Explainable machine-learning predictions for the prevention of hypoxaemia during surgery

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Although anaesthesiologists strive to avoid hypoxaemia during surgery, reliably predicting future intraoperative hypoxaemia is not possible at present. Here, we report the development and testing of a machine-learning-based system that predicts the risk of hypoxaemia and provides explanations of the risk factors in real time during general anaesthesia. The system, which was trained on minute-by-minute data from the electronic medical records of over 50,000 surgeries, improved the performance of anaesthesiologists by providing interpretable hypoxaemia risks and contributing factors. The explanations for the predictions are broadly consistent with the literature and with prior knowledge from anaesthesiologists. Our results suggest that if anaesthesiologists currently anticipate 15% of hypoxaemia events, with the assistance of this system they could anticipate 30%, a large portion of which may benefit from early intervention because they are associated with modifiable factors. The system can help improve the clinical understanding of hypoxaemia risk during anaesthesia care by providing general insights into the exact changes in risk induced by certain characteristics of the patient or procedure.

Over 300 million surgeries are performed worldwide every year. Although an integral part of healthcare, surgery and anaesthesia pose considerable risk of complications and death. Studies have shown a perioperative mortality rate of 0.4–0.8% and a complication rate of 3–17%, just in industrialized countries. Fortunately, half of these complications are preventable. With an increase in the adoption of electronic medical record systems, high-fidelity heterogeneous data are being captured during surgery and anaesthesia care, but these data are rarely used to improve patient safety and quality of care. There is untapped potential for data science to utilize perioperative data to positively impact surgical and anaesthesia care. To address this unmet need we leverage recent advances in perioperative informatics and present new machine-learning methods to predict harmful physiological events and to inform anaesthesiologists.

Hypoxaemia, or low arterial blood oxygen tension, is an unwanted physiological condition known to cause serious patient harm during general anaesthesia and surgery. Hypoxaemia is associated with cardiac arrest, cardiac arrhythmias, post-operative infections and wound healing impairments, decreased cognitive function and delirium, and cerebral ischaemia through a number of metabolic pathways. Despite the advent and use of pulse oximetry to continuously monitor blood oxygen saturation (SpO2) during general and regional anaesthesia, hypoxaemia can neither be reliably predicted nor prevented at future time points. Real-time blood oxygen monitoring through pulse oximetry only allows anaesthesiologists to take reactive actions to minimize the duration of hypoxic episodes after their occurrence. Decision support systems that process electronic medical record data have been shown to help increase adherence to guidelines, but remain primarily reactive rather than predictive in nature, see ref. for a full review. If hypoxaemia can be predicted or anticipated before it occurs, then actions can be taken by anaesthesiologists to proactively prevent hypoxaemia and minimize patient harm.

Machine-learning techniques use statistical methods to infer relationships between patient attributes and outcomes in large datasets and have been successfully applied to predict adverse events in health care settings, such as sepsis or patient deterioration in the intensive care unit. Yet machine-learning techniques to predict adverse events such as hypoxaemia in a considerably more complex setting such as the operating room are lacking at present. Moreover, although previous complex machine-learning approaches provide good prediction accuracy, their application in an actual clinical setting is limited because their predictions are difficult to interpret and hence not actionable. Interpretable methods explain why a certain prediction was made for a patient, that is, the specific patient characteristics that led to the prediction. This lack of interpretability has thus far limited the use of powerful methods such as deep learning and ensemble models in medical decision support.

We present an ensemble-model-based machine-learning method, Prescience, that predicts the near-term risk of hypoxaemia during anaesthesia care and explains the patient- and surgery-specific factors that led to that risk (Fig. 1). We believe this is an important step forwards for machine learning in medicine because although machine-learning models have significantly improved the ability to predict the future condition of a patient, the inability to explain the predictions from accurate, complex models is a serious limitation. Understanding what drives a prediction is important for determining targeted interventions in a clinical setting. For this reason, machine-learning methods employed in clinical applications avoid using complex, yet more accurate, models and retreat to simpler interpretable (for example, linear) models at the expense of accuracy. To address this problem, some approaches have achieved interpretability by carefully limiting the complexity of the
Prescience integrates many data sources into a single risk, which is explained through a succinct visual summary. A wide variety of data sources were used to build a predictive model of hypoxaemia events. An explanation (overlaid) is then built for each prediction. Pink features have values that increased risk, whereas green features decreased hypoxaemia risk. The combination of the impacts of all features is the predicted Prescience risk; in this case, the odds are 2.4 times higher than normal. Each feature impact value represents the change in risk when the value of that feature is known versus unknown. Qualitative terms such as ‘low’ or ‘high’ are based on the distribution of a feature value in our dataset.

