Research and Applications

Privacy-preserving model learning on a blockchain network-of-networks

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ABSTRACT

Objective: To facilitate clinical/genomic/biomedical research, constructing generalizable predictive models using cross-institutional methods while protecting privacy is imperative. However, state-of-the-art methods assume a “flattened” topology, while real-world research networks may consist of “network-of-networks” which can imply practical issues including training on small data for rare diseases/conditions, prioritizing locally trained models, and maintaining models for each level of the hierarchy. In this study, we focus on developing a hierarchical approach to inherit the benefits of the privacy-preserving methods, retain the advantages of adopting blockchain, and address practical concerns on a research network-of-networks.

Materials and Methods: We propose a framework to combine level-wise model learning, blockchain-based model dissemination, and a novel hierarchical consensus algorithm for model ensemble. We developed an example implementation HierarchicalChain (hierarchical privacy-preserving modeling on blockchain), evaluated it on 3 healthcare/genomic datasets, as well as compared its predictive correctness, learning iteration, and execution time with a state-of-the-art method designed for flattened network topology.

Results: HierarchicalChain improves the predictive correctness for small training datasets and provides comparable correctness results with the competing method with higher learning iteration and similar per-iteration execution time, inherits the benefits of the privacy-preserving learning and advantages of blockchain technology, and immutable records models for each level.

Discussion: HierarchicalChain is independent of the core privacy-preserving learning method, as well as of the underlying blockchain platform. Further studies are warranted for various types of network topology, complex data, and privacy concerns.

Conclusion: We demonstrated the potential of utilizing the information from the hierarchical network-of-networks topology to improve prediction.

Key words: blockchain distributed ledger technology, privacy-preserving predictive modeling, hierarchical network, clinical information systems, decision support systems

INTRODUCTION

Background and significance

Cross-institutional predictive modeling can accelerate clinical, genomic, and biomedical research1–5 by learning more generalizable models from the increased number of patient records (Figure 1A). Aiming at protecting privacy of the patients, several centralized privacy-preserving algorithms6–9 were developed based on the principle of exchanging the models instead of disseminating Protected Health Information (PHI) data directly (Figure 1B). Although these approaches ensured prediction correctness while honoring patients’ privacy, the...
fact that only a central server manages the entire model training process creates problems yet to be solved: (1) imbalance in the compute resource allocation can occur, where the central server can potentially assign work to a participating healthcare institution unfairly while letting the other institutions remain idle most of the time; (2) model parameters in the central server can be modified obliviously during model training; (3) the provenance (i.e., source institution) of models may be changed by the central server in an undetectable way by local sites; and (4) information gaps about disseminated models occur between the central servers with full visibility about generated models and non-central-servers with only partial visibility. These potential concerns can become hurdles when adopting privacy-preserving learning algorithms among multiple institutions.

To mitigate these risks, a plausible solution is to adopt blockchain, the underlying technology of modern fully decentralized crypto-currencies (Figure 1C). (1) As a peer-to-peer architecture, blockchain can serve as a distributed ledger for the institutions to exchange machine learning models without a central server thus re...
moving the concern of a single-point-of-control and potential computational unfairness. (2) The design of the consensus protocols in blockchain makes the change of the ledger extremely difficult and therefore ensures the immutability of the models stored on the blockchain.12 (3) Blockchain also preserves provenance of the ledger, which makes the source of the models verifiable.13 (4) Blockchain is transparent (ie, "everyone can see everything"),13 and therefore all models are visible to every participating healthcare institution. It should be noted that although there are many other existing decentralized architecture (eg, gossiping algorithms14–16) blockchain contains the above-mentioned technical features (ie, immutability, provenance, and transparency) and has been adopted for critical financial applications for an extended period of time.11 Therefore, we utilize blockchain as the decentralized architecture to alleviate the concerns of centralization.

However, the state-of-the-art blockchain-based learning methods10,17,18 assume that the network has a “flattened” topology, as shown in Figure 1A. In the real world, the structure of research networks can be more complicated than a simple topology. For example, PCORNet,19 a major initiative to support an effective, sustainable national research infrastructure, includes 13 clinical data research networks (CDRNs). Specifically, pSCANNER,20 one of the CDRN subnetworks in PCORNet led by the University of California San Diego (UCSD), includes subnetworks with data from diverse sites, and is part of this multi-level “network-of-networks” (ie, pSCANNER includes SCANNER21 and UCREX22 subnetworks, with each containing multiple sites).

