

Gated recurrent units classify amyloidogenic proteins from sequence alone

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Abstract

The evolution of amyloidogenic proteins narrowly balances function with disorder. Amyloid fibrils confer diverse utility across domains of life— curli fibers secreted by enterobacteria such as *Escherichia coli* scaffold cell adhesion and biofilm formation (1, 2), while silk fibers spun by *Bombyx mori* protect the silkworm during metamorphosis. Both proteins aggregate in the spleens of mice upon intravenous injection; a pathology characteristic of amyloidosis (3). Evidence of heterologous proteins cross-seeding amyloid fibrillization (2–5) outlines the need to investigate environmental factors that may contribute to amyloidoses. Here, an artificial neural network model composed of gated recurrent units (GRU) is introduced to the amyloid classification task, rivalling current methods with 95% specificity, 81% sensitivity and 87% accuracy, after 10-fold cross-validation; performance is retained upon external validation. The feature extraction process used to train the model is minimal, emphasizing a data-driven paradigm to mitigate biases in feature selection. Performance, model architectures, and feature sets are compared against other amyloid prediction approaches. The model is applied to predict 310,485 amyloidogenic proteins within the Gut Phage Database, a novel collection of ~ 7.5 million bacteriophage proteins across 142,809 species (6).

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Introduction

In 1854, Rudolph Virchow first characterized the term “amyloid” in medical literature by describing insoluble, macroscopic tissue abnormalities that stained positive in starch-iodine assays (7–9). The nature of the assay led Virchow to assume these proteins were identical to starch, and named these aggregates *corpora amylacea*— a term rooted in *amylum*, the Latin word for “starch” (8, 9). In 1959, electron microscopy found that amyloids were morphologically diverse proteins, composed of fibrils ranging from 70 to 140 Å in width, and up to 16,000 Å in length (10). X-ray diffraction studies during the 1960s revealed that amyloid fibrils are composed of common cross- β motifs (11), a secondary structure derived from β -sheet proteins. Throughout the 2000s, conformational and thermodynamic studies of amyloid fibrillation found intrinsically disordered proteins and partially unfolded globular proteins driving aggregation (12–17). These proteins leverage mutations, overexpression of amyloid precursors, and free energy disruptions to seed amyloidogenesis. Aggregates remain partially unfolded, yet highly-ordered and extremely stable, arranging into cross- β structures characteristic of amyloid fibrils (12, 13).

Amyloids have long been associated with disease, but many functional roles were outlined in recent years; particularly in bacteria, fungi, and humans (1, 2, 18). In a microbiome environment, functional amyloids may leak from the gut and cross-seed with the host, driving neurodegeneration (16, 17, 19, 20), type 2 diabetes (17), and inflammatory disorder (19). The microbiome (17, 19, 21–23) and central nervous system (24) have established roles in modulating intestinal permeability, and are probable effectors of cross-seeding. Long-term transfer of healthy gut microbiota via faecal transplantation ameliorated A β deposition, tau pathology, reactive gliosis and memory impairment in a murine model (25), though etiology is intermingled with the downstream consequences of systemic inflammation. More evidence of cross-seeding via the intestinal barrier is needed to validate this hypothesis (16, 17, 26).

A nucleated growth mechanism drives amyloid aggregation through sigmoidal kinetics, composed of three stages; a lag phase, growth phase, and final plateau (13, 27). This classical nucleation theory offers consensus on macroscopic kinetics, but several models postulate different physicochemical mechanisms for self-catalyzed fibrillization (27–30). Amyloids lie on a spectrum of virulence; while all aggregates may be transmissible (31), readily infectious aggregates, such as those found in transmissible spongiform encephalopathies, are sub-classified as prions (32).

Chapter 1

Finding amyloids in the proteome

1.1 Published methods

Amino acid composition strongly indicates amyloidogenic propensity (33–36). Sequence-based prediction is a productive approach; many algorithms trained upon abstract physicochemical features classify amyloids with useful accuracy, but have notable variances in specificity and sensitivity. Several of these are focused on prion classification, such as PAPA (37), pWALTZ (28), PLAAC (38), prionW (39), and pRANK (33). These classifiers are biased towards glutamine/asparagine-enriched prion peptides within yeast, impairing validity in human disease contexts (33, 38). Recent approaches are generalized towards amyloid classification, and will be the focus of comparison in this study.

