Evaluation of domain generalization and adaptation on improving model robustness to temporal dataset shift in clinical medicine

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Temporal dataset shift associated with changes in healthcare over time is a barrier to deploying machine learning-based clinical decision support systems. Algorithms that learn robust models by estimating invariant properties across time periods for domain generalization (DG) and unsupervised domain adaptation (UDA) might be suitable to proactively mitigate dataset shift. The objective was to characterize the impact of temporal dataset shift on clinical prediction models and benchmark DG and UDA algorithms on improving model robustness. In this cohort study, intensive care unit patients from the MIMIC-IV database were categorized by year groups (2008–2010, 2011–2013, 2014–2016 and 2017–2019). Tasks were predicting mortality, length of stay, sepsis and invasive ventilation. Feedforward neural networks were used as prediction models. The baseline experiment trained models using empirical risk minimization (ERM) on 2008–2010 (ERM[08–10]) and evaluated them on subsequent year groups. DG experiment trained models using algorithms that estimated invariant properties using 2008–2016 and evaluated them on 2017–2019. UDA experiment leveraged unlabelled samples from 2017 to 2019 for unsupervised distribution matching. DG and UDA models were compared to ERM[08–16] models trained using 2008–2016. Main performance measures were area-under-the-receiver-operating-characteristic curve (AUROC), area-under-the-precision-recall curve and absolute calibration error. Threshold-based metrics including false-positives and false-negatives were used to assess the clinical impact of temporal dataset shift and its mitigation strategies. In the baseline experiments, dataset shift was most evident for sepsis prediction (maximum AUROC drop, 0.090; 95% confidence interval (CI), 0.080–0.101). Considering a scenario of 100 consecutively admitted patients showed that ERM[08–10] applied to 2017–2019 was associated with one additional false-negative among 11 patients with sepsis, when compared to the model applied to 2008–2010. When compared with ERM[08–16], DG and UDA experiments failed to produce more robust models (range of AUROC difference, − 0.003 to 0.050). In conclusion, DG and UDA failed to produce more robust models compared to ERM in the setting of temporal dataset shift. Alternate approaches are required to preserve model performance over time in clinical medicine.

The wide-spread adoption of electronic health records (EHRs) and the enhanced capacity to store and perform computation with large amounts of data have enabled the development of highly performant machine learning models for clinical outcome predictions¹. The utility of these models critically depends on sustained performance to maintain safety², end-users’ trust, and to outweigh the high cost of integrating each model into the clinical workflow³. However, this is hindered in the non-stationary healthcare environment by temporal dataset shift due to mismatch between the data distribution on which models were developed and the distribution to which models were applied⁴.

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There has been limited research on the impact of temporal dataset shift in clinical medicine. Recent approaches largely relied on maintenance strategies consisting of performance monitoring, model updating and calibration over certain time intervals. Another approach grouped clinical features into their underlying concepts to cope with a change in the record-keeping system. Generally, these approaches either require detection of model degradation or rely on explicit knowledge or assumptions about the underlying cause of the shift. Complementary to these approaches would be ones that attempt to proactively produce robust models that incorporate relatively few assumptions on the nature of the shift.

The past decade of machine learning research offered numerous algorithms that learn robust models by using data from multiple environments to identify invariant properties. These algorithms were often developed for domain generalization (DG) and unsupervised domain adaptation (UDA). In the DG setting, the goal is to learn models that generalize to new environments unseen at training time. In the UDA setting, the goal is to adapt models to target environments using labeled samples from the source environment as well as a limited set of unlabeled samples from the target environment. If we consider EHR data across discrete time windows as related but distinct environments, then both DG and UDA settings may be suitable to combat the impact of temporal dataset shift. To date, these approaches have not been evaluated on improving model robustness to temporal dataset shift for clinical prediction tasks. Therefore, the objective was to benchmark learning algorithms for DG and UDA on mitigating the impact of temporal dataset shift on machine learning model performance in a set of clinical prediction tasks.