As shown in Figure 1D, such a real-world network-of-networks topology can imply practical issues such as (1) small data size for rare diseases/conditions (eg, Kawasaki Disease23,24), (2) each site may prefer to prioritize their own model (eg, UCSD may tend to put more weight on the model learned from local data, while still including models from other institutions to increase model generalizability) while still considering its data size; and (3) subnetwork model maintenance (eg, the pSCANNER network may prefer to retain the aggregated model from its own participating networks in parallel with the model learned from the whole network such as PCORNet). Without utilizing the information of the hierarchical topology, the learning method designed for a flattened topology10,17,18 could not address these practical issues effectively. As a result, the attempt to improve the correctness of the cross-institutional predictive modeling while preserving patient privacy may not be feasible due to the insufficiency of the existing methods regarding small data size, model prioritization, and model maintenance for subnetworks.

Therefore, a hierarchical approach (Figure 1E) that considers the network-of-networks information is critical to address these issues. By computing, recording, and combining the models from each level with different weights based on data size, we anticipate the hierarchical method to (1) improve predictive correctness with small data; (2) prioritize local data for each site while considering the number of records, and (3) retain consensus for each subnetwork. Also, the benefits of the methods designed for flattened network (eg, the property of fair compute loads for every site of GloreChain10) and the advantages of adopting a decentralized architecture (ie, no single point of control, immutable data/records, data provenance, and complete visibility10) should be inherited.

Objective

We aim at developing a hierarchical modeling framework with 3 goals: (a) inherit the benefits of the privacy-preserving learning methods designed for flattened network, (b) retain the advantages of adopting a decentralized architecture, and (c) address practical data size, local model, and subnetwork consensus issues on a real-world clinical, genomics and biomedical research network-of-networks.

MATERIALS AND METHODS

To achieve the first goal of inheriting the benefits of the state-of-the-art learning method at every level of the hierarchical network-of-network topology (Figure 1E), we adopt the learning methods designed for a flattened network. With this design, we ensure that at each level the predictive correctness and fair compute loads properties are preserved. Next, to retain the advantages of a decentralized architecture, we utilized peer-to-peer blockchain technology11,13,25–35 to disseminate models and therefore avoid concerns such as single point of control. Finally, to tackle the practical issues on a real-world research network-of-networks, we propose to leverage the hierarchical topology information to store and combine the models from each level. Figure 2 demonstrates the concept of the proposed hierarchical consensus learning using an example of pSCANNER, which consists of 2 subnetworks (SCANNER and UCREX).

We developed HierarchicalChain to evaluate our proposed framework. HierarchicalChain contains the following 3 main components: 1) a level-wise learning method which is originally designed for a flattened network, 2) a blockchain network and its on-chain data structure, and 3) a hierarchical consensus learning algorithm. These components are introduced in the next 3 subsections, followed by the implementation details, datasets, and experiment settings.

Level-wise GloreChain decentralized model learning

Our proposed general framework can adopt both online and batch decentralized learning algorithms. Online methods, such as ModelChain17 and ExplorerChain,18 focus on efficient retraining (ie, updated model for new data without a complete retrain of the model). In contrast, batch methods, such as GloreChain,10 emphasize effective prediction results (ie, learn mode using all data at once to achieve higher correctness).

For HierarchicalChain, we selected the batch method GloreChain because we aim at achieving high predictive correctness. GloreChain also provides an additional advantage of having fair compute loads for each participating site.10 GloreChain is based on GLORE,7 a centralized privacy-preserving learning method. We adapted GloreChain to a level-wise method, such that the consensus models can be trained at each level of the hierarchical topology (eg, a total of 7 models learned from 3 levels as shown in Figure 2). We denote the adapted method as GloreChain-LevelWise.

The blockchain network and on-chain data structure

HierarchicalChain utilized a permissioned blockchain network, in which only authorized sites (eg, member institutions of consortia like PCORNet19) can participate. Such a permissioned network improves privacy protection by prefiltering participants. The incentives for each site and each subnetwork are the improved model generalizability and thus predictive correctness, as well as the immutably recorded models for each level of the network-of-networks. Although the network structure is hierarchical, we use only one blockchain network to disseminate all models. This simple design can reduce the maintenance cost.