Various amyloids such as islet amyloid polypeptide (40), silkworm chorion protein (41) and *Sup35* yeast prion (42) depend on small interchangeable motifs to aggregate at a significant rate, ranging from five to seven amino acids in length (40–42). Because of this observation, some amyloid classifiers limit feature extraction to hexapeptides (35, 36, 43, 44). It is important to note that local motifs are not the sole drivers of amyloidogenesis. Global sequence specificity is directly proportional to the amyloid fibrillization rate of hen lysozyme, emphasizing that long-range interactions are relevant (34). Moreover, gatekeeper residues often flank amyloidogenic domains and neutralize electric charge (45, 46) or bind allosteric factors (47), modulating aggregation rate and further defining the spectrum of amyloidogenesis. More comprehensive models can be developed by leveraging elusive features in the protein sequence. The three most performant algorithms to-date, RFAmyloid (48), PredAmyl-NLP (49), and iAMY-SCM (50), extract distal features within the greater protein sequence and will be compared against the current model.

RFAmyloid was developed in 2018, and utilizes random forest architecture (48) based upon decision trees to classify amyloid. Two feature extraction methods are applied; SVMProt-188D extracts amino acid frequencies across the first twenty dimensions, while eight physicochemical features fill the remaining 168 dimensions: hydrophobicity, normalized Van der Waals volume, polarity, polarizability, charge, surface tension, secondary struc-

ture, and solvent accessibility (51). The 20 amino acids are placed into one of three abstract categories, on a per-feature basis. Three within-category descriptors are applied to each physicochemical feature— composition (C) captures amino acid frequency, transition (T) notes the frequency of changes to physicochemically different amino acids in the sequence, and distribution (D) describes emergent physicochemical properties with respect to the first 25, 50, and 75% of amino acids in the sequence, normalized by peptide length. The complete feature space is described in Equation 1.1 (51).

$$20 \text{ amino acids} + (C + T + D) \times 8 \text{ chemical properties} = 188 \text{ dimensions.} \quad (1.1)$$

Pse-in-one recovers a broader sequence profile by embedding positional features via pseudo components, which represent a set of feature vectors embodying the composition of amino acids within a peptide (48, 52). Short-range pseudo components are calculated using the occurrence frequencies of k nearest residues with respect to a single amino acid, while long-range psuedo components capture autocorrelation across di- and tri-peptides within the greater sequence (52).

The only shared feature between RFAmyloid and the current model is amino acid composition. Despite this, sensitivity, specificity and accuracy are highly comparable, which demonstrates that a neural network architecture can learn the amyloid classification problem without imperatively assigning a feature set. A performance trade-off is observed with the current model; RFAmyloid has over 15% lesser specificity, while sensitivity is 10% greater. Table 1.1 compares these predictors across the amyloid classifiers discussed here. The RFAmyloid training dataset filters out peptides shorter than 50 amino acids, which is a notable departure from previous methods trained exclusively on hexapeptides. The current model does not apply a minimum length restriction; the shortest amyloidogenic protein sequence in training data, physalaemin, is 11 amino acids long.

The group behind RFAmyloid introduced another tool in 2020, PredAmyl-MLP, marking an iteration upon their previous work (49). This tool applies a multilayer-perceptron neural network architecture trained on the same dataset used for RFAmyloid, now filtered using CD-HIT (53) to cluster highly similar sequences; emphasizing canonical features and reducing noise (51, 53). As in RFAmyloid, SVM-Prot188-D is applied to extract physicochemical and higher-order compositional features across the protein into 188 dimensions, but features are also filtered using the maximum relevant maximum distance method (49, 54). Psuedo components are more focused, only utilizing tri-peptide composition, and the binomial distribution method (55) is applied to find optimal tri-peptides. Model sensitivity and specificity increased by 3-5% compared to RFAmyloid, closely competing with the current model.

iAMY-SCM is the most recent amyloid classification algorithm to emerge in 2021, applying a novel score card method to classify amyloid (50). Di-peptide composition is extracted and normalized from the protein sequence, lending a 400-dimensional feature set for training (50). Di-peptide scores range from 0 to 1000, with 1000 indicating residues critical to amy-

loidogenic proteins. Notably, this method exposes specific amyloidogenic regions within a protein via dipeptide arrangements, lending more interpretability than a single classification produced by RFAmyloid (48) and the current model. Tables 1.1 and 1.2 compare accuracy, sensitivity, specificity and Matthews correlation across these models during cross-validation and external validation, respectively.

