On the Tradeoff between Energy, Precision, and Accuracy in Federated Quantized Neural Networks

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Abstract—Deploying federated learning (FL) over wireless networks with resource-constrained devices requires balancing between accuracy, energy efficiency, and precision. Prior art on FL often requires devices to train deep neural networks (DNNs) using a 32-bit precision level for data representation to improve accuracy. However, such algorithms are impractical for resource-constrained devices since DNNs could require execution of millions of operations. Thus, training DNNs with a high precision level incurs a high energy cost for FL. In this paper, a quantized FL framework, that represents data with a finite level of precision in both local training and uplink transmission, is proposed. Here, the finite level of precision is captured through the use of quantized neural networks (QNNs) that quantize weights and activations in fixed-precision format. In the considered FL model, each device trains its QNN and transmits a quantized training result to the base station. Energy models for the local training and the transmission with the quantization are rigorously derived. An energy minimization problem is formulated with respect to the level of precision while ensuring convergence. To solve the problem, we first analytically derive the FL convergence rate and use a line search method. Simulation results show that our FL framework can reduce energy consumption by up to 53% compared to a standard FL model. The results also shed light on the tradeoff between precision, energy, and accuracy in FL over wireless networks.

I. INTRODUCTION

The emergence of federated learning (FL) ushered in a new era of distributed inference that can alleviate data privacy concerns \[^1\]. In FL, massively distributed mobile devices and a central server (e.g., a base station (BS)) collaboratively train a shared model without requiring devices to share raw data. Many FL algorithms employ complex deep neural networks (DNNs) to achieve a high accuracy by allocating many bits for the precision level in data representation \[^2\]. DNN structures, such as convolutional neural networks (CNNs), can have tens of millions of parameters and billions of multiply-accumulate (MAC) operations \[^3\]. In practice, the energy consumed for computation and memory access is proportional to the level of precision \[^4\]. Hence, computationally intensive neural networks with a conventional 32 bits full precision level may not be suitable for deployment on energy-constrained mobile and Internet of Things (IoT) devices. In addition, a DNN may increase the energy consumption of transmitting a training result due to the large model size. To design an energy-efficient FL scheme, one could reduce the level of precision to decrease the energy consumption for the computation and transmission. However, the reduced precision level could introduce quantization error that degrades the accuracy and the convergence rate of FL. Therefore, deploying real-world FL frameworks over wireless systems requires one to balance precision, accuracy, and energy efficiency – a major challenge facing future distributed learning frameworks.

Remarkably, despite the surge in research on the use of FL, only a handful of works in \[^5\]–\[^10\] have studied the energy efficiency of FL from a system-level perspective. A novel analytical framework that derived energy efficiency of FL algorithms in terms of the carbon footprint was proposed in \[^5\]. Meanwhile, in \[^6\], the authors formulated an energy minimization problem under heterogeneous power constraints of mobile devices. The work in \[^7\] investigated a resource allocation problem to minimize the total energy consumption considering the convergence rate. In \[^8\], the energy consumption of FL was minimized by controlling workloads of each device, which has heterogeneous computing resources. The work in \[^9\] proposed a quantization scheme for both uplink and downlink transmission in FL and analyzed the impact of the quantization on the convergence rate. The authors in \[^10\] considered a novel FL setting, in which each device trains a binary neural network so as to improve the energy efficiency of transmission by uploading the binary parameters to the server.

However, the works in \[^5\]–\[^9\] did not consider the energy efficiency of their DNN structure during training. Since devices have limited computing and memory resources, deploying an energy-efficient DNN will be a more appropriate way to reduce the energy consumption of FL. Although the work in \[^11\] considered binarized neural networks during training, this work did not optimize the quantization levels of the neural network to balance the tradeoff between precision and energy. To the best of our knowledge, there is no work that jointly considers the tradeoff between precision, energy, and accuracy.

