PREDICTIVE PRECOMPUTE WITH RECURRENT NEURAL NETWORKS

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ABSTRACT

In both mobile and web applications, speeding up user interface response times can often lead to significant improvements in user engagement. A common technique to improve responsiveness is to precompute data ahead of time for specific activities. However, simply precomputing data for all user and activity combinations is prohibitive at scale due to both network constraints and server-side computational costs. It is therefore important to accurately predict per-user application usage in order to minimize wasted precomputation ("predictive precompute"). In this paper, we describe the novel application of recurrent neural networks (RNNs) for predictive precompute. We compare their performance with traditional machine learning models, and share findings from their large-scale production use at Facebook. We demonstrate that RNN models improve prediction accuracy, eliminate most feature engineering steps, and reduce the computational cost of serving predictions by an order of magnitude.

1 INTRODUCTION

The relationship between application latency and user engagement is well-known; improving the responsiveness of an application by just a few seconds can result in significant increases in user engagement due to the limited attention span of users (Palmer, 2016).

In modern applications, the most common source of latency is data fetching. To serve the content required for a particular activity, one might first make a network request, retrieve the raw content from a database, and then apply some additional processing — each of which may take a significant amount of time. One common strategy to improve responsiveness from the user’s perspective is to precompute the results (prefetching them ahead of time) so that it is immediately available for the user.

For any individual activity, the simplest precompute strategy is to just always perform a precompute at application startup. However, this causes issues both client-side and server-side:

- On the client side (especially on mobile clients), aggressive precompute adversely affects cellular data usage, application startup time, and battery usage, all of which negatively impact user engagement.

- On the server side, the computational cost of the data fetches becomes significant at scale, especially during peak hours where computational resources are relatively scarce.

To minimize computational costs, one solution is to use an approach we call predictive precompute: we can predict the probability that a user will access a particular activity given the current application state and their historical access logs. We then only precompute data when the probability surpasses a certain threshold, significantly reducing the proportion of wasted precompute.

The key to this approach is accurately predicting user access probabilities. Estimating probabilities translates well into a standard machine learning problem, where existing research offers some solutions: for example, (Wang et al., 2015) describe a system to precompute links embedded in Twitter posts using linear regression on various content-specific features and (Sarker et al., 2019) demonstrate the effectiveness of decision tree models for context-aware smartphone usage predictions. However, it is often difficult to “feature engineer” a user’s historical access logs into a form that traditional machine learning methods can use.

We propose the use of deep learning models, particularly those based on recurrent neural networks (RNNs), as a novel improvement over previous methods. We demonstrate through offline experiments that RNNs are able to make effective use of historical data with minimal feature engineering to achieve superior prediction accuracy over traditional models. We prove these benefits in production through an online experiment where RNNs yield a 7.81% increase in successful precompute over a traditional model.

Finally, we highlight the benefits of the RNN computation model from a systems perspective. By eliminating the time-based aggregations used in traditional models in favor of a single hidden state, the overall computational cost of serving predictions is reduced by a factor of 10x.
2 RELATED WORK

Existing literature describes relatively simple models to estimate access probabilities in the context of precompute:

- (Wang et al., 2015) describe a system to prefetch links embedded in social network feeds using linear regression on mostly content-based features.
- (Parate et al., 2013) describe using the CDF of the “time since last use” as the probability estimate.
- (Sarker et al., 2019) showcase the effectiveness of decision tree models for context-aware smartphone usage prediction.

However, there are a few common limitations in approaches like the ones mentioned above:

- Making full use of historical access logs is difficult. Most classical machine learning models operate on fixed-length feature vectors, so a common approach is to compute aggregate functions based on the timestamps of previous accesses (e.g. time since last access, average time between historical accesses, number of accesses within a certain time window.) However, the choice of aggregations must be manually chosen through trial-and-error (“feature engineering”).
- Incorporating contextual features from historical access logs adds another dimension of difficulty, as any time-based aggregation can be combined with any subset of context dimensions (e.g. what was the number of accesses associated with this application surface?).
- The act of performing the predictions themselves is resource-constrained: care must be taken to ensure that model serving must not become as computationally expensive as the precomputation itself. For example, computing and serving aggregation features like the examples mentioned above may require specialized infrastructure to remain efficient at scale.