| Model | Accuracy | Specificity | Sensitivity | MCC |
|--------------|--------------|--------------|--------------|--------------|
| GRU | 0.869 | 0.952 | 0.812 | 0.751 |
| RFAmyloid | 0.892 | 0.781 | 0.927 | 0.739 |
| PredAmyl-MLP | 0.916 | 0.836 | 0.950 | 0.798 |
| iAMY-SCM | 0.895 | 0.757 | 0.954 | 0.750 |

Table 1.1: Comparison of accuracy, sensitivity, specificity, and Matthews correlation coefficients between models, on 10-fold cross-validation (48–50). Heuristics for the current model are in bold. "NA" abbreviates "not available", indicating that the heuristic was not published. "MCC" abbreviates "Matthews correlation coefficient".

| Model | Accuracy | Specificity | Sensitivity | MCC |
|--------------|--------------|-------------|--------------|--------------|
| GRU | 0.889 | 1.00 | 0.818 | 0.797 |
| RFAmyloid | 0.897 | 0.818 | 0.932 | 0.757 |
| PredAmyl-MLP | NA | NA | NA | NA |
| iAMY-SCM | 0.827 | 0.606 | 0.922 | 0.570 |

Table 1.2: Comparison of accuracy, sensitivity, specificity, and Matthews correlation coefficients between models, on external validation (48–50). Heuristics for the current model are in bold. "NA" abbreviates "not available", indicating that the heuristic was not published. "MCC" abbreviates "Matthews correlation coefficient".

1.2 Model architecture & dataset collection

The current study is the first known approach using a recurrent neural network to classify amyloids using protein sequence alone. A stacked bi-directional, gated recurrent unit (GRU) architecture was applied to find amyloidogenic likelihood, by traversing tokenized amino acids of a given protein. Two stacked GRUs increase non-linearity, enabling deeper representations of abstract features in the protein sequence. A bi-directional recurrent neural network highlights features dependent on protein sequence direction with respect to N- and C- termini, and reliably predicts secondary structure prediction concerning long-range motifs (56).

Two pre-existing amyloid and prion datasets were constructed into 180 positive examples, from the AmyPro amyloid database (57) and the set of yeast prions used to train PLAAC (38). Protein sequences obtained from AmyPro are limited to broad amyloid-forming regions, as defined in literature (57). The negative dataset was constructed with protein sequences not

known to be amyloidogenic, from UniprotKB (58). A balanced proportion of peptides was collected from each species in the positive dataset, totalling 180 negative examples. The negative dataset has not been validated for amyloidogenic activity, so contamination may be present.

1.3 Feature extraction & training

A minimal set of features was extracted from protein sequences. Proteins were tokenized by amino acid into a predetermined, 1-hot-like embedding layer of 64 dimensions, for each of the 20 amino acids and a wildcard character. No inferences were made upon the protein sequence; all abstract features were left for the model to learn, in favor of a data-driven approach. This is in stark contrast to existing methods, which extract physicochemical properties, position-specific information, and 3-D structural features. Performance compares favorably against these methods using the data-driven paradigm. The model is penalized using smooth L_1 -loss, illustrated in Equation 1.2.

$$L_{1;smooth} = \begin{cases} |x| & \text{if } |x| > \alpha \\ \frac{1}{|\alpha|}x^2 & \text{if } |x| \leq \alpha \end{cases} \quad (1.2)$$

The smooth L_1 -loss function utilizes hyperparameter α to determine if the output is large or small. This loss function maintains steady gradients over large and small outputs by squaring the loss at a certain threshold, mitigating the effect of outliers in the dataset.