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Second, Prescience explains why a prediction was made, regardless of the complexity of the machine-learning model used to make the prediction. Significant progress has been made recently in integrating predictive machine-learning solutions into medical care15–17,19. However, accurately and intuitively conveying to doctors why a prediction was made remains a key challenge. For example, a numeric representation of risk is useful (for example, the 2.4 odds ratio in Fig. 1). However, a more detailed presentation that shows the risk is due to the body mass index (BMI), current tidal volume and pulse rate of a patient is more clinically meaningful as some factors may be modifiable and result in clinical changes mitigating that risk (Fig. 1). Typically, understanding why a prediction was made requires limiting the complexity of the model15, but Prescience enables explanations for models of arbitrary complexity. The feature impact values computed by Prescience essentially represent the change in the predicted risk of the model when we observe a feature (such as the weight of a patient) versus when we do not observe the feature (such as not knowing the weight of a patient). This change in the output prediction of a model when a feature is observed indicates its importance for the prediction. Feature importance does not imply a causal relationship and so do not represent a complete diagnosis of hypoxaemia in a patient. However, they do enable an anaesthesiologist to better formulate a diagnosis by knowing which attributes of the patient and procedure contributed to the current risk predicted by the machine-learning model.

Results

To demonstrate the value of the explained predictions made by Prescience and gain insight into factors that affect intraoperative hypoxaemia, we present the following results: (1) a comparison of Prescience hypoxaemia predictions against anaesthesiologists’ predictions with and without the aid of Prescience; (2) an example of how Prescience explains hypoxaemia risk at a specific time-point during a surgical procedure; (3) a comparative summary of relevant AIMS data features for hypoxaemia prediction chosen by Prescience...
and by anaesthesiologists and (4) a detailed presentation of key risk factors for hypoxaemia identified by Prescience.

Prescience overview. Based on the World Health Organization recommendations and for the purposes of prediction, we defined hypoxaemia as the decrease in SpO₂, that is, arterial blood oxygen saturation as measured by pulse oximetry, to a threshold value of 92% or lower (see Methods; Supplementary Fig. 1). From the AIMS data, we extracted 3,797 static extracted features for each patient from more than 20 original static sources and an expanded super-set of 3,905 real-time and static extracted features for each time point during anaesthesia care from the more than 20 original static sources as well as 45 different real-time data sources (see Methods; Supplementary Table 1). Features such as words from text data get directly mapped to the display in Fig. 1, whereas sets of features from a single time series (such as tidal volume) are combined in Fig. 1. We excluded select cases (heart transplant, lung transplant, tracheostomy and coronary artery bypass surgeries) in which SpO₂ and other hemodynamic parameters can be significantly affected by non-physiological measurements, for example, during cardiopulmonary bypass. All the experiments were performed after appropriate Institutional Review Board approval (see Methods), with clinical data summarized in Fig. 2.

We trained a gradient boosting machine model to solve the following two types of prediction problems. The first is the initial prediction, at the start of a procedure the risk of hypoxaemia at any time during a procedure is predicted on the basis of the static extracted features. The second is the real-time prediction, predicting hypoxaemia in the next 5 min at various points of the operative period based on real-time and static extracted features collected up to that time point. We chose this 5 min window as it would be long enough to allow an anaesthesiologist enough time to intervene but also short enough so that it represents near-term risks that would benefit from immediate attention. For initial prediction we used 42,420 procedures (each a single surgical case) as training samples to train the gradient boosting machine, 5,649 procedures as validation samples to choose the tuning parameters for the gradient boosting machine (and other prediction models for comparison) and 5,057 as test samples for comparing across different prediction models (Supplementary Fig. 2). For real-time prediction we used 8,087,476 per-minute time points as training samples, 1,053,629 as validation samples and 963,674 as test samples, for which all time points from the same procedure were included in the same sample set and no missing data imputation was performed (Supplementary Fig. 3).

Fig. 2 | Patient and procedure characteristics. Histograms summarizing the basic properties of the anaesthesia procedures used for training (y axes represent normalized procedure counts). Prescience was trained and evaluated using data from 53,126 procedures recorded at two hospitals over two years (representing 36,232 unique patients). In our dataset, 37.4% of adults aged 20 or over have a BMI of 30 or more, which is a close match to the US obesity rate of 37.9%

Dividing the time points by procedure is important as samples from the same procedure are not independently and identically distributed but have some time dependency. To ensure that there was no bias towards the final test set, the test data was initially compressed and left compressed until method development was completed.

