

ML Lab Research Proposal

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Project: HF-Bench — Classical ML vs. Deep Learning for 30-Day Heart Failure Readmission Prediction on MIMIC-IV

Dataset: MIMIC-IV v2.2 (PhysioNet, public credentialed access)

Target compute: 1x AMD Instinct MI300X (deep learning arms); CPU-only for classical ML arms. Est. total wall-clock: 2–3 days.

Target journal: npj Digital Medicine / PLOS ONE / JAMIA Open

1. Problem Statement

Heart failure (HF) is one of the leading drivers of preventable hospital readmission in the United States, with roughly 20–25% of patients returning within 30 days of discharge and associated costs exceeding \$26 billion annually. Accurate risk stratification at the point of discharge could enable targeted interventions — closer follow-up, medication adjustment, home health referrals — that reduce readmissions and improve patient outcomes.

A substantial body of literature applies machine learning to this problem, with most published models relying on classical approaches: logistic regression, random forest, or gradient-boosted trees (XGBoost/LightGBM). Deep learning methods — particularly attention-based architectures (TabTransformer, FT-Transformer) and purpose-built tabular DL models (TabNet) — have shown promise on general tabular benchmarks but remain underexplored in the clinical HF readmission setting.

Critically, most existing comparisons are conducted on single-institution datasets, rarely report subgroup performance across demographic groups, and do not systematically evaluate the interpretability-performance tradeoff that clinicians actually care about. This project fills that gap: we conduct the first systematic head-to-head benchmark of classical ML versus deep learning for 30-day HF readmission prediction on MIMIC-IV, with explicit evaluation of fairness across demographic subgroups and clinical interpretability via SHAP attribution.

2. Proposed Approach

We benchmark five model families on an identical MIMIC-IV HF cohort, held to a strict train/validation/test split (temporal, to simulate real deployment):

Model	Family	Hardware	Why included
Logistic Regression	Classical	CPU	Clinical standard; maximally interpretable baseline
XGBoost	Classical (GBDT)	CPU	Current SOTA on tabular EHR; sets the bar DL must beat
LightGBM	Classical (GBDT)	CPU	Faster GBDT alternative; cross-checks XGBoost results
TabNet	Deep Learning	MI300X	Attention-based tabular DL with built-in feature selection

FT-Transformer	Deep Learning	MI300X	Feature Tokenizer + Transformer; SOTA on tabular DL benchmarks
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All models are trained on identical feature sets. Hyperparameters are tuned via Optuna (100 trials per model) on the validation set. Feature importance is assessed for all models using SHAP — TreeExplainer for classical models, GradientExplainer for DL models — enabling a direct interpretability comparison across paradigms.

Subgroup analyses by race/ethnicity, sex, and age group (≥ 65 vs < 65) are conducted for all five models. This surfaces not only which model performs best overall, but which model performs most equitably across patient populations — a contribution largely absent from prior benchmarks.

The core research questions are: (1) Does deep learning outperform classical ML on this task? (2) If so, at what cost to interpretability and fairness? (3) Which model a clinician should actually deploy?

3. Dataset

All data comes from MIMIC-IV v2.2, a publicly available de-identified EHR dataset from Beth Israel Deaconess Medical Center, accessed via PhysioNet credentialed access (no DUA beyond PhysioNet credentialing, typically approved in 1–7 days).

Cohort definition:

- Adults (≥ 18 years) with primary ICD-10 diagnosis of heart failure (I50.x)
- Discharged alive following an inpatient hospitalization
- At least one prior admission in MIMIC-IV (required for historical feature computation)
- Estimated cohort: ~15,000–25,000 admissions
- Outcome (label): readmitted within 30 days of discharge — binary (1 = readmitted, 0 = not)

MIMIC-IV module	Features extracted
admissions / patients	Age, sex, race/ethnicity, insurance type, admission source, length of stay
diagnoses_icd	Comorbidities: diabetes, CKD, COPD, AFib, hypertension (one-hot); Charlson Comorbidity Index
labevents	BNP/NT-proBNP, creatinine, sodium, hemoglobin, eGFR at discharge + missingness flags
prescriptions	Loop diuretic dose, ACE inhibitor/ARB/ARNI, beta-blocker, total medication count
chartevents	Discharge weight, systolic BP, heart rate, O2 saturation
Engineered features	Admissions in prior 6 months, BNP delta (admission→discharge), days since last admit

Preprocessing: median imputation for missing labs + binary missingness indicators, one-hot encoding for categoricals, StandardScaler normalization (fit on train only, applied to val/test), SMOTE oversampling on training fold only. Temporal train/val/test split: 70% / 15% / 15% by admission date.

4. Evaluation Metrics

Experiment	Metric
Primary benchmark (all 5 models)	AUROC, AUPRC — primary performance metrics; target AUROC ≥ 0.75
Clinical operating point	Sensitivity + specificity at 80% specificity threshold; PPV, NPV
Calibration	Brier score + calibration plot (reliability diagram) for all models
Fairness / subgroup	AUROC stratified by race/ethnicity, sex, age ≥ 65 vs <65 — for all 5 models
Interpretability	Mean SHAP feature ranking; top-10 feature table; SHAP summary plot per model
DL vs classical gap	Delta AUROC (FT-Transformer – XGBoost); paired bootstrap significance test
Ablation	All models trained without engineered features vs full feature set; delta AUROC

Seeds: 42, 123, 7, 0, 256. All metrics reported as mean $\pm$ 95% CI across 5 seeds. Hyperparameter tuning performed on validation set only; final numbers reported on held-out temporal test set.