A 70:30 split was applied to the full dataset; 15% of the data is kept within an independent set for external validation. The model is cross-validated 10-fold during training. Regularization is applied via dropout layers of 30% probability in-between the stacked directional GRUs, and at the final layer as outputs are concatenated across both directions. Model architecture is illustrated in Supplemental Data 1. The Pytorch (59) and SKLearn (60) libraries were applied during implementation, outlined in Supplemental Data 2 as an IPython notebook. Supplemental Data 3 provides a confusion matrix, receiver operating characteristic curve, and loss plots. Tables 1.3 and 1.4 contextualize empirical performance upon 10-fold cross-validation and external validation, respectively.

| Fold | Accuracy | Specificity | Sensitivity | MCC |
|------|----------|-------------|-------------|-------|
| 1 | 0.869 | 0.952 | 0.812 | 0.751 |
| 2 | 0.853 | 0.909 | 0.810 | 0.712 |
| 3 | 0.867 | 0.938 | 0.814 | 0.742 |
| 4 | 0.866 | 0.931 | 0.818 | 0.740 |
| 5 | 0.856 | 0.943 | 0.798 | 0.727 |
| 6 | 0.849 | 0.950 | 0.786 | 0.717 |
| 7 | 0.869 | 0.952 | 0.812 | 0.751 |
| 8 | 0.846 | 0.949 | 0.782 | 0.712 |
| 9 | 0.866 | 0.938 | 0.814 | 0.742 |
| 10 | 0.860 | 0.882 | 0.839 | 0.720 |

Table 1.3: Comparison of accuracy, sensitivity, specificity, and Matthews correlation coefficient of the current model, on cross-validation. "NA" abbreviates "not available", indicating that the heuristic was not published. "MCC" abbreviates "Matthews correlation coefficient".

| Fold | Accuracy | Specificity | Sensitivity | MCC |
|------|----------|-------------|-------------|-------|
| 1 | 0.870 | 0.955 | 0.813 | 0.753 |
| 2 | 0.852 | 0.913 | 0.807 | 0.712 |
| 3 | 0.870 | 0.955 | 0.813 | 0.754 |
| 4 | 0.870 | 0.917 | 0.833 | 0.745 |
| 5 | 0.889 | 1.00 | 0.818 | 0.798 |
| 6 | 0.870 | 0.955 | 0.813 | 0.754 |
| 7 | 0.870 | 0.955 | 0.813 | 0.753 |
| 8 | 0.870 | 1.00 | 0.794 | 0.767 |
| 9 | 0.870 | 0.955 | 0.813 | 0.754 |
| 10 | 0.796 | 0.786 | 0.808 | 0.592 |

Table 1.4: Comparison of accuracy, sensitivity, specificity, and Matthew correlation coefficient of the current model, on external validation. "NA" abbreviates "not available", indicating that the heuristic was not published. "MCC" abbreviates "Matthews correlation coefficient".

Chapter 2

Amyloid in gut bacteriophages

2.1 Current evidence

Bacteriophages are the dominant viral constituents of the microbiome, and are concentrated in the gut (61–64). Most phage-derived peptides are not well-represented in reference databases (6). Discussion of bacteriophages in disease etiology has risen in literature, with the gut-brain axis being a prominent concern (63–65). Studies have observed the presence of bacteriophage communities within cerebrospinal fluid (CSF) donated by healthy individuals, challenging prior assumptions of CSF sterility in the absence of disease (66, 67).

Tetz and Tetz (68) marks the first in-depth investigation of amyloidogenic propensity in phagobiota. With a focus on the prion amyloid subclass, Tetz and Tetz (68) examined 370,617 bacteriophage protein sequences available on UniprotKB (58) for prion-like domains using PLAAC (38): a hidden–Markov model that profiles sequence similarity to yeast prions, via the maximum likelihood estimation (38, 68). PLAAC found 5040 putative prions with a log likelihood ratio greater than 0.003; representing 1.35% of phage peptide sequences available on UniprotKB. Tetz and Tetz (68) characterized cell attachment and DNA injection as the most abundant functional groups of proteins matching these amyloid predictions. Several pathogenic amyloid proteins in humans readily interact with nucleic acids (69–71), indicating the possibility of cross-seeding.

Interestingly, filamentous phage capsid protein g3p binds A β in a manner inducing disaggregation, through a general amyloid–interaction motif (GAIM) dependent on hydrophobic interactions (72). In a murine model, intraperitoneal injection of a fusion protein containing this GAIM reduced A β deposition, phospho-tau pathology, atrophy, and improved cognition in murine models of Alzheimer’s (73). While the g3p protein disrupts amyloidogenesis, it stands to reason that other phage peptides have a converse effect on amyloid aggregation, possibly those sharing sequence profile similarity with yeast prions (68).

