

# Mitigating Bias in Radiology Machine Learning: 1. Data Handling

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Minimizing bias is critical to adoption and implementation of machine learning (ML) in clinical practice. Systematic mathematical biases produce consistent and reproducible differences between the observed and expected performance of ML systems, resulting in suboptimal performance. Such biases can be traced back to various phases of ML development: data handling, model development, and performance evaluation. This report presents 12 suboptimal practices during data handling of an ML study, explains how those practices can lead to biases, and describes what may be done to mitigate them. Authors employ an arbitrary and simplified framework that splits ML data handling into four steps: data collection, data investigation, data splitting, and feature engineering. Examples from the available research literature are provided. A Google Colaboratory Jupyter notebook includes code examples to demonstrate the suboptimal practices and steps to prevent them.

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Machine learning (ML) applications in radiology have resulted in more than 8000 publications worldwide from 2000 to 2018 (1). Nonetheless, mitigation of possible mathematical bias (hereafter called bias) remains a critical concern for adopters (2,3). Although bias in medical research may be *random*, such as sampling variability or measurement precision, *systematic bias* produces consistent and reproducible differences between observed and expected performance (4). Unrecognized bias may contribute to suboptimal results.

To mitigate bias, researchers should carefully design and implement a pipeline of data handling, model development, and performance evaluation (5). Each of these steps may introduce systematic or random bias. Systematic biases can reduce the fairness of ML systems; such biases must be recognized and, ideally, eliminated. Table 1 describes several systematic biases that can arise in the data handling phase of ML system development.

Suboptimal quality of clinical data often limits the performance of ML algorithms (6,7). Therefore, developers must handle data accurately when performing data sampling, de-identification, annotation, labeling, or managing missing values. Although several guidelines exist for the proper development of ML systems, discussion of proper data handling is often insufficient (8,9). This report highlights common suboptimal data handling practices that may lead to systematic biases and briefly introduces common techniques to address them.

In line with Kocak et al (5), we define *data handling* as all data-related processes following the initial planning for an ML study up to model development and training. Although there are many ways to define data handling (10), we define four steps: data collection, data investigation,

data splitting, and feature engineering (Fig 1). These steps focus on the aspects of data handling that ML researchers can control. We avoid the terms *data preprocessing* and *data wrangling* as these terms may encompass more than one step. This report, intended for radiologists interested in ML who are not expert data scientists, focuses on computer vision examples; however, most of the principles are applicable to other ML applications such as natural language processing or tabular data analysis. A Google Colaboratory Jupyter notebook is provided with code examples for each step ([https://colab.research.google.com/drive/1c4G2b\\_ynikPsf2J5ExNS2D0or4uHX1dy?usp=sharing](https://colab.research.google.com/drive/1c4G2b_ynikPsf2J5ExNS2D0or4uHX1dy?usp=sharing)).

## Step 1: Data Collection

Data collection typically follows study design and usually requires researchers to access private or public data pools, query desired data types (eg, imaging, text, tabular), and transfer data to research storage. Depending on data properties, institutional policies, and study design, data de-identification may be done in this step or during feature engineering. Researchers may also access pre-existing human-labeled data, such as annotations for segmentation or classification models, or algorithm-generated data, such as synthetic images or text.

### Improper Identification of the Dataset

Careful selection of data is critical for ML system development. Collecting all available data (instead of a chosen subset) is not always feasible, and feeding redundant data may hinder training by adding unnecessary complexity (11). Conversely, failing to collect all available data predisposes trained models to additional biases. An example of improper dataset identification occurs when

## Abbreviations

DICOM = Digital Imaging and Communications in Medicine, EDA = exploratory data analysis, ML = machine learning

## Summary

This report describes 12 major practices during data handling that predispose machine learning systems to mathematical bias.

## Key Points

- Systematic biases in machine learning systems produce consistent and reproducible differences between the observed and expected performance from those systems, which limit the application of these systems to real-world scenarios.
- Although proper data handling is gaining more attention, guidelines to address the correct management of big data are scarce.
- Strategies are introduced to prevent different mathematical biases resulting from poor data collection, data investigation, data splitting, and feature engineering of machine learning models.