To address these problems, we are able to draw inspiration from research in the recommendation domain, where historical user behavior is similarly predictive. In recent times, deep learning-based recommendation systems have become extremely widespread due to their effectiveness at handling complex relationships in large datasets (Zhang et al., 2017). Of particular interest are recommendation systems based on recurrent neural networks (RNNs) due to their innate ability to model sequential data:

- (Beutel et al., 2018) describe the use of RNN-based recommender systems for video recommendations based on user actions. Of special note is the improved handling of “contextual features” (e.g. time, location, interface) which we find to be very relevant in the domain of predictive precompute as well.
- (Soh et al., 2017) describe the use of RNNs (specifically, gated recurrent units) to personalize recommendations in user interfaces based on a sequence of past user interactions, which is a very similar domain to the work in this paper.
- (Katevas et al., 2017) have also demonstrated the effectiveness of RNN architectures to predict user responses to notifications based on a sequence of mobile sensor data.

Our findings indicate that the results from the works above transfer well to the domain of predictive precompute. Using several test datasets, we compare RNN models with simpler models similar to the ones mentioned above and demonstrate the benefits of RNNs through online and offline experimentation.

3 DEFINING PREDICTIVE PRECOMPUTE

In this paper, we will focus on precomputation of individual activities within large-scale applications, where we have datasets of access logs from previous user sessions.

To be more specific, we would like to have a function that estimates the probability that a user accesses a particular activity within an application session, based on the current session context and past access logs as inputs. We will estimate the access probability at the beginning of each session and then choose to trigger precomputation at that point if the probability is greater than some fixed threshold.

3.1 Definitions

Sessions are defined as discrete time windows where the user is actively using the application; a session typically starts when the user opens the application. For simplicity, we consider each session to have a fixed length, e.g. 20 minutes, to avoid having to precisely measure when a session ends.

Context describes session-specific information that may be predictive of user behavior. Examples of context include:

- the current timestamp (including the hour of the day, day of the week, etc.)
- the current application surface
- indicators visible to the user, e.g. a “badge count” indicating the number of unseen notifications

Access logs are a sequential record of past application sessions, keyed by a user identifier. For each session we will record the context at the start of the session, as well as an additional Boolean access flag indicating if the activity was accessed within that application session or not. Access logs will be used as the training dataset, where access flags are used as ground truth labels and contexts are used to extract features.
3.2 Problem Statement

For any given user, assume that we have access to logged data for \( n - 1 \) previous user sessions, where \( C_i \) denotes the context and \( A_i \in \{0, 1\} \) denotes the access flag for the \( i \)th session. If we refer to the current session as session \( n \), we would like to estimate the probability of an access, \( P(A_n) \), given all known information past and present:

\[
P(A_n | C_1, A_1, C_2, A_2, ..., C_{n-1}, A_{n-1}, C_n)
\]

The remainder of this paper will primarily focus on how to best estimate \( P(A_n) \). We will train machine learning models treating each session as an individual data point, with the recorded value of \( A_n \) as the ground truth (label) and \( C_1, A_1, ..., C_{n-1}, A_{n-1}, C_n \) as the basis for features.

3.2.1 Timeshifted Precompute

An interesting related problem occurs in the case where we wish to enable precomputation of data prior to the start of an application session (e.g. several hours in advance). We refer to this modified problem as timeshifted precompute.

The primary motivation of precomputing data further in advance is to shift computational cost from peak hours to off-peak hours. At scale, being able to shift a meaningful amount of computational cost to off-peak hours smooths out the peak/off-peak power curve and can reduce overall capacity requirements.

In this scenario we do not have access to any session-specific context and instead must rely on existing access logs alone to predict the probability that an access will occur within a pre-defined peak hours window of a particular day. If we denote access during peak hours of day \( d \) as \( PA_d \in \{0, 1\} \), then we can state the probability estimate as:

\[
P(PA_d | C_1, A_1, ..., C_n, A_n)
\]

Each training example corresponds to one user × peak window pair, with the ground truth label being the presence of an access within the peak window.

4 Datasets

In this paper we analyze two real-world datasets where predictive precompute is being employed at Facebook, each consisting of a random sample of \( 10^6 \) users over 30 days. As a publically available baseline, we also make use of the Mobile Phone Use dataset published by (Pielot et al., 2017).

4.1 Mobile Tab Access (MobileTab)

Upon startup of the Facebook mobile application, we may choose to prefetch data for certain sections (“tabs”) if we can predict that the user is likely to access them.

