply this gradient, at the risk of slowing down or even completely preventing convergence [40, 88]. The Standard FL algorithm (FedAvg [53]) simply drops stale gradients. However, even if computed on a stale model, the gradient may incorporate potentially valuable information. Moreover, in FLEET, the gradient computation may drain energy directly from the battery of the phone, thus making the result even more valuable. Therefore, ADASGD utilizes even stale gradients without jeopardizing the learning process, by multiplying

 $^{^5}$ Not to be confused with the model of the profiler (θ_{prof}).

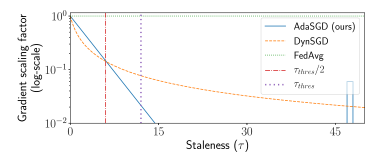


Fig. 6. Gradient scaling schemes of SGD algorithms. ADASGD, proposed in this article, dampens stale gradients with an exponentially decreasing function ($\Lambda(\tau)$) based on the expected percentage of non-stragglers ($\tau_{\rm thres} := s$ -th percentile of staleness values), and boosts the gradient of the straggler ($\tau = 48$) due to its low similarity ($sim(x_i)$).

each gradient with an additional weight to the learning rate. This weight consists of (1) a dampening factor based on the staleness ($\Lambda(\tau_i)$) and (2) a boosting factor based on the user's data novelty $(\frac{1}{\sin(r_i)})$, which we describe in the following.

Staleness-based dampening. AdaSGD builds on prior work on staleness-aware learning that has shown promising results [40, 88]. To accelerate learning, AdaSGD relies on a system parameter: the expected percentage of non-stragglers (denoted by s%). We highlight that this value is not a hyperparameter that needs tuning for each ML application but a system parameter that solely depends on the computing and networking characteristics of the workers, while it can be adapted dynamically [65, 66]. We define the staleness-aware dampening factor $\Lambda(\tau) = e^{-\beta \tau}$, with β chosen s.t. $\frac{1}{\frac{\tau_{\text{thres}}}{2}+1} = e^{-\beta \frac{\tau_{\text{thres}}}{2}}$ (i.e., the inverse dampening function [40] intersects with our exponential dampening function in $\frac{\tau_{\text{thres}}}{2}$), where τ_{thres} is the s-th percentile of past staleness values. Figure 6 shows the dampening factor of AdaSGD compared to the inverse dampening function (employed by DynSGD [40]). Our hypothesis is that the perturbation to the learning process introduced by stale gradients increases exponentially and not linearly with the staleness. We empirically verify the superior performance of our exponential dampening function compared to the inverse in Section 3.3.

Similarity-based boosting. In the presence of stragglers with large delays (comparing to the mean latency), staleness can grow and drive $\Lambda(\tau)$ close to 0 (i.e., almost neglect the gradients of these stragglers). Nevertheless, these gradients may contain valuable information. In particular, they may be computed on data that are not similar to the data used by past gradients. Hence, ADASGD boosts these gradients by using the following similarity value:

$$sim(\mathbf{x}_i) = BC(LD(\mathbf{x}_i), LD_{global}),$$
 (4)

where BC denotes the Bhattacharyya coefficient [81], and LD the label distribution, which captures the importance of each gradient. For instance, given an application with four distinct labels and a local dataset (x_i) that has one example with label 0 and two examples with label 1:

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 $LD(x_i) = [\frac{1}{3}, \frac{2}{3}, 0, 0]$. The global label distribution (LD_{global}) is computed on the aggregate number of previously used samples for each label. We highlight that LD is not specific to classification ML tasks; for regression tasks, LD would involve a histogram, with the length of the LD vector being equal to the number of bins instead of the number of classes. We choose BC for two main reasons: (1) it is suitable for comparing two probability distributions (in our case, the Label Distribution), and (2) $sim(x_i)$ should belong to the interval [0,1]. We also mention that this coefficient is similar but more general compared to the (more standard) Euclidean distance: BC is a generalization of Mahalanobis distance, which is the Euclidean distance after scaling to unit variance.

The similarity value essentially captures how valuable the information of the gradient is. For instance, if a gradient is computed on examples of an unseen label (e.g., a very rare animal), then its similarity value is less than 1 (i.e., has information not similar to the current knowledge of the model). For the similarity computation, the server needs only the indices of the labels of the local datasets without any semantic information (e.g., label 3 corresponds to "dogs").

Computational complexity. I-Prof poses negligible overhead (O(1)) to the overall training procedure, given the handful of features used. Similarity computation costs an extra O(L), where L is the total amount of labels. This can also be treated as O(1) when compared to the complexity of SGD (that depends on the number of model parameters and the number of examples). Overall, the computation complexity of FLEET is asymptotically the same as the underlying SGD complexity which in turn depends on (1) the number of iterations, (2) model parameters, and (3) the number of training samples.

2.4 Implementation

The server of FLEET is implemented as a web application (deployed on an HTTP server) and the worker as an Android library. The server transfers data with the workers via Java streams by using Kryo [27] and Gzip. In total, FLEET accounts 26,913 Java LoC, 3,247 C/C++ LoC, and 1,222 Python LoC.

Worker runtime. We design the worker of our middleware (FLEET) as a library and execute it only when the overlying ML application (see Figure 2) is running in the foreground for two main reasons. First, since Android is a UI-interactive operating system, background applications have low priority so their access to system resources is heavily restricted and they are likely to be killed by the operating system to free resources for the foreground running app. Therefore, allowing the worker to run in the background would make its performance very unpredictable and thus impact the predictions of I-Prof. Second, running the worker in the foreground alleviates the impact of collocated (background) workload.

We build our main library for CNNs in C++ on top of FLEET. We employ (1) the Java Native Interface (JNI) for the server, (2) the Android NDK for the worker, (3) an encoding scheme for transferring C++ objects through the java streams, and (4) a thread-based parallelization scheme for the independent gradient computations of the worker. On recent mobile devices that support NEON [5], FLEET accelerates the gradient computations by using SIMD instructions. We also port a popular deep learning library (DL4J [22]) to FLEET, to benefit from its rich ecosystem of ML algorithms. However, as DL4J is implemented in Java, we do not have full control over the resource allocation.

FLEET relies on the developer of the overlying ML application to ensure the performance isolation between the running application and the worker runtime. The worker can execute in a window of low user activity (e.g., while the user is reading an article) to minimize the impact of the overlying ML application on the predictive power of I-Prof.