The main contribution of this paper is a novel energy-efficient quantized FL framework that can represent data with a finite level of precision in both local training and uplink transmission. In our FL model, each device trains a quantized neural network (QNN), whose weights and activations are quantized with a finite level of precision, so as to decrease en-

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energy consumption for computation and memory access. After training, each device quantizes the result with the same level of precision used in the local training and transmits it to the BS. The BS aggregates the received information to generate a new global model and broadcasts it back to the devices. To quantify the energy consumption, we propose rigorous energy model for the local training based on the physical structure of a processing chip. We also derive the energy model for the uplink transmission considering the quantization. To achieve a high accuracy, FL requires a high level of precision at the cost of increased total energy consumption. Meanwhile, although, a low level of precision can decrease the energy consumption per iteration, it will decrease the convergence rate to achieve a target accuracy. Thus, there is a need for a new approach to analyze and optimize the tradeoff between precision, energy, and accuracy. To this end, we formulate an optimization problem by controlling the level of precision to minimize the total energy consumption while ensuring convergence with a target accuracy. To solve the problem, we first analytically derive the convergence rate of our FL framework and use a line search method to numerically find the local optimal solution. Simulation results show that our FL model can reduce the energy consumption up to 53% compared to a standard FL network.

Fig. 1: An illustration of the quantized FL model over wireless network.

In our model, each device trains a QNN of identical structure using \( n \) bits of precision for quantization. We can express data more precisely if we increase \( n \) at the cost of more energy usage. We can represent any given number in fixed-point format such as \([\Omega, \omega]\), where \( \Omega \) is the integer part and \( \omega \) is the fractional part of the given number. Here, we use one bit to represent the integer part and \( (n-1) \) bits for the fractional part. Then, the smallest positive number we can present would be \( \kappa = 2^{-n+1} \), and the possible range of numbers with \( n \) bits will be \([-1, 1 - 2^{-n+1}]\). Note that a QNN restricts the value of weights to \([-1, 1] \). We consider a QNN structure whose weights and activations are quantized in fixed-point format rather than conventional 32-bit floating-point format. A. Quantized Neural Networks

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Algorithm 1: Quantized FL Algorithm

Input: $K$, $I$, initial model $w_0$, $t = 0$, target accuracy $\epsilon$

1 repeat
2 The BS randomly select a subset of devices $N_t$ and broadcasts the $w_t$ to the selected devices;
3 Each device $k \in N_t$ trains $w^k_t$ by running $I$ steps of SGD as (6);
4 Each device $k \in N_t$ transmits $d^{Q,k}_{t+1}$ to the BS;
5 The BS generates a new global model $w_{t+1} = w_t + \frac{1}{K} \sum_{k \in N_t} d^{Q,k}_{t+1}$;
6 $t \leftarrow t + 1$;
7 until target accuracy $\epsilon$ is satisfied;

where $\eta$ is the learning rate and $\xi$ is a mini-batch for the current update. Then, we restrict the values of $w^k_t$ to $[-1, 1]$ as $w^k_t \leftarrow \operatorname{clip}(w^k_t, -1, 1)$, where $\operatorname{clip}(\cdot, -1, 1)$ projects each input to 1 (-1) for any input larger (smaller) than 1 (-1), or returns the same value as the input. Otherwise, $w^k_t$ can become very large without a meaningful impact on quantization [11]. After each training, $w^k_t$ are quantized as $w^{Q,k}_t$.

B. FL model

For learning, without loss of generality, we adopt FedAvg [2] to solve problem (3). At each global iteration $t$, the BS randomly selects a set of devices $N_t$ with $|N_t| = K$ and broadcasts the current global model $w_t$ to the scheduled devices. Each device in $N_t$ trains its local model based on the received global model by running $I$ steps of SGD on its local loss function as below

$$w_{t+1}^{k} = w_{t}^{k} - \eta_t \nabla E_k(w_{t}^{Q,k}, \xi_{t}), \forall \tau = 1, \ldots, I,$$  \hspace{1cm} (7)