Camarillo-Guerrero et al. (6) presents the Gut Phage Database (GPD), a massive collection of 7,581,807 phage-derived proteins derived from human gut microbiota, ranging across 142,809 bacteriophage species; of which the majority are uncharacterized to-date. Less than

1% of canonical sequences overlap with bacteriophages represented in the NCBI RefSeq database (6, 74).

2.2 Results

This study iterates upon the prior work of Tetz and Tetz (68) by applying a highly-specific model to classify aggregation-prone proteins, within the greater amyloid class. The current model predicts 310,947 phage proteins have amyloidogenic propensity, totalling 4.1% of the GPD.

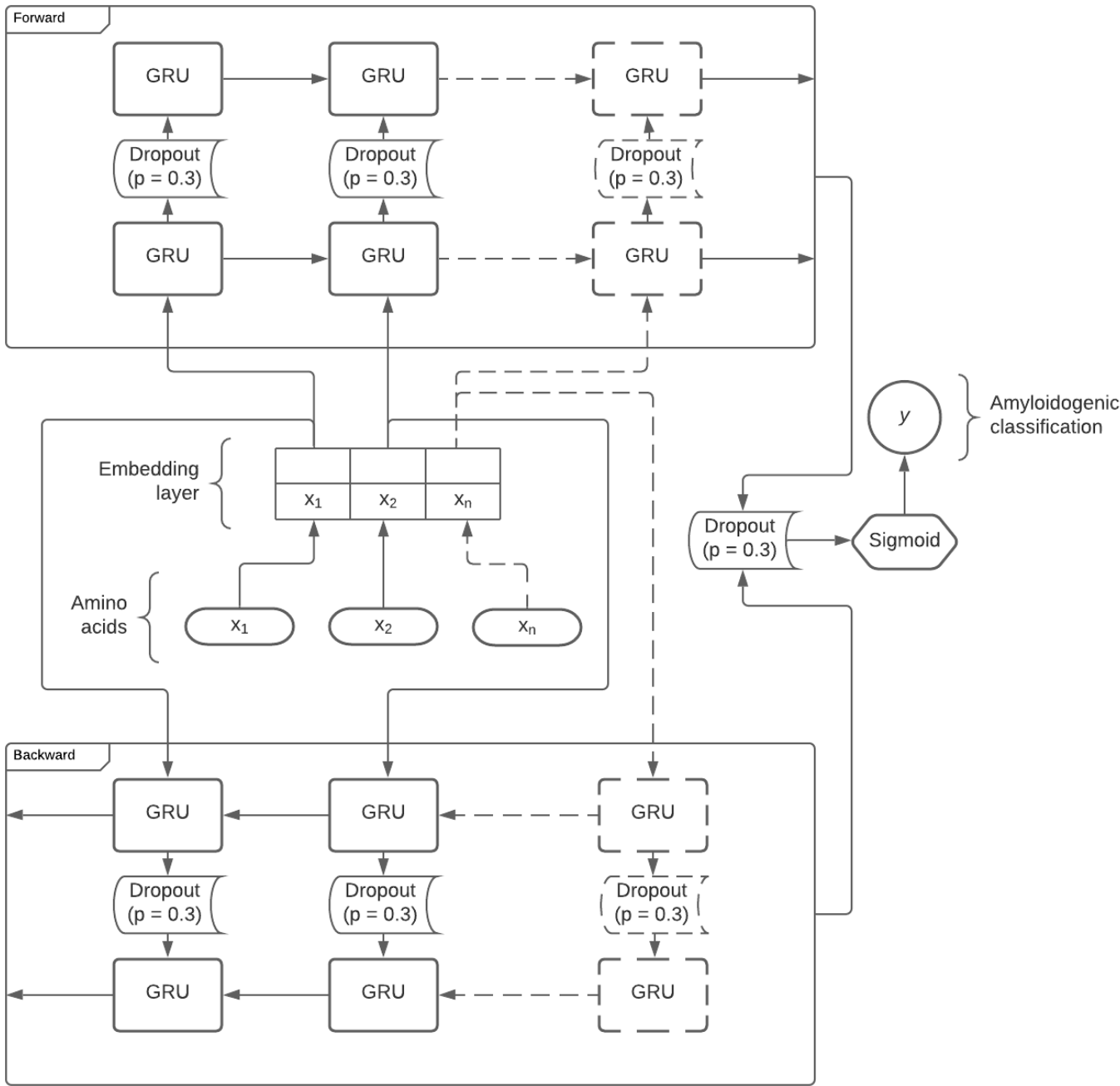
Chapter 3

Conclusions and Future Work

A gated recurrent unit architecture was introduced to classify amyloid protein sequences with high specificity, offering competitive results despite minimal feature extraction. Inclusion of higher-order features, such as tri-peptide composition and physicochemical properties, may close the gap in sensitivity with existing amyloid classifiers. These features can also improve biological interpretability by outlining regions-of-interest within the predicted amyloid sequence.