Resource allocation. Allocating system resources is a very challenging task given the latency and energy constraints of mobile devices [24, 57]. Our choice of employing only stock Android without root access means that we can only control which cores execute the workload on the worker, with no access, for instance, to low-level advanced tuning. Given this limited control and the inherent mobile device heterogeneity, we opt for a simple yet effective scheme for allocating resources.

This scheme schedules the execution only on the "big" cores for ARM big.LITTLE architectures and on all of the cores otherwise. In the case of computationally intensive tasks (e.g., the learning tasks of FLEET), big cores are more energy efficient than LITTLE cores because they finish the computation much faster [32]. Regarding ARMv7 symmetric architectures with two and four cores that equip older mobile devices, the energy consumption per workload is constant regardless of the number of cores: a higher level of parallelism will consume more energy, but the workload will execute faster. For this reason, our allocation policy relies on all available cores so that we can take advantage of the embarrassingly parallel nature of the gradient computation tasks. For such tasks, we empirically show (Section 3.5) that this scheme outperforms more complex alternatives [57].

Controller thresholds. In practice, the service provider can adopt various approaches to define the size and similarity thresholds of the controller (see Figure 2). One option is A/B testing along with the gradual increase of the thresholds. In particular, the system initializes the thresholds to zero and divides the users into two groups. The first group tests the impact of the mini-batch size and the second the impact of the label similarity. Both groups gradually increase the thresholds until the impact on the service quality is considered acceptable. The server can execute this A/B testing procedure periodically (i.e., reset the thresholds after a time interval). We empirically evaluate the impact of these thresholds on prediction quality in Section 3.6.

3 EVALUATION

Our evaluation consists of two main parts. First, in Sections 3.1 and 3.2, we evaluate the claim that Online FL holds the potential to deliver better ML performance than Standard FL [8] for applications that employ data with high temporality (Section 1). Second, we evaluate in more detail the internal mechanisms of FLEET, namely ADASGD (Section 3.3), I-PROF (Section 3.4), the resource allocation scheme (Section 3.5), and the controller (Section 3.6).

We deploy the server of FLEET on a machine with an Intel Xeon X3440 with four CPU cores, 16 GiB of RAM, and 1-Gb Ethernet, on Grid5000 [33]. For the Twitter hashtag recommender (Section 3.1), we deploy the worker on a Raspberry Pi 4 as our hashtag recommender is implemented on TensorFlow that does not yet support training on Android devices. For the MovieLens recommender (Section 3.2), we deploy the worker on four different Android devices in our lab. For all other experiments, the workers are deployed on a total of 40 different mobile phones from the AWS Device Farm [7] (Oregon, USA).

3.1 Online vs Standard FL: The Twitter Hashtag Recommender

We compare Online with Standard FL on a Twitter hashtag recommender. The goal of this recommender is to assist Tweeter users in their hashtag selection, by proposing them hashtags suitable for the Tweet they want to publish. Tweepy [77] enables us to collect around 2.6 million tweets located on the West Coast of the United States over a period of 13 consecutive days. We preprocess these tweets (e.g., remove automatically generated tweets, remove special symbols) based on the work of Dhingra et al. [23]. We then divide the data into shards, each spanning 2 days, and divide each shard into chunks of 1 hour. We finally group the data into mini-batches based on the user id.

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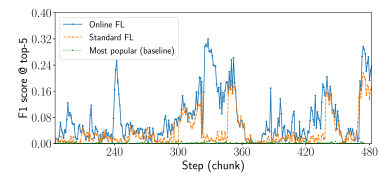


Fig. 7. Online FL boosts Twitter hashtag recommendations by an average of 2.3× comparing to Standard FL.

Our training and evaluation procedure follows an Online FL setup. Our model is a basic recurrent neural network implemented on TensorFlow with 123,330 parameters [30], which classifies each Tweet to 100 classes representing the 100 most popular hashtags in the previous timeframe. The model training consists of successive gradient descent operations, with each gradient derived from a single mini-batch (i.e., sent by a single user). Every day includes 24 mini-batches and thus 24 distinct gradient computations, whereas each model update uses 1 gradient. For the Online FL setup, the 24 daily updates are evenly distributed (one update every hour). Training uses the data of the previous hour and testing uses the data of the next hour. For the Standard FL setup, the 24 daily updates occur all at once (at the end of the day). Training uses the data of the previous day and testing uses the data of the next day. We highlight that under this setup, the two approaches employ the same number of gradient computations and the difference lies only in the time they perform the model updates. We also compare against a baseline model that always predicts the most popular hashtags [43, 64]. We evaluate the model on the data of each chunk and reset the model at the end of each shard.

Quality boost. For assessing the quality of the hashtag recommender, we employ the F1-score @ top-5 [28, 43] to capture how many recommendations were used as hashtags (precision) and how many of the used hashtags were recommended (recall). For each Tweet, the predicted probabilities (on the 100 classes) are sorted before we compare the top-5 hashtags with the actual hashtags of the tweet, and derive the F1-score. Figure 7 shows that Online FL outperforms Standard FL in terms of F1-score, with an average boost of 2.3×. Online FL updates the model in a more timely manner (i.e., soon after the data generation time) and can thus better predict (higher F1-score) the new hashtags than Standard FL. The performance of the baseline model is quite low, as the nature of the data is highly temporal [46].

Energy impact. We measure the energy impact of the gradient computation on the Raspberry Pi worker. The Raspberry Pi has no screen; nevertheless, recent trends in mobile/embedded processor design show that the processor is dominating the energy consumption, especially for compute-intensive workloads such as the gradient computation [35]. We measure the power consumption of every update of Online FL by executing the corresponding gradient computation 10 times and by taking the median energy consumption. We observe that the power depends on the batch size and increases from 1.9 W (idle) to 2.1 W (batch size of 1) and to 2.3 W (batch size of 100). The computation latency is 5.6 seconds for batch size of 1 and 8.4 for batch size of 100. Across all of the updates of Online FL (that employ various batch sizes and result in the quality boost shown in Figure 7), we measure the average, median, 99th percentile, and maximum values of the daily energy

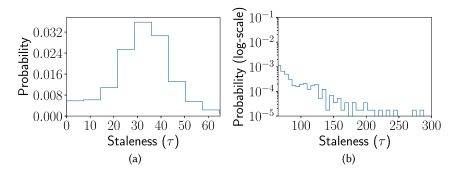


Fig. 8. Staleness distribution of collected tweets follows a Gaussian distribution (τ < 65) with a long tail (τ > 65).

consumption as 4, 3.3, 13.4, and 44 mWh, respectively. Given that most modern smartphones have battery capacities over 11,000 mWh, we argue that Online FL imposes a minor energy consumption overhead for boosting the prediction quality.