The transaction metadata of blockchain was exploited to store the models and related information, as shown in Figure 3. The transaction...
amount are all zeroes because we adopt blockchain as a pure ledger for data dissemination instead of coin transferring. The details of the data stored on-chain are explained in Table 1. Compared to GloreChain, we added 4 new fields ("Hierarchy," "Record," "Level," and "Type") in HierarchicalChain to incorporate information from the hierarchical topology. The space complexity of the on-chain transaction metadata is $O(M^2 + H)$, where $M$ is the number of covariates and $H$ is the number of the level of the hierarchy. By only disseminating partially learned models (ie, aggregated parameters) on-chain and keeping all observation-level PHI data off-chain, the privacy of the patient can be preserved.

The Proof-of-Hierarchy consensus learning algorithm

We developed Proof-of-Hierarchy (PoH), a new algorithm to learn the consensus predictive model on a hierarchical network-of-networks. First, the consensus models of each level are learned using GloreChain-LevelWise method. These models are stored on the single shared blockchain network and thus can be accessed by every site freely. Finally, to predict the outcome of a new patient data record, a site computes the prediction scores using all models and combines the scores to generate the final prediction result.

To combine the scores, we adopted an ensemble approach, which has been utilized often in biomedical informatics research, such as in medical information extraction on clinical notes and early detection of breast cancer using X-ray images. In the PoH algorithm, we exploited 2 simple weighted-average ensemble methods: horizontal ensemble and vertical ensemble. The horizontal ensemble (Figure 4A) combines Level 1 models, weighted by their training data size, with the intuition that a large institution may prefer to emphasize their own model for the prediction. The vertical ensemble (Figure 4B) combines all levels of models related to the local site weighted by the size of each level of network. The intuition of this method is to consider both the specificity from the small/local data and the generalizability from the remote/subnetwork data. Note that although these 2 ensemble methods combine the results...
Table 1. An example of on-chain data of HierarchicalChain. In this example, $M = 2$ is the number of the covariates in the dataset and $H = 3$ is the number of levels of the hierarchical topology. The partial model of GloreChain-LevelWise contains both “Model Mean” and “Model Covariance,” while the final model is the consensus mean vector. The “Flag” is TRANSFER, representing the submission of a model from 1 site to another via the blockchain. In this round, the model is transferred from the UCReX Site $s_2$ (“From Site”) to the SCANNER Site $s_1$ (“To Site”) at "2019-06-28 10: 53: 26” (“Time”). The “Hierarchy” of pSCANNER, UCReX, and UCReX Site $s_2$ represents the subnetworks of the local site (“UCReX Site $s_2$”), and the number of records on the local site is 30 (“Record”). The “Level” of 3 shows the current learning process happening on Level 3, and the “Type” of the model is single-level (“SINGLE”). The “Iteration” of 16 is the number of learning iterations at current level, and the “Result” indicates the value of the evaluation metric for correctness (eg, the full area under the receiver operating characteristic curve [AUC]).