The current model was applied to predict novel amyloidogenic proteins in phagobiota; these predictions cover 4.10% of the Gut Phage Database. Analysis is ongoing to characterize the broader functions of these proteins, and possible etiologies in human disease.

Supplemental Data 1

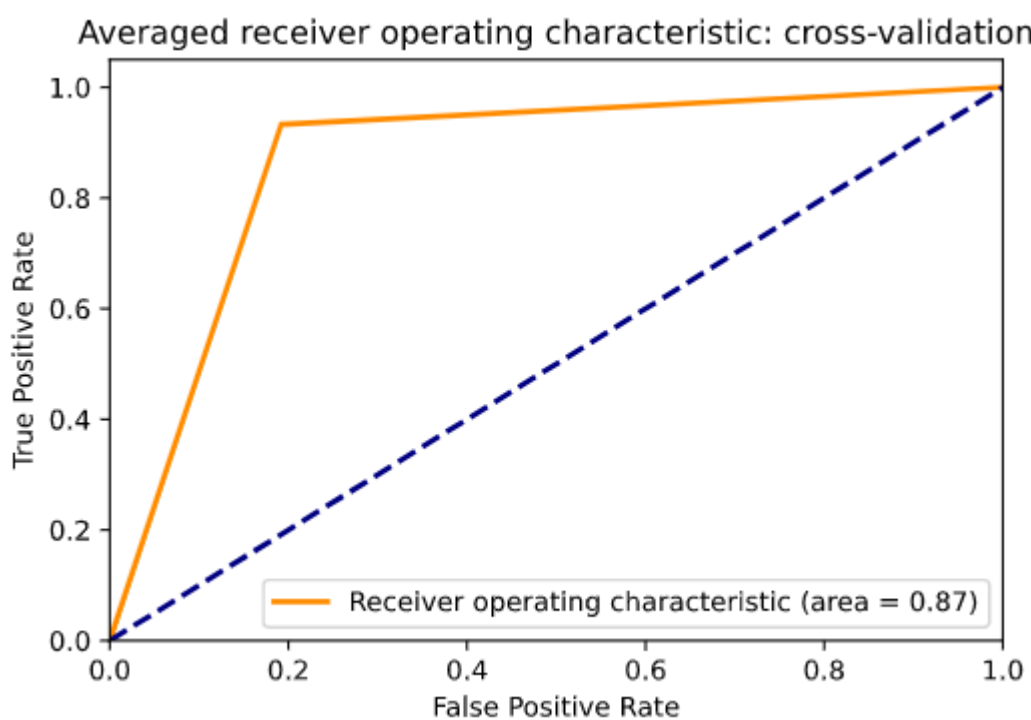
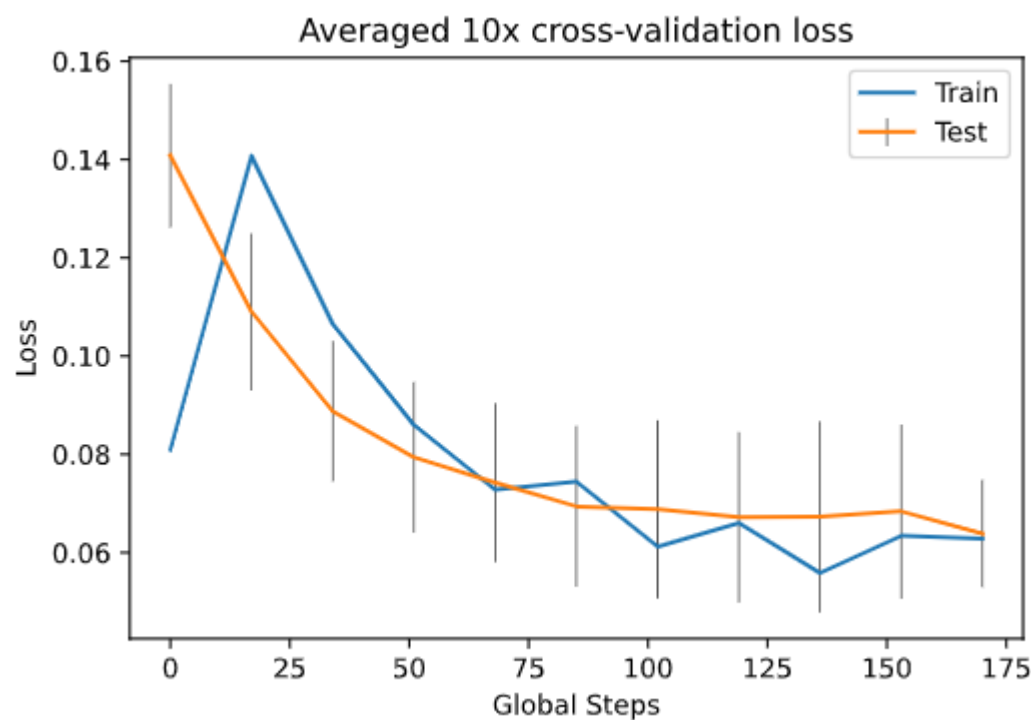