Staleness distribution. We study the staleness distribution of the updates on our collected tweets to set our experimental setup for evaluating ADASGD (Section 3.3). We assume that the round-trip latency per model update (gradient computation time plus network latency) follows an exponential distribution (as commonly done in the distributed learning literature [3, 25, 49, 59]). The network latency for downloading the model (123,330 parameters) and uploading the gradients is estimated to 1.1 second for 4G LTE and 3.8 seconds for 3G HSPA+ [38]. We then estimate the average computation latency to be 6 seconds, based on our latency measurements on the Raspberry Pi. Therefore, we choose the exponential distribution with a minimum of 6 + 1.1 = 7.1 seconds and a mean of $\frac{(6+1.1)+(6+3.8)}{2} = 8.45$ seconds. Given the exponential distribution for the round-trip latency and the timestamps of the tweets, we observe in Figure 8 that the staleness follows a Gaussian distribution with a long tail (as assumed in the work of Zhang et al. [88]). The long tail is due the presence of certain peak times with hundreds of tweets per second.

It is noteworthy that for applications such as our Twitter hashtag recommender where the user activity (and therefore the per-user data creation) is concentrated in time, the difference between Online and Standard FL in terms of communication overhead (due to the gradient-model exchange) is negligible. For applications where the per-user data creation is spread across the day, the communication overhead of Online FL grows, as each user communicates more often with the server.

3.2 Online vs Standard FL: The MovieLens Recommender

To better quantify the concrete benefits of FLEET for online recommenders, we have also used a MovieLens dataset [34] consisting of 1 million ratings from 6,000 users on 4,000 movies. We defined a simple scenario in which users continuously feed an online service with movie ratings while consuming movie recommendations (Figure 9(a)). The experiment distributes data based on user IDs and uses standard matrix factorization to provide recommendations. Following the FL philosophy, each user repeatedly computes gradients locally on their own (private) data, and uses these gradients to update a central model. Time (*x*-axis of Figure 9(a)) is simulated by splitting the MovieLens dataset in an initial training set of 100k and 45 rolling sets of 20k entries. A rolling set is initially used to evaluate the prediction accuracy and then it is appended to the training set. In Standard FL ("offline" curve), the model is trained every night with the data generated the day before, whereas Online FL ("online" curve) trains the model 10 times per day. Figure 9(a) charts the

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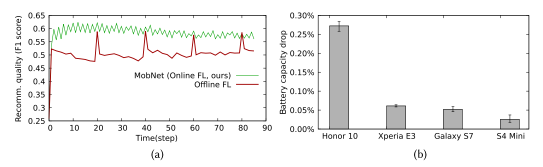


Fig. 9. Recommendation quality vs energy consumption for a movie recommender: recommendation quality of FLEET vs traditional FL on MovieLens 1M (higher is better) (a) and energy consumption (as percentage of the battery capacity) per FLEET gradient update (lower is better) (b). One can observe that the energy consumption is fairly heterogeneous even among recent phone generations.

F1-score from these two training approaches over time. The figures clearly shows that the online approach delivers scores up to 23% higher than those of the Standard FL, because it is able to incorporate new user inputs on the fly (see Figure 1).

Energy impact. Although the results of Figure 9(a) are encouraging, mobile devices may compute model gradients at any time, including when they are running on battery. To assess the impact of FLEET in terms of energy, Figure 9(b) represents the average and the variance for the total energy consumption over all gradient updates used to derive Figure 9(a). The experiment is performed using four off-the-shelf Android devices (Honor 10, Sony Xperia E3, Samsung Galaxy S7, and Samsung Galaxy S4 mini) connected over a WiFi network. Figure 9(b) shows that even if all phones compute identical workloads, there is an order of magnitude difference between the most energy efficient and the least efficient device. The daily energy consumption may vary depending on the device type, employed MF algorithm, and policy, but with a policy of 10 gradient updates per day, we observed a daily energy consumption ranging from 0.25% to 2.7% (the values in Figure 9(b) multiplied by 10).

3.3 ADASGD Performance

We now dissect the performance of ADASGD via an image classification application that involves CNNs. We choose this benchmark due to its popularity for the evaluation of SGD-based approaches [2, 12, 41, 53, 88, 89]. We employ multiple scenarios involving various staleness distributions, data distributions, and a noise-based differentially private mechanism.

Image classification setup. We implement the models shown in Table 1 in FLEET⁸ to classify handwritten characters and colored images. We use three publicly available datasets: MNIST [48], E-MNIST [15], and CIFAR-100 [44]. MNIST consists of 70,000 examples of handwritten digits (10 classes), whereas E-MNIST consists of 814,255 examples of handwritten characters and digits (62 classes). CIFAR-100 consists of 60,000 color images in 100 classes, with 600 images per class. We perform min-max scaling as a preprocessing step for the input features.

 $^{^6}$ To put this number into context, in 2009, Netflix awarded \$1,000,000 to a research team that was able to improve their recommendation algorithm with 10.06%.

⁷For this experiment, the average energy spent on network communication is around 55% of the total energy consumption. ⁸We implement the CNN for E-MNIST on DL4J and the rest on our default CNN library.

Dataset	Parameters	Input	Conv1	Pool1	Conv2	Pool2	FC1	FC2	FC3
MNIST	Kernel size	$28 \times 28 \times 1$	$5 \times 5 \times 8$	3×3	$5 \times 5 \times 48$	2×2	10	-	-
	Strides		1×1	3×3	1×1	2×2			
E-MNIST	Kernel size	$28 \times 28 \times 1$	$5 \times 5 \times 10$	2×2	$5 \times 5 \times 10$	2×2	15	62	-
	Strides		1×1	2×2	1×1	2×2			
CIFAR-100	Kernel size	$32 \times 32 \times 3$	$3 \times 3 \times 16$	3 × 3	$3 \times 3 \times 64$	4×4	384	192	100
	Strides		1×1	2×2	1×1	4×4			

Table 1. CNN Parameters

We split each dataset into training/test sets: 60,000/10,000 for MNIST, 697,932/116,323 for E-MNIST, and 50,000/10,000 for CIFAR-100. Unless stated otherwise, we set the aggregation parameter K (Section 2.3) to 1 (for maximum update frequency), the mini-batch size to 100 examples [61], the PA parameter ϵ to 0.1, and the learning rate to 15 * 10⁻⁴ for CIFAR-100, 8 * 10⁻⁴ for E-MNIST, and 5 * 10⁻⁴ for MNIST.