As shown in Supplementary Figs. 2 and 3, the gradient boosting machine outperforms alternative prediction models previously used for similar problems, particularly for the primary task of real-time prediction.

We used 198 and 523 test samples to evaluate the performance of anaesthesiologists for initial and real-time prediction tasks, respectively (see below; Fig. 3). Prescience outputs the risk prediction and its explanations (Figs. 1 and 4a), which show a set of features that increased (pink) and decreased (green) the risk.

We developed an efficient, theoretically justified machine-learning technique based on recent advances for interpreting models to estimate the importance of each feature in a prediction made for a single patient. This drives real-time explanations (Fig. 4) for the Prescience model. We verified the quality of the explanations given to the anaesthesiologists (in the experiments described below) by comparing the explanations with the change in model output when a feature is perturbed (Supplementary Fig. 4). We also developed effective visualizations of these explanations that encode them in a compact visual form for anaesthesiologists (Fig. 1; Supplementary Figs. 5–7) and a more detailed visualization that highlights the relevant contributing features (Fig. 4; see Methods for details).

Prescience improves the ability of an anaesthesiologist to predict hypoxaemia. To test the potential of Prescience to aid hypoxaemia prediction we replayed pre-recorded intraoperative data from test sample procedures in a web-based visualization to five practising anaesthesiologists (Supplementary Figs. 5–7). Each anaesthesiologist was given both types of prediction tasks, initial prediction (198) and real-time prediction (523). For each prediction task, anaesthesiologists were asked to provide a relative risk of hypoxaemia compared to a normal acceptable risk, for example, 0.01 for one-hundredth of the normal risk or 3.4 for 3.4 times the normal risk. These relative risks were then used to calculate standard receiver operating characteristic (ROC) curves averaged over five anaesthesiologists as shown in Fig. 3, which plots the true positive rate (that is, the percentage of correctly predicted desaturations) on the y axis against the false positive rate (that is, the percentage of incorrectly predicted non-desaturations) on the x axis. Note that ROC curves depend only on the order of the relative risk values among
predictions from a single anaesthesiologist. This eliminates the need to choose a threshold and the need to separately calibrate risk scores between anaesthesiologists.

Figure 3a,b shows that for both types of prediction tasks, the predictions made by Prescience (pink) are considerably more accurate than the predictions made by anaesthesiologists (green). The prediction accuracy of anaesthesiologists markedly improved when the anaesthesiologists were given Prescience’s risk prediction and its explanations in addition to the original procedure data (Supplementary Figs. 5–7). A clear separation between the performance of anaesthesiologists with and without the aid of Prescience is observed for both initial prediction (Fig. 3a, P < 0.0001) and real-time prediction (Fig. 3b, P < 0.0001). This suggests that Prescience can enhance the assessment of future risk made by anaesthesiologists and their ability to proactively anticipate hypoxaemia events. Interestingly, the prediction performance of anaesthesiologists with Prescience explanations was slightly lower than direct predictions from Prescience. This means that when the anaesthesiologists adjust their risk estimate for a patient away from what Prescience originally predicted they are more likely to be wrong than right.

To avoid the scenario in which an anaesthesiologist is tested twice on the same prediction task—one with and the other without Prescience, we created replicate test sets by dividing the prediction tasks into two groups of similar size: tasks for initial prediction and tasks for real-time prediction. The procedures shown to anaesthesiologists were receive Prescience’s assistance in one of these two replicate test sets and their ability to proactively anticipate hypoxaemia events. Note that the false positive rate (FPR; x axis) measures how many points without upcoming hypoxaemia were incorrectly predicted to have upcoming hypoxaemia. The true positive rate (TPR; y axis) measures the fraction of hypoxaemic events that were correctly predicted. P values were computed using bootstrap resampling over the tested time points while measuring the difference in area between the curves. If we instead resample over anaesthesiologists we observe bootstrap P values of 0 and Student’s t-test P < 0.001 for Prescience improvements. See Supplementary Fig. 8 for plots of the statistical separation between the mean ROC curves across all false positive rates.

The anaesthesiologists consulted had experience after residency ranging from 3 to 26 years (median = 7 yr) and were all actively practising at University of Washington Medical Center, VA Puget Sound Health System or Seattle Children’s hospital. The extensive experience level of the anaesthesiologists who participated in our study may not represent the typical experience level of anaesthesia providers, especially when nurse anaesthetists and residents in training provide anaesthesia care.