A session begins when the user starts the application and ends after a fixed window of 20 minutes. For this dataset, we selected a tab with moderate usage and recorded an access for every session where an interaction occurred with the tab within the time window.

Context for this dataset includes the current time, unread notification count displayed over the tab icon (0-99), and the name of the active application tab at startup. Access logs consisting of the three context features and the access flag are stored over a 30-day period.

Table 1 illustrates an example sequence of sessions for an individual user in the MobileTab dataset.

<table>
<thead>
<tr>
<th>TIMESTAMP</th>
<th>ACCESS FLAG</th>
<th>UNREAD</th>
<th>ACTIVE TAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1564642800</td>
<td>1</td>
<td>3</td>
<td>Home</td>
</tr>
<tr>
<td>1564642900</td>
<td>0</td>
<td>0</td>
<td>Home</td>
</tr>
<tr>
<td>1564643000</td>
<td>0</td>
<td>1</td>
<td>Messages</td>
</tr>
</tbody>
</table>

4.2 Timeshifted Data Queries (Timeshift)

On the Facebook website, data queries that are relatively static can be precomputed and cached several hours ahead of time (as described in the Timeshifted Precompute problem statement). During off-peak hours of the day, we predict if a user will require a particular data query result in a session during peak hours on the following day.

A session begins upon the start of a website load and ends after a fixed 20-minute window. We selected a moderately used data query and recorded an access for every session where the data query was used.

Context for this dataset includes only the session timestamp and a flag indicating whether or not the session occurred during peak hours. Any additional context quickly loses relevance by prediction time, where no context is available.

4.3 Mobile Phone Use (MPU)

(Pielot et al., 2017) have generously published a dataset containing data traces for 279\(^1\) mobile phone users over the course of four weeks. We borrow heavily from the work in (Katevas et al., 2017) and also attempt to predict the probability that a user opens the app associated with a notification when it is received. In the context of predictive precompute, the OS could conceivably preload the app in the background for notifications with a high probability of interaction.

\(^1\)There are 342 users in the full dataset, but only 279 have an Android version with full support for notification tracking.
For MPU we define each session to start with the appearance of a notification, with a fixed length of 10 minutes. An access occurs if the user opens the app associated with the notification. To provide a reasonable comparison against the previous datasets we ignore the auxiliary “sensors” and focus on only the access logs associated with previous notifications.

We derive four context variables for each notification: the current time, the current screen state (on/off/unlocked), the application ID the notification was associated with, and the last opened application ID.

4.4 Data Statistics

Table 2 displays summary statistics for each dataset. In MobileTab and MPU each labeled example corresponds to a single session, while in Timeshift an example corresponds to a single peak period (one peak period per day and 30 days per user, for a total of 30M unique training examples). Although the MPU dataset has a very small sample of users, there is much more data per user — on average over 8,000 notification events per user — which still yields a sufficient amount of data for the purposes of model training.

Figure 1 displays the CDF of access rates for each dataset. Note that for MobileTab and Timeshift a significant percentage of users (36% and 42% respectively) have no recorded accesses at all in 30 days; this is typical of real-world scenarios, where not all users may access a particular activity.

Table 2. Summary of each dataset.

<table>
<thead>
<tr>
<th>DATA SET</th>
<th>POSITIVE RATE</th>
<th>SESSIONS</th>
<th>USERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>MobileTab</td>
<td>11.1%</td>
<td>60.8M</td>
<td>1M</td>
</tr>
<tr>
<td>Timeshift</td>
<td>7.1%</td>
<td>38.5M</td>
<td>1M</td>
</tr>
<tr>
<td>MPU</td>
<td>39.7%</td>
<td>2.34M</td>
<td>279</td>
</tr>
</tbody>
</table>

5 Baseline Models

In this section we will describe a selection of simpler machine learning models for comparison against the recurrent neural network model.

5.1 Percentage-Based Model

A very simple baseline model is to return the current access percentage based on all historical sessions for each user, disregarding additional context information. We can seed the percentage estimate for each user with the globally averaged access percentage across all sessions ($\alpha \in (0, 1)$):

$$P(A_n) = \frac{\alpha + \sum_{i=1}^{n-1} A_i}{n}$$

For Timeshift the calculation is similar, except we average over accesses at peak rather than individual sessions:

$$P(PA_d) = \frac{\alpha + \sum_{i=1}^{d-1} PA_i}{d}$$

5.2 Prelude: Feature Engineering

In order to make use of traditional models we must first convert all the available raw context for each session into a fixed-length numerical vector. We use common feature engineering techniques to accomplish this.