Since the training data present on mobile devices are typically collected by the users based on their local environment and usage, both the size and the distribution of the training data will typically heavily vary among users. Given the terminology of statistics, this means that the data are **not independent and identically distributed (non-IID)**. Following recent work on FL [79, 80, 87, 89], we employ a non-IID version of MNIST. Based on the standard data decentralization scheme [53], we sort the data by the label, divide them into shards of size equal to $\frac{60000}{2 \text{ * number of users}}$, and assign two shards to each user. Therefore, each user will contain examples for only a few labels.

Staleness controller. To be able to precisely compare ADASGD with its competitors, we experimentally control the staleness of the updates produced by FLEET, beyond the natural variability of the phones at our disposal. We use the pseudo-code shown in Algorithm 1, called τ Controller, to inject a predefined staleness distribution into FLEET. By allowing a fine control of staleness values, τ Controller can be used for the design and evaluation of any staleness-aware ML algorithm on a predefined staleness distribution (with arbitrary large values), independently of the underlying hardware and infrastructure setup (e.g., number of mobile devices, device type, network).

```
ALGORITHM 1: \tauController
```

```
Input: n: mini-batch size, \gamma_t: sequence of learning rates, \tau_{min}, \tau_{max}: max and min staleness, K: model updates
            aggregation window size
                                                                                                    // List of latest models (\theta_i)
    // Model update := \langle \text{gradient } (q), \text{ model step } (s), \text{ label distribution } (LD) \rangle
                                                                  // Map: model step \rightarrow list of computed model updates
2 G = {}
                                                                                        // Model version to be sent to client
3 Priority = 0
   Server
4 Function Descent(\mathcal{G}^{aggr}, \Theta):
                                                                                        // \mathcal{G}^{aggr}: K aggregated model updates
         t = |\Theta|
                                                                                                                        // Current step
         \theta^{(new)} = \Theta[t] - \gamma_t \sum_{(g, s, LD) \in \mathcal{G}} aggr \min \left(1, \Lambda(t-s) \cdot \frac{1}{sim(LD)}\right) \cdot g
                                                                                                                        // Equation (3)
         return \theta^{(new)}
   Function Pull():
         lock(\Theta)
         \theta = \Theta[Priority]
10
         unlock(\Theta)
         return \theta, Priority
12
```