When using Prescience predictions to generate early warning alarms in the operating room, it is important to minimize the false alarm rates. This can be accomplished by adjusting the trade-off between precision (the positive predictive value) and recall (the sensitivity). High precision means a low false alarm rate (which is 1 – precision), however, it comes at the cost of low recall. Supplementary Figure 10 plots the precision and recall trade-off for Prescience on the full set of test time points. Given that the performance of the complex model in Prescience improves with larger datasets, we also included results from a model trained on
an expanded dataset of 175,000 procedures to measure the benefit of using more data to train Prescience. The larger dataset resulted in notably better performance and could capture 9% of all minutes with upcoming hypoxaemia at 70% precision (or 44% of all minutes with upcoming hypoxaemia at a precision of 30%, if the threshold for precision-recall trade-off is selected for higher recall).
Precisions are strikingly higher than those we project that anaesthesiologists would achieve on the full test dataset (Supplementary Fig. 10). We also note that the predictive accuracy can be further improved by shortening the predictive window to less than five minutes (Supplementary Fig. 9).

Anaesthesiologists must not only decide when to act to prevent hypoxaemia, but also when not to act. To assist in this, Prescience can predict not only when hypoxaemia will occur, but also when it will not occur. Prescience can predict when hypoxaemia will not occur for 60% of all time points while maintaining a precision of 99.9% (Supplementary Fig. 11).

Explained risks reveal both procedure and time specific effects. An explanation from Prescience represents the effects of interpretable groups of extracted patient features (see Figs. 1 and 4a), where each group corresponds to the set of extracted patient features from a single input feature in the AIMS dataset, such as the SpO2 monitor time series. These effects explain why the model predicted a specific risk and thus allow an anaesthesiologist to plan appropriate interventions.

In Fig. 1 only the most significant features contributing to hypoxaemia risk are shown for quick reference. However, in Fig. 4 the relative contributions of all patient and case features (that is, attributes) towards hypoxaemia risk can be seen at every sample time point during a procedure (Fig. 4b). Without a meaningful explanation, the sudden increase in risk shown at the time point marked ‘Now’ might be hard to interpret; however, by representing the predicted risk as a cumulative effect of contributing patient and procedure features, the reason for the increase becomes clear (Fig. 4a).
The increase in the risk of hypoxaemia in the next five minutes shown in Fig. 4 is driven by a set of features capturing both static attributes, such as patient height and weight, and dynamic parametric values, such as tidal volume (that is, the volume of gas exhaled per breath) and the administration of drugs. The risk explanation bar in Fig. 4a has pink features that push the risk higher (to the right) and green features that push the risk lower (to the left). Each group of features is sorted by the magnitude of their impact and the features with the greatest impact are labelled. Through this representation we can see that many of the 3,905 real-time extracted features have only a small impact and the risk for this time point is predominantly driven by a few features. The choice of features provided to the model was driven by the data recorded in the AIMS system and hence available for training. Rather than only provide the model with features we believed important, we let the model use any feature it chose. This means that it may find features we would not initially expect to be predictive of hypoxaemia. For some of these features it is helpful to tag them with indicators of how they relate to hypoxaemia risk before final deployment in an operating room. This can help anaesthesiologists to quickly see non-obvious connections with patient physiology, such as how the muscle relaxant succinylcholine in Fig. 4 does not represent a direct causal impact on hypoxaemia, but rather is a proxy that captures the risk from a potentially difficult airway or full stomach (in the hospital system we considered, succinylcholine is given to patients with a high risk of a difficult airway during intubation). Figure 4b shows the trend in the Prescience risk predictions over the course of the procedure. The plot in Fig. 4b is equivalent to rotating the feature explanation in Fig. 4a by 90° and then stacking the explanations for each time point horizontally. We can see from the trend in Fig. 4b that the large increase in risk at the current time was driven by tidal volume (there was a recent drop in this patient's tidal volume). The future SpO2 measurements confirm that the patient did indeed progress to hypoxaemia. Not only does Prescience alert anaesthesiologists when the patient risk of hypoxaemia is high, but it also provides information on the factors driving the risk and their relative contributions. This informed risk prediction enables anaesthesiologists to plan an appropriate course of action to avoid hypoxaemia.