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Possible Values</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model Mean</td>
<td>The mean vector of the GloreChain-LevelWise partial model</td>
<td>A numerical vector with its length equals $M + 1$</td>
<td>[0.676883, 1395.043314, 55.376925 ]</td>
</tr>
<tr>
<td>Model Covariance</td>
<td>The variance-covariance matrix of the GloreChain-LevelWise partial model</td>
<td>A numerical $(M+1) \times (M+1)$ symmetric matrix</td>
<td>[ 2.164606, 913.081988, 60.973002 ] , [ 913.081988, 1524148.948051, 17102.681580 ], [ 60.973002, 17102.681580, 3334.172388 ]</td>
</tr>
<tr>
<td>Flag</td>
<td>The type of action a site has taken to the model</td>
<td>UNKNOWN, HIERARCHY, INITIALIZE, UPDATE, EVALUATE, TRANSFER, CONSENSUS, COMPLETE, TEST, CLEAR</td>
<td>TRANSFER</td>
</tr>
<tr>
<td>From Site</td>
<td>The site that has submitted the model</td>
<td>A unique name or identifier representing the site</td>
<td>UCReX Site $s_2$</td>
</tr>
<tr>
<td>To Site</td>
<td>The site which will receive the model</td>
<td>A unique name or identifier representing the site</td>
<td>SCANNER Site $s_1$</td>
</tr>
<tr>
<td>Time</td>
<td>The time that the site submitted the model</td>
<td>A timestamp</td>
<td>2019-06-28 10: 53: 26</td>
</tr>
<tr>
<td>Hierarchy *</td>
<td>The subnetworks that the local site belong to</td>
<td>A string vector with its length equals $H$ and contains unique names or identifiers of each level of the hierarchy</td>
<td>pSCANNER, UCReX, UCReX Site $s_2$</td>
</tr>
<tr>
<td>Record *</td>
<td>The number of the records of the local site</td>
<td>A non-negative integer</td>
<td>30</td>
</tr>
<tr>
<td>Level *</td>
<td>The current level of hierarchy for learning (“1” for ensemble models)</td>
<td>A non-negative integer</td>
<td>3</td>
</tr>
<tr>
<td>Type *</td>
<td>The type of the model, either single-level (&quot;SINGLE&quot;) or ensemble (&quot;HORIZONTAL&quot; or &quot;VERTICAL&quot;)</td>
<td>UNKNOWN, SINGLE, HORIZONTAL, VERTICAL</td>
<td>SINGLE</td>
</tr>
<tr>
<td>Iteration</td>
<td>The current iteration of the learning process at current level</td>
<td>A non-negative integer</td>
<td>16</td>
</tr>
<tr>
<td>Result</td>
<td>The value of the evaluation metric when the learning process completes</td>
<td>A numerical value between 0 and 1</td>
<td>0.921604</td>
</tr>
</tbody>
</table>

from the models of each level of the hierarchical topology, those models are stored immutably on the blockchain and can be retrieved at any time as needed.

The details of the PoH algorithm are described in Supplementary Algorithms A.1, A.2, and A.3 in Appendix A. The main PoH (Supplementary Algorithm A.1) contains both level-wise model learning (Supplementary Algorithm A.2) and horizontal/vertical ensemble (Supplementary Algorithm A.3). We assume the topology of the network-of-network is a perfect tree (eg, the network-of-networks contains 2 sub-networks and each sub-network contains 2 sites; that is, the number of levels is 3, and the total number of participating sites is 4). The 5 hyperparameters of PoH includes the polling time period $\Delta$, the waiting time period $\Theta$, the maximum per-level iteration $\Omega$, the total number of participating sites $N$, and the number of levels $H$.

The implementation of HierarchicalChain

The system architecture of HierarchicalChain is shown in Figure 5. We implemented PoH, Blockchain-Connector, and GloreChain-LevelWise in Java. HierarchicalChain only uses the patient data to compute the models (in the GloreChain-LevelWise component) without disseminating the data to the blockchain network. We adopted MultiChain16,41 as our blockchain platform, because it is both a system built on top of the well-known Bitcoin Blockchain11 and a permissioned blockchain network for general-purpose ledgering10,32,18. The default consensus protocol, Mining Diversity,27,41 is adopted with default parameters for MultiChain. The system was developed in the UCSD campus Amazon Web Services (AWS)43,44 and evaluated on the integrating Data for Analysis, Anonymization, and SHaring (iDASH) 2.0 cloud network,45,46 a private cloud network also based on AWS45 and compliant with the Health Insurance Portability and Accountability Act (HIPAA) requirements. In both cloud networks, we used Linux-based Virtual Machines (VMs) to simulate 4-site scenario, and the type of each VM is Amazon EC2 T2 Large (ie, 2 virtual CPUs and 8GB of RAM) with 100GB of storage.47

Datasets

To evaluate the algorithms and models of HierarchicalChain, we adopted the following 3 datasets, each with one binary outcome:
(1) Edinburg Myocardial Infarction (Edin): this dataset includes 9 covariates and 1253 observations with the purpose of predicting the presence of disease (class distribution: 0.219 positive and 0.781 negative); (2) Cancer Biomarkers (CA): there are 2 covariates and 141 observations to predict the cancer (class distribution: 0.638 positive and 0.362 negative); and (3) Total Hip Arthroplasty (THA): the data contains 34 covariates and 960 observations aiming at predicting extended hospital stay (ie, hospital length of stay for total hip arthroplasty surgery > 3 days). For the THA dataset, an IRB Exemption Category 4 (Project Number 190385XX) was certified by the UCSD Human Research Protections Program (HRPP) on March 20, 2019.