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```
13 Function Push(q, s, LD):
         lock(\Theta)
14
         G[s].append(\langle g, s, LD \rangle)
15
          // Cold-start phase
         if |\Theta| < \tau_{max} and |\mathcal{G}.values()| > K then
16
              G^{aggr} = G.getRandomValues(K)
                                                                                                     // Get K model updates
17
               \theta^{(new)} = \text{Descent}(\mathcal{G}^{aggr}, \Theta)
18
              \Theta.append(\theta^{(new)})
              Priority++
20
         end
21
          // On-demand staleness phase
         for \tau \in [\tau_{min}, \tau_{max}] do
22
              if \tau \notin G.keys() or |G[|\Theta| - \tau]| < K then
23
                    Priority = |\Theta| - \tau
                                                                    // This model version requires more updates
                    unlock(\Theta)
                    return
               end
         end
28
          // Operation phase
         for \tau \in [\tau_{min}, \tau_{max}] do
29
              G^{aggr} = []
                                                 // Aggregates the K model updates necessary for Descent
30
              for K iterations do
31
                    \hat{\tau} = \text{Gaussian}(\mu = \frac{\tau_{max} - \tau_{min}}{2}, \sigma = \frac{\tau_{max} - \tau_{min}}{6}) // \hat{\tau} \in [\tau_{min}, \tau_{max}] with 99.7% prob
32
                    \hat{\tau} = min(\tau_{max}, max(\tau_{min}, \hat{\tau}))
                                                                                                              //\hat{\tau} \in [\tau_{min}, \tau_{max}]
                    upd = \mathcal{G}[|\Theta| - \hat{\tau}]
                    G^{aggr}.append(upd)
                    G[|\Theta| - \hat{\tau}].remove(upd)
                                                                // Successful due to On-demand staleness phase
               end
               \theta^{(new)} = Descent(\mathcal{G}^{aggr}, \Theta)
38
               \Theta.append(\theta^{(new)})
39
              \Theta.remove(0)
                                                                                                      // Remove oldest model
40
         end
41
          // Drop unused gradients
         for \tau \in G.keys() do
42
              if (|\Theta| - \tau) > \tau_{max} then
43
                  G.remove(|\Theta| - \tau)
              end
45
         end
46
         unlock(\Theta)
47
         return
48
```

 τ Controller executes on the server side of FLEET. For each client request, the function "Pull()" (lines 8–12) is invoked first and the appropriate model (θ) alongside with its corresponding version number (Priority) is sent to the client. The client then locally computes the gradient (g) and pushes it back to the server by invoking "Push(g, s, LD)" (lines 13–48), where s is the local model version that the gradient is computed on (i.e., Priority received from "Pull()") and LD is the label distribution needed to compute the similarity factor (Section 2.3). The function "Push()" executes

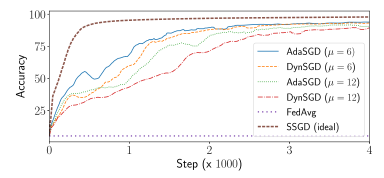


Fig. 10. Impact of staleness on learning.

in three phases. In the cold-start phase (lines 16–21), the server performs the descent operation and saves each model version for $\tau_{max} - \tau_{min}$ times before proceeding to the on-demand staleness phase. The purpose of the on-demand staleness phase (lines 22–28) is to accumulate K gradients for each of the possible outputs of the staleness distribution. This phase obtains each missing staleness value τ by sending the corresponding saved model to the client (set by using the *Priority* counter). In the operation phase (lines 29–41), the server performs the descent operation by choosing K accumulated model updates, the staleness of which is the output of the staleness distribution. Inevitably, the on-demand staleness method discards updates that are too stale to be selected (lines 42–46).

Staleness-aware learning. Based on the work of Zhang et al. [88] and the shape of the staleness distribution shown in Figure 8, we employ Gaussian distributions for the staleness with two setups $(D1 := \mathcal{N}(\mu = 6, \sigma = 2))$ and $D2 := \mathcal{N}(\mu = 12, \sigma = 4)$ to measure the impact of increasing the staleness. We set the expected percentage of non-stragglers (s%) to 99.7% (i.e., $\tau_{\text{thres}} = \mu + 3\sigma$). We evaluate the SGD algorithms on FLEET by using commercial Android devices from AWS.

We evaluate the performance of ADASGD against three learning algorithms: (1) DYNSGD [40], a staleness-aware SGD algorithm employing an inverse dampening function ($\Lambda(\tau) = \frac{1}{\tau+1}$), which ADASGD builds upon (Section 2.3); (2) the standard SGD algorithm with synchronous updates (SSGD) that represents the ideal (*staleness-free*) convergence behavior; and (3) FedAvg [53], the standard *staleness-unaware* SGD algorithm that is based on gradient averaging.

Staleness-based dampening. Figure 10 shows that ADASGD outperforms the alternative learning schemes for the non-IID version of MNIST. As expected, the staleness-free scenario (SSGD) delivers the fastest (ideal) convergence, whereas the staleness-unaware Fedauges. The comparison between the two staleness-aware algorithms (DynsGD and AdasGD) shows that our solution (AdasGD) better adapts the dampening factor to the noise introduced by stale gradients (Section 2.3). AdasGD reaches 80% accuracy 14.4% faster than DynsGD for D1 and 18.4% for D2. Figure 10 also depicts the impact of staleness on DynsGD and AdasGD. We observe that the larger the staleness, the slower the convergence of both algorithms. The advantage of AdasGD over DynsGD grows with the amount of staleness, as the larger amount of noise gives more leeway to AdasGD to benefit from its superior dampening scheme.

Similarity-based boosting. We evaluate the effectiveness of the similarity-based boosting property of ADASGD (Section 2.3) in the case of long tail staleness (see Figure 8). We employ the non-IID MNIST dataset, D1 (thus, τ_{thres} is 12), and set the staleness to $4 \cdot \tau_{\text{thres}} = 48$ for all gradients computed on data with class 0. This setup essentially captures the case where a particular label is only

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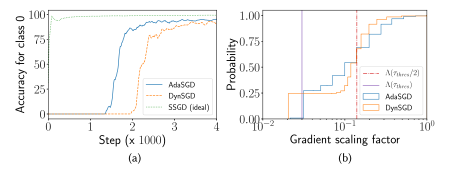


Fig. 11. Impact of long tail staleness on learning.

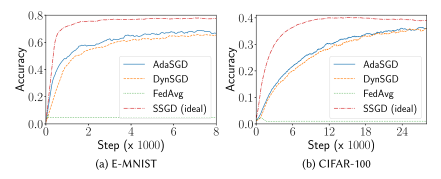


Fig. 12. Staleness awareness with IID data.

present in stragglers. Figure 11(a) shows that ADASGD incorporates the knowledge from class 0 much faster than DYNSGD.

Figure 11(b) shows the CDF for the dampening values used to weight the gradients of Figure 11(a). We mark the two points of interest regarding the $\tau_{\rm thres}$ by vertical lines (as also shown in Figure 6). If ADASGD had no similarity-based boosting, all updates related to class 0 would almost not be taken into account, as they would be nullified by the exponential dampening function, therefore leading to a model with poor predictions for this class. Given the low class similarity of the learning tasks involving class 0, ADASGD boosts their dampening value. The second vertical line denotes the staleness value ($\frac{\tau_{\rm thres}}{2}$ = 6) for which ADASGD and DYNSGD give the same dampening value (0.14). The slope of each curve at this point indicates that the dampening values for DYNSGD are more concentrated, whereas the ones for ADASGD are more spread around this value.

IID data. Although data are more likely to be non-IID in an FL environment, the data collected on mobile devices might in some cases be IID. We thus benchmark AdaSGD under two additional datasets (E-MNIST and CIFAR-100) with the staleness following D2. Figure 12 shows that our observations from Figure 10 hold also with IID data. As with non-IID data, Fedavg diverges also in the IID setting, and AdaSGD performs better than DynSGD on both datasets.