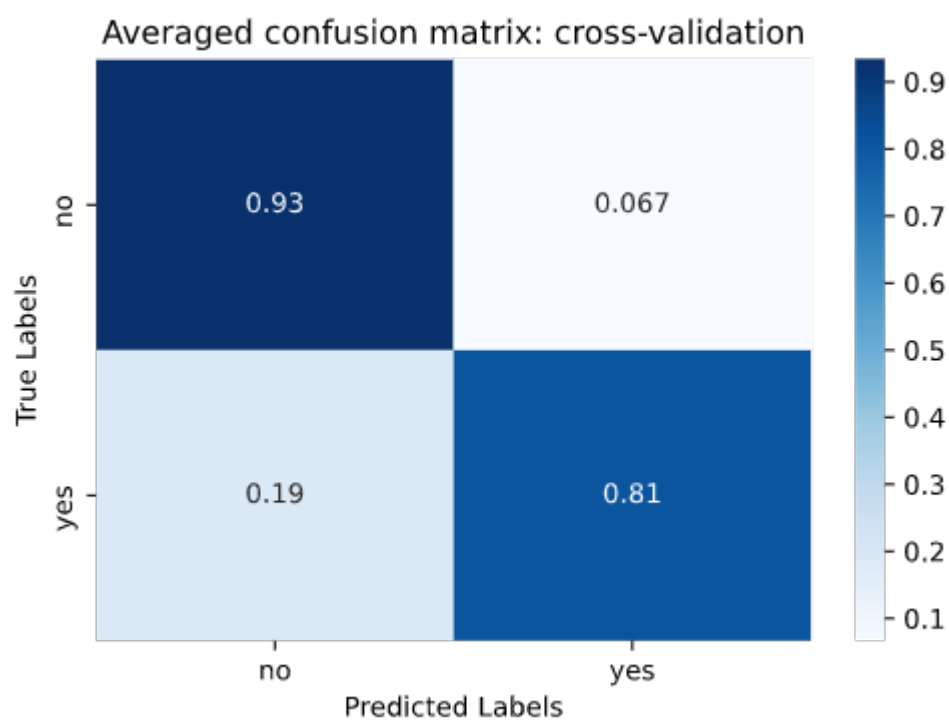
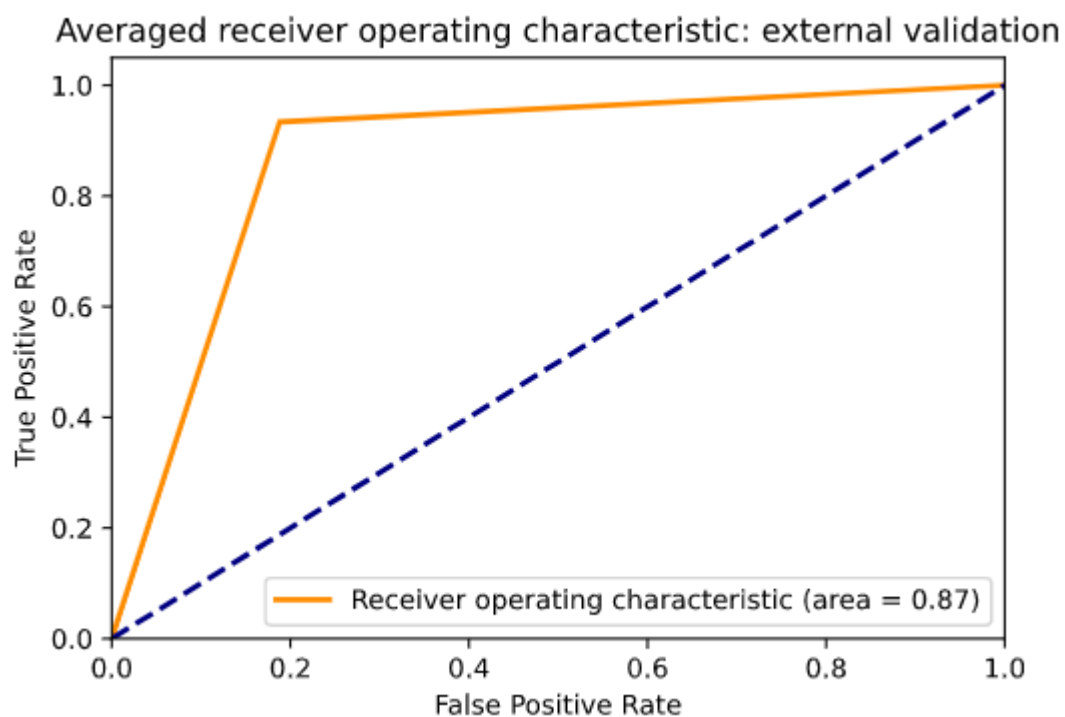
Supplemental Data 2