**Experiment settings**

Our goal of experiment is to evaluate whether HierarchicalChain can improve the prediction correctness for practical issues—especially for small training data—by using the hierarchical topology information, prioritizing local data, and retaining consensus models from each level in the hierarchy. We compare the horizontal and vertical ensemble methods of HierarchicalChain (ie, Hierarchical-Chain-Horizontal and Hierarchical-Chain-Vertical) with Glore-Chain, the state-of-the-art blockchain-based decentralized learning method designed for flattened network topology.

For both HierarchicalChain and GloreChain methods, the precision of the convergence criterion was $10^{-6}$ with the following same hyperparameters based on previous studies, the network latency of the cloud networks, and the various sizes and splitting methods of the data: the polling time period $\Delta = 1$ (second), the waiting time period $\Theta = 5$ (seconds), the maximum per-level iteration $\Omega = 100$. The latest 4 transactions with the size of the transaction metadata > 20 were checked to identify new transactions on the blockchain network.

We used the abovementioned 3 datasets (ie, Edin, CA, and THA) to evaluate the methods. For the hierarchical topology, we set the total number of participating sites to $N = 4$, and the total level of hierarchy to $H = 3$ (ie, the same topology as shown in Figure 2). That is, we simulated the network-of-networks by splitting each of the 3 datasets into 4 sites in a hierarchical topology.

To simulate the real-world scenario, we tested 2 different ways to split the training data among sites: balanced (ie, the number of records on each site is even) and imbalanced (ie, the number of records on each site is uneven). Therefore, we split each dataset randomly with (1) balanced ratio of 25% for each site, and (2) imbalanced ratios of 10%, 20%, 30%, and 40% for each of the 4 sites, respectively.

For the Edin and THA datasets, each dataset is already small (141 records). Therefore, we randomly sampled the training data from only 0.5 to 1.0, in increments of 0.1. For each training and test dataset, including the sampled training data, we randomly sampled the training data from 0.1 to 1.0 (using the full training data), in increments of 0.1, for the Edin and THA datasets. Since the original size of the CA dataset is already small (141 records), we randomly sampled the training data from only 0.5 to 1.0, in increments of 0.1. For each training and test dataset, including the sampled training data, we preserved a class distribution similar to the original dataset, and kept at least 1 positive and 1 negative record. Our evaluation metric is the full area under the receiver operating characteristic curve (AUC). We calculated weighted-average AUC on the test data with the data ratio as the weights (eg, 10%, 20%, 30%, and 40% for imbalanced data splitting) accounting for both balanced and imbalanced data-splitting scenarios. We measured consensus iterations and execution times as well.

The abovementioned process (ie, data splitting, predictive modeling, and weighted-average test AUC computing) was repeated 30 times to collect the results. For the configuration with the smallest training sizes, we further conducted a Wilcoxon signed-rank test to examine whether the 2 ensemble methods (ie, horizontal
and vertical) of HierarchicalChain perform with statistical significant difference when compared with GloreChain in terms of predictive correctness. We reset the blockchain network for each trial to collect a more accurate execution time.

RESULTS

Predictive correctness
The predictive correctness results on the small training datasets are shown in Figure 6A. In general, both ensemble methods of HierarchicalChain outperformed GloreChain. Especially for the balanced split Edin and THA datasets and the imbalanced split THA datasets, the differences in AUC were statistically significant (with $P$ value < .05). For the CA dataset, vertical ensemble performed better on data with both types of splitting methods, and horizontal ensemble performed better for the balanced split data; however, all of the results have $P$ value $\geq$ .05. Also, in almost all cases (except the horizontal ensemble on CA dataset), HierarchicalChain showed smaller standard deviation when compared with GloreChain. The results for different training data ratio are depicted in Figure 6B. In general, HierarchicalChain-Vertical performed similar to GloreChain with a larger training data size, while HierarchicalChain-Horizontal provided worse correctness results and was less stable. The lower performance of HierarchicalChain-Horizontal may be due not using high-level models (ie, models from Level 2 and Level 3).

Learning iteration
The results of learning iteration on the small training data are shown in Table 2. In general, HierarchicalChain required 2–10 times of iterations when compared with GloreChain. The results for different size of training data are illustrated in Figure 6C. Note that the iterations for HierarchicalChain (computing Level 1, Level 2, and Level 3) were expected to be around 3 times the iterations of GloreChain (computing only Level 3). While this was true for the full-sized training data, on smaller training data the variation became larger.