Differential privacy. Differential privacy [26] is a popular technique for privacy-preserving FL with formal guarantees [9]. We thus compare ADASGD against DYNSGD in a differentially private setup by perturbing the gradients as in the work of Abadi et al. [2]. We keep the previous setup (IID data with D2) and employ the MNIST dataset. Based on the work of Wu et al. [82], we fix the probability $\delta = 1/N^2 = \frac{1}{60000^2}$ and measure the privacy loss (ϵ) with the moments accountant

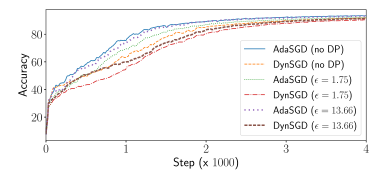


Fig. 13. Staleness awareness with differential privacy.

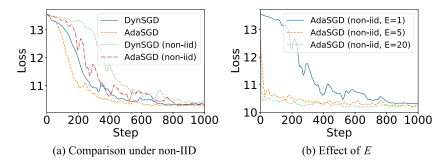


Fig. 14. Non-IID data impacts convergence. Larger E mitigates this impact.

approach [2] given the sampling ratio $(\frac{\text{mini-batch size}}{N} = \frac{100}{60000})$, the noise amplitude, and the total number of iterations.

Figure 13 demonstrates that the advantage of ADASGD over DynSGD also holds in the differentially private setup. A better privacy guarantee (i.e., smaller ϵ) slows down the convergence for both staleness-aware learning schemes.

Local updates. FL enables local updates—that is, multiple gradient computations on the same mobile device that result in a single global model update. *E* defines number of local updates [53]. We follow the same setup as for the experiment shown in Figure 14(a), and test ADASGD under non-IDD and different number of local updates for each worker (by employing the local updater of FLEET (Section 2)).

Figure 14(b) depicts that increasing *E* to 5 introduces a significant acceleration (in terms of steps) for the learning. Each update step involves five local updates that explain the speedup. Increasing the number of local updates to 20 provides a relatively smaller boost.

Mini-batch size variability. We empirically demonstrate the trade-off between a variable minibatch size and the convergence speed for the E-MNIST dataset. To isolate the effect of mini-batch size, we employ only one mobile device and perform synchronous updates (SSGD). We use two settings to compare the effect of a variable mini-batch size. In the first setting, we fix the minibatch size to n, whereas in the second setting, we sample the size from a Gaussian distribution $\mathcal{N}(\mu = \frac{n-1}{2}, \sigma = \frac{n-1}{6})$, based on the distribution of the output of I-Prof (see Figure 16(d)).

Figure 15(a) shows that the fixed scenario results in faster convergence as expected. The main reason is that a higher mini-batch size leads to a more robust gradient estimation. The SGD

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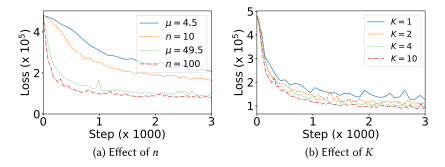


Fig. 15. Larger n or K results in faster (in terms of steps) and more stable convergence.

convergence theory also justifies this observation as the convergence time is inversely proportional to \sqrt{n} [88].

Aggregation size. The parameter K denotes the number of responses that ADASGD aggregates for each global model update (Section 2.3). To isolate the effect of K, we also deploy FLEET with a mobile device and perform synchronous updates (SSGD) for the E-MNIST dataset. Figure 15(b) shows that a larger K leads to a faster convergence and less noise in the learning curve w.r.t. loss, as ADASGD increases the robustness of each update by increasing K.

Choosing the optimal value for the parameter K involves a trade-off between the robustness of each update and the update frequency (as we discuss in Section 1). Increasing the value of K triggers a decrease in the model update throughput. Therefore, the model converges faster in terms of steps but slower in terms of time. The staleness in the accumulated gradients diminishes this increase in the robustness.

3.4 I-Prof Performance

We compare I-Prof against the profiler of MAUI [18], a mobile device profiler aiming to identify the most energy-consuming parts of the code and offload them to the cloud. MAUI predicts the energy by using a linear regression model (similar to the global model of I-Prof) on the number of CPU cycles ($\hat{E} = \theta_0 \cdot n$), to essentially capture how the size of the workload affects the energy (as in the work of Mittal et al. [60]). We adapt the profiler of MAUI to our setup by replacing the CPU cycles with the mini-batch size for two main reasons. First, our workload has a static code path so the number of CPU cycles on a particular mobile device is directly proportional to the mini-batch size. Second, measuring the number of executed CPU cycles requires root access that is not available on AWS.

We bootstrap the global model of I-Prof and the model of MAUI by pretraining on a training dataset. To this end, we use 15 mobile devices in AWS (that are different from the ones used for the rest of the experiments), assign them learning tasks with increasing mini-batch size until the computation time becomes two times the SLO, and collect their device information for each task. We rely on the same methodology to evaluate energy consumption but use only 3 mobile devices in our lab, as AWS prohibits energy measurements.

For testing, we use a different set of 20 commercial mobile devices in AWS, each performing requests for the image classification application (on MNIST), starting at different timestamps (login events) as shown in Figure 16(a). To ensure a precise comparison with MAUI, we add a *round-robin dispatcher* to the profiler component that alternates the requests from a given device between I-Prof and MAUI.

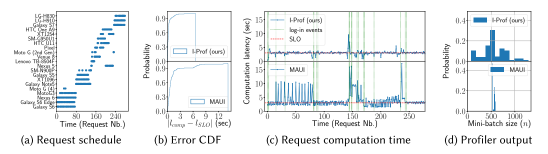


Fig. 16. I-Prof outperforms MAUI and drives the computation time closer to the SLO.

Computation time SLO. Figure 16(b) shows that I-Prof largely outperforms MAUI in terms of deviation from the computation time SLO. A total of 90% of approximately 280 learning tasks deviate from an SLO of 3 seconds by at most 0.75 seconds with I-Prof and 2.7 seconds with MAUI. This is the direct outcome of our design decisions. First, I-Prof adds dynamic features (e.g., the temperature of the device) to train its global model (Section 2.2). As a result, the predictions are more accurate for the first request of each user. Second, I-Prof uses a personalized model for each device that reduces the error (deviation from the SLO) with every subsequent request (Figure 16(c)). Figure 16(d) shows that the personalized models of I-Prof are able to output a wider range of minibatch sizes that better match the capabilities of individual devices. On the contrary, MAUI relies on a simple linear regression model that has acceptable accuracy for its use case but is inefficient when profiling heterogeneous mobile devices.

Energy SLO. To assess the ability of I-Prof to also target the energy SLO, we use the same setup as for the computation time, except on five mobile devices. We configure I-Prof with a significantly smaller error margin, $\epsilon=6*10^{-5}$ (Equation (2)), because the linear relation (capture by α as defined in Section 2.2) is significantly smaller for the energy than for the computation time (as shown in Figure 4). Figure 17 shows that I-Prof significantly outperforms MAUI in terms of deviation from the energy SLO. A total of 90% of 36 learning tasks deviate from an SLO of 0.075% battery drop by at most 0.01% for I-Prof and 0.19% for MAUI. The observation that I-Prof is able to closely match the latency SLO, whereas MAUI suffers from huge deviations, holds for the energy as well. The PA personalized models are able to quickly adapt to the state of the device as opposed to the linear model of MAUI that provides biased predictions.

Sensitivity analysis. The parameter ϵ regulates the adaptability of the personalized models of I-Prof. Figure 18 depicts the effect of the choice of ϵ on the learning procedure of the PA algorithm with a total of five mobile devices in AWS Device Farm. A larger value (Figure 18(b)) increases the error margin (see Equation (2)) and thereby leads to fewer updates. This results in a larger error (deviation from SLO) compared to a tuned value for ϵ (Figure 18(a)).

3.5 Resource Allocation