where $\eta_t$ is the learning rate at global iteration $t$. Note that unscheduled devices do not perform local training. Then, $K$ devices calculates the model update $d_{t+1}^{Q,k} = w_{t+1}^{k} - w_{t}^{k}$, where $w_{t+1}^{k} = w_{t+1}^{k,I}$ and $w_{t}^{k} = w_{t}^{k,0}$ [9]. Typically, $d_{t+1}^{Q,k}$ has a millions of elements. It is not practical to send $d_{t+1}^{Q,k}$ with full precision for energy-constrained devices. Hence, we apply the same quantization scheme used in QNNs to $d_{t+1}^{Q,k}$ and denote its quantization result as $d_{t+1}^{Q,k}$. Then, $K$ devices transmit their model update to the BS. The received model updates are averaged by the BS, and the next global model will be generated as below

$$w_{t+1} = w_{t} + \frac{1}{K} \sum_{k \in N_t} d^{Q,k}_{t+1}.$$  \hspace{1cm} (8)

The FL system repeats this process until the global loss function converges to a target accuracy constraint $\epsilon$. We summarize the aforementioned algorithm in Algorithm 1.

Next, we propose the energy model for the computation and the transmission for our FL system.

C. Computing and Transmission model

1) Computing model: We consider a typical two dimensional processing chip for CNNs as shown in Fig. 2 [4]. This chip has a parallel neuron array, $p$ MAC units, and two levels of memory: a main and a local buffer. A main buffer stores the current layers’ weights and activations, while a local buffer caches currently used weights and activations. From

![Fig. 2: An illustration of the two dimensional processing chip.](image)

[13], we use the energy model of a MAC operation for $n$ levels of precision $E_{MAC}(n) = A(n/n_{max})^a$, where $A > 0$, $1 < \alpha < 2$, and $n_{max}$ is the maximum precision level. Here, a MAC operation includes the operation of a layer such as output calculation, batch normalization, activation, and weight update. Then, the energy consumption for accessing a local buffer $E_l$ can be modeled as $E_{MAC}(n)$, and the energy for accessing a main buffer is $E_m = 2E_{MAC}(n)$ [4].

The energy consumption of device $k$ for one iteration of local training is given by $E_{C,k}^{C,k}(n)$ when $n$ bits are used for the precision level in the quantization. Then, $E_{C,k}^{C,k}(n)$ is the sum of the computing energy $E_C(n)$, the access energy for fetching weights from the buffers $E_W(n)$, and the access energy for fetching activations from the buffers $E_A(n)$, as follows [13]:

$$E_{C,k}^{C,k}(n) = E_C(n) + E_W(n) + E_A(n),$$
$$E_C(n) = E_{MAC}(n)N_c + 4O_s E_{MAC}(n_{max}),$$
$$E_W(n) = E_mN_c + E_lN_c\sqrt{n/p_{max}},$$
$$E_A(n) = 2E_mO_s + E_lN_c\sqrt{n/p_{max}},$$

where $N_c$ is the number of MAC operations, $N_s$ is the number of weights, and $O_s$ is the number of intermediate outputs throughout the network. For $E_C$, in a QNN, batch normalization, activation function, gradient calculation, and weight update are done in full-precision $n_{max}$ to each output $O_s$ [11]. Once we fetch weights from a main to a local buffer, they can be reused in the local buffer afterward as shown in $E_W(n)$. In Fig. 2 a MAC unit fetches weights from a local buffer for computation. Since we are using a two dimensional MAC array of $p$ MAC units, they can share fetched weights with the same row and column, which has $\sqrt{p}$ MAC units respectively. In addition, a MAC unit can fetch more weights due to the quantization with $n$ bits compared with when weights are represented in $n_{max}$ bits. Thus, we can reduce the access to a local buffer by the amount of $\sqrt{n}/p_{max}$. A similar process applies to $E_A$ since activations are fetched (stored) from (to) the main buffer.