- **One-hot encoding of categorical variables.** For context variables such as the unread/notification counts, active tab, and application names, we use the standard technique of one-hot encoding. Note that for the tab and application name features we first limit the number of distinct values to a reasonable range by hashing and taking the remainder modulo 97.

- **Time-based features.** Given the raw timestamp, we additionally calculate the hour of day (0 - 23) and day of week (1 - 7) and apply a one-hot encoding to them.

- **Time-based aggregations.** We can track the number of accesses, number of sessions, and their ratio (the access percentage) for each user across a variety of time windows. In our comparisons we use the last 28 days, 7 days, 1 day, and 1 hour as time windows. We can also filter past accesses to those whose contexts match the current session context, e.g. having the same active tab or notification count (or both). To maximize coverage, we calculate aggregations based on all (time window) × (matching subset of context) combinations.

- **Time elapsed.** We can calculate the time difference (in seconds) from both the last access and the last session. As with the aggregation features, we also condition these to past events with a matching context subset.

\(^2\)https://en.wikipedia.org/wiki/One-hot
Table 5 illustrates the importance of thorough feature engineering with GBDT models; evaluation metrics drop sharply once aggregation and time-elapsed features are removed.

5.3 Logistic Regression (LR)

As a first step we use logistic regression (LR) on the features described above, treating each session as an individual data point. To give the aggregation-based features adequate warm-up time, only sessions from the latest 7 days of each dataset are used for training; of these, we take 90% of the users as the training set and leave 10% as a test set for evaluation. Details are explained in Section 8. An additional feature pre-processing step is to bucketize \( \text{time elapsed} \) features into 50 buckets and one-hot encode them, due to the uneven distribution. To do so, we take \( \lfloor \frac{50}{t} \rfloor \) where \( t \) is the time difference in seconds; note that the largest possible \( t \) (30 days) is about \( e^{14.76} \) seconds.

To train a logistic regression model, we use the scikit-learn (Pedregosa et al., 2011) LogisticRegression\(^3\) API with the saga solver and default settings.

5.4 Gradient Boosted Decision Trees (GBDT)

Gradient boosted decision trees (GBDT) are a popular model type that provides solid results with minimal tuning. Training is similar to the logistic regression approach, but we skip the one-hot encoding step for time-elapsed features and also some categorical features (e.g. the time of day and the day of the week).

We use XGBoost\(^4\) 0.90 (Chen & Guestrin, 2016) to train a decision tree model with mostly default settings, except for the tree depth hyperparameter. To determine the optimal tree depth, we split off 10% of the users from the training set as validation set. We then use a simple exhaustive search over all possible depths in the range \([1, 10]\) to minimize the log loss objective over the validation dataset.

With the manually engineered numerical features, we find that GBDTs are very hard to beat. We tested simple neural network architectures (e.g. a multi-layer perceptron) and could not obtain significant gains over GBDT models.

6 Recurrent Neural Networks

Whereas traditional models treat the access prediction for individual sessions as independent events, the innovation of recurrent neural networks (RNNs) is to process events in a sequential manner while introducing a persistent hidden state to carry over information from previous events.

An RNN model accepts both the current feature vector and the hidden state as inputs, and produces both a prediction and an updated hidden state for the subsequent prediction.

The primary benefit of RNN architectures is to obsolete the tedious and manual feature engineering steps described in section 5.2 — specifically the aggregation and time difference features. Instead, the RNN hidden state has the ability to automatically capture features based on past events.

Traditionally, RNN architectures have been popularized in text and audio domains, as they are well suited to handle regular sequences of information like characters or audio samples (Graves et al., 2013). In this section we describe various technical considerations when applying RNNs in the domain of predictive precompute.

6.1 Sequence Modeling

Recall from section 3.2 that for each user we have a sequence of \( n \) logged historical sessions, with contexts \( C_1, \ldots, C_n \) and access activity \( A_1, \ldots, A_n \). Let \( t_1, \ldots, t_n \) also denote the UNIX timestamp of each session where \( t_1 < t_2 < \cdots < t_n \). To model this using RNNs, we first introduce some preliminary concepts:

- **Feature extraction.** Each step of the RNN model must receive a fixed-length feature vector. While we can omit all of the aggregation features described in Section 5.2, we still must construct a feature vector from each \( C_i \) consisting of the one-hot encoded categorical context variables (e.g. notification count) and time-based features (hour of day, day of week). Let \( f_i \) denote the feature vector of context \( C_i \).