Execution time
The execution time results for the small training data are shown in Tables 3 and 4. While the total execution times (Table 3) are roughly in proportion to the learning iterations (Table 2), the per-iteration execution times (Table 4) demonstrate similar results for HierarchicalChain and GloreChain. The per-iteration execution time results for different sizes of training data are depicted in Figure 6D. In general, the per-iteration time on smaller training data was also shorter, because the same overhead (eg, initialization time)
was shared by a larger number of learning iterations. Also, HierarchicalChain had lower per-iteration time, because the same overhead was shared by 3 levels of computation (GloreChain only computed for 1 level). According to a previous study, most of the execution time was used for waiting/synchronization, and only a small portion of the time (eg, < 0.2 second) was used for actual computation of the models.

**DISCUSSION**

**Findings**

According to the results, HierarchicalChain, using vertical ensemble to combine the models and prioritize the local ones, outperforms GloreChain for small training data—especially for the THA dataset collected from UCSD Health. Also, in general, the prediction correctness of HierarchicalChain is comparable to GloreChain. Despite the increased learning iterations, the per-iteration execution time of HierarchicalChain remains at the same level as the one for GloreChain. Additionally, HierarchicalChain inherits the benefits of GloreChain (eg, fair compute loads) and advantages of blockchain (eg, no single point of control), and can record models for each level on chain immutably. Finally, HierarchicalChain is more generalizable than GloreChain. That is, HierarchicalChain can be deployed on a flattened network, and, in this case, it becomes exactly the same as GloreChain.

HierarchicalChain, based on GloreChain, can adopt any privacy-preserving learning algorithms, including both batch and online methods. HierarchicalChain overcomes blockchain confidentiality issues by exchanging models from each level without transferring patient-level data; avoids the blockchain scalability issue, because the per-iteration learning time (5–30 seconds per iteration) is way longer than the average transaction time of a blockchain (< 1 second); and mitigates the blockchain 51% attack issue because of the permissioned network nature.

HierarchicalChain can also adopt different underlying blockchain platforms. That is, in our experiment, we adopted MultiChain with its low energy-consuming Mining Diversity consensus protocol; however, any other blockchain platform can also serve as the peer-to-peer infrastructure. The size of the model in our experiments is about 5 KB, which is smaller than the default size limit (2 MB) of MultiChain or other mainstream blockchain platforms. Also, the iDASH 2.0 cloud network provides an additional layer of security protection beyond the permissioned blockchain network.

Figure 6. The results on data with different training data ratio, including 3 datasets (Edin, CA, and THA) as well as 2 data-splitting methods (balanced and imbalanced). We compared 2 ensemble methods (horizontal and ensemble) of HierarchicalChain with the state-of-the-art GloreChain. The data are split to balanced or imbalanced ratios among the sites. A. The predictive correctness results on small training data. The top header represents dataset name (data split ratio). The models are trained using only small portions of the training data. The evaluation metrics is the weighted-average AUC and the P values are computed using the Wilcoxon signed-rank test. B. Prediction correctness, measured in weighted-average test AUC for different training data ratio. C. Learning iterations for different training data ratios. D. Per-iteration execution time measured in seconds for different training data ratios.
Limitations

The limitations for this work include: (1) Topology. Hierarchical-Chain was not tested on nonperfect tree topologies, which can contain different site numbers in various levels and may impact the performance of the correctness, iterations, and execution time; (2) Data. We did not evaluate on data with a large number of covariates, missing/nonrepresentative data, and highly different data distribution among the sites or the levels; (3) Advanced privacy concerns. Although this study focused on protecting patient privacy by having healthcare institutions exchange aggregated machine learning model without disseminating patient-level data, more advanced privacy concerns, such as institutional privacy and differential privacy (ie, the model may still reveal some information for the institution) and differential privacy (ie, the patient-level data may be inferred under certain...
Table 2. Learning iteration results on small training data, including both mean and standard deviation (SD). Note that the learning iteration of HierarchicalChain were the sum of the iterations for learning models on Level 1, Level 2, and Level 3, and the computation of vertical and horizontal ensemble did not contribute to the number of iterations. Therefore, only one result per data/split combination was reported for HierarchicalChain. It should also be noted that the maximum per-level iteration \( \Omega \) in our experiments is set to 100, and therefore the upper limit of the iterations for GloreChain is 100 (iterations) \( \times \) 3 (levels) = 300