2) Transmission Model: We use orthogonal frequency domain multiple access (OFDMA) to transmit a model update to the BS. The achievable rate of device $k$ is given by

$$r_k = B \log_2 \left(1 + \frac{P_{h_k}}{N_0 B}\right),$$  \hspace{1cm} (10)

where $B$ is the allocated bandwidth, $h_k$ is the channel gain between device $k$ and the BS, $P$ is the transmit power of each device, and $N_0$ is the power spectral density of white noise.
After local training, device $k$ will transmit $d_t^{Q,k}$ to the BS at given global iteration $t$. Then, the transmission time $T_k$ for uploading $d_t^{Q,k}$ is given by
\[
T_k(n) = \frac{||d_t^{Q,k}||}{r_k} = \frac{||d_t^k||_n}{r_k n_{\text{max}}}. \tag{11}
\]
Note that $d_t^{Q,k}$ is quantized with $n$ bits of precision while $d_t^k$ is represented with $n_{\text{max}}$ bits. Then, the energy consumption for the uplink transmission is given by
\[
E^{U,L,k}(n) = T_k(n) \times P = \frac{P||d_t^k||_n}{B \log_2 \left(1 + \frac{P h}{N_0 B}ight) n_{\text{max}}}. \tag{12}
\]

In the following section, we formulate an energy minimizing problem based on the derived energy models.

III. PROPOSED APPROACH FOR ENERGY-EFFICIENT FEDERATED QNN

We formulate an energy minimization problem while ensuring convergence under a target accuracy. A tradeoff exists between the energy consumption and the convergence rate with respect to $n$. Hence, finding the optimal $n$ is important to balance the tradeoff and achieve the target accuracy. We propose a numerical method to solve this problem.

We aim to minimize the expected total energy consumption until convergence under the target accuracy as follows:

\[
\min_n \mathbb{E} \left[ \sum_{t=1}^{T} \sum_{k \in N_t} E^{U,L,k}(n) + IE^{C,k}(n) \right] \tag{13a}
\]

s.t. $n \in [1, \ldots, n_{\text{max}}]$, \quad \mathbb{E}[F(w_T) - F(w^*)] \leq \epsilon, \tag{13b}

where $T$ is the number of local iterations, $\mathbb{E}[F(w_T)]$ is the expectation of global loss function after $T$ global iteration, $F(w^*)$ is the minimum value of $F$, and $\epsilon$ is the target accuracy.

Since $K$ devices are randomly selected at each global iteration, we can derive the expectation of the objective function of (13a) as follows
\[
f_E(n) = \mathbb{E} \left[ \sum_{t=1}^{T} \sum_{k \in N_t} E^{U,L,k}(n) + IE^{C,k}(n) \right] = \frac{KT}{N} \sum_{k=1}^{N} \left\{ E^{U,L,k}(n) + IE^{C,k}(n) \right\}. \tag{14}
\]

To represent $T$ with respect to $\epsilon$, we assume that the loss function is $L$-smooth, $\mu$-strongly convex and that the variance and the squared norm of the stochastic gradient are bounded by $\sigma_k^2$ and $G$ for device $k$, $\forall k \in N$, respectively. Before we present the expression of $T$, in the following lemma, we will first analyze the quantization error of the stochastic quantization in Sec. II.

Lemma 1. For the stochastic quantization $Q(\cdot)$, a scalar value $w$, and a vector $w \in \mathbb{R}^d$, we have
\[
\mathbb{E}[Q(w)] = w, \quad \mathbb{E}[(Q(w) - w)^2] \leq \frac{1}{2n^2}, \tag{15}
\]
\[
\mathbb{E}[Q(w)] = w, \quad \mathbb{E}[(Q(w) - w)^2] \leq \frac{d}{2n^2}. \tag{16}
\]

Proof. We first derive $\mathbb{E}[Q(w)]$ as
\[
\mathbb{E}[Q(w)] = \left[ w + \frac{\kappa - w}{\kappa} \right] = w. \tag{17}
\]
Similarly, $\mathbb{E}[(Q(w) - w)^2]$ can be obtained as
\[
\mathbb{E}[(Q(w) - w)^2] = \left[ w - w \right]^2 \frac{\kappa + w - w}{\kappa} = \frac{\kappa^2}{4} = \frac{1}{2n^2}. \tag{18}
\]

From Lemma 1, we can see that our quantization scheme is unbiased as its expectation is zero. However, the quantization error can still increase for a large model. We next leverage the results of Lemma 1, and so as to derive $T$ with respect to $\epsilon$ in the following proposition.