**Methods**

**Data source.** We used the MIMIC-IV database, which contains deidentified EHRs of 382,278 patients admitted to an intensive care unit (ICU) or the emergency department at the Beth Israel Deaconess Medical Center (BIDMC) between 2008 and 2019. For this cohort study, we considered ICU admissions sourced from the clinical information system MetaVision at the BIDMC, in which records from 53,150 patients were made available in the latest version of MIMIC-IV 1.0.

**Data access.** Data from MIMIC-IV was approved under the oversight of the Institutional Review Boards of BIDMC (Boston, MA) and the Massachusetts Institute of Technology (MIT; Cambridge, MA). Because of deidentification of protected health information, the requirement for individual patient consent was waived by the Institutional Review Boards of BIDMC and MIT. Data access was credentialled under the oversight of the data use agreement through PhysioNet and MIT. All experiments were performed in accordance with institutional guidelines and regulations.

**Cohort.** Each patient’s timeline in MIMIC-IV is anchored to a shifted (deidentified) year with which a year group and age are associated. The year group reflects the actual 3-year range (for example 2008–2010) in which the shifted year occurred, and age reflects the patient’s actual age in the shifted year. There are four available year groups in MIMIC-IV: 2008–2010, 2011–2013, 2014–2016 and 2017–2019. We included patients who were 18 years or older and randomly selected one ICU admission that occurred in the year group for each patient. As a result, each patient is represented once in our dataset and is associated with a single year group. We excluded ICU admissions less than 4 h in duration.

**Outcomes.** We defined four clinical outcomes. For each outcome, the task was to perform binary predictions over a time horizon with respect to the time of prediction, which was set as 4 h after ICU admission. Long length of stay (LOS) was defined as ICU stay greater than three days from the prediction time. Mortality corresponded to in-hospital mortality within 7 days from the prediction time. Invasive ventilation corresponded to initiation of invasive ventilation within 24 h from the prediction time. Sepsis corresponded to the development of sepsis according to the Sepsis-3 criteria within 7 days from the prediction time. For invasive ventilation and sepsis, we excluded patients with these outcomes prior to the time of prediction. Further details on each outcome are presented in the Supplementary Methods online.

**Features.** Our feature extraction followed a common procedure and obtained six categories of features including diagnoses, procedures, labs, prescriptions, ICU charts and demographics. Demographic features included age, biological sex, race, insurance, marital status and language. Clinical features were extracted over a set of time-intervals defined relative to the time of ICU admission as follows: 0–4 h after ICU admission, 0–7 days prior, 7–30 days prior, 30–180 days prior, and 180 days-any time prior. For each time interval, we obtained counts of unique concept identifiers for diagnoses, procedures, prescriptions and labs with the exception that identifiers for diagnoses and procedures were not obtained in the 0–4 h interval after admission as they were not available. We also obtained measurements for lab tests for each time interval, and measurements for chart events in the 0–4 h interval after admission. In addition, we mapped each measurement variable in each time interval to the patient-level mean, minimum and maximum. Number of extracted features for each category and time interval are listed in the Supplementary Methods online.

Feature preprocessing pruned features that had less than 25 patient observations, replaced non-zero count values with 1, encoded measurement features to quintiles, and one-hot encoded all but count features. This process resulted in binary feature matrices that were extremely sparse. All feature preprocessing procedures were fit on the training set (e.g., to determine the boundaries of each quintile) and were subsequently applied to transform the validation and test sets.
Model and learning algorithms. In all experiments, we leveraged fully connected feedforward neural network (NN) models for prediction, as they enable flexible learning for differentiable objectives across algorithms. The standard algorithm to learn models is the empirical risk minimization (ERM) algorithm in which the objective is to minimize average training error without considerations of environment annotations (year groups in our case).