5. Compute Requirements

Classical ML arms (logistic regression, XGBoost, LightGBM) run entirely on CPU and require no special hardware. Deep learning arms (TabNet, FT-Transformer) use the MI300X for training and hyperparameter search.

Phase	Hardware	Estimated wall-clock
MIMIC-IV data pull + preprocessing (mimic-code SQL + pandas)	CPU	~2–4 hours (one-time)
Classical ML training + Optuna tuning (100 trials each)	CPU	~1–2 hours total
TabNet training + Optuna tuning (100 trials, 5 seeds)	1x MI300X	~4–6 hours
FT-Transformer training + Optuna tuning (100 trials, 5 seeds)	1x MI300X	~6–8 hours
SHAP attribution (all 5 models, test set)	CPU / MI300X	~1–2 hours
Subgroup analysis + figure generation	CPU	~1 hour
Total	1x MI300X node	~1.5–2 days wall-clock with buffer

GPU memory: TabNet and FT-Transformer on a ~20,000-patient tabular dataset fit comfortably within 16 GB VRAM. The MI300X (192 GB HBM3) is far above what is strictly required, but allows large Optuna search spaces and full-batch training without minibatch approximation artifacts.

Storage: ~5–10 GB for relevant MIMIC-IV tables + preprocessed .parquet cache. Well under any reasonable budget.

Software / dependencies (all pip-installable):

- Python 3.10+, pandas, numpy, scipy, scikit-learn, xgboost, lightgbm
- pytorch-tabnet, pytorch-frame (FT-Transformer implementation)
- shap, optuna, imbalanced-learn, matplotlib, seaborn
- ROCm 6.x + official PyTorch ROCm wheel (for MI300X)
- mimic-code SQL scripts for cohort extraction (MIT-licensed)
- No Docker required; standard Python venv. No CUDA-only kernels.

6. Expected Deliverables

By end of the two-week window:

- Reproducible cohort extraction pipeline (SQL + Python) from MIMIC-IV
- Trained and tuned models for all 5 model families with full cross-validation results
- Head-to-head benchmark table: AUROC, AUPRC, Brier score, sensitivity, specificity for all models
- Subgroup fairness table: AUROC by race/ethnicity, sex, and age group for all 5 models
- SHAP summary plots and top-10 feature tables for all models (classical TreeExplainer + DL GradientExplainer)
- ROC and precision-recall curves overlaid for all 5 models
- Draft methods + results sections formatted for npj Digital Medicine

Longer-term (post-sprint):

- Full manuscript submitted to npj Digital Medicine, PLOS ONE, or JAMIA Open
- Abstract submitted to AMIA Annual Symposium or AHA Scientific Sessions
- Public GitHub repo with fully reproducible pipeline, MIT-licensed

I am intentionally not promising that deep learning beats XGBoost — a negative result (classical ML matches or outperforms DL on this task) is equally publishable and arguably more clinically useful. The goal is a clean, honest, reproducible benchmark with equity findings.

7. Risk & Mitigation

Risk	Likelihood	Mitigation
MIMIC-IV PhysioNet credentialing delayed	Medium	Apply immediately (typically 1–7 days). Use MIMIC-IV demo (100-patient public subset, no credentialing) to build and validate full pipeline in parallel. No time lost.
DL models underperform XGBoost on this task	Medium	This is a valid, publishable finding — 'classical ML matches DL on tabular EHR data with better interpretability' is a useful clinical result. Frame as the honest benchmark it is. XGBoost arm is the safety net.

FT-Transformer or TabNet fails to converge / hyperparameter sensitivity	Medium	Isolate DL training behind unit-tested modules. Fall back to MLP (simpler DL baseline) if attention-based models misbehave. ROCm compatibility tested in smoke run before full training.
SHAP GradientExplainer is slow or unstable on DL models	Medium	Use SHAP KernelExplainer as fallback (model-agnostic, slower but reliable). Limit explainability analysis to a random 500-patient subset of the test set if runtime is prohibitive.
Subgroup sample sizes too small for reliable AUROC estimates	Low	Report CIs explicitly. Collapse granular race categories to broader groups if $n < 100$. Flag small- n subgroups rather than omitting them — reviewers accept this with honest reporting.
Class imbalance (~20% positive) degrades minority-class recall	Low	SMOTE on training fold + <code>class_weight='balanced'</code> for classical models + <code>pos_weight</code> in DL loss. Report AUPRC alongside AUROC. Tune threshold to clinical operating point rather than defaulting to 0.5.

I will check in at the 1-week mark with classical ML results and Stage-1 DL training numbers so we can decide together whether to push for the full 5-model benchmark or trim to the strongest 3.