Proposition 1. For learning rate $\eta_t = \frac{\beta}{4 + \gamma}$, $L < \frac{2n^2}{2n^2 + 1}$, $\beta > \frac{1}{\mu}$, and $\gamma > 0$, we have
\[
\mathbb{E}[F(w_T) - F(w^*)] \leq \frac{L}{2} \frac{v}{T + \gamma}, \tag{20}
\]
where $v$ is
\[
v = \sum_{k=1}^{N} \frac{\sigma_k^2}{G^2} \left[ 1 + \frac{d}{2n^2} \right] + 4(1 - I)^2 G^2 + \frac{4(N - K)}{K(N - 1)} I^2 G^2. \tag{21}
\]

Proof. The complete proof is omitted due to space limitations. Essentially, proposition 1 can be proven by using Lemma 1 the convergence result with a quantized model update [9], and replacing the SGD weight update in [14] with [7].

From Proposition 1, we let (20) be upper bounded by $\epsilon$ in (13c) as follows
\[
\mathbb{E}[F(w_T) - F(w^*)] \leq \frac{L}{2} \frac{v}{T + \gamma} \leq \epsilon. \tag{22}
\]

We then take equality in (22) to obtain $T = Lv/(2\epsilon) - \gamma$ and approximate the problem as
\[
\min_n \frac{K}{N} \left[ \frac{L v}{2 \epsilon} - \gamma \right] \sum_{k=1}^{N} \{ E^{U,L,k}(n) + IE^{C,k}(n) \} = f_E(n) \tag{23a}
\]

s.t. $n \in [1, \ldots, n_{\text{max}}]$. \tag{23b}

Note that any optimal solution $n^*$ from problem (23a) can satisfy problem (13a) [13]. For any feasible $T$ from (22), we can always choose $T_0 > T$ such that $T_0$ satisfies (13c).
Now, we relax \( n \) as a continuous variable, which will be rounded back to an integer value. From (9), (12), and (21), we can observe that, as \( n \) increases, \( E_{UL,k}(n) \) and \( E_{C,k}(n) \) becomes larger while \( T \) decreases. Hence, we can know that a local optimal \( n^* \) may exist for minimizing \( f_E(n) \). Since \( f_E(n) \) is differentiable with respect to \( n \) in the given range, we can find \( n^* \) by solving \( \partial f_E(n)/\partial n = 0 \) from Fermat’s Theorem [16]. Although it is difficult to derive \( n^* \) analytically, we can obtain it numerically using a line search method. Hence, we can find a local optimal solution, which minimizes the total energy consumption under the given target accuracy.

IV. Simulation Results

For our simulations, we uniformly deploy \( N = 50 \) devices over a square area of size \( 100 \text{ m} \times 100 \text{ m} \) serviced by one BS at the center, and we assume a Rayleigh fading channel with a path loss exponent of 2. We also use MNIST dataset. Unless stated otherwise, we use \( P = 100 \text{ mW}, B = 10 \text{ MHz}, N_0 = -100 \text{ dBm}, K = 30, I = 5, n_{\text{max}} = 32 \text{ bits}, \epsilon = 0.01, \beta = 5, L = 1, \mu = 1, \gamma = 1, \sigma_k = 1, \) and \( G = \sqrt{4Lc}, \forall k = 1, \ldots, N \) [17]. For the computing model, we set \( A = 3.7 \text{ bit} \) and \( \alpha = 1.25 \) as done in [13], and we assume that each device has the same architecture of the processing chip. All statistical results are averaged over 10000 independent runs.