In this study, we largely employ DG and UDA algorithms that learn to produce representations that exhibit certain invariances across environments by leveraging the training data and their corresponding year groups. One exception is Group distributionally robust optimization (GroupDRO)\textsuperscript{17}, which does not “learn” invariances but instead minimizes training error in the worst-case training environment by increasing the importance of environments with larger errors. DG algorithms that learn invariant representations include Invariant risk minimization (IRM)\textsuperscript{18} and distribution matching. IRM aims to learn a latent representation (i.e., hidden layer activations) where the optimal classifier leveraging that representation is the same for all environments. Distribution matching algorithms include correlation alignment (CORAL)\textsuperscript{19}, which seeks to match the mean and covariance of the distribution of the data encoded in the latent space across environments; maximum mean discrepancy (MMD)\textsuperscript{20}, which minimizes distribution discrepancy between predictions belonging to different environments; and domain adversarial learning (AL)\textsuperscript{21,22}, which matches the distributions using an adversarial network and an objective that minimizes discriminability between environments. The adversarial network used in this study is a NN model with one hidden layer of dimension 32. Since distribution matching algorithms (CORAL, MMD, and AL) do not require outcome labels, they can be leveraged for DG as well as UDA.

Model development. We conducted baseline, DG and UDA experiments. The baseline experiment consisted of several aspects. First, to characterize temporal dataset shift on model performance, we trained models with ERM on the 2008–2010 group (ERM[8–10]) and evaluated these models in each subsequent year group. Next, to describe the extent of temporal dataset shift, we compared the performance of ERM[8–10] in each subsequent year group with models trained using ERM on that year group. Difference in performance in the target year group (2017-2019) between ERM[8–10] and models trained and evaluated on the target year group (ERM[17–19]) described the extent of temporal dataset shift in the extreme scenario in which models were developed on the earliest available data and were never updated. All models in the baseline experiment used ERM.

For DG and UDA experiments, model training was performed on 2008–2016, with UDA also incorporating unlabelled samples from the target year group. Performance of DG and UDA models were compared with ERM models trained using 2008–2016 (ERM[8–16]) as these are the fairest ERM comparators for DG and UDA models. For models in the DG and UDA experiments, we focused on their performance in the target year group, but also described their performance in 2008–2016.

Data splitting procedure. Data splitting procedure was performed separately for each task and experiment (see Fig. 1). The baseline experiment split each year group into 70% training, 15% validation and 15% test sets. In DG and UDA experiments, the training set included 85% data from 2008 to 2010 and 2011–2013, and 45% of data from 2014 to 2016. The validation set included 35% of data from 2014 to 2016 (chosen because of its temporal proximity to the target year group). The test set included the same 15% from each year group as the baseline experiment, which allowed us to compare model performance across experiments and learning algorithms on the same patients. For UDA, training year groups were combined into one group, and unlabeled samples of various sizes (100, 500, 1000, and 1500) from the target year group (2017–2019) were leveraged for unsupervised distribution matching.

Model training. We developed NN models on the training sets of each experiment for each task and selected hyperparameters based on performance in the validation sets. DG and UDA models used the same model hyperparameters as ERM[8–16], but involved an additional search over the algorithm-specific hyperparameter that modulated the impact of the algorithm on model learning. For all experiments, we trained 20 NN models using the selected hyperparameters for each combination of outcome, learning algorithm and experiment-specific characteristic (for example, year group in baseline experiment and size of unlabeled samples for UDA). Further details on the models, learning algorithms, as well as hyperparameter selection and model training procedures are presented in the Supplementary Methods online.

Model evaluation. We evaluated models on the test sets of each experiment. Model performance was evaluated using area-under-receiver-operating-characteristic curve (AUROC), area-under-precision-recall curve (AUPRC) and the absolute calibration error (ACE)\textsuperscript{23}. ACE is a calibration measure similar to the integrated calibration index\textsuperscript{24} in that it assesses overall model calibration by taking the average of the absolute deviations from an approximated perfect calibration curve. The difference is that ACE uses logistic regression for approximation instead of locally weighted regression such as LOESS.

To aid clinical interpretation of the impact of temporal dataset shift and its mitigation strategies, we translated change in performance to interpretable threshold-based metrics (including sensitivity and specificity) across clinically reasonable threshold levels. We chose the task with the most extreme temporal dataset shift. Next, we set up a scenario with 100 hypothetical ICU patients and estimated the number of patients with and without a positive label using average prevalence from 2008 to 2019. We then estimated the number of false-positive (FP) and false-negative (FN) predictions using the average sensitivity and specificity for: (1) ERM[8–10] in 2008–2010, illustrating the results of initial model development with training and test sets in 2008–2010, and representing performance anticipated by clinicians applying the model to patients admitted in 2017–2019 if the model is not.
updated; (2) ERM[8–10] in 2017–2019, illustrating the actual performance of the earlier model on patients, or the impact of temporal dataset shift; (3) ERM[8–16] in 2017–2019, illustrating the ERM comparators for DG and UDA models; (4) models trained using a representative approach from DG or UDA; and (5) ERM[17–19] in which training and test sets are both using 2017–2019 data.