## Keywords

Data Handling, Bias, Machine Learning, Deep Learning, Convolutional Neural Network (CNN), Computer-aided Diagnosis (CAD)

retrieving Digital Imaging and Communications in Medicine (DICOM) files. Researchers may lose access to nonimaging data attributes available in DICOM metadata (eg, patient age and sex) if these files are converted to other file formats upon retrieval. Such attributes could be used to check study inclusion and exclusion criteria and to assure appropriate demographic representation (10).

To identify and collect the appropriate datasets, it is vital to estimate the types, attributes, and size of the data required. First, an in-depth review of available clinical and technical literature, accompanied by insights from medical experts, helps to determine the data types or attributes that are critical (12). Second, statistical power estimation techniques and knowledge of similarly developed ML systems can help determine the minimum dataset size needed to show an effect and also ensure the generalizability of the trained model (10,13). Finally, researchers may have access to more than one data type (eg, imaging from different modalities, different clinical datasets, or even human-labeled annotations for other ML purposes). Training a model on multiple data types may improve the performance of ML systems (14).

### Single Source of Data

Models may not generalize well if training data are collected from a single source (15). A model may perform well with data from the pool it was trained on, but there is a substantial risk of failing to perform well on data from other sources. Data from a single source may not sample the full data distribution in the real world. For example, chest radiograph classifiers trained on publicly available datasets may have multiple racial, sex, and socioeconomic biases that reflect the patient distribution in their training data (16), resulting in poor generalizability.

Another less recognized “single source” issue is collection of data from a single point in time. Hardware and software scanner upgrades may alter image appearance. Disease incidence and

appearance change with time, and new treatments may affect the appearance. Although the input data to ML systems may remain homogeneous over time, potential data drift should be anticipated (17).

Although it can be difficult to estimate how much expansion or diversification of training data will ensure generalization, there are multiple strategies to improve the heterogeneity of data sources. One strategy is to collect data from multiple institutions with different patient compositions (7). Although this strategy is not always feasible due to technical or patient privacy issues, the emergence of data de-identification tools, federated learning, and cloud data storage may help address this challenge (10,18–20). A second strategy is to collect data from different vendors (eg, imaging devices or electronic health records) within a single institution. Different makes and models of devices, including older devices, will help, although this option is likely inferior to collecting multi-institutional data (21). A third strategy is to use public datasets. Several nonmedical and medical datasets have been released publicly (22). These datasets may include human- or machine-generated labels, but even a dataset without labels may still help train self-supervised models. Such models are trained on tasks that do not need ground truth labels but still learn meaningful features from their training data, which can facilitate the training of other supervised models through a process called *transfer learning*. Any external data from other institutes or public sources should be checked in advance to avoid implicit biases (see below) (23). Finally, estimating the exact training data qualities needed to minimize distribution shift issues in real clinical practice is not always feasible. However, the training data should at least be sampled in a way that is representative of the population data.

### Unreliable Source of Data

Data should be collected from reliable sources: Users should understand unambiguously how the data have been retrieved, processed, and transferred. For example, developers may be unable to determine if protected health information embedded in DICOM files was removed appropriately in the absence of de-identification protocols. Human-labeled annotations (eg, segmentation masks or bounding boxes) require a clear protocol and measurement of intra- and interannotator reliability to reduce recall, observer, or measurement biases (24).

It is easier to remove any ambiguity in data before collection. Access to live clinical data from picture archiving and communication system and electronic health records is usually restricted to staff physicians, technologists, and data managers. Therefore, developers may need to collaborate with clinical stakeholders to communicate needs and clarify any ambiguities (10). Public datasets may be prone to biases and may not represent or report the age, race, or sex of the original population (25). It is also critical to make sure that there is no overlap of specific patients and images in public datasets, the so-called Frankenstein problem, where one public set is used to train a model and another is used to validate, but the presence of the same case in both sets means the real-world performance is poorer than measured (26).

**Table 1: A List of Different Systematic Data Handling Biases of Machine Learning Studies, Their Definitions, and Predisposing Suboptimal Practices**

Name of Bias	Definition	Predisposing Suboptimal Practices
Selection bias (also called sampling bias)	Collecting data that is not representative of the target population	Improper identification of the dataset Single source of data Inadequate EDA Unrepresentative datasets
Exclusion bias	Deleting valuable data that was thought to be unimportant	Inadequate EDA EDA with no domain expertise Failing to observe actual data Mismanagement of missing data
Measurement bias	Systematically favoring a particular result when observing or measuring variables in data	Unreliable source of data EDA with no domain expertise Failing to observe actual data Leakage between datasets Overfitting to hyperparameters Improper feature removal Improper feature scaling Mismanagement of missing data
Recall bias	Labeling similar types of data inconsistently	Unreliable source of data
Survey bias	Substantial missing, incomplete, and inconsistent responses to surveys, questionnaires, or interviews used to collect data	Improper identification of the dataset Inadequate EDA Mismanagement of missing data
Observer bias (also called confirmation bias)	Favoring information that does not contradict the researcher's desire or previous beliefs	Unreliable source of data EDA with no domain expertise Failing to observe actual data
Prejudice bias (also called human bias)	Training data includes (human) biases containing implicit racial, gender, or ideological prejudices	Unreliable source of data Inadequate EDA Mismanagement of missing data
Algorithmic bias	ML algorithm creating or amplifying the bias over the training data	Mismanagement of missing data

Note.—EDA = exploratory data analysis, ML = machine learning.

## Step 2: Data Investigation

Developers should investigate collected data from multiple perspectives to discover properties that may assist in detecting potential data issues, a process known as *exploratory data analysis* (EDA) (27). The main goals of EDA are to (a) organize and summarize the raw data, (b) discover important features and patterns in the data and flag any deviations, and (c) interpret findings in the context of the problem. The analytic operations of an EDA (eg, filtering, aggregation, and visualization) are done sequentially, where the result of one operation often denotes the next operation (28).

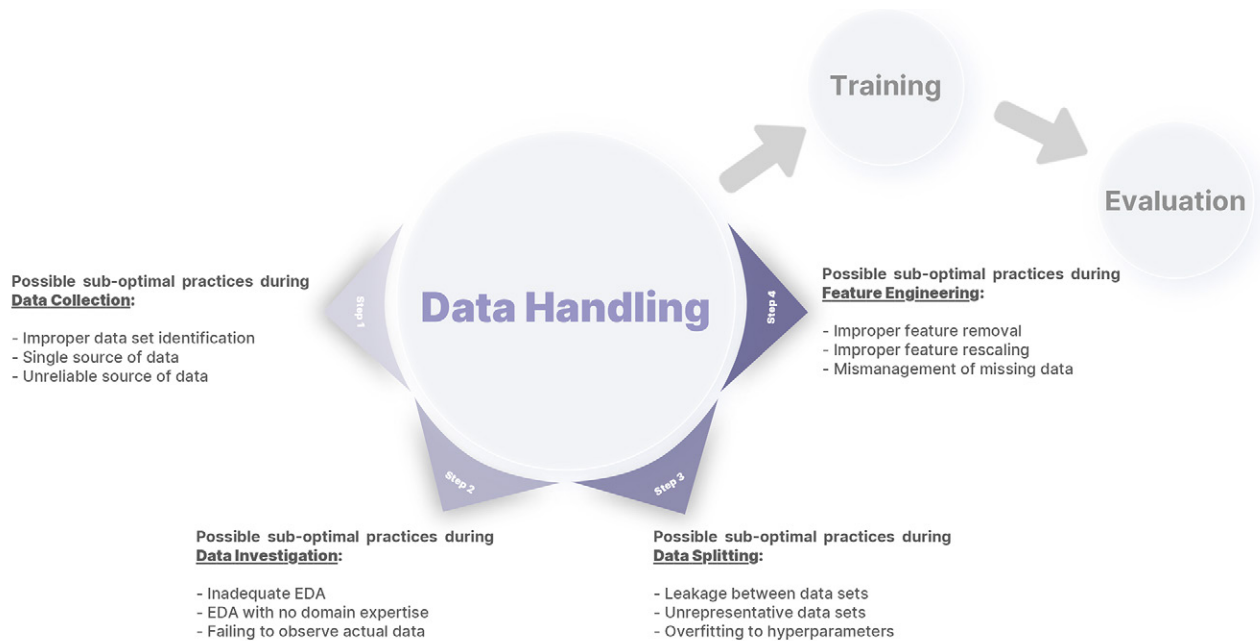
### Inadequate EDA

An inadequate EDA may be challenging to detect, but there may be no EDA at all if the collected data appear trivial. Many statistical aspects of the data should be investigated during an EDA, including measures of frequency, central tendency and spread, the shape of data distribution, missing data, and outlier data (27). One should investigate the interdependency within the data, such as if there is more than one imaging examination for some patients. Table 2 provides recommendations for EDA in a medical ML study.

EDA tools allow data visualization in many formats, including graphs, plots, or tables. Most ML applications are coded in Python, and many packages, such as Pandas and Matplotlib, provide basic EDA tooling (29–31). DataPrep and AutoAIViz perform EDA automatically for different data types (28,32,33). Although helpful, none of these tools will replace the need for in-depth and targeted analysis of data by clinical and data science experts.

### EDA with No Domain Expertise

Incorporation of domain expertise can enrich EDA (34). In fact, the quality of an EDA depends on the ability of the developers to ask the right questions and detect patterns within the data based on their cognitive ability, domain expertise, and experience in data interpretation (35). To detect bias, EDA should be conducted as a joint effort of ML statisticians, data scientists, and clinicians or other domain experts. Knowledge sharing, which can facilitate collaboration between developers and domain experts, involves identifying related domain experts, obtaining information from them using formal or informal tools (eg, meetings or interviews), and processing and



**Figure 1:** An arbitrary framework for defining data handling, consisting of four different steps: data collection, data investigation, data splitting, and feature engineering. Different errors introduced in this report for each step are also summarized. EDA = exploratory data analysis.

storing the collected information in repositories, such as question banks or manuals, for further use. Knowledge sharing is facilitated by many communication standards (36,37).

### Failing to Observe Actual Data

Although it seems trivial, personally reviewing the collected data is of utmost importance for ML developers and clinicians. While we already described several statistical measures to evaluate input data, observing the data itself (and not its statistical properties) often provides new insights. For example, in a study for developing an ML model to segment brain tumors from brain MRI studies, initial observation of a random sample from the input data may reveal varying sequences and quality of MRI studies. Such findings indicate a need to check the frequency of available sequences, as well as vendor and scan protocols. If feasible, all data points in datasets should be reviewed, and if not, researchers should draw a representative sample and inspect that subset. For example, imaging data could be visualized by programmatically building a mosaic photograph of individual imaging instances (38,39). Despite the lower resolution of each of these images (tile), they may be helpful for comparison and anomaly detection purposes (Fig 2). Similarly, conventional DICOM viewers may be used to inspect images and any human-labeled annotations. Tools like ITK-SNAP can overlay segmentation masks on underlying images to help check the quality of annotations (40). It is also necessary to review data after the feature engineering step (see below) to detect unwanted or undesirable modifications to the data.

### Step 3: Data Splitting

The data are partitioned into *training*, *validation*, and optionally *testing* sets. The training set is used to fit an ML model during training. The validation set is used to track the learning

process during training but is not used directly to train the model. The test set is used to evaluate the performance and generalizability of the ML model after the training is complete (Fig 3A). Critical data handling errors may occur, particularly when working with medical data.

### Leakage between Datasets

There should be no leakage between the training and test sets. In other words, to assure that models can distinguish meaningful signals from noise, the ML model should not “see” any of the test data during training (41). Data may “leak,” even if developers ensure that no data are repeated in both training and test sets because medical data are usually clustered at different levels. For example, a patient with a liver tumor may have four different liver MRI studies, each with more than one series, and each series with several images. In such a scenario, a random train-test split at the image level will result in biased training. Individual images are not necessarily independent, and images from the same series could go to both training and testing sets. Although the ML model will not see the same section in both the training and test sets, it will likely find very similar features from adjacent sections. This will help the model learn their inherent similarities. In such a case, both training and validation loss would decrease, but the model would not have learned general features and will subsequently perform poorly if applied to other patients. Even splitting based on series or study levels is not sufficient to prevent data leakage. Different scans from the same patient will have similarities, and the model’s performance may still be overestimated when applied to validation or test sets.

The standard way to prevent data leakage is to split medical data at the patient level. When training and test sets consist of data from different patients, developers can be more

**Table 2: Recommendations for Performing an Exploratory Data Analysis in a Computer Vision Machine Learning Study with Available Imaging and Nonimaging Medical Data**

Recommendation	Example(s)
Check the data type heterogeneity	<p>Check if imaging data are available as DICOMs or in other formats</p> <p>Check the heterogeneity of imaging data between patients and studies (eg, are all available MRI scans from the same sequence for all patients, or is there any heterogeneity)?</p> <p>If DICOMs are available, see what attributes (tags) are accessible in their metadata</p> <p>Check what clinical variables and text data are provided along with the images</p>
Analyze the frequency of data	<p>Count the frequency of patients for whom the data are provided</p> <p>Count the frequency of available imaging studies, series, and scans</p> <p>Count the frequency of available nonimaging data variables</p> <p>Plot frequency tables or graphs for imaging and nonimaging data</p> <p>Find out how the data availability is similar or different between patients. Are there any patients with too many or too few available data points of any kind?</p> <p>Count the number of missing data points</p>
Perform univariate analysis for nonimaging data	<p>Check the measures of central tendency and spread for each of the variables</p> <p>Plot the distribution of each variable in a graphical chart</p> <p>Determine outliers for each variable</p>
Perform multivariate analysis for nonimaging data	<p>Check for significant differences of mean, correlation, and covariance between the variables</p> <p>See if missing or outlier values for each variable are following any specific distribution with respect to other variables</p>
Explore the timing of studies	<p>Find out the date and time intervals for available studies</p> <p>Find out the general trajectory for each patient over time</p>
Evaluate the dimension (aspect ratio) of imaging data	<p>Plot the frequency of values for different dimensions (height, width, and the number of sections) of available imaging; see if any outlier values exist</p> <p>Check if any resizing or resampling is needed to make the available imaging more homogeneous</p>
Analyze the pixel (or voxel) values	<p>Calculate the measures of central tendency and spread for pixel (voxel) values</p> <p>See if any clipping is needed to remove or replace outlier intensities</p>
Observe the actual imaging data	<p>Visualize imaging data as single images or mosaic pictures and try to identify any unexpected characteristics</p> <p>Overlay any available labels (eg, segmentation masks or detection bounding boxes) on imaging data and check if they are clinically and technically acceptable</p>

Note.—DICOM = Digital Imaging and Communications in Medicine.

assured that their models will not have a chance to find similar noise between the sets. Splitting at the patient level is not always easy, as patients may have a different number of data points available, and multiple variables must be balanced between the sets. Python packages such as scikit-learn provide algorithmic tools that can speed up data splitting while grouping by a patient-level variable, such as the *PatientID* attribute of DICOM files, and stratifying the data based on other variables (see the notebook, [https://colab.research.google.com/drive/1c4G2b\\_ymikPsf2J5ExNS2D0or4uHX1dy?usp=sharing](https://colab.research.google.com/drive/1c4G2b_ymikPsf2J5ExNS2D0or4uHX1dy?usp=sharing)) (42). Finally, there are instances where data split at levels higher than the patient level may be beneficial. For example, if there are consistent differences in how a hospital handles patient scans, it may be valuable to separate data at the institution level.

### Imbalanced Datasets

The training and test sets should both represent real-world data. This condition is not easily met in medical domains for two important reasons: (a) the size of medical datasets is substantially smaller than most nonmedical datasets (eg, ImageNet), and (b) even the available data may have imbalanced distribution based on several variables. The causes of this imbalance go beyond ML and may pertain to population-level frequencies or patient recruitment protocols. Suppose developers want to train an ML model on hip radiographs from 10 000 patients who have undergone total hip arthroplasty to predict the risk of a complication. If the incidence of the complication in their institution is 2%, they would have 9800 patients with no complications versus 200 with complications, yielding a highly imbalanced dataset. As the conventional train-test split

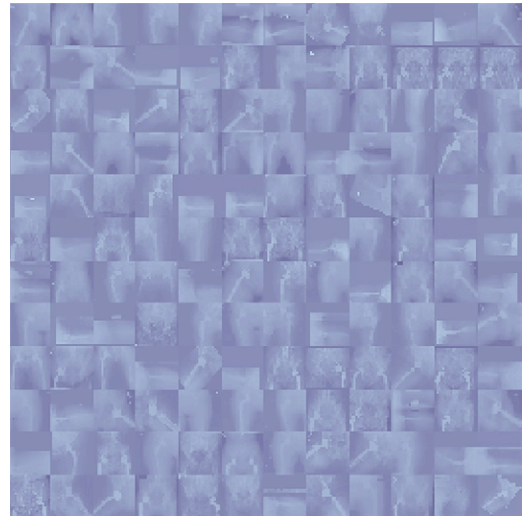
often holds out 10%–20% of data for testing, the test set will contain the data for only 20–40 patients with complications. In addition to the many training challenges that such a dataset will cause, there is a substantial risk that the few patients with complications from the test set are not representative of the population. Therefore, developers cannot rely on the reported model performance applied to their test set, as it may not generalize to the real-world data.

Another data splitting technique, *k-fold cross-validation* (Fig 3B), splits the data into *k* nonoverlapping folds (usually five or 10) (43). Each fold is used once as the test set, while all other folds are used as a training dataset. A total of *k* models are trained (one for each test set), and the mean performance is reported. Using *k*-fold cross-validation in our example, developers can ensure that the data from each of the 200 patients with complications have been used once to evaluate the model’s performance. Also, as the *k* models have been trained on different data, it is more likely that their mean performance is generalizable to real-world data, particularly if the variance is low. Using *k*-fold cross-validation is typically better than using the conventional train-test data splitting, specifically when dealing with small and imbalanced datasets. If used, we strongly recommend reporting not only the mean but also the range and the CIs of model performance to assess reliability.

### Overfitting to Hyperparameters

Overfitting is a known issue of training ML models when they do not perform as well on the test set as on the training data, denoting the lack of generalizability of the model. Although overfitting is generally a training issue, a particular case of it

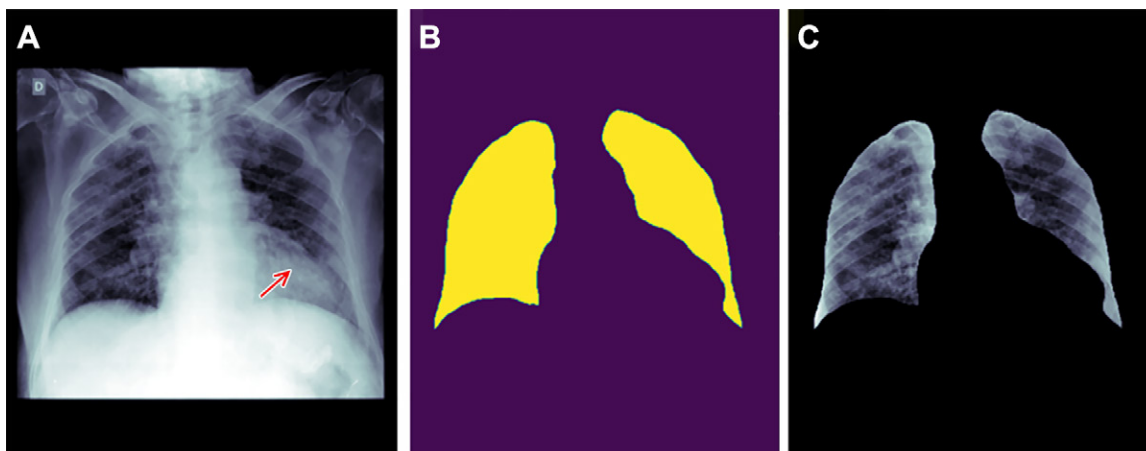
may be traced to data splitting: overfitting to hyperparameters or the test set. While training ML models, developers use various approaches like grid searching to choose the best hyperparameters. To compare each set of hyperparameters, one



**Figure 2:** A mosaic photograph of random radiographs was collected from our institutional dataset of patients who underwent total hip arthroplasty. Despite the reduced resolution of individual images, a quick look at this photograph reveals valuable insights for developers who desire training models on this dataset; for example, radiographs have different views, not all radiographs have prostheses, radiographs are from different sexes (the anatomy of pelvis is different between male and female patients), different prosthesis brands are available in the data, some radiographs have outlier intensities (presenting darker or brighter than expected), and so forth.



**Figure 3:** Schematic description of (A) traditional train-validation-test splitting, (B) fivefold cross-validation ( $k = 5$ ; where  $k$  is the number of folds), and (C) fivefold nested cross-validation ( $k = m = 5$ ; where  $k$  is the number of folds in the first-level cross-validation, and  $m$  is the number of folds in the second-level cross-validation).



**Figure 4:** Example of how improper feature removal from imaging data may lead to bias. **(A)** Chest radiograph in a male patient with pneumonia. **(B)** Segmentation mask for the lung, generated using a deep learning model. **(C)** Chest radiograph is cropped based on the segmentation mask. If the cropped chest radiograph is fed to a subsequent classifier for detecting consolidations, the consolidation that is located behind the heart will be missed (arrow, **A**). This occurs because primary feature removal using the segmentation model was not valid and unnecessarily removed the portion of the lung located behind the heart.

might evaluate their effect on model performance on the test set. Therefore, there is a risk that developers may gradually find the hyperparameters that work best on their test set but are not necessarily generalizable to other unseen data (44). To avoid this issue in the conventional train-test split, a distinct subset of the training set (aka, validation set) would be used for hyperparameter tuning (and no more for training), and the test set will be used in neither of these tasks.

We noted above that cross-validation can be useful for imbalanced and small datasets. Nested cross-validation is a variant of the  $k$ -fold cross-validation technique that tries to address the issue of overfitting to the hyperparameters (45). In this technique (Fig 3C), the  $k$ th fold will be held out, and data from the  $k - 1$  other folds, which were previously regarded as the training set, will collectively be split into  $m$  distinct folds using a second (inner-layer)  $m$ -fold cross-validation. The resulting  $m$  folds will then be used to train  $m$  models and find the best hyperparameter settings. The best model will finally be applied to the holdout  $k$ th fold, and its performance will be reported. In summary,  $m \times k$  models will be trained in a nested cross-validation approach, while no hyperparameter tuning happens on the holdout sets. It should be noted that both the  $k$ -fold cross-validation and nested cross-validation approaches are prone to methodologic biases and may not be always feasible; therefore, specific guidelines must be followed to obtain their most reliable performance (46).

#### Step 4: Feature Engineering

Feature engineering transforms or removes features from the input data before fitting an ML model (46). The idea is to change the input data so that ML models see more meaningful and less redundant features. Many techniques could be incorporated into feature engineering, including techniques that aim to reduce noise in the data (eg, denoising, kernel-based transformation, organ windowing followed by normalization, and cropping followed by zero padding of imaging) before feeding to ML models (47). Because of the complicated nature

of feature engineering, suboptimal practices can occur during feature engineering and make the ML models prone to underfitting and bias.

#### Improper Feature Removal

One routine question in feature engineering is “What features can be removed from the input data to make the learning easier for ML models?” The underlying assumption for this question is that there is noise in the data that, if removed, enables easier fitting of an ML model. Whether this assumption is valid depends on the input data, the algorithm, and the task at hand. Suppose some developers would like to train a deep learning classifier for differentiating COVID-19 pneumonia from other viral pneumonia using chest radiographs. Model performance may not necessarily improve if they use another model to first segment the lungs from the original chest radiograph and then train the classifier only on cropped lung regions. While noise may be reduced, the classifier could look at nonlung regions of the chest radiograph (eg, the heart) and find additional helpful signals. Another problem with that assumption is that ML models do not necessarily learn as humans do. What is considered noise to developers may be a source of valuable signal data for models.

Evaluating all scenarios in separate experiments is the best way to understand if removing features will cause the model to fit better or worse (thus reducing or increasing bias). Although such a solution could increase training costs, developers should consider it whenever possible while also relying on common sense or implicit clinical knowledge and experience. Figure 4 demonstrates an example of how improper feature removal may lead to subsequent bias in a model’s performance.

#### Improper Feature Scaling

Feature scaling means applying arithmetic operations on feature values so that most (if not all) of the features from the input data are brought to a similar scale. This way, features will not have dramatically different magnitudes, and the learn-

ing process (eg, the back propagation in gradient descent algorithms) is more efficient. Many techniques are available for feature scaling, but two common ones are (a) normalization, which changes the feature values to a common scale (eg, 0–1) and (b) standardization (or z score normalization), which scales feature values to have zero mean and unit SD.

Although these techniques are applied to improve model fit and generalizability, they can result in poorer fit, and even bias, if misapplied. Seeking consultation from expert statisticians may help choose the appropriate scaling techniques for the data and algorithms. For example, standardization is more effective if the feature values have an almost Gaussian distribution, and the ML algorithm assumes the distribution of data to be normal. Normalization, on the other hand, does not have this limitation. Therefore, while normalization is more appropriate for algorithms like k-nearest neighbors and artificial neural networks, standardization works better for regression algorithms. As with other examples, normalization may amplify the noise in features with almost constant values, and standardization can significantly distort data values in the presence of outliers (48).

There are several tips to ensure feature scaling results in more benefit than harm for the input data. First, feature scaling must be deemed necessary. For example, ML algorithms such as tree-based models do not consider the scale of the data and often do not need feature scaling. One should also check the current scale and distribution of data and apply the appropriate feature scaling technique. For example, it is needed to remove outlier values in data before applying standardization. To do so, voxel values in medical imaging data like radiographs or MRI can be clipped to a specific value range (eg, between 5% and 95% percentile of intensity distribution or between the lower and upper narrow tails of a histogram obtained from probability distribution function of the image intensity) (47).

### Mismanagement of Missing Data

All real-world data are likely to have missing data. In fact, some ML repositories may have more than 40% of data missing (49). While discussing why missing data exists in ML is outside the scope of this report, how missing data are handled does affect bias and fairness. Previous research has shown that missing data may not be evenly distributed, leading to unwanted effects on the fairness of the data and the resulting models (50). For example, underrepresented groups may be reluctant to provide sensitive information on medical or demographic questionnaires and are more likely to miss follow-up medical visits (51). They may even receive more or fewer diagnostic and treatment options than necessary (52). Therefore, the medical data for such underrepresented groups may be less available and ignored (or interpreted mistakenly) by ML systems that have not addressed missing values properly during training, resulting in prejudice and selection biases.

The simplest (but not recommended) way to deal with missing data during ML development is to remove all patients for whom data points are missing. This approach can result in bias and fairness issues, as (a) removal of cases may exacerbate the problem of small or imbalanced datasets, and (b) the missing data are not random and more commonly affects a specific

population of patients; thus, the final model's performance may be significantly worse on underrepresented populations (53,54). Moreover, various studies have shown how the performance of models may be degraded when missing data are discarded. A better strategy is to substitute missing data with synthetic values (so-called *imputation*). Imputation may be as simple as replacing the missing values with the mean or median of the available data; however, it could also be done using more sophisticated techniques and even by using ML to predict the missing values (55). While some imputation techniques (especially those that use predictive ML) may amplify algorithmic bias in the data, they generally improve the model's performance while maintaining its fairness (50). Exhaustive discussions of the pros and cons of different imputation methods are beyond the scope of this report but are available elsewhere (53–55).

### Conclusion

ML has been widely deployed, from image reconstruction and hypothesis testing to improving diagnostic, prognostic, and monitoring tools (56). This report discussed how ML tools could be susceptible to biases in their data handling phase and how developers might prevent those biases. Our report covered items 7–13 of the Checklist for Artificial Intelligence in Medical Imaging and items 1–5 of the Guiding Principles for Good Machine Learning Practice for Medical Device Development, recently introduced by the U.S. Food and Drug Administration, Health Canada, and the United Kingdom's Medicines and Healthcare products Regulatory Agency (24,57).

The recommendations here should be interpreted and applied with caution. First, the discussion is based on an arbitrary data handling framework of four successive steps: data collection, investigation, data splitting, and feature engineering. This framework is oversimplified, and we acknowledge that ML systems are not developed as a linear process. Second, this report has considered only the mathematical forms of bias. Social and ethical forms of bias can harm underrepresented groups or benefit those with more resources. Addressing such biases often needs political decisions and time-consuming changes in stakeholders' behaviors. Third, ML bias is an active field of research, and this report is not an exhaustive list of problems or solutions. Additionally, bias mitigation strategies are problem specific, and certain strategies described above may be unnecessary or even inappropriate in specific situations. Finally, not all bias-prone practices in ML can be easily replaced or improved. Limitations do exist in ML studies, and ML systems may be biased in different ways. Depending on the data and task at hand, such biases could be tolerated or even considered helpful (58). However, developers must identify erroneous practices during ML development, mitigate as many of them as possible, and explicitly report the remaining limitations of their ML systems. In further reports, we will focus on biases that exist in two other phases of ML studies, namely *model development* and *performance evaluation*.

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