In this section, we evaluate the low-level execution of our resource allocation scheme (Section 2.4) and compare it against CALOREE [57], which is a state-of-the-art resource manager for mobile devices.

⁹AWS prohibits energy measurements, so we only rely on devices available in our lab, listed in their log-in order: Honor 10, Galaxy S8, Galaxy S7, Galaxy S4 mini, and Xperia E3.

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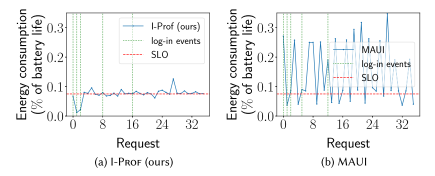


Fig. 17. I-PROF outperforms MAUI and drives the energy closer to the SLO.

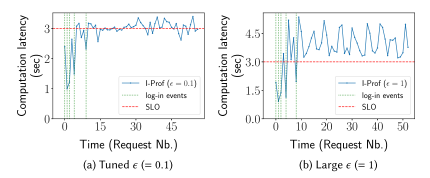


Fig. 18. Large aggressiveness (ϵ) prevents the convergence of the personalized models of I-Prof.

Table 2. Low-Level Performance Counters for SGD Computation on a Batch
Size of 10,000 Examples Executed on a Samsung Galaxy S7 Mobile Device

Monitored Event	Big Cores	All Cores		
Total branch loads	28,867,459,781.00	30,741,621,368.00		
Branch load misses	145,237,513.00	195,482,991.00		
Total L1 cache loads	223,053,108,577.00	242,798,977,320.00		
L1 cache misses	676,120,047.00	676,389,172.00		
Total L2 cache loads	1,357,014,880.00	1,500,271,296.00		
L2 cache misses	69,308,983.00	120,932,664.00		
Memory bus accesses	203,816,034.00	398,014,352.00		
Executed instructions	360,784,601,285.00	361,105,214,999.00		
Total CPU cycles	311,574,013,167.00	384,924,861,807.00		
Execution time (ms)	37,844	30,923		

Low-level big.LITTLE performance. For the ARM big.LITTLE architectures, our resource allocation scheme (Section 2.4) relies only on the big cores. In Table 2, we compare the low-level performance counters collected during the gradient computation on a batch size of 10,000 examples, executed on all eight cores of a Samsung Galaxy S7 device compared to only the four big cores. We can observe that even if gradient computation finishes 7 seconds faster by employing all cores, the execution efficiency is much lower: around 25% more CPU cycles for about the same number

Table 3.	Performance of CALOREE [57]
	on New Devices

Running Device	Deadline Error (%)
★Galaxy S7	1.4
Galaxy S8	9
Honor 9	46
Honor 10	255

Note: Galaxy S7, marked with a star (\star) , was used for training.

of instructions. This experiment highlights one of the reasons behind the superior performance of our resource allocation scheme.

Comparison with CALOREE. The goal of CALOREE is to optimize resource allocation for the workload execution to meet its predefined deadline while minimizing the energy consumption. To this end, CALOREE profiles the target device by running the workload with different resource configurations (i.e., number of cores, core frequency). Since FLEET executes on non-rooted mobile devices, we can only adapt the number of big/little cores (but not their frequencies). By varying the number of cores allocated to our workload (i.e., gradient computation), we obtain the energy consumption of each possible configuration. From these configurations, CALOREE only selects those with the optimal energy consumption (the lower convex hull) that are packed in the so-called **performance hash table (PHT)**.

CALOREE on new devices. In their thorough evaluation, the authors of CALOREE used the same device for training and running the workloads. Therefore, we first benchmark the performance of CALOREE when running on new devices. We employ Galaxy S7 to collect the PHT and set the mini-batch size to that returned by I-Prof for a latency SLO of 3 seconds (Section 3.4). We then run this workload with CALOREE on different mobile devices, as shown in Table 3.

The performance of CALOREE degrades significantly when running on a different device than the one used for training. The first line of Table 3 shows the baseline error when running on the same device. The error increases more than $6\times$ for a device with similar architecture and the same vendor (Galaxy S8) and more than $32\times$ for a device of similar architecture but different vendor (Honor 9 and 10). This significant increase for the error is due to the heterogeneity of the mobile devices that make PHTs not applicable across different device models.

CALOREE vs FLEET. We evaluate the resource allocation scheme of FLEET by comparing it to the ideal environment for CALOREE—that is, training and running on the same device (a setup nevertheless difficult to achieve in FL with millions of devices). Following the setup used for the energy SLO evaluation (Section 3.4), we employ five devices and fix the size of the workload (mini-batch size) based on the output of I-Prof. In particular, we set the mini-batch size to 280, 4,320, 6,720, 5,280, and 1,200 for the devices shown in Figure 19, respectively. We set the deadline of CALOREE either equal or double that of the computation latency of FLEET. We take 10 measurements and report on the median, 10th, and 90th percentile.

Figure 19 shows the fact that in the ideal environment for CALOREE and even with double the time budget (giving more flexibility to CALOREE), FLEET has comparable energy consumption. Since gradient computation is a compute-intensive task with high temporal and spacial cache locality, the configuration changes performed by CALOREE negatively impact the execution time and cancel any energy saved by its advanced resource allocation scheme. In addition, the fewer configuration knobs available on non-rooted Android devices limit the full potential of CALOREE.

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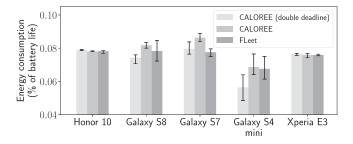


Fig. 19. Resource allocation of FLEET vs CALOREE.

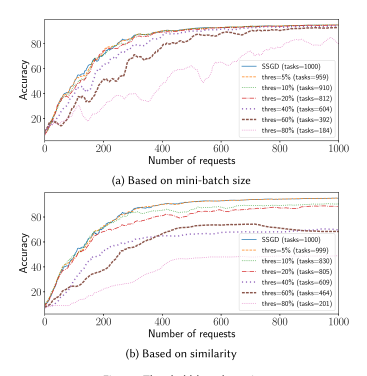


Fig. 20. Threshold-based pruning.

3.6 Learning Task Assignment Control

The controller of FLEET employs a threshold to prune learning tasks and thus control the trade-off between the cost of the gradient computations and the model prediction quality. This threshold can be based either on the mini-batch size or on the similarity values (see Figure 2). To evaluate this trade-off, we employ non-IID MNIST with the mini-batch size following a Gaussian distribution $\mathcal{N}(\mu=100,\sigma=33)$ (based on the distribution of the output of I-Prof shown in Figure 16(d)), and set the threshold to the n-th percentile of the past values. Figure 20 illustrates that a threshold on the mini-batch size is more effective in pruning the less useful gradient computations than a threshold on the similarity. Figure 20(a) shows that even dropping up to 39.2% of the gradients (with the smallest mini-batch size) has a negligible impact on the accuracy (less than 2.2%). Figure 20(b) shows that one can drop 17% of the most similar gradients with an accuracy impact of 4.8%.

Application	Image Classification			
Compression	E-MNIST	MNIST	CIFAR-100	
Original	52 (KiB)	49 (KiB)	2.2 (MiB)	
GZIP + Kryo	42 (KiB)	41 (KiB)	2 (MiB)	

Table 4. Transfer (Download/Upload) Size Per Request

3.7 Impact of FLEET on Mobile Devices

The usability of FLEET greatly depends on the fact that the impact on the user device is low enough such that a standard usage of the device is guaranteed. The impact may concern the CPU usage, the memory requirements, or the network bandwidth.

Impact on the CPU. To measure the impact of FLEET workers on other collocated apps, we employ a benchmark (called BaseApp) that performs counter increment operations (CTROPS). We then deploy FLEET along with BaseApp on Honor 9 and execute 20 requests for the E-MNIST and CIFAR-100 datasets. We observe that the throughput of BaseApp drops only by 2.9% and 3.1%, respectively. This shows that FLEET can be executed with a minimal impact even on a CPU-intensive collocated application.

Memory requirements. We measure the memory requirements of FLEET with on Honor 9. The memory consumption before the learning task is 14.5 MiB (maximum of 4 GiB available). Performing the ML task raises the memory usage to 22 MiB for MNIST and 135 MiB for CIFAR-100, thus highlighting the impact of the model size.