**Statistical analysis.** For each combination of outcome, experiment-specific characteristic (e.g., year group in the baseline experiment), and evaluation metric (AUROC, AUPRC and ACE), we reported the median and 95% confidence interval (CI) of the distribution over mean performance (across 20 NN models) in the test set obtained from 10,000 bootstrap iterations. To compare models (for example, learned using IRM vs. ERM[8–16]) in the target year group, metrics were computed over 10,000 bootstrap iterations and the resulting 95% confidence interval of the differences were used to determine statistical significance.

Model training was performed on an Nvidia V100 GPU. Analyses were implemented in Python 3.8, Scikit-learn 0.24 and Pytorch 1.7. The code for all analyses is open-source and available at https://github.com/sungrsear/ mimic4ds_public.

**Results**

Cohort characteristics for each year group and outcome are presented in Table 1. Figure 2 shows performance measures (AUROC, AUPRC and ACE) of ERM[8–10] models in each year group vs. models trained on that year group. Largest temporal dataset shift was observed for sepsis predictions in 2017–2019 (drop in AUROC, 0.090; 95% CI 0.080–0.101).

Figure 3 illustrates change in the performance measures of DG and UDA models in the target year group (2017–2019) relative to ERM[8–16]. In addition, change in performance measures of ERM[8–10] and ERM[17–19] are plotted in grey for comparison. ERM[8–16] performed better than ERM[8–10] (largest gain in AUROC, 0.049; 95% CI 0.041–0.057; see Supplementary Table S1 online), but performed worse than ERM[17–19] (worst drop in AUROC, 0.071; 95% CI 0.062–0.081) with some exceptions in mortality and invasive ventilation predictions (see Supplementary Table S2 online). Performance of DG and UDA models was similar to ERM[8–16] and while some models performed significantly better than ERM[8–16], others performed significantly worse with all differences being relatively small in magnitude (see Supplementary Table S3). Correspondingly, shifts in the cumulative distribution of predicted probabilities across year groups were not reduced by DG and UDA relative to ERM (Supplementary Fig. S1), and model selection for DG and UDA algorithms tended to select small algorithm-specific hyperparameter values with little impact on model training (Supplementary Methods). However, increasing the magnitude of these hyperparameters did not result in performance gains (see Supplementary Fig. S2, S3, and S4).
Table 2 illustrates a clinical interpretation of temporal dataset shift in sepsis prediction using a scenario of 100 consecutively admitted patients to the ICU between 2017 and 2019 with a risk threshold of 10% and an estimated outcome prevalence of 11% (see Supplementary Table S5 for results across thresholds from 5 to 45%).

ERM[8–10] applied to 2017–2019 was associated with one additional FN among 11 patients with sepsis and 7 additional FP among 89 patients without sepsis when compared to the model applied to 2008–2010. FN with AL, as the representative mitigation approach, was similar to ERM[8–16].

Table 1. Cohort characteristics by year group. pos: positive labels; SD: standard deviation.

Table 2 illustrates a clinical interpretation of temporal dataset shift in sepsis prediction using a scenario of 100 consecutively admitted patients to the ICU between 2017 and 2019 with a risk threshold of 10% and an estimated outcome prevalence of 11% (see Supplementary Table S5 for results across thresholds from 5 to 45%). ERM[8–10] applied to 2017–2019 was associated with one additional FN among 11 patients with sepsis and 7 additional FP among 89 patients without sepsis when compared to the model applied to 2008–2010. FN with AL, as the representative mitigation approach, was similar to ERM[8–16].

Discussion
Our results revealed heterogeneity in the impact of temporal dataset shift across clinical prediction tasks, with the largest impact on sepsis prediction. When compared to ERM[8–16], DG and UDA algorithms did not substantially improve model robustness. In some cases, DG and UDA algorithms produced less performant models than ERM. We also illustrated the impact of temporal dataset shift and the effect of mitigation approaches for clinical audiences so that they can determine whether the extent of dataset shift precludes utilization in practice.

The heterogeneity of impact by temporal dataset shift as characterized by our baseline experiment echoes the mixed results in model deterioration across several studies that made predictions of clinical outcomes in various
populations. This calls for careful investigation of potential model degradation due to temporal dataset shift at both the population and task level. In addition, these investigations should translate model degradation, typically measured as change in AUROC, into change in clinically relevant performance measures or utility in allocation of resources, and place its impact in the context of clinical decision making and downstream processes.

This study is one of the first to benchmark the capability of DG and UDA algorithms on EHR data across multiple clinical prediction tasks to mitigate the impact of temporal dataset shift. Our findings align with recent empirical evaluations of DG algorithms demonstrating that they do not outperform ERM under distribution shift across data sources in real-world clinical datasets as well as non-clinical datasets. The reasons underlying the failure of DG algorithms are topics of active research, with several recent works offering theory to explain why models derived with IRM and groupDRO are typically not more robust than ERM in practice. Furthermore, other work has demonstrated that UDA objectives based on distribution matching, such as AL, CORAL, and MMD, failed to improve generalization to the target domain under shifts in the outcome rate or in the association between the outcome and features. These findings highlight the difficulty of improving robustness to dataset shift with methods that estimate invariant properties without explicit knowledge of the type of dataset shift.

Future research should continue to explore methods that have demonstrated success in improving model robustness outside of clinical medicine and identify the types of shift for which these methods might be effective. For instance, models adapted from foundation models pre-trained on diverse datasets have displayed impressive performance gains in addition to robustness to various types of dataset shifts. Domain adaptation methods that leverage distinct latent domains in the target environment, data augmentation including the use of adversarial examples, similarities between the source and target environments, and semi-supervised learning that alternatively generate pseudo labels and incorporate them into re-training the model have all demonstrated success in various domains. In addition, domain knowledge as to which causal mechanisms are likely to be stable or change across time (for example, the causal effect of a disease on patient outcome versus the policy used to prescribe a specific drug) can be incorporated into learning robust models. We observed heterogeneity across tasks in the impact of temporal dataset shift on model performance. While investigating the cause of the heterogeneity is beyond the scope of this work, it is an important issue to investigate in future research.

Strengths of this study include the use of multiple clinical outcomes and the illustration of temporal dataset shift and its mitigations using more clinically relevant metrics. There are several limitations in this study. First,
the coarse characterization of temporal dataset shift did not offer insight about the rate at which model performance deteriorated. This was due to the deidentification in the MIMIC-IV database that left year group as the only time information that followed a correct chronological order across patients. Second, using data from the target year group to estimate best-case models is not realistic in real-world deployment as such data might not be available. Third, our assessment of clinical implications did not consider clinical use-cases in which the model alerts physicians of patients with the highest risks (i.e., acting on a threshold that is adaptively selected)\textsuperscript{50}. In those scenarios, the amount of agreement in the ranking of risks between models need to be additionally considered. Finally, our choice of methods for DG and UDA algorithms were limited to those that estimate invariant properties across training data environments.

In conclusion, DG and UDA using methods that estimate invariant properties across environments failed to produce more robust models compared to ERM in the setting of temporal dataset shift. Alternate approaches are required to preserve model performance over time in clinical medicine.

Figure 3. Difference in mean performance of DG and UDA approaches relative to ERM\textsuperscript{[8–16]} in the target year group (2017–2019). Performance of ERM\textsuperscript{[8–10]} (train set 2008–2010 and test set 2017–2019, dashed line) and ERM\textsuperscript{[17–19]} (train and test sets 2017–2019, solid line) models are also shown for comparison. Error bars indicate 95% confidence interval obtained from 10,000 bootstrap iterations. Here, we show results from three of the four experimental conditions using differing number of unlabelled samples for UDA—we did not observe meaningful differences across the number of unlabelled samples evaluated. Numerical representation of the performance measures relative to ERM\textsuperscript{[8–16]} are presented in Supplementary Table S3. LOS: length of stay; ERM: empirical risk minimization; IRM: invariant risk minimization; AL: adversarial learning; GroupDRO: group distributionally robust optimization; CORAL: correlation alignment; MMD: maximum mean discrepancy; AUROC: area under the receiver operating characteristics curve; AUPRC: area under the precision recall curve; ACE: absolute calibration error; domain generalization: DG; unsupervised domain adaptation: UDA.
There are 100 consecutive patients admitted to the ICU between 2017 and 2019. Management will differ depending on whether the risk of sepsis is greater than 10% at 4 h after admission. The table below illustrates the anticipated performance metrics of ERM[8–10] based on their performance in 2008–2010 (first column), the implications of dataset shift on ERM[8–10] indicated by their actual performance in 2017–2019 (second column), results of updating the model ERM[08–16] and mitigation by a representative UDA approach (third and fourth columns), and the ERM[17–19] (last column).

| Training set | 2008–2010 | 2008–2010 | 2008–2016 + 1500 unlabelled samples from 2017 to 2019 | 2017–2019 |
|--------------|------------|------------|-------------------------------------------------|------------|
| Test set     | 2008–2010  | 2017–2019  | 2017–2019                                       | 2017–2019  |
| Learning Algorithm | ERM      | ERM      | AL (UDA)                  | ERM        |

| Diagnostic properties in test set | Sensitivity | Specificity | PPV | NPV | False positives for 89% patients who did not develop sepsis | False negatives for 11% patients that developed sepsis |
|-----------------------------------|-------------|------------|-----|-----|---------------------|---------------------|
|                                   | 0.65        | 0.64       | 0.22| 0.94| 25                   | 4                   |
|                                   | 0.57        | 0.64       | 0.16| 0.92| 32                   | 5                   |
|                                   | 0.76        | 0.68       | 0.22| 0.95| 29                   | 3                   |
|                                   | 0.75        | 0.68       | 0.22| 0.95| 28                   | 3                   |
|                                   | 0.61        | 0.74       | 0.23| 0.94| 23                   | 4                   |

Table 2. Clinical interpretation of temporal dataset shift in sepsis prediction. Sepsis was chosen as this task exhibited the most discrimination deterioration due to temporal dataset shift. ERM: empirical risk minimization; AL: domain adversarial learning; UDA: unsupervised domain adaptation. *10% threshold for sepsis was chosen as a clinically reasonable value although results across thresholds from 5 to 45% are shown in Supplementary Table S5. †ERM[8–16] models are the fair ERM comparators to DG and UDA models given they were trained using 2008–2016. ‡Outcome prevalence was estimated based on average sepsis prevalence from 2008 to 2019. The table illustrates the results of initial model development with training and evaluation in the earliest period or 2008–2010 (first column), which represents performance anticipated by clinicians applying the model to patients admitted in 2017–2019 if the model is not updated. The second column shows actual performance of that model on their patients, or the impact of temporal dataset shift. In other words, the first two columns illustrate the clinical impact of temporal dataset shift for the task with the most extreme dataset shift, namely sepsis. It shows that for the 11 patients who developed sepsis, the false negative rate increased by 1 patient. The table also shows the impact of retraining with the more updated data (third column), and one approach to mitigate dataset shift, namely AL (UDA) (fourth column). Results of AL (UDA) was almost identical to third column (ERM[8–16]). For illustrative purposes, it also shows the ERM[17–19] in which training and test sets are both 2017–2019 data.

Data availability
The MIMIC-IV dataset analyzed during the current study is available from https://mimic.mit.edu/docs/iv/. The code for all analyses is open-source and available at https://github.com/sungreresearch/mimic4ds_public.

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Author contributions

All authors conceptualized and designed the study. L.L.G., L.S., and A.J. acquired the data. L.L.G. and S.R.P. analyzed the data. All authors interpreted the data, drafted and revised the manuscript critically for important
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**Competing interests**
The authors declare no competing interests.

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