Figure 3 shows the total energy consumption of the FL system until convergence for varying levels of precision \( n \). In Fig. 3 we assume a QNN structure with two convolutional layers: 32 kernels of size \( 3 \times 3 \) with one padding and three of strides and 32 kernels of size \( 3 \times 3 \) with one padding and two of strides, each followed by 2 \times 2 max pooling. Then, we have one dense layer of 220 neurons and one fully-connected layer. In this setting, we have \( N_\ell = 20.64 \times 10^6, N_s = 0.18 \times 10^6 \), and \( O_s = 1354 \). From this figure, we can see that the total energy consumption decreases and then increases with \( n \). This is because when \( n \) is small, quantization error becomes large as shown in Lemma 1 which slows down the convergence rate in (20). However, as \( n \) increases, the energy consumption for the local training and transmission also increases. Hence, a very small or very high \( n \) may induce undesired large quantization error or unnecessary energy consumption due to a high level of precision. From this figure, we can see that \( n = 10 \) can be optimal for minimizing energy consumption for our system.

Figure 4 shows the optimal level of precision \( n^* \) when varying the number of local iterations \( I \). We use the same CNN architecture in Fig. 3. We can observe that \( n^* \) increases with \( I \). This is because, as \( I \) increases, the local models converge to the local optimal faster as SGD averages out the effect of quantization error [11]. Hence, a lower \( n \) can be chosen by leveraging the increased \( I \) to minimize the total energy consumption. We can observe that only \( n = 7 \) is required at \( I = 20 \) while we need \( n = 10 \) at \( I = 3 \).

Figure 5 presents the optimal level of precision \( n^* \) for varying model size \( d \). Note that \( d \) equals to the number of model parameters \( N_s \). To scale the number of MAC operations for increasing \( d \) accordingly, we set \( N_c = 0.5 \times 10^3 N_s \). From Fig. 5, we can see that \( n^* \) increases with \( d \). From Lemma 1 the quantization error accumulates as \( d \) increases. This directly affects the convergence rate in (20) resulting in both increased global iterations and the total energy consumption. Therefore, to mitigate the increasing quantization error from increasing \( d \), a larger level of precision may be chosen.

Figure 6 shows the total energy consumption and \( n^* \) when varying a target accuracy \( \epsilon \). For these results, we use the same CNN architecture as Fig. 3. We can see that a higher accuracy level requires larger total energy consumption and
more bits for data representation to mitigate the quantization error. In addition, the FL system needs a more number of global iterations to achieve $\epsilon$ from (22). As $\epsilon$ becomes looser, a lower $n_*$ can be chosen. From Fig. 6 we can see that an additional 127 J energy is required to increase $\epsilon$ from 0.01 to 0.001 while one additional bit of precision is needed.

Figure 7 compares the total energy consumption until convergence for varying the size of CNN models with the baseline that uses standard 32 bits for data representation. Case 1 is assumed to be CNN of two layers and have $d = 25.6 \times 10^6$ and $N_c = 1.8 \times 10^9$. Case 2 and 3 are assumed to be CNN of five layers. They have $61.6 \times 10^6$ and $83.5 \times 10^6$ number of parameters and $0.35 \times 10^6$ and $83.5 \times 10^6$ number of MAC operations, respectively. Case 4 is 7 layers of CNN with $d = 115.1 \times 10^6$ and $N_c = 10.4 \times 10^9$. Lastly, we assume Case 5 is 9 layers of CNN with $d = 138.6 \times 10^6$ and $15.5 \times 10^9$. The corresponding $n_*$ are 13, 14, 15, 15, respectively. We can see that our FL scheme is more effective for CNN models with a large model size. Note that Case 5 has 138.8 M weights while Case 1 has 25.6 M weights. In particular, for Case 5, we can reduce the total energy consumption up to 53% compared to the baseline.

V. CONCLUSION

In this paper, we have studied the problem of energy-efficient quantized FL over wireless networks. We have presented the energy model for the quantized FL based on the physical structure of a processing chip and the convergence rate. Then, we have formulated an energy minimization problem that considers a level of precision in the quantized FL. To solve this problem, we have used a line search method. Simulation results have shown that our model requires much less energy than a standard FL model for convergence. The results particularly show significant improvements when the local models rely on large neural networks. In essence, this work provides the first holistic study of the tradeoff between energy, precision, and accuracy for FL over wireless networks.

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