- **Representing different time intervals.** Traditional RNN models usually operate on regular sequences where elements in the sequence represent fixed time steps or consecutive characters. However, for a sequence of user sessions, some sessions may be seconds apart, while others may be hours or days apart. To allow the network to react effectively to different timescales we input \( \Delta t_i = t_i - t_{i-1} \) to the recurrent network at each step, where \( \Delta t_1 = 0 \). We find in our datasets that the distribution of \( \Delta t \) tends to be power-law distributed, so we apply the log/bucketing transform described in 5.2 here as well, denoted \( T(\Delta t_i) \).

- **Hidden states.** Each user starts with an initial hidden state \( h_0 \), an all-zero vector. At the end of session \( i \), the RNN model produces an updated \( h_i \) based on the previous hidden state \( h_{i-1} \) and inputs \( f_i, A_i, \) and \( \Delta t_i \).

- **Update delays.** To model real-world behavior accurately we must take into account two sources of delays: (1) that the ground truth \( A_i \) cannot be determined until the session ends (recall that each session has a fixed length, e.g. 20 minutes), and (2) that obtaining \( h_i \) is not
Figure 2. Modeling sequences of access logs with recurrent neural networks. Multilayer perceptron (MLP) units produce output probabilities at time \( t \), while hidden state updates occur through the gated recurrent unit (GRU) at time \( t + \delta \) due to the delay. Note that because \( t_5 \) occurs before \( t_2 + \delta \) it cannot make use of \( h_2 \) and uses \( h_1, t_1 \) as inputs instead.

Instantaneous (i.e. it takes some time, \( \epsilon \)). To address this we define a lag parameter, \( \delta \), equal to the session length plus \( \epsilon \).

- Functions for hidden updates and predictions. Abstractly, an RNN can be separated into two functions: an updater \( RNN_{update} \) which produces new hidden states, and a feed-forward network \( RNN_{predict} \) which produces predictions as output. It is often convenient to combine the two into a single model that produces both outputs simultaneously, but this separation is required in order to properly model the lag \( \delta \).

Putting everything together, we can define a sequence of hidden states \( h_0 = 0, h_1, ..., h_n \), with a recurrence relation:

\[
h_i = RNN_{update}(h_{i-1}, [f_i; A_i; T(\Delta t_i)])
\]

To obtain a prediction for \( P(A_i) \), we use \( RNN_{predict} \) with the latest known hidden vector accounting for update lag, denoted \( h_k \), where \( k \) is the maximum \( k \) such that \( t_k < t_i - \delta \) (if no such \( k \) exists, then we let \( k = 0 \) and \( t_i - t_k = 0 \)):

\[
P(A_i) = RNN_{predict}(h_k, [f_i; T(t_i - t_k)])
\]

For \textit{timeshifted precompute (3.2.1)}, predictions do not have access to \( f \) or \( t \) and instead can only use \( start_d \) and \( h_k \), where \( start_d \) marks the start of the peak period on day \( d \) and \( k \) is the maximum index such that \( t_k < p_1 - \delta \):

\[
P(PA_d) = RNN_{predict}(h_k, [T(start_d - t_k)])
\]

Intuitively, we can think of the hidden vectors \( h \) as an encoding of the sequences of contexts \( C \), accesses \( A \), and timestamps \( t \). Many of the feature engineering techniques in section 5.2 can be seen as attempts to do this in a manual way. However, through training a recurrent neural net we hope to learn the optimal way to encode the entire sequence into a single vector rather than relying on manual tuning and heuristics. Another significant benefit is the incremental nature of hidden updates: with manual feature engineering we need to store and retrieve the entire sequence, but with RNNs we only need the last known hidden vector.

### 6.2 Model Architecture

A few options are available for hidden state updates (\( RNN_{update} \)). For this paper, we evaluated three options: a basic \textit{tanh}-based recurrent unit, a gated recurrent unit (GRU) and a long short-term memory (LSTM) unit. (Chung et al., 2014) found that GRU and LSTM units result in comparable performance on a number of example datasets, while \textit{tanh} performance lags behind.