Averaged feature-importance estimates broadly align with a survey of prior expectations. To gain an understanding of the general impact of features across all procedures, we computed the average importance of each feature in the Prescience model. In contrast to the explanations shown in Figs. 1 and 4a, which are specific to a single prediction at a particular time point, these average feature-importance estimates are over many procedures and time points. Estimates of average feature importance are shown in Fig. 5 for both initial prediction and real-time prediction.

To estimate which clinical features anaesthesiologists use to estimate hypoxaemia risk, we first performed a survey before using Prescience, asking four anaesthesiologists to list the most important factors they consider when assessing the risk of hypoxaemia, both before (for initial prediction) and during (for real-time prediction) a procedure. Their responses were then aggregated into a single ranked list of features (Supplementary Tables 2 and 3). Figure 5 shows the rankings chosen by anaesthesiologists next to the feature-importance estimates derived by Prescience for the initial (Fig. 5a) and real-time predictions (Fig. 5b). The ranking of features by anaesthesiologists appears to correspond well with the ranking by Prescience.

As another way to measure which features anaesthesiologists think contribute to hypoxaemia, we learned from their behaviour by training a separate gradient-boosting-machine model on the basis of their predictions. This allows a direct comparison between the anaesthesiologists and Prescience on the same set of features. We fit this model to all of the anaesthesiologist relative-risk predictions using ten-fold cross-validation. We then computed the feature-importance estimates for this model that was trained to mimic the behaviour of anaesthesiologists. Given the smaller set of training examples used to train the model (198 initial predictions and 523 real-time predictions), we used bootstrapping to estimate the variability of the feature-importance estimates (Fig. 5; right).

In general, there is reasonable agreement between the Prescience feature-importance estimates and those identified by the anaesthesiologists. However, there are important differences that may stem from the comprehensive nature of the Prescience analysis, whereas anaesthesiologists necessarily focus on what they consider the most likely causes for hypoxaemia. One striking difference is the reduced role of current SpO2 levels in the predictions of anaesthesiologists. Although anaesthesiologists are clearly influenced by the recent patterns of patient SpO2 levels, Prescience strongly depends on these patterns, whereas anaesthesiologists appear to be equally influenced by other factors, such as end-tidal CO2 (the amount of CO2 exhaled by the patient) and peak ventilation pressure. The second and fourth ranked features by anaesthesiologists for initial prediction were lung disease and asthma, respectively, which did not show up as important features for Prescience. This is potentially because they must be extracted from preoperative text notes and only about 1% of the procedures recorded the term chronic obstructive pulmonary disease, for example, and only 3% of case notes mention asthma.

Our study used data from two hospitals and the initial hypoxaemia predictions were driven by a bias between the two hospitals. This is perhaps unsurprising as one hospital is a Level-1 trauma centre and a significant proportion of its surgical cases involve trauma patients who are more susceptible to hypoxaemia. However, it is interesting to note that the importance of hospital as a risk factor became insignificant for the intraoperative real-time predictions, presumably because the risk differences in each hospital were captured by the real-time features.

Among the static features, BMI and age were significant risk factors. These features are well understood in the medical literature as risk factors that can increase the chances of hypoxaemia. Among the real-time (intraoperative) features, SpO2 was also a significant intraoperative feature identified by Prescience as predictive of hypoxaemia. Lower values may indicate inadequate ventilation or airway obstruction, which can in turn increase the risk of hypoxaemia.
also determined that hypotension (systolic blood pressure below 80 mm Hg) increases the risk of hypoxaemia. On the other hand, moderately higher FiO₂ (inspired O₂ concentration) and positive pressure ventilation can reduce the risk of hypoxaemia, as expected by anaesthesiologists.

Prescience’s estimated importance of individual features for hypoxaemia risk highlight important clinical relationships. Three important features for both initial and real-time predictions were chosen to illustrate how the Prescience model modifies hypoxaemia risk on the basis of changes to feature characteristics (Fig. 6). Although many such relationships are present for the various features, Fig. 6 shows a representative selection that demonstrates informative risk relationships that are captured in the Prescience model.

Among the static features, we find that patient BMI has a clear effect on the risk of hypoxaemia. When the BMI is greater than 26, the risk of hypoxaemia increases linearly until it has more than doubled when the BMI is greater than 50. Although a qualitative association between hypoxaemia and body weight is well established in the field of anaesthesia[27,28], Prescience quantifies this relative risk.

Prescience shows that patients with higher ASA physical-status codes have a higher risk of intraoperative hypoxaemia. This is not surprising as higher ASA codes represent increased severity of the physical condