Interpretable Data-Based Explanations for Fairness Debugging

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ABSTRACT
A wide variety of fairness metrics and eXplainable Artificial Intelligence (XAI) approaches have been proposed in the literature to identify bias in machine learning models that are used in critical real-life contexts. However, merely reporting on a model’s bias, or generating explanations using existing XAI techniques is insufficient to locate and eventually mitigate sources of bias. In this work, we introduce Gopher, a system that produces compact, interpretable and causal explanations for bias or unexpected model behavior by identifying coherent subsets of the training data that are root-causes for this behavior. Specifically, we introduce the concept of causal responsibility that quantifies the extent to which intervening on training data by removing or updating subsets of it can resolve the bias. Building on this concept, we develop an efficient approach for generating the top-k patterns that explain model bias that utilizes techniques from the ML community to approximate causal responsibility and uses pruning rules to manage the large search space for patterns. Our experimental evaluation demonstrates the effectiveness of Gopher in generating interpretable explanations for identifying and debugging sources of bias.

1 INTRODUCTION
Machine learning (ML) is being increasingly applied to decision-making in sensitive domains such as finance, healthcare, crime prevention and justice management. Although it has the potential to overcome undesirable aspects of human decision-making, concern continues to mount that its opacity perpetuates systemic biases and discrimination reflected in training data [3, 4, 47]. This concern gives rise to increasing regulations and demands for generating human understandable explanations for ML algorithms behavior, an issue addressed by the rapidly growing field of eXplainable Artificial Intelligence (XAI); see [67] for a recent survey.

The development of XAI methods is motivated by technical, social and ethical objectives [12, 24, 46, 48, 56] including: (1) providing users with actionable insights to change the results of algorithms in the future, (2) facilitating the identification of sources of harms such as bias and discrimination, and (3) providing the ability to debug ML algorithms and models by identifying errors or biases in training data that result in adverse and unexpected behavior.

Most XAI research to date has focused on generating feature-based explanations, which quantify the extent to which input feature values contribute to an ML model’s predictions. In this direction, methods based on feature importance quantification [5, 24, 30, 57, 59, 60, 66], surrogate models, causal and counterfactual methods [60, 75, 76], and logic-based approaches [23, 40, 81] are designed to reveal the dependency patterns between the input features and output of ML algorithms. These methods differ in terms of whether they address correlational, causal, counterfactual or contrastive patterns. While such explanations satisfy the aforementioned objective (1) and (2), if we only consider the test data as a source of bias, they fall short in generating diagnostic explanations that let users trace unexpected or discriminatory algorithmic behavior back to its training data. Hence, they fail to satisfy objective (3) if the training data is the source of the bias and do not fulfill objective (4). Indeed, information solely about output-input feature dependency is insufficient to explain discriminatory and unexpected algorithmic decisions in terms of data errors and biases introduced during data collection and preparation. While a feature-based explanation can identify which features of a test data point are correlated with a misclassification or bias, it does not explain why the model exhibits this bias. To do this, we must trace the bias back to the data used to train the model. For example, feature-based approaches cannot generate explanations of the form: “The main source of gender bias for this classifier, which decides about loan applications, is its training data, which exhibits bias against the credit scores of unmarried females who are house owners.”

In this paper, we take the first step toward developing a framework for generating diverse, compact and interpretable training data-based explanations that relate an unexpected and discriminatory behavior of an ML algorithm to its training data, not its input features. Specifically, we introduce Gopher, a system that explains ML model bias by evaluating its training data to find cohesive patterns of data points that, when eliminated, remove the bias. This means that if the ML algorithms had been trained on the modified training data, it would not have exhibited the unexpected or undesirable behavior or would have exhibited this behavior to a lesser degree. Explanations generated by our framework, which complement existing approaches in XAI, are crucial for helping system developers and ML practitioners to debug ML algorithms for data errors and bias in training data, such as measurement errors and misclassifications [35, 42, 94], data imbalance [27], missing data and selection bias [29, 62, 63], covariate shift [74, 82], technical biases introduced during data preparation [85], and poisonous data points.
Figure 1: An overview of explanations generated by XAI approaches for an ML algorithm built using the UCI Adult dataset (see Section 6 for details), which predicts if an individual earns ≥ 50K/year. (a) Feature-based explanations rank attributes in descending order of importance toward model behavior. (b) Instance-based explanations output a list of data points most responsible for model behavior. Gopher generates two kinds of explanations: (c) top-k explanations identifying the patterns most causally responsible for model bias, and (d) updates to the top-k explanations that reduce the model’s bias by the most.

Injected through adversarial attacks [36, 43, 65, 83]. It is known in the algorithmic fairness literature that information about the source of bias is critically needed to build fair ML algorithms because no current bias mitigation solution fits all situations [27, 31, 36, 82, 94].

We use the example shown below to illustrate the difference between feature- and data-based techniques, that explain bias based on the contribution of a subset of the training data.

Example 1.1. Consider a classifier that predicts whether individuals described by their attributes (such as gender, education level, marital status, working hours, etc.) earn more than 50K a year. A system developer uses the classifier to predict the income of individuals and, when analyzing the results, finds an unexpected negative result (earning less than 50K) for a female user:

Based on her education (education = Bachelors), marital status (marital = Never-married), (age = 40), and other features, this user should be classified as earning more than 50K. Upon closer examination, the developer realizes that the algorithm violated the commonly used fairness metric of statistical parity with respect to the protected class gender = Female. Statistical parity requires that the probability of being classified as the positive class (earning more than 50K in our example) is the same for persons from the protected class as it is for persons from the privileged class (we present fairness definitions in Section 2).

The developer uses an existing XAI package, such as LIME [75] or SHAP [60], to explain the algorithm’s output for the user and learns that the user’s gender and unmarried status were most responsible for the faulty prediction. We show examples of such explanations in Figure 1(a) and (b). These explanations, however, do not help the developer identify the source of the model’s bias. In contrast, Gopher identifies cohesive subsets of the training data that are most responsible for the bias and compactly describes the subsets using patterns that are conjunctions of predicates. Figure 1(c) shows the top-3 explanations Gopher produces. Pattern \( \phi_2 \) states that a sub-population that strongly impacts model bias is male spouses who work in the federal government for less than 40 hours per week. Indeed, in the Adult Income dataset (see Section 6), this subpopulation is the major source of gender bias, because it reports household income; hence, married males become highly correlated with high-income, due to the bias in data collection. In addition to producing such explanations, Gopher can also identify homogeneous updates for such sub-populations that can maximally improve model bias. Figure 1(d) shows one such update (with bold outline). For instance, model bias would significantly decline if we changed relationship status from “Spouse” to “Not married” and the Workclass from federal employee to self-employed in the subset of the training data corresponding to \( \phi_2 \). Such updates mean that our approach not only identifies which subsets of training data are responsible for the bias, but also which features of the data points in this subset are responsible.

Root causes and responsibility. Our first contribution is to formalize the concepts of root causes of bias and causal responsibility of a subset of training data on bias of an ML model. Responsibility is measured in terms of the difference between the bias of the original model and a new model trained on modified training data, obtained by either removing the subset or updating it. Our system, then, generates data-based explanations of the top-k subsets of training data that have the highest causal responsibility for model bias.

Pattern-based explanations. Such “raw explanations” based on subsets of data points may overwhelm rather than inform end-users. More compact and coherent descriptions are needed. Furthermore, sources of bias and discrimination in training data are typically not randomly distributed across different sub-populations; rather they manifest systematic errors in data collection, selection, feature engineering, and curation [29, 35, 42, 62, 63, 70, 94]. That is, more often than not, certain cohesive subsets of training data are responsible for bias. Therefore, our system generates explanations in terms of compact patterns expressed as first-order predicates that describe homogenous subsets of training data. By identifying commonalities within training data subsets, which are main contributors to model bias, such explanations unearth root causes of the bias. To ensure the
We introduce a novel quality metric which is expensive. Second, it is computationally prohibitive to explore all possible subsets of training data to identify the top-k explanations. To address the first challenge, we trace the model’s predictions through its learning algorithm back to the training data. To do so, we approximate the change in model parameters by using either influence functions [22] or by assuming that the updated model parameters are computed as a single gradient step from the original model. First-order (FO) approximations of influence functions have recently been proven useful in ML model debugging [20] by identifying the top-k training data instances responsible for model mis-predictions. However, they do not effectively capture the ground truth influence of coherent subsets of training data because of correlations between data instances. To better approximate the change in model parameters, we utilize second-order (SO) influence functions, which were shown to better correlate with ground truth subset influence [9]. While even SO influence functions can provide relatively poor estimates of the influence of large portions of the training data, the approximation error of SO influence functions is typically much better for coherent sets of data points as described by our patterns. To address the second challenge, we develop a lattice-based search algorithm based on ideas from frequent itemset mining [6] to discover frequent patterns. Our algorithm identifies coarse-grained subsets of training data that are influential and successively refines them to discover smaller influential subsets. We introduce a novel quality metric for explanations and rules to further prune the search space and generate the top-k explanations.

Update-based explanations. Gopher is unique in that it in addition to reaches explanations in terms of data removal we also consider explanations based on updating data points. This is motivated by the observation that not all features of a data point are erroneous and responsible for bias. We generate explanations based on updates (potentially on a subset of features) to reduce bias. Specifically, we search for a homogeneous perturbation to an explanation (data described by a pattern) that maximizes the reduction in model bias, where we update data instances in the subset by the same perturbation vector (i.e., all data points described by the pattern are uniformly perturbed). We formulate the problem of computing update-based explanations as a constrained optimization problem that searches for a homogeneous update to a subset of training data in order to maximize bias reduction, and we solve it using the projected gradient descent [15].

We make the following contributions:

- We propose a principled approach for generating top-k explanations based on frequent pattern mining and use pruning techniques to reduce the search space. (Section 4.2).
- We introduce update-based, actionable explanations. These explanations identify not only which parts of the training data are responsible for the bias but how to reduce or “repair” the bias. We formalize the problem of finding the best explanations as a constrained optimization problem that maximizes bias reduction. We propose a new algorithm based on projected gradient descent to determine the best homogeneous update for an explanation (Section 5).
- We conducted extensive experiments using real-world datasets to evaluate: (1) the end-to-end performance of Gopher, (2) the accuracy of influence estimation techniques, and (3) the quality and interpretability of explanations (Section 6). We show that Gopher generates explanations that are consistent with insights from existing literature.

2 BACKGROUND AND ASSUMPTIONS

We now present pertinent background information on classification and algorithmic fairness.

Classification. In this paper, we consider the problem of binary classification, the focus of most literature on algorithmic fairness. Consider a training dataset of $n$ examples $D = \{(x_i, y_i)\}_{i=1}^n$, with domain $\text{Dom}(X) \times \text{Dom}(Y)$, drawn from an unknown data distribution $D$, where $X$ denotes a set of discrete and continuous features and $Y = \{0, 1\}$ is a binary label to be predicted. Here, the goal is to find a classifier $f : \text{Dom}(X) \rightarrow \hat{Y}$ that associates each data point $x$ with a predicted label $\hat{y} \in \{0, 1\}$ that maximizes accuracy $E_{x,y \sim D}[\hat{y}(f(x) = y)]$, i.e., the expectation of the fraction of data points from the unknown data distribution $D$ that are correctly predicted by $f$. Note that since we have access to only a sample $D$ of $D$, model performance is determined over the training data as a substitute for calculating $E_{x,y \sim D}[\hat{y}(f(x) = y)]$. Typically, $\hat{y}(\cdot)$ is a member of a family of functions $f_\theta$ that are parameterized by $\theta$ with domain $\Theta$. A learning algorithm $\mathcal{A}$ uses $D$ and learns parameters $\theta^* \in \Theta$ that maximize the empirical accuracy $\frac{1}{n} \sum_{i=1}^n \mathbb{1}[f_\theta(x_i) = y_i]$. Learning algorithms typically use some loss function $L(z, \theta)$ and minimize the empirical loss $\frac{1}{n} \sum_{i=1}^n L(z_i, \theta)$. We denote the optimal parameters of a classifier $\theta^*$ as $\theta$ when it is clear from the context. Throughout this paper, we consider learning algorithms that use a loss function $L(\theta)$ that is twice-differentiable. This covers classification algorithms such as logistic regression [72], support vector machines (SVM) [72], and feed forward neural networks [88].

Algorithmic fairness. Consider a binary classifier $f_\theta$ with output $\hat{Y}$ and a protected attribute $S \in X$, such as gender or race. Without loss of generality, we interpret $\hat{Y} = 1$ as a favorable (positive) prediction and $\hat{Y} = 0$ as an unfavorable (negative) prediction. To simplify the exposition, we assume $\text{Dom}(S) = \{0, 1\}$, where $S = 1$ indicates a privileged and $S = 0$ indicates a protected group (e.g., males and non-males, respectively). Algorithmic fairness aims to ensure that the classifier $f$ makes fair predictions devoid of discrimination wrt. the protected attribute(s). Many fairness definitions have been proposed to quantify the bias of a binary classifier (see [91] for a recent survey). The best-known ones are based on associative relationships between the protected attribute and the classifier’s outcome. These
are unlike causal notions of fairness, which incorporate background knowledge about the underlying data-generative process and define fairness in terms of the causal influence of the protected attribute on the classifier’s outcome [49, 53, 69, 79]. We focus on three of the most widely used associational notions of fairness.

Statistical parity requires an algorithm to classify both the protected and privileged group with the same probability:

$$\Pr(\tilde{Y} = 1 | S = 1) = \Pr(\tilde{Y} = 1 | S = 0).$$

Equal opportunity requires both the protected and privileged group to have the same true positive rate:

$$\Pr(Y = 1 | Y = 1, S = 1) = \Pr(Y = 1 | Y = 1, S = 0).$$

Predictive parity requires that both protected and privileged groups have the same predicted positive value (PPV), i.e., correct positive predictions have to be independent of the protected attribute:

$$\Pr(Y = 1 | \tilde{Y} = 1, S = 1) = \Pr(Y = 1 | \tilde{Y} = 1, S = 0).$$

In practice, these fairness notions are often evaluated by estimating the probabilities on a test dataset $D_{\text{test}}$. Note that our techniques also work for other associational and causal notions of fairness.

3 PROBLEM STATEMENT

This section introduces concepts of root causes and data-based explanations of the bias of a classifier. For now, we focus on generating explanations based on removal of data points from training data. In Section 5, we extend our framework to support explanations obtained by updating data points, not removing them.

Consider training dataset $D$, a learning algorithm $\mathcal{A}$, a classifier $\theta_0$ trained using $\mathcal{A}$ on $D$, and a fairness metric (cf. Section 2) $\mathcal{F} : \theta, D_{\text{test}} \rightarrow \mathbb{R}$ which we refer to as bias. The bias quantifies the fairness violation of the classifier on a testing data $D_{\text{test}}$ that is unseen during the training process, such that if $\mathcal{F}(\theta, D_{\text{test}}) > 0$, the classifier is biased, and it is unbiased otherwise. For instance, for statistical parity we may define $\mathcal{F}$ as $\Pr(\tilde{Y} = 1 | S = 1) - \Pr(\tilde{Y} = 1 | S = 0)$, where $\Pr(\tilde{Y} = 1 | Y = s)$, for $s \in \{0, 1\}$, is estimated on $D_{\text{test}}$ using empirical probabilities.

Intervening on training data. We evaluate the effect of the subset of the training data $S \subseteq D$ on the bias of a classifier $\theta_0$ using an intervention that removes $S$ from $D$. We then assess whether the removal reduces the bias of a new classifier trained on the adjusted (“intervened”) training data. More specifically, let $D_S = D \setminus S$ be the intervened training data and $\theta_S$ be the new model trained on $D_S$ (using the same learning algorithm $\mathcal{A}$). The effect of $S$ on the bias of $\theta_0$ can be measured by comparing the bias of the original and updated classifiers, i.e., by comparing $\mathcal{F}(\theta, D_{\text{test}})$ and $\mathcal{F}(\theta_S, D_{\text{test}})$.

Definition 3.1 (Root cause of bias). A subset $S \subseteq D$ of the training data $D$ is a root cause of the bias of a classifier $\theta_0$ if:

$$0 \leq \mathcal{F}(\theta_S, D_{\text{test}}) < \mathcal{F}(\theta, D_{\text{test}}),$$

where $\theta_S$ is a classifier trained on $D_S = D \setminus S$.

Next, we introduce a metric to quantify the causal responsibility of a subset of training data for classifier bias.

Definition 3.2 (Causal responsibility). We measure the causal responsibility of a subset of training data $S \subseteq D$ for the bias $\mathcal{F}$ of a classifier $\theta_0$ w.r.t. a fairness metric as

$$\mathcal{R}_S(\mathcal{F}) = \frac{\mathcal{F}(\theta, D_{\text{test}}) - \mathcal{F}(\theta_S, D_{\text{test}})}{\mathcal{F}(\theta, D_{\text{test}})}.$$

Intuitively, causal responsibility captures the relative difference between the bias of the original and new models obtained by intervening on training data. As such, it quantifies the degree of contribution $S$ has on model bias. Note that $-\infty < \mathcal{R}_S(\mathcal{F}) < 1$. If $S$ is a root cause of bias, then $\mathcal{F}(\theta, D_{\text{test}}) - \mathcal{F}(\theta_S, D_{\text{test}}) > 0$; hence, $0 < \mathcal{R}_S(\mathcal{F}) < 1$, and the larger the $\mathcal{R}_S$, the greater the responsibility of $S$ for the bias. If $\mathcal{R}_S(\mathcal{F}) < 0$, then removing $S$ from training data either does not change the bias or further exacerbates it.

Causal responsibility forms the basis for defining data-based explanations. To formalize the notion of pattern-based explanations, we present the following definitions.

Definition 3.3 (Pattern). A pattern $\phi$ is a conjunction of predicates $\phi = \bigwedge_i \phi_i$ where each $\phi_i$ is of the form $[X \ op \ c]$, where $X \in X$ is a feature, $c$ is a constant and $op \in \{=, <, \leq, >, \geq\}$.

Patterns represent training data subsets. For example, the pattern $\phi = (\text{gender} = \text{Female}) \land (\text{age} < 45)$ describes data instances where gender is ‘Female’ and age is less than 45. We use $D_{\phi}$ to denote the set of all patterns defined over the training data $D$ and denote the set of data instances satisfying pattern $\phi$ by $D(\phi)$.

Definition 3.4 (Support of a pattern). The support of pattern $\phi$ is denoted by $\text{Sup}(\phi)$ and is defined as the fraction of data instances that satisfy $\phi$, i.e.,

$$\text{Sup}(\phi) = \frac{|D(\phi)|}{|D|}.$$

Definition 3.5 (Interestingness of a pattern). Given a biased classifier $\theta_0$ and a pattern $\phi$, we define the interestingness of $\phi$ to explain the bias of a classifier as:

$$U(\phi) = \frac{\mathcal{R}_S(\mathcal{F}(\phi))}{\text{Sup}(\phi)}.$$

The intuition behind our definition of interestingness is that if two patterns result in the same reduction in model bias, we are more interested in the pattern that requires fewer changes to the data (less support). Thus, interestingness measures the average bias reduction per data point covered by a pattern.

In addition to finding patterns with high interestingness scores, we also aim to return a diverse set of patterns that have little overlap in terms of the data points that they contain. Diversity helps us avoid returning patterns that are too similar to each other, i.e., that differ only in minor details. This is a desirable property because patterns with large overlap convey similar information and, thus, lead to redundancy. We capture the notion of diversity through a containment score as defined next.

Definition 3.6 (Containment score). Given two patterns $\phi$ and $\phi'$, the containment score $C$ is a measure of their intersection:

$$C(\phi, \phi') = \frac{|D(\phi) \cap D(\phi')|}{|D(\phi)|}.$$
Furthermore, for a pattern \( \phi \) and set of patterns \( \Phi \), we define
\[
C(\phi, \Phi) = \max_{\phi' \in \Phi} C(\phi, \phi').
\]

The containment score quantifies the overlap between the data represented by a pair of patterns. Similar patterns have a higher fraction of overlapping data instances (a containment score close to 1) and largely convey the same information differently. Dissimilar or diverse patterns, have a containment score closer to 0.

We are now ready to formalize the problem of generating the top-\( k \) diverse and interesting data-based explanations:

**Definition 3.7 (Top-\( k \) explanations).** Generating the top-\( k \) data-based explanations for a bias \( F \) of a classifier requires finding the top-\( k \) most responsible and diverse explanations. Formally, given a containment threshold \( c \), the goal is to compute a set \( \text{TOP-}k \subseteq \Phi_D \) of explanation candidates as defined below.

\[
\text{TOP-}1 = \arg\max_{\phi \in \Phi_D} U(\phi)
\]

\[
\text{TOP-}i + 1 = \arg\max_{\phi \in \Phi_D \setminus C(\phi, \text{TOP-}i) \leq c} U(\phi), \quad 1 < i \leq k.
\]

Intuitively, \( \text{TOP-}k \) is computed by iteratively including into the current result set the next candidate with the highest score, skipping any candidate patterns whose overlap with one of the explanations in the result we have computed so far exceeds the threshold \( c \). Note that \( \text{TOP-}k \) is not well-defined if multiple patterns have the same score. We impose an arbitrary order over \( \Phi_D \) to break ties.

Two major challenges complicate efforts to efficiently compute top-\( k \) explanations. First, computing the causal responsibility of a pattern is computationally intensive: we must train a new classifier on the intervened training data obtained by removing the subset of data points covered by the pattern. Second, it is computationally prohibitive to exhaustively explore the space of all candidate explanations, which is in the number of features.

## 4 COMPUTING TOP-k EXPLANATIONS

This section describes our methods for addressing the challenges of computing top-\( k \) explanations for the bias of an ML model. First, we describe how to efficiently approximate the causal responsibility of a training data subset, without having to retrain classifiers (Section 4.1). Then, we develop a lattice-based search algorithm which utilizes the approximation methods for estimating causal responsibility, to efficiently generate top-\( k \) explanations (Section 4.2).

### 4.1 Approximating Causal Responsibility

We now describe two methods for approximating the causal responsibility of a subset of the training data \( D \) on classifier bias without needing to train a new classifier. The first is a method based on influence functions (Section 4.1.1). The second is a simple, yet effective, approximation based on gradient descent (Section 4.1.2).

#### 4.1.1 Influence function approximation

Recall from Section 2 that the optimal model \( \theta^* \) is the element of the set of possible models \( \Theta \) minimizing the empirical risk, i.e.,

\[
\theta^* = \arg\min_{\theta \in \Theta} \mathcal{L}(\theta) = \arg\min_{\theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \mathcal{L}(z_i, \theta).
\]

Let \( \nabla_\theta \mathcal{L}(\theta) \) and \( \mathcal{H}_\theta = \nabla^2_\theta \mathcal{L}(\theta) = \frac{1}{n} \sum_{i=1}^{n} \nabla^2_\theta \mathcal{L}(z_i, \theta) \) be the gradient and the Hessian of the loss function, respectively. Under the assumption that the empirical risk \( \mathcal{L}(\theta) \) is twice-differentiable and strictly convex, it is guaranteed that \( \mathcal{H}_\theta \) exists and is positive-definite, which further implies \( \mathcal{H}_\theta^{-1} \) exists.\(^1\)

The influence of a data instance \( z \in D \) is measured in terms of the effect of its removal from \( D \) on model parameters \( \theta^* \). We first measure the effect of up-weighting \( z \) by some small \( \varepsilon \); as we will see later, removing \( z \) is equivalent to up-weighting it by \( \varepsilon = -\frac{1}{n} \), where \( n \) is the number of data instances in \( D \). Up-weighting \( z \) by \( \varepsilon \) leads to a new set of optimal model parameters:

\[
\theta^*_\varepsilon = \arg\min_{\theta \in \Theta} \mathcal{L}(\theta) + \varepsilon \mathcal{L}(z, \theta).
\]

We are interested in estimating how model parameters change due to an \( \varepsilon \) change in \( z \). Let \( \Delta_\varepsilon = \theta^*_\varepsilon - \theta^* \) denote the change in model parameters. Note that because \( \theta^* \) is independent of \( \varepsilon \), we can capture the change in model parameters in terms of \( \Delta_\varepsilon \):

\[
\frac{d\theta^*_\varepsilon}{d\varepsilon} = \frac{d\Delta_\varepsilon}{d\varepsilon}.
\]

**Influence of a single data point.** Since \( \theta^*_\varepsilon \) minimizes the updated training loss (Equation (2)), the gradient of the loss function at \( \theta^*_\varepsilon \) should be zero, i.e.,

\[
\nabla_\theta \mathcal{L}(\theta^*_\varepsilon) + \varepsilon \nabla_\theta \mathcal{L}(z, \theta^*_\varepsilon) = 0.
\]

Let us denote the gradient of the loss function as \( g(\varepsilon, \theta) \); hence, the LHS of Equation (4) reads \( g(\varepsilon, \theta^*_\varepsilon) \). The key idea behind influence functions is that up-weighting a data point \( z \) by a very small \( \varepsilon \) does not significantly change the optimal parameters, i.e., if \( \varepsilon \rightarrow 0 \), then \( \theta^*_\varepsilon \rightarrow \theta^* \). Therefore, using the first principles, \( g(\varepsilon, \theta^*_\varepsilon) \) can be effectively approximated by Taylor’s expansion of \( g(\varepsilon, \theta) \) at \( \theta^* \), the optimal parameters of the original model.\(^2\) By plugging this approximation into the LHS of Equation (4), we obtain:

\[
\nabla_\theta \mathcal{L}(\theta^*) + \varepsilon \nabla_\theta \mathcal{L}(z, \theta^*) \big|_{\varepsilon = 0} = 0.
\]

Solving for \( \Delta_\varepsilon \), we get

\[
\Delta_\varepsilon = -\nabla_\theta \mathcal{L}(\theta^*) \big|_{\varepsilon = 0} \nabla_\theta \mathcal{L}(z, \theta^*)^{-1}.
\]

From Equation (1), since \( \theta^* \) minimizes \( \mathcal{L}(\theta) \), we obtain \( \nabla_\theta \mathcal{L}(\theta^*) = 0 \). Ignoring higher orders of \( \varepsilon \),

\[
\Delta_\varepsilon = -\nabla_\theta \mathcal{L}(z, \theta^*) \nabla_\theta \mathcal{L}(\theta^*)^{-1} \varepsilon.
\]

From Equations (3) and (7), as \( \varepsilon \rightarrow 0 \), the influence of up-weighting a data instance \( z \in D \) by \( \varepsilon \) on the model parameters is computed as:

\[
I_\theta(z) = \left. \frac{d\theta^*_\varepsilon}{d\varepsilon} \right|_{\varepsilon = 0} = -\nabla_\theta \mathcal{L}(\theta^*)^{-1} \nabla_\theta \mathcal{L}(z, \theta^*) = -\mathcal{H}_\theta^{-1} \nabla_\theta \mathcal{L}(z, \theta^*) \quad \text{for } \varepsilon = -\frac{1}{n}.
\]

To remove data instance \( z \), we consider up-weighting it by \( \varepsilon = -\frac{1}{n} \). The change in model parameters due to removing \( z \) can, therefore, be linearly approximated by computing \( d\theta^*_\varepsilon \approx -\frac{1}{n} I_\theta(z) \).

**Influence of subsets.** Using first-order (FO) influence function approximation (Equation (8)), the effect of removing a subset of

\[^1\]See please [50] for a discussion on relaxing these assumptions on the loss function.

\[^2\]Recall that Taylor’s expansion of function \( u \) about the point \( t \) with increment \( h \) is given by \( u(t + h) = u(t) + u'(t)h + u''(t)h^2/2 \). Hence, in our case \( g(\varepsilon, \theta^*_\varepsilon) = g(\varepsilon, \theta^* + \Delta_\varepsilon) = [\nabla_\theta \mathcal{L}(\theta^*) + \varepsilon \nabla_\theta \mathcal{L}(z, \theta^*)] + [\nabla^2_\theta \mathcal{L}(\theta^*) + \varepsilon \nabla^2_\theta \mathcal{L}(z, \theta^*)] \Delta_\varepsilon \).
data instances $S \subseteq D$ on model parameters can be obtained by:

$$I_\theta^{(1)}(S) = \sum_{z \in S} I_\theta(z). \quad (9)$$

The approximation shown in Equation (9) is quite accurate when the updated model’s parameters are close to the original model’s parameters. That is because we estimate the effect of removing a set of data points by summing their individual effects; this essentially means that we assume that the effect of removing one data point on the model is independent of the effect of removing any other data point. Put differently, when approximating the influence of a data point, we assume that the removal of other data points will not affect the model. While in general this assumption does not hold, it leads to an acceptable approximation if a small fraction of training data points is removed, but accuracy will decline when a larger fraction of training data is removed. For such large model perturbations, using higher order terms can reduce approximation errors significantly. The issue can be alleviated by computing the effect of uniformly up-weighting data instances in a subset of training data by some small $\epsilon$, using the same idea as influence function but considering higher-order optimality criteria [9]. Specifically, in the derivation of influences, we do not ignore second-order terms of $\epsilon$ for an even more accurate estimation of the group influence of up-weighting a subset of training data instances on model parameters, and we obtain the influence as:

$$I_\theta^{(2)}(S) = \left(\frac{1}{|D| - |S|}\right) I_\theta^{(1)}(S) + \left(\frac{|S|}{|D| - |S|}\right) I_\theta(S), \quad (10)$$

where

$$I_\theta(S) = \left(1 - \sum_{z \in S} \frac{1}{|S|} \frac{\partial^2 \mathcal{L}(\theta, \mathcal{D})}{\partial \theta \partial \theta} \right) I_\theta^{(1)}(S) \quad (11)$$

We refer readers to [9] for detailed derivations of the above expression for the group influence of a subset. The second term ($I_\theta^{(2)}$) in the second-order (SO) approximation captures correlations among data instances in $S$ through a function of gradients and Hessians of the loss function at the optimal model parameters. Thus, when training data instances are correlated, the SO group influence function is more informative and captures the ground truth influence more accurately. We empirically compare the accuracy of first- and SO influence function bias approximations in Section 6.

### Causal responsibility

Influence functions can be adopted to approximate causal responsibility using the chain rule of differentiation, as follows. We can estimate the effect of up-weighting data instances in $S$ on a function $f(\theta)$ as:

$$I_f(S) = \frac{df(\theta^*_S)}{d\theta} \bigg|_{\theta=0} = \frac{df(\theta^*_S)}{d\theta} \bigg|_{\theta=0} = \nabla_\theta f(\theta^*)^T I_\theta(S), \quad (11)$$

where $I_\theta(S) = I_\theta^{(1)}(S)$ or $I_\theta^{(2)}(S)$ depending on whether the influence is computed using first- or second-order group influence function approximations, respectively.

Given a fairness metric $F$, $I_F(S)$ captures the effect of intervening on $S$ on the bias of the algorithm and is used as the numerator in Definition 3.2 to compute the responsibility of $S$.

#### 4.1.2 Gradient-based approximation

As shown in [50], FO influence function approximations can be used to update individual training data points to maximize test loss. However, they deviate from ground truth influence for a group of data points because they do not capture data correlations among points in a group [10]. SO influence functions capture correlations but have only been explored for subset removal [10].

For the task of generating update-based explanations, we introduce an alternative approach to approximate causal responsibility. We assume that the updated model parameters are obtained through one step of gradient descent which we use in Section 5 and empirically find explanations that are more accurate than even SO influence approximations (details in Section 6.3).

Note that as in Equation (1), the updated model parameters when a subset of data points is removed is given by:

$$\theta_S = \argmin_{\theta \in \Theta} \left(\mathcal{L}(\mathcal{D}, \theta) - \frac{1}{|S|} \sum_{z \in S} \mathcal{L}(z, \theta)\right). \quad (12)$$

The preceding equation can be solved using gradient descent by taking repeated steps in the direction of steepest descent of the loss function (for data points in $D\setminus S$). However, inspired by the existing literature in adversarial ML attacks [44], which assumes that model parameters do not change significantly when a small subset of data points is poisoned, we also assume minimal change in model parameters. Thus, the change in model parameters is approximated in terms of a single step of gradient descent, as follows:

$$\theta_S = \theta - \eta \left(\frac{\nabla_\theta \mathcal{L}(\mathcal{D}, \theta^*)}{n} - \frac{1}{n} \sum_{i=1}^{n} \nabla_\theta \mathcal{L}(z_i, \theta)\right), \quad (13)$$

where $\eta$ is the learning rate for the gradient step.

The effect of removing $S$ on bias is then computed as the difference in test bias before and after $S$ is removed: $I(S) = F(\theta_S, D_{test}) - F(\theta, D_{test})$. While our single-step gradient descent approach can be used to estimate subset influence, this may not be a good idea for learning algorithms use more efficient techniques than gradient descent. We, therefore, use this approach mainly for generating update-based explanations in Section 5.

#### 4.2 Lattice-Based Search

Having discussed several techniques for efficiently approximating influence, we now introduce our algorithm for finding top-k explanations according to Definition 3.7. As discussed in Section 1, the naive approach for computing the top-k explanations by evaluating all possible patterns is exponential in the number of attributes, and, thus, is infeasible even when we estimate the influence of patterns. Toward this objective, we propose ComputeCandidates, an algorithm that takes as input training data and generates a list of candidate patterns. ComputeCandidates generates explanations starting with patterns with a single predicate each. The algorithm then iteratively generates patterns with $i$ predicates by merging two patterns with $i-1$ predicates that differ only in one predicate. For instance, we can merge patterns $\phi_1 = ((\text{hours} < 40) \land (\text{married} = \text{Married}))$ and $\phi_2 = ((\text{married} = \text{Married}) \land (\text{gender} = \text{Male}))$ into a pattern $\phi = ((\text{hours} < 40) \land (\text{gender} = \text{Male}) \land (\text{married} = \text{Married}))$. The idea is similar to frequent itemset mining [6, 39] in
Figure 2: An overview of the lattice structure and search.

data mining where itemsets with n items are generated by successively merging candidate itemsets of smaller size. Figure 2 shows an example of part of the lattice search space for patterns.

By itself, building patterns bottom-up does not reduce the size of the search space. For that we propose two heuristics. The first heuristic is that we assume as input a support threshold τ and only consider patterns whose support is above τ. The rationale for this heuristic is that patterns with low support describe only a small portion of the training data and, thus, are unlikely to identify systematic issues. In Section 6, with τ as small as 1%, we did not observe patterns with low support to reduce bias by much. Even for τ = 1%, much larger patterns (with ~ 27% data) are dominating the top-k. For a candidate pattern φ generated by merging two patterns φ₁ and φ₂, we have sup(φ) ≤ sup(φ₁) and sup(φ) ≤ sup(φ₂). Thus, pruning φ₁, for example, prunes the entire sublattice whose root is φ₁. In Figure 2, pruning the pattern φ₁ = {(gender = Male) ∧ (marital = Married) ∧ (hours < 40)} also prunes other patterns including φ = {(hours < 40) ∧ (marital = Married) ∧ (gender = Male)} formed by merging φ₁ with another pattern.

Our second heuristic is to prune patterns during merging. Consider a pattern φ generated by merging patterns φ₁ and φ₂. We only consider φ if its responsibility exceeds the responsibility of the two patterns it is generated from: R(φ) > R(φ₁) and R(φ) > R(φ₂). Note that if this is the case, then U(φ) > max(U(φ₁), U(φ₂)), because U(φ) = \frac{Sup(φ)}{Sup(φ₁) + Sup(φ₂)} - and as mentioned above sup(φ) ≤ min(Sup(φ₁), Sup(φ₂)). In Figure 2, the pattern φ = {(gender = Male) ∧ (marital = Married)} is generated from patterns φ₁ = {gender = Male} and φ₂ = {marital = Married} if U(φ) > U(φ₁) and U(φ) > U(φ₂). The rationale for this heuristic is that patterns with more predicates are harder to interpret. Thus, an increase in the number of predicates should be justified by an increased impact on the bias of the model.

We now discuss our algorithm in more detail (pseudo code is shown in Algorithm 1). We start by generating all possible single predicate patterns (Line 1) by iterating through all features X ∈ X and each possible value val for the feature and generate three types of comparisons: X < val, X = val, and X > val. Note that for features with a large number of possible values, we can apply binning techniques to reduce the number of candidate patterns. Additionally, this has the advantage of avoiding the generation of almost identical explanations (e.g., hours < 40 and hours < 42).

We then iteratively create patterns (Line 7) of size i by merging two patterns of size i – 1 that only differ in one predicate to generate a candidate patterns of size i. Iteration finishes once no more candidate have been produced in the previous iteration. For each generated candidate φ we test if its support is larger than the threshold τ and that its responsibility estimated using influence is larger than the influence of both patterns it was derived from. Pattern φ is only included in Φ_i, the set of candidate patterns of size i, if both conditions are fulfilled. Finally, we return all candidate explanations (the union of all sets Φ_i generated so far).

To determine the support of a pattern, we have to evaluate a query over the training data. If the patterns φ₁ and φ₂ we are attempting to merge are conflicting, then φ₁ ∪ φ₂ has zero support, and we can avoid running this query. Two patterns φ₁ and φ₂ are conflicting if they both contain a predicate on an attribute X and the conjunction of these predicates is unsatisfiable. For example, in Figure 2, patterns φ₁ = {marital = Married} and φ₂ = {marital = Unmarried} are conflicting.

Diversity of explanations. We use Algorithm 2 to compute a diverse set of top-k explanations as defined in Definition 3.7. We first use ComputeCandidates to generate a set of candidate explanations. This set is then sorted based on U and we are iterating over this set in sort order and including patterns into the result, skipping patterns whose overlap with any of the previously added patterns exceeds the threshold c. Note that we are interested in patterns that, when modified or removed, reduce bias maximally without significantly affecting model accuracy. Instead of directly optimizing for minimal accuracy loss, we penalize patterns with high support (low interestingness score). As an extreme example, optimizing for bias reduction and accuracy loss simultaneously is an interesting direction for future work.
Algorithm 2: Generate Top-k Explanations

Input: Training dataset D, support threshold r, containment threshold c, desired number of top explanations k

Output: Top-k explanations

1. \( \phi_{\text{cand}} \leftarrow \text{ComputeCandidates}(D, r) \)
2. \( \Phi_{\text{top-k}} \leftarrow \emptyset \) = Set of top-k explanations
3. for \( \phi \in \text{sort-by-score}(\phi_{\text{cand}}) \) do
4. \( \text{if } \exists \phi_j \in \Phi_{\text{top-k}} \colon C((\phi, \phi_j)) > c \) then
5. \( \Phi_{\text{top-k}} \leftarrow \Phi_{\text{top-k}} \cup \phi_j \)
6. if \( |\Phi_{\text{top-k}}| = k \) then
7. \( \text{return } \Phi_{\text{top-k}} \)

that Gopher’s explanations describe training data subsets that may have potential data quality errors or point out to historical biases reflected in training data or bias that was introduced during data collection. These errors are otherwise not detectable using standard data cleaning algorithms such as outlier detection. Gopher can be complemented with existing error detection mechanisms or external sources of information about data provenance to expose the errors in the identified subsets.

5 UPDATE-BASED EXPLANATIONS

In this section, we formalize the problem of generating explanations based on updating training data. Given an influential subset of training data obtained through methods proposed in Section 4, our goal is to find a homogenous update that can lead to maximum bias reduction. We first discuss an approach for approximating the influence of updating or perturbing a subset of training data on bias reduction. We first discuss an approach for approximating causal responsibility of training data subsets.

Consider a classifier with optimal parameters \( \theta^* \) trained on a training dataset D. Let \( S = \{z_i = (x_i, y_i)\}_{i=1}^m \) be a subset of D consisting of \( m \) data points. Also, let \( S^p = \{z_i^p = (x_i^p, y_i^p)\}_{i=1}^m \) be the result of updating each instance in S using a perturbation vector \( \delta = \{\delta_1, \ldots, \delta_N\} \) to uniformly update the attribute values of data point \( z \in S \) as \( z^p = z + \delta \), where the \( j \)-th attribute of \( z \) is updated by \( \delta_j \). For example, if the working hours for an individual is 32, then an update of 8 will change it to 40. Uniform perturbation means the working hours for all individuals in a subset is updated by 8 hours. Our goal is to compute the optimal parameters \( \theta_p \) of a new model trained on \( D^p = (D \setminus S) \cup S^p \). We approximate \( \theta_p \) using a single step of gradient descent (cf. 4.1.2), as follows:

\[
\theta_p = \theta^* - \eta \left( \frac{1}{n} \sum_{i=1}^{m} \nabla_g L(z_i, \theta^*) + \frac{1}{m} \sum_{i=1}^{m} \nabla_g L(x_i^p, \theta^*) \right),
\]

where \( \eta \) is the learning rate for the gradient step.

Therefore, the influence of an update to \( S \) on the bias of the model can be measured using the chain rule, as follows:

\[
\Delta F(\theta^*, \theta_p, D_{\text{test}}) = \nabla_{\theta} F(\theta^*, D_{\text{test}})^\top \left( \theta_p - \theta^* \right).
\]

Now, given a subset of training data, \( S \subseteq D \), our goal is to obtain an update on \( S \) that leads to the greatest bias reduction. Formally, we aim to optimize the problem

\[
\delta^* = \arg \max_{\delta} \Delta F(\theta^*, \theta_p, D_{\text{test}}).
\]

Using Equations (14) and (15), Equation (16) becomes:

\[
\delta^* = \arg \min_{\delta} \nabla_{\theta} F(\theta^*, D_{\text{test}})^\top \nabla_{\theta} L(S^p, \theta^*).
\]

We solve Equation (17) using the gradient descent algorithm to obtain the perturbation vector:

\[
\delta_{k+1} = \delta_k - \eta \nabla_{\theta} F(\theta_{\text{test}}, \theta^*)^\top \nabla_{\theta} L(S^p + \delta_k, \theta^*),
\]

where \( \eta \) is the learning rate of the gradient ascent step.

The updated data point is obtained as \( z^p = z + \delta_{k+1} \). Note that the preceding formulation can result in perturbations that lie outside the input domain. We add domain constraints to prevent this from happening. In particular, the updates should change an attribute of a data point from one value to another in the input domain, i.e., \( z, z^p \in \text{Dom}(X) \times \text{Dom}(Y) \). We solve this constrained optimization problem in Equation (18) using projected gradient descent [20], which works as follows: if the updated data point violates the domain constraint, i.e., \( z^p \notin \text{Dom}(X) \times \text{Dom}(Y) \), then project it back to the input domain \( \text{Dom}(X) \times \text{Dom}(Y) \) as:

\[
z_{up} = \arg \min_{z' \in \text{Dom}(X) \times \text{Dom}(Y)} \| z' - z^p \|
\]

The projection ensures that \( z_{up} \) is the data point in the input domain \( \text{Dom}(X) \times \text{Dom}(Y) \) that is closest to the actual perturbation \( z^p \).

6 EXPERIMENTS

In our experimental evaluation of Gopher, we aim to address the following questions: Q1: How effective are the proposed techniques for approximating causal responsibility of training data subsets? Q2: What is the end-to-end performance of Gopher in generating interpretable and diverse data-based explanations? Q3: What is the quality of the update-based explanations? Q4: How effective is our approach at detecting data errors responsible for ML model bias?

6.1 Datasets

We use standard datasets from the fairness literature. The data and code for the experiments can be found at the project page.

German Credit Data (German) [26]. The personal, financial and demographic information (20 attributes) of 1,000 bank account holders. The prediction task classifies individuals as good/bad credit risks.

Adult Income Data (Adult) [26]. Demographic information, level of education, occupation, working hours, etc., of 48,000 individuals (14 attributes). The task predicts whether the annual income of an individual exceeds $50K.

Stop, Question, and Frisk Data (SQF) [1]. Demographic and stop-related information for 72,548 individuals stopped and questioned (and possibly frisked) by the NYC Police Department (NYPD). The classification task is to predict if a stopped individual will be frisked.

6.2 Setup

We considered three ML algorithms: logistic regression, support vector machines, and a feed-forward neural network with 1 layer and 10 nodes (we provide details about the hyper-parameters for the algorithms in the shared code). We used PyTorch [71] or sklearn [72] implementation of these algorithms. In accordance with existing literature on evaluation of fairness of ML algorithms on these datasets [19], the sensitive attribute attributes are: gender (Adult),

3https://drive.google.com/drive/folders/1_jQhBg3Y1us48HNYbE_QfZc4WngFjBE?usp=sharing
We split each dataset into training and test data, trained an ML model (one pattern are color-coded). We use the logistic regression model, in both kinds of explanations, predicates that occur in more than one pattern are color-coded. We identify the 5 modes up to a depth of 3 having the maximum combined influence across data points and report their paths to the root node. For example, to generate top-5 explanations having up to 3 predicates, we identify the 5 modes up to a depth of 3 having the maximum FO influence and report the conjunction of predicates that form the path from the root for each node.

### 6.3 Causal Responsibility Approximations

In this experiment, we evaluated the quality of the proposed approaches for approximating influence, and thereby causal responsibility. The ground truth influence of subsets is computed by retraining the model after removing the subset from the training data. In this set of experiments, we evaluate the proposed approximation methods. To estimate subset influence using FO approximations, we summed the individual influences, whereas the SO subset influence is computed using Equation (10). We also compared the approximations to the one-step gradient descent approach described in Section 4.1.2, that we use for update explanations.

**Effectiveness.** In Figures 3a to 3c, we report the absolute deviation of the influence estimated by the different methods from the ground truth (y-axis) for the fairness definitions (x-axis) defined in Section 2. We trained different classifiers on the German dataset. We observed that, as expected, FO influence approximations and one-step gradient descent deviated more from ground truth influence than SO influence. First-order influence approximation exhibits larger errors because it does not account for possible correlations among data points in the subset [10]; one-step gradient descent influence approximation is not accurate because it uses a single step of gradient descent instead of iterating until the model parameters converge. This behavior is especially evident when the size of the group is large, which usually corresponds to large influence (bins on either side for each method). The key takeaway from this experiment is that second-order influence functions closely approximate ground truth influence especially when model parameters do not change substantially (the middle bins).

**Efficiency.** In Figure 4, we report the average time taken by each method when subsets of varying sizes are removed from the training data. The brute force approach – retraining the model after removing the subset – is typically more than two orders of magnitude slower than even the most expensive method, especially when the fraction of removed training data points is small (e.g., influence functions are up to 4 orders of magnitude faster when 0%-10% of the training dataset is deleted). We also observe that the one-step gradient descent approach, with the exception for SO influence functions on neural networks, while effective in estimating ground truth influence (as shown in Figure 3a, Figure 3c), is significantly slower than influence functions for estimating subset influence. Note that the time cost for retraining is close to that of one-step
gradient descent because we used the initial model parameters to speed up convergence during retraining. In practice, we have to adjust the learning rate for one-step gradient descent over multiple iterations, which makes it even more expensive. We conclude that retraining the model and using the one-step gradient descent approach are not feasible for generating the top-k explanations. The one-step gradient descent, however, is useful for generating update-based explanations as described in Section 4.1.2: influence functions are not applicable to the optimization problem we have to solve for updates, and retraining is also not an option.

6.4 End-to-end Performance

Next, we evaluate the performance of Gopher for generating the top-k explanations for ML algorithms trained on different datasets.

**German.** This dataset is biased toward older individuals and considers them less likely to be characterized as high credit risks. In Table 1, we report the top-3 explanations up to 4 predicates generated by Gopher (with their support and ground truth influence) sorted by their interestingness score. We observe that one subset of 5% of data points explains more than half of the model bias, whereas another subset of ~ 6% of data points reduces bias by ~ 36%. These explanations highlight fractions of training data that may have potential errors and hence, need attention. On inspection, we found that these explanations correspond to training data points where older individuals are primarily labeled as low credit risks. By removing these individuals, the probability of an individual being classified as a high/low credit risk is uniformly distributed across the sensitive attribute age. As a result, the model’s dependency on age is reduced, thus reducing overall model bias. Note that the top-2 explanations consist of predicates with the sensitive attribute for this dataset, signifying its importance in bias reduction. In comparison, we made the following observations about the explanations generated by FO-tree: Entirely different regressor trees, and hence different explanations, were generated depending upon whether sklearn or PyTorch was used to fit the model. While the sensitive attribute (age) formed the root node for the FO-tree generated on the PyTorch model, a non-intuitive attribute (installment rate) was the root in the FO-tree for the sklearn model. The top-2 explanations from FO-tree over the PyTorch model were consistent with our explanations whereas those generated from FO-tree over the sklearn model were less compact (consisting of 4 predicates each). Their (support, bias reduction) were (6.13%, 32.3%), (5.63%, 33.1%) and (12.9%, 8%), respectively.

**Adult.** This dataset has been at the center of several studies that analyze the impact of gender [78, 87] and has been shown to be inconsistent: income attributes for married individuals report household income. The dataset has more married males, indicating a favorable bias toward males. As seen in Table 2, the sensitive attribute gender plays an important role in all of the explanations. In this set of
In these experiments on neural networks, we observed that second-order influence functions did not estimate the model parameters accurately and greatly underestimated the ground truth influence. This observation was consistent with the analysis provided in [10] for neural networks where the influence of a group of data points has low correlation with ground truth influence, and second-order influences underestimate ground truth influence. Our approach hinges on the applicability of influence functions to correctly estimating the model parameters around the optimal parameters— an assumption that might not hold for neural networks. However, Gopher still identified patterns that reduce model bias to some extent. In comparison, the top-3 patterns returned by FO-tree had higher support and lower ground truth influence: (10.0%, 9.8%), (13.2%, 10.8%) and (5.9%, 11%), respectively. Note that even though the dataset has single predicates (.marital → Married, ~ 47% data) that remove bias almost completely, these predicates do not rank in the top-k explanations because of their low interestingness scores.

**SQF.** This dataset highlighted that the practices of NYPD in stopping, questioning and frisking blacks (and latinos) more often compared to whites were unconstitutional and violating Fourth Amendment rights [2]. In Table 3, our top-3 explanations identify patterns consisting of the protected group that were frisked and the privileged group that were not frisked. Because of these data points, the model learns that data points belonging to the privileged (protected) group are less (more) likely to be frisked, and therefore, cause bias in the model. By removing these data points, the privileged group becomes less correlated with the ‘no frisk’ outcome, thus reducing model bias. The topmost explanation generated by the FO-tree [(location=Outside) ∧ (race=White) ∧ (build<>>Thin) ∧ (does not fit a relevant description)] had similar support (13.2%) and bias reduction (27%) as our topmost explanation. The other two patterns had lower bias reduction (one of them had 0.15% reduction in bias) and greater support, and hence were less interesting.

### 6.5 Update-based Explanations

In these experiments, we generated updates for the top-k explanations. For each explanation, we provide the perturbation that would result in the maximum bias reduction.

**German.** As seen in Table 4, updates typically involve perturbing the protected attribute (age). For example, the first explanation suggests that by updating data points satisfying the pattern such that after the update they are in the protected instead of the privileged group, and by changing the gender to increase the chance of a positive outcome, bias is reduced by ~42%. In this case, we found an update for the pattern that would reduce model bias but not by as much as it would reduce by deleting those data points. Similarly, in explanation 2, older individuals that have a good credit history are considered low credit risks. By changing their age group to the protected group and credit history to a worse level, we now associate younger individuals with good credit risk and reduce model bias (albeit by an amount smaller than if the group was removed). The key takeaway here is that we can reduce model bias by updating the training data points referring to these patterns instead of removing them altogether.

**Adult.** We report the update-based explanations for this dataset in Table 5. As mentioned before, this subset is biased toward married individuals and males. In explanation 1, individuals that were not married had lower income. By changing their marital status to married, we were able to reduce model bias by at least as much as would have been achieved were those patterns deleted. Note, however, that in explanation 2, even by changing the gender and marital status to the preferred attribute values, we could not find an appropriate update that would reduce the bias. After marital status, we found that education accounts for most of the bias [78]―individuals with a higher level of education are associated with higher incomes. In explanation 2, by changing the education level
Table 5: Update-based explanations for the top-3 explanations for Adult (τ = 5%). Updates to the original explanation (top) are shown in bold outline (bottom). Change in bias reduction Δbias is represented by ▼ (decrease) or ▲ (increase). Average time taken to update each point = 0.24s.

| Pattern | Support | Δbias |
|---------|---------|-------|
| Marital = Divorced/Separated/Widowed ∧ (Age ≥ 45) | 9.1% | 13.3% |
| Marital = Married-spouse-civ/AF/absent ∧ (Age ≥ 45) | 13.8% ▲ | |
| Gender = Female ∧ Marital = Never married ∧ (Relationship / Wife, Husband) | 14.3% | 19.1% ▲ |
| Gender = Male ∧ Marital = Married-spouse-civ/AF/absent ∧ (Relationship / Wife, Husband) | 0.6% ▼ | |
| Gender = Female ∧ (Education = Bachelors) ∧ (Relationship / Wife, Husband) | 7.6% | 9.1% ▲ |
| Gender = Female ∧ (Education = Assoc-std) ∧ (Relationship / Wife, Husband) | 9.5% ▲ | |

Table 6: Update-based explanations for the top-3 explanations for SQF (τ = 5%). Updates to the original explanation (top) are shown in bold outline (bottom). Change in bias reduction Δbias is represented by ▼ (decrease) or ▲ (increase). Average time taken to update each point = 0.5s.

| Pattern | Support | Δbias |
|---------|---------|-------|
| Race = Black ∧ (Fits a relevant description=No) ∧ Location=Outside ∧ Age< 25 | 16.9% | 25.6% |
| Race = Black ∧ (Fits a relevant description=No) ∧ Location=Outside ∧ Age∈[25, 45) | 14.3% ▼ | |
| Race = Black ∧ (Fits a relevant description=No) ∧ Location=Outside ∧ Age∈[25, 45) | 13.0% | 13.7% |
| Race = White ∧ (Fits a relevant description=Yes) ∧ Location=Inside ∧ Age∈[25, 45) | 8.6% ▼ | |
| Race = White ∧ Engaging in a violent crime=No ∧ (Casing a victim=Yes) ∧ (Proximity to scene of offense=No) | 7.0% | 8.2% |
| Race = White ∧ Engaging in a violent crime=No ∧ (Casing a victim=No) ∧ (Proximity to scene of offense=Yes) | 13.7% ▲ | |

6.6 Scalability Analysis

To evaluate the scalability of influence computations, we report in Figure 5 the effect of dataset size on the time taken to compute influence of a subset by the approaches described in Section 4.1. We replicated the German dataset to increase its size by a factor of 50 to 1,600, yielding up to 1.6M training data points. Since we wanted to evaluate how dataset sizes affect influence computation runtimes, we fixed the size of the subset for which we compute the subset influence to 5% (we chose this threshold because it reflects our problem setting where we are interested in small fractions of training data that need attention). We observed that both first-order and second-order influence computations scale well in the dataset size, and achieve speed-ups of several orders of magnitude over retraining the model or using one-step gradient descent. Note that Gopher has an upfront cost of pre-computing the gradients of the loss function and the Hessian. However, once these computations are done, the time taken to compute subset influence is negligible (as is also seen in Figure 5). In contrast, model retraining to compute subset influence can be quite expensive. For example, using the feed-forward network on Adult, we observed that the pre-computations took ~1,800s and the top-3 explanations were generated in 56s for a total time cost of ~1,856s. In comparison, retraining the model after removing one subset took > 10s. We see benefit in using Gopher when a relatively large number of candidates are being considered to generate top-k explanations (which is the case with the datasets used in this paper).
We found that the outlier detection mechanism supported by scikit-learn is an interesting direction for future work. To test the viability of detecting data errors, we applied our implementation as a detection mechanism for errors injected by adversarial attacks.

### Data errors injected by adversarial attacks
In this experiment, we tested whether Gopher’s explanation generation using the lattice structure has good scalability (runtimes of < 25 min) for explanations with fewer predicates. In comparison, our filtering mechanism that accounts for diversity of explanations takes negligible amount of time (although considering more candidates does take longer to generate diverse explanations).

#### Table 7: Scalability in the number of candidate patterns.

| Level | execution (s) | filtering (ms) | #candidates |
|-------|---------------|----------------|-------------|
| 1     | 0.03          | 36             | 29          |
| 2     | 0.24          | 45             | 371         |
| 3     | 1.59          | 70             | 2,770       |
| 4     | 17.5          | 75             | 13,625      |
| 5     | 26.9          | 76             | 36,704      |
| 6     | 147.2         | 90             | 62,955      |

In Table 7, we report the time taken to generate the top-5 explanations on the German dataset when we allow more predicates (indicated by the level in the lattice structure) in an explanation, and hence consider greater number of candidate explanations. We observe that Gopher’s explanation generation using the lattice structure has good scalability (runtimes of < 25 min) for explanations with fewer predicates. In comparison, our filtering mechanism that accounts for diversity of explanations takes negligible amount of time (although considering more candidates does take longer to generate diverse explanations).

### 6.7 Detecting data errors
To test the viability of detecting data errors, we applied our implementation as a detection mechanism for errors injected by adversarial attacks.

#### Data errors injected by adversarial attacks
In this experiment, we tested whether Gopher can detect errors introduced by data poisoning attacks [52, 64]. The objective was to develop techniques to detect attacks that have superior performance (measured in terms of accuracy and fairness) on training data and are targeted at exacerbating model performance on test data. The state of the art in safeguarding against data poisoning attacks is to detect anomalies that do not conform to the rest of the data. However, anomaly detection fails in the presence of sophisticated attacks that are targeted at deteriorating model accuracy and/or fairness [36, 43, 65, 83].

We performed experiments where we injected poisoned data points into the training data using non-random anchoring attacks [64]. We found that the outlier detection mechanism supported by scikit-learn [72], LocalOutlierFactor, was not able to detect any of the poisoned data points, because they follow a similar distribution as the original training data points. In comparison, when the data was clustered (using k-means or Gaussian mixture models clustering) and clusters were ranked in decreasing order of estimated SHAP influence, we observed that the top-2 clusters contained ~ 70% of poisoned points. While these results are promising, a detailed study demonstrating the effectiveness of second-order influence functions in detecting adversarial attacks is out of the scope of this work and is an interesting direction for future work.

### 7 RELATED WORK
Our work relates to the following lines of research.

#### Feature-based explanations
XAI research focuses on explaining ML models in terms of patterns and dependencies between input features and their outcomes. Its methods are based on feature attribution, such as the SHAP value, to quantify the responsibility of input features for ML model predictions [5, 24, 30, 57, 59, 60, 66, 86]. Methods based on surrogate explainability approximate ML models using a simple, interpretable model (such as linear regression) [60, 75, 76]. Contrastive and causal XAI methods explain ML model predictions in terms of minimal interventions or perturbations on input features that change the prediction [11, 32, 45, 55, 61, 68, 89, 90, 93].

Logic-based methods use tools from logic-based diagnosis that operate on logical representations of ML algorithms [23, 40, 81] to compute minimal sets of features that are sufficient and necessary for ML model predictions. These approaches fall short in generating diagnostic explanations that help users trace an ML algorithm’s unexpected or discriminatory behavior back to its training data.

#### Explanations based on training data
These XAI methods attribute ML model prediction to specific parts of its training data [14]. All current approaches rank individual training data points based on their influence on ML model predictions [10, 33, 34, 51, 83] using influence functions [22]. Influence functions are a classic technique from robust statistics that measure how optimal model parameters depend on training data instances. Based on first-order influence functions, [95] introduced an approach for identifying training data points that are responsible for user constraints specified by an SQL query. There are other methods than influence functions, however, e.g. [98] develops an approach for identifying and ranking training data points based on their influence on predictions of neural networks, and [96] develops an approach for incremental computation of the influence of removing subset of training data points. Furthermore, other recent work argues for the use of data Shapley values to quantify the contribution of individual data instances [33, 34, 54]; these approaches are computationally expensive because each data instance requires the model to be retrained. Unlike prior methods, our method generates: (1) explanations for fairness of an ML model, (2) interpretable explanations based on first-order predicates that pinpoint a subset of training data responsible for model bias, and (3) update-based explanations that reveal data errors in certain attributes of a training data subset. It is worth mentioning here that reducing the bias in the training data can be done not only by removing subsets of training data but also by adding certain appropriate training samples. Searching for in-distribution data points that explain away bias upon insertion into training data is a challenging problem that we plan to explore in the future. Please note, however, that our update-based explanations can act as a combination of removing existing data points and adding new ones.

#### Debugging ML models
Much work examines the debugging of ML models for fairness and bias. fair-DAGs [97] and mlinspect [37] regard class imbalance as a cause of discrimination and track the distribution of sensitive attributes along ML pipelines to address bias. These approaches cannot highlight problematic parts of training data, i.e., those responsible for biased outputs. MLDebugger [58] applies a method that automatically identifies one or more minimal causes of unsatisfactory performance in ML pipelines using provenance of the previous runs. [7, 8] develop methods to identify regions of attribute space not adequately covered by training data. [99] develops an approach for debugging training data by making the smallest set of changes to training data labels so the ML model trained on the updated data can correctly predict labels of a trusted set of items in test data; since the focus of their approach is on updating training labels, it is not applicable to our setting. Another related line of work concerns finding large data slices in which
the model performs poorly [21, 73, 77]; the slices are discovered based on their association with model error and do not capture the causal effect of interventions. These techniques are also not directly applicable in the context of fairness, where, unlike model error, bias due to a subset is not additive. Moreover, none of the preceding interventions update data instances.

**Adversarial ML.** Our work shares similarities with adversarial machine learning literature, which explores ways in which ML can be compromised by adversarial attacks. The most relevant classes of adversarial attacks are based on data poisoning [18, 84], where polluting training data compromises model fairness or predictions [43, 65, 83]. Data poisoning attacks on fairness aim to inject a minimum set of synthetic poisonous data points in training data that compromise the fairness of a model trained on the contaminated data. This is in reverse to the goal of our research. Note that many data poisoning attacks methods rely on influence functions.

**Machine unlearning.** Our research also relates to the nascent field of machine unlearning [13, 38, 41, 80], which addresses the “right to be forgotten” provisioned in recent legislation, such as the General Data Protection Regulation (GDPR) in the European Union [92] and the California Consumer Privacy Act in the United States [25]. Current methods typically designed for particular classes of ML models, e.g., [80], support efficient unlearning requests for decision trees. However, the techniques in this sub-field could be integrated into our framework for efficient computation of causal responsibility. We defer this investigation to future study.

**Bias detection and mitigation.** Prior work explores ways to detect and mitigate bias in ML models. These approaches can be categorized into pre-, post- and in-processing (see [17] for recent surveys). Our goal is to pre-process data to reduce bias which is different from post-processing that deals with model output to handle bias, and in-processing that is aimed at building fair ML models. The idea is to remove bias and discrimination signals from training data by repairing or pre-processing it (see, e.g., [16, 28, 79]). Pre-processing methods in fairness are independent of the downstream ML model and usually are not interpretable; hence, they are insufficient in generating explanations that reveal the potential source of bias. While bias mitigation is not a direct focus of our research, the approach we propose could be useful for developing bias mitigation algorithms that are interpretable and take into account the downstream ML model; hence, they would incur minimal information loss and generalize better. We defer this research to future work.

8 CONCLUSIONS

This work presented a new approach for debugging bias in machine learning models by identifying coherent subsets of training data that are responsible for the bias. We introduce Gopher, a principled framework for reasoning about the responsibility of such subsets and develop an efficient algorithm that produces explanations, which compactly describe responsible sets of training data points through patterns. We demonstrate experimentally that Gopher is efficient and produces explanations that are interpretable and correctly identify sources of bias for datasets where ground truth biases are well understood. In future work, we plan to expand our approach beyond supervised ML algorithms with differentiable loss functions to support a wider range of ML algorithms such as tree-based ML models and clustering algorithms. Moreover, we plan to leverage database techniques for incremental maintenance, for efficient computation of causal responsibility, as opposed to approximating it. Another interesting future direction is to integrate Gopher with database provenance to formalize the notion of provenance of ML model decisions that trace ML model outcomes all the way back to decisions made in the ML pipeline that might explain the bias and unexpected behavior of the model.

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