Empirically we find that GRUs provide the best performance over all of the datasets (at least without significant tuning).

The primary hyperparameter available when using RNN units is the dimensionality of the hidden vectors. Empirically, \( d = 128 \) seems to be a good dimensionality for all datasets. Another possible modification is to stack multiple...
GRU or LSTM units on top of each other; however, we report similar findings to (Beutel et al., 2018), where the addition of multiple GRU units did not provide a meaningful improvement over a single unit.

For $R N N_{\text{predict}}$, a simple architecture where the input vector and hidden vector are concatenated and passed into a feed-forward multilayer perceptron (MLP) provides good performance. Inspired by (Beutel et al., 2018), we find that an element-wise multiplication of the hidden vector with a latent factor derived from the context provides a meaningful improvement:

$$h'_{i} = h_{k} \circ (1 + L([f_{i}; T(t_{i} - t_{k})]))$$

where $k$ is the latest known index as described previously and $L$ is a linear transformation matrix.

As for the MLP layer, we find that a single hidden layer with 128 neurons combined with a rectified linear unit (ReLU) layer seems to be sufficient for the best performance; adding more layers does not lead to meaningful improvements. We can therefore summarize the formulation as:

$$P(A_{i}) = \sigma(b_{2} + W_{2} \cdot \text{ReLU}(b_{1} + W_{1}[h'_{i}; f_{i}; T(t_{i} - t_{k})]))$$

Here $\sigma$ denotes the sigmoid function while $b_{1}$, $b_{2}$ represent constant bias vectors and $W_{1}, W_{2}$ represent linear transformation matrices.

### 6.3 Loss Functions

For binary classification problems, log loss is the standard loss measurement function. For an individual access prediction $P(A_{i})$ the log loss is defined as:

$$- [A_{i} \cdot \log(P(A_{i})) + (1 - A_{i}) \cdot \log(1 - P(A_{i}))]$$

For traditional tasks, training is often optimized over the log loss averaged over all points in the training dataset. However, we find that this is suboptimal when comparing evaluation metrics over later days; it over-weights errors from predictions early in the sequence (when only a small number of access logs have been included in the hidden state). On the other hand, optimizing directly on later days (e.g. the last 7 days) appears to be suboptimal as well, possibly because the gradient is less stable for later elements in the sequence. Empirically, we consistently find it is best to train on the log loss for the last 21 days out of the 30 days available for each user. We did not find weighing the loss with an exponential time decay to be of significant benefit.

Each session is weighted equally, which does mean that users with more active sessions have greater influence over the model’s performance. Although this is desirable when considering computational costs as a whole, different weighting schemes can be explored if we wish to give more weight to inactive users as well.

### 7 RNN Training

RNN models are trained using PyTorch v1.1\(^6\) using the Adam optimizer with a learning rate of $1e-3$. We also include a dropout layer in the middle of the MLP set to 20% to prevent overfitting. Figure 3 displays sample PyTorch code for the key model definitions.

```python
import torch.nn as nn
class RNNClassifier(nn.Module):
    # i_n = feature vector dimensions
    # h_n = hidden dimensions
    # w_n = number of hidden neurons
    def __init__(self, i_n, h_n, w_n):
        self.L = nn.Linear(i_n, h_n)
        self.W_1 = nn.Linear(i_n + h_n, w_n)
        self.W_2 = nn.Linear(w_n, 1)
        self.Dropout = nn.Dropout(0.2)
        self.GRU = nn.GRUCell(i_n + 1, h_n)

    # Returns a new hidden vector $h_{i+1}$.
    # f_i includes both the feature vector
    # and the encoded time difference.
    def hidden_forward(self, h_i, f_i, A_i):
        cross_h_i = self.L(f_i)
        cross_h_i = h_k * (1 + self.L(f_i))
        mlp = self.W_2(torch.cat((cross_h_i, f_i), 1), h_i, 1)
        mlp = torch.relu(self.Dropout(mlp))
        return torch.sigmoid(self.W_2(mlp))

    # Predicts $P(A_{i+1})$.
    def forward(self, h_k, f_i):
        h_i = self.L(f_i)
        mlp = self.W_1(torch.cat((h_i, f_i), 1), h_i, 1)
        mlp = torch.relu(self.Dropout(mlp))
        return mlp

Figure 3. Sample PyTorch code for key model definitions.
```

\(^6\)https://github.com/pytorch/pytorch/releases/tag/v1.1.0

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Predictive Precompute with Recurrent Neural Networks

Figure 4. Log loss vs. number of sessions processed (based on the full cross-validated dataset). Each vertical line represents the end of one epoch, with 8 epochs in total.

Figure 5. Distribution of MPU session counts.

7.1 Minibatch Training

For MobileTab and Timeshift we can significantly increase training speed by using minibatch training with batches of 10 users. Note that for MPU this becomes ineffective due to the small number of users versus high number of sessions per user, and we fall back to processing users individually.

For each minibatch we compute predictions for the last 21 days and then calculate the average log loss over all prediction/label pairs. The loss gradient of each minibatch is then back-propagated to complete one training iteration. For the larger datasets, training converges in just one epoch, but for the MPU dataset a total of 8 epochs are required for convergence. Figure 4 displays the training log loss curves.

We find that a key optimization to speed up training is to evaluate minibatches via custom parallelism. While a standard approach is to pad user histories to a uniform length before evaluating them in batch, in practice the distribution of access history lengths has a very long tail (Figure 5). This results in an excessive amount of operations wasted on padding values. Instead, we can evaluate predictions and calculate gradients for each user on a separate thread and then accumulate the gradients afterwards. Models train twice as quickly with this approach versus the padded batch approach.

A final consideration for the MPU dataset is to truncate user histories to the most recent 10,000 sessions. This limits the effect of long tail users on training time without having a noticeable impact on model quality.

8 Evaluation Results

Unless otherwise noted, evaluation metrics are reported on the test dataset for each model. We are careful to use the same train / test split for all models (for MPU, the same 4-fold validation sets). We split based on users because it allows more historical data per user at training time. Another key consideration is the timeframe used for evaluation: evaluating on the full 30-day period does not accurately reflect real-world performance because the majority of users already have a full 30 days of history. For example, on any given day in the MobileTab dataset, less than 1% of sessions have no previous history in the previous 29 days. In practice, only 1% or fewer users do not already have previous logged history over the past 30 days. Therefore, we evaluate predictions on the last 7 days of testing data in each dataset to get a better estimate of performance in production.

As for the evaluation metric itself, the most important metrics in the predictive precompute domain are precision and recall. Precision corresponds to the percentage of precomputations that were followed by an actual access, while recall corresponds to the percentage of accesses that were successfully precomputed. In practice, improvements in recall are almost linearly correlated to reductions in application latency. Figure 6 shows the full precision-recall curve across all tested models for MobileTab.

To obtain a single numerical metric for model comparison we use the area under the precision-recall curve (PR-AUC): (Davis & Goadrich, 2006) show that PR-AUC tends to be an effective measure when dealing with highly skewed datasets (MobileTab and Timeshift). Table 3 compares the PR-AUC across all tested models and datasets. When applying models in practice, we typically select a threshold that keeps wasted precomputations within an acceptable range (i.e. maximizing recall while constraining on precision; for example constraining precision to 50%). Table 4 compares

Figure 6. Precision-recall curve comparison for MobileTab.

the recall for each model at a fixed 50% precision, where the difference between models becomes more apparent for MobileTab and Timeshift.

9 ONLINE EXPERIMENTATION

While the offline experiments described above show clear improvements, we also have results from online experiments to verify that they carry over to production environments. For the MobileTab dataset, we productionized the RNN model to replace an existing production GBDT model as follows:

- The most recent hidden state for each user (a 128-element floating point vector) and session timestamp are stored in a real-time data store similar to Redis8.
- TorchScript9 versions of the MLP and GRU models are made available in a remote execution environment.
- At session startup time, the most recent hidden state along with the current context variables are retrieved and sent through the MLP part of the model to calculate an access probability $p$. We eagerly precompute and retrieve the tab contents if $p$ is greater than a fixed threshold, chosen to target a precision of 60%. This corresponds to a recall of about 51.1% in the RNN model vs. 47.4% in the GBDT model. This comes out to a 7.81% increase in “successful prefetches” (i.e. accesses that were successfully prefetched).
- Context variables are sent to a stream processing system similar to Apache Kafka10, tagged by a unique session ID. Tab accesses are also sent to the same system with a matching session ID. Events are buffered by session ID, and after a timer corresponding to the session length fires, the context $C_i$ and access flag $A_i$ are computed. We then retrieve the most recent hidden state for the user $h_i$ and execute the GRU part of the model to calculate and store a new hidden state.

We report several observations after monitoring the behavior of the productionized model over a period of about 90 days:

Relative production resources. RNN models are indeed more resource intensive — empirically the TorchScript model is about 9.5x more computationally intensive than a GBDT model. However, in practice, the most compute-intensive component is actually the serving of aggregate access percentages and time elapsed features, which requires about two orders of magnitude more compute than the model computation itself. One approach is to retrieve the 30-day access logs for each user to compute aggregations on the fly, but some users have hundreds or thousands of past accesses which makes this impractical. Instead, aggregations are computed using a stream processing service in combination with a key-value store. However, we still need to keep track of every combination of context values in order to serve context-dependent aggregations, which may

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8https://redis.io
10https://kafka.apache.org
result in thousands of unique keys per user. For example, MobileTab requires about 20 aggregation feature lookups for every individual prediction.

In contrast, with the RNN model we only need to make one key-value lookup to retrieve a 128-dimensional (512-byte) hidden vector for each prediction. By decreasing both the storage footprint and request volume, this reduces the overall serving computational cost by about 10x in practice. Furthermore, if necessary, hidden states allow for very fine-grained control over resource usage via the hidden state dimensionality. In more resource-constrained environments, we can easily train a model that has fewer hidden dimensions to trade off model quality for a smaller storage footprint per user. Neural network quantization methods can also be applied to store single bytes instead of floating-point numbers for each dimension.

**Cold start behavior.** In our online experiment, we compared two groups of users starting with an empty history to compare the warmup behavior between the GBDT and RNN models. We find that it takes about 14 days for the RNN model to stabilize, and that it is consistently superior than the GBDT model. Figure 7 displays the online PR-AUC for the first 30 days of the online experiment.

**Long-term model quality and stability.** Despite the fact that the training data only spans 30 days, we see that the empirical precision and recall are consistent with the results obtained from offline experiments, and continue to maintain the same level of quality (with no sign of degradation) over a 90-day period. This suggests that the hidden states produced by the RNNs are stable over long-term periods.

**Retraining the model.** While not tested in production, it should be possible to train newer versions of the RNN model using the existing hidden states as the value for $h_0$, thus providing a path to replacing the production model without invalidating all existing hidden states. Another approach may be to preserve the GRU parameters and hidden states and retrain only the MLP portion of the model, which is significantly faster to retrain.

**Tradeoffs.** While RNNs have a clear advantage in model performance and significantly reduce serving computational costs, the primary drawbacks are 1) the increased training time and 2) the increased amount of data required to train an effective model. The MPU dataset is a realistic baseline for the amount of data required, with $2 \times 10^6$ sessions, and takes about 10 hours to complete 8 training epochs. In contrast, the aggregation-based features described in Section 5.2 are very generically applicable to any predictive precompute use case, and GBDT models can be trained in minutes with just $10^4$ data points. Finally, hidden states are almost entirely “black-box” and are not easily explainable, whereas aggregation functions are easily human-interpretable.

**10 Conclusion**

We present a review of existing techniques that can be applied to predictive precompute problems as well as a selection of real-world datasets for comparison.

We demonstrate the novel use of recurrent neural network (RNN) models to achieve state-of-the-art results in this domain. In addition to achieving superior precision and recall metrics, RNNs significantly reduce the need for manual feature engineering due to the automatic encoding of historical information into hidden states.

We show that these advantages carry over to an online production environment, where models maintain consistent performance over an extended 90-day period. We highlight how RNN models can help decrease the computational cost of serving models by an order of magnitude by encoding all prior history into a compact hidden state.

In closing, we hope that the techniques described in this paper make it easier for other applications to utilize predictive precompute.

**10.1 Future Work**

**Reusable models.** The very simple percentage-based model described in 5.1 in some sense acts as a “universal model” that works as a solid baseline across all use cases with almost no training. In a similar vein it may be possible to create a generic RNN-based model that uses only past session timestamps and their access labels to produce high-quality estimates without any pre-training.

**Interpretable hidden states.** The hidden state model is practically a black box. Extracting interpretable relations from the hidden state could suggest ways of feature engineering the dataset to enable the use of simpler models.
REFERENCES