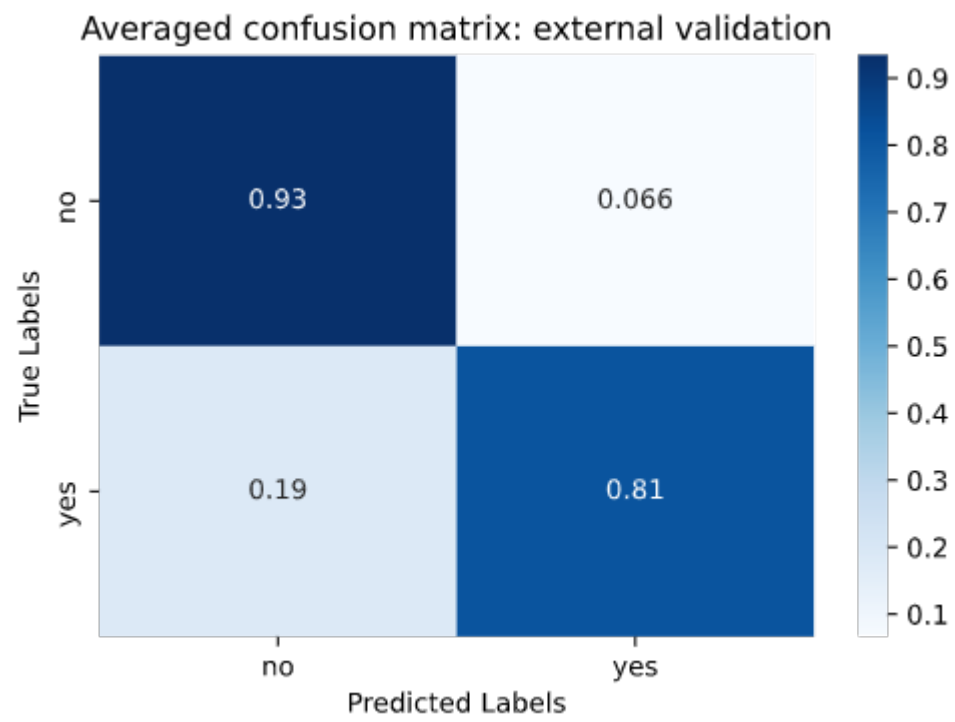
Output data and an IPython notebook are maintained at this GitHub repository:

<https://github.com/mocklee/AgGRU-Check>

Supplemental Data 3







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