<table>
<thead>
<tr>
<th>Dataset (Training Data Ratio)</th>
<th>Edin (0.1)</th>
<th>CA (0.5)</th>
<th>THA (0.1)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Iterations</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Balanced Data Splitting</td>
<td>GloreChain</td>
<td>59,300 (22.968)</td>
<td>5.133 (2.871)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>143,983 (28.953)</td>
<td>30.042 (19.370)</td>
</tr>
<tr>
<td>Imbalanced Data Splitting</td>
<td>GloreChain</td>
<td>47,967 (20.388)</td>
<td>4.800 (3.818)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>131,842 (27.006)</td>
<td>40.608 (25.022)</td>
</tr>
</tbody>
</table>

Abbreviations: CA, cancer biomarkers; Edin, Edinburg myocardial infarction; THA, total hip arthroplasty.

Table 3. Total execution time results on small training data, including both mean and standard deviation (SD). The measurements are in seconds and are averaged over 4 sites. The time for HierarchicalChain includes the computation of the model on Level 1, Level 2, and Level 3, as well as the calculation of the horizontal and vertical ensembles

<table>
<thead>
<tr>
<th>Dataset (Training Data Ratio)</th>
<th>Edin (0.1)</th>
<th>CA (0.5)</th>
<th>THA (0.1)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Execution Time (Second)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Balanced Data Splitting</td>
<td>GloreChain</td>
<td>451.292 (152.107)</td>
<td>89.200 (29.019)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>1142.308 (227.684)</td>
<td>522.142 (213.102)</td>
</tr>
<tr>
<td>Imbalanced Data Splitting</td>
<td>GloreChain</td>
<td>382.483 (140.145)</td>
<td>92.733 (26.047)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>1094.283 (228.273)</td>
<td>645.392 (284.194)</td>
</tr>
</tbody>
</table>

Abbreviations: CA, cancer biomarkers; Edin, Edinburg myocardial infarction; THA, total hip arthroplasty.

Table 4. Per-iteration execution time results (total execution time in Table 3 divided by the average iterations, which are shown in Table 2)

<table>
<thead>
<tr>
<th>Dataset (Training Data Ratio)</th>
<th>Edin (0.1)</th>
<th>CA (0.5)</th>
<th>THA (0.1)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Execution Time (Second)</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Balanced Data Splitting</td>
<td>GloreChain</td>
<td>7.610 (2.565)</td>
<td>17.377 (5.653)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>7.934 (1.581)</td>
<td>17.381 (7.094)</td>
</tr>
<tr>
<td>Imbalanced Data Splitting</td>
<td>GloreChain</td>
<td>7.974 (2.922)</td>
<td>19.319 (5.426)</td>
</tr>
<tr>
<td></td>
<td>HierarchicalChain</td>
<td>8.300 (1.731)</td>
<td>15.893 (6.998)</td>
</tr>
</tbody>
</table>

Abbreviations: CA, cancer biomarkers; Edin, Edinburg myocardial infarction; THA, total hip arthroplasty.

CONCLUSION

By training the predictive models using level-wise methods, disseminating the models using a blockchain network, and combining models using a novel hierarchical consensus learning algorithm, our privacy-preserving learning framework: 1) improves prediction correctness especially for the use case of having a small training dataset for rare diseases/conditions; 2) keeps similar per-iteration execution time; 3) inherits benefits from decentralized learning and blockchain technology; and 4) records models of each level, immutably. Although such an improvement may not have clinical significance and more learning iterations are needed, we demonstrated the potential of utilizing the information from the hierarchical network-of-networks topology to improve the prediction. With further evaluations and enhancements, our proposed framework can create more generalizable predictive models to support clinical/genomic/biomedical studies within real-world research networks.

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AUTHOR CONTRIBUTIONS
T-TK contributed in conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, resources, software, validation, visualization, supervision, and writing (original draft). JK contributed in validation, visualization, and writing (review and editing). RAG contributed in data curation and writing (review and editing).

SUPPLEMENTARY MATERIAL
Supplementary material is available at Journal of the American Medical Informatics Association online.

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CONFLICT OF INTEREST STATEMENT
None declared.

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