Bandwidth consumption. We measure the size of the data that Honor 9 receives from the server (i.e., the model parameters) and sends back to the server (i.e., the gradients). We also employ GZIP to compress the KRIO objects received and sent, and compare with a setting without any compression. Table 4 depicts that compression reduces the transfer size by 19% for E-MNIST, 16% for MNIST, and 10% for CIFAR-100. The difference in the original size among the three datasets is primarily due to the difference in the amount of model parameters, and the difference in the compression reduction is due to the data structures employed to host model parameters.

4 RELATED WORK

Distributed ML. Adam [12] and TensorFlow [1] adopt the parameter server architecture [50] for scaling out on high-end machines and typically require cross-worker communication. FLEET also follows the parameter server architecture by maintaining a global model on the server. However, FLEET avoids cross-worker communication, which is impractical for mobile workers due to the device churn.

A common approach for large-scale ML is to control the amount of staleness for boosting convergence [19, 68]. In Online FL, staleness cannot be controlled, as this would impact the model update frequency. The workers perform learning tasks asynchronously with end-to-end latencies that can differ significantly (due to device heterogeneity and network variability) or even become infinite (user disconnects).

Petuum [84] and TensorFlow handle faults (worker crashes) by checkpointing and repartitioning the model across the workers whenever failures are detected. In a setting with mobile devices, such failures may appear very often, thus increasing the overhead for checkpointing and repartitioning. FLEET does not require any fault-tolerance mechanism for its workers, as from a global learning perspective, they can be viewed as stateless.

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Federated learning. To minimize the impact on mobile devices, Standard FL algorithms [8, 39, 53, 74] require the learning task to be executed only when the devices are idle, plugged in, and on a free wireless connection. However, in Sections 3.1 and 3.2, we have shown that these requirements may drastically impact the performance of some applications. It is worth noting that techniques for reducing the communication overhead [39] or increasing the robustness against adversarial users [20, 21] are orthogonal to the online characteristic so they can be plugged into FLEET. Recently, other research works have started to investigate semi-synchronous and asynchronous learning algorithms adapted to heterogeneous federated environments. Xie et al. [83] propose an asynchronous algorithm for federated optimization that uses a weighted average to update the global model, where the mixing weight is set dynamically as a function of the staleness. Shi et al. [73] propose a similar aggregation algorithm based on model timestamp where the weights assigned to model updates decreases as the staleness value increase. In the same vein, Zhou et al. [90] propose a caching mechanism with a staleness-based weighted aggregation algorithm. However, previous work computes the weights of stale model updates based on a linear relationship to the staleness. ADASGD relies on an exponential dampening function that adapts to past staleness values and finds the most appropriate weights for the stale model updates. In addition, ADASGD proposes a similarity-based boosting scheme that increases the weight of gradients computed on diverse and rare data.

Performance prediction for mobile devices. Estimating the computation time or energy consumption of an application running on a mobile device is a very broad area of research. Existing approaches [10, 13, 14, 36, 45, 86] target multiple applications generally executing on a single device. They typically benchmark the device or monitor hardware and operating system level counters that require root access. In contrast, FLEET targets a single application executing in the same way across a large range of devices. I-Prof poses a negligible overhead, as it employs features only from the standard Android API to enable Online FL, and requires no benchmarking of new devices. I-Prof is designed to make predictions for *unknown* devices.

Neurosurgeon [41] is a scheduler that minimizes the end-to-end computation time of inference tasks (whereas FLEET focuses on training tasks), by choosing the optimal partition for a neural network and offloading computations to the cloud. The profiler of Neurosurgeon only uses workload-specific features (e.g., number of filters or neurons) to estimate computation time and energy, and ignores device-specific features. By contrast, mobile phones, as targeted by I-Prof, ¹⁰ exhibit a wide range of device-specific characteristics that significantly impact their latency and energy consumption (see Figure 4).

Systems such as CALOREE [57] and LEO [58] profile mobile devices under different system configurations and train an ML model to determine the ones that minimize the energy consumption. They rely on a control loop to switch between these configurations such that the application does not miss the preset deadline. Due to the restrictions of the standard Android API, the available knobs are limited in our setup. For our application (i.e., gradient computation), we show that a simple resource allocation scheme (Section 2.4) is preferable even in comparison with an ideal execution model.

5 CONCLUDING REMARKS

This article presented FLEET, the first system that enables *online* ML at the edge. FLEET employs I-Prof, a new *ML-based profiler* that determines the ML workload that each device can perform

¹⁰In their in-depth experimental evaluation the authors of [41] consider a single hardware platform and not Android mobile devices

within predefined energy and computation time SLOs. FLEET also makes use of ADASGD, a new *staleness-aware learning* algorithm that is optimized for Online FL. We showed the performance of I-Prof and ADASGD on commercial Android devices with popular benchmarks. I-Prof can be used for any iterative computation with embarrassingly parallel CPU-bound tasks, whereas ADASGD is specific to gradient descent and thus ML applications. In our performance evaluation, we do not focus on network and scalability aspects that are orthogonal to our work and addressed in existing literature. We also highlight that transferring the label and device information (see Figure 2) poses a negligible network overhead compared to transferring the relatively large FL learning models. Finally, addressing biases is an important problem even in cloud-based online ML (not only FL) that we also do not address in this work. For Online FL, we arguably need to keep some of these biases (e.g., recommend more politics to people that wake up earlier). Of course, diversity is also crucial.

Although we believe FLEET to represent a significant advance for online learning at the edge, there is still room for improvement. First, for the energy prediction, I-Prof requires access to the CPU usage that is considered as a security flaw on some Android builds and thus not exposed to all applications. In this case, I-Prof requires a set of additional permissions that belong to services from Android Runtime. Second, the transfer of the label distribution from the worker to the server introduces a potential privacy leakage. However, we highlight that the server has access only to the indices of the labels and not their values. In this article, we focus on the protection of the input features and mention the possibility to deactivate the similarity-based boosting feature of AdaSGD in the case that this leakage is detrimental. We plan to investigate noise addition techniques for bounding this leakage [26] in our future work. Finally, theoretically proving the convergence of AdaSGD is non-trivial due to the unbounded staleness and the non-IID datasets among the workers. In this respect, a dissimilarity assumption similar to that of Li et al. [51] may facilitate the derivation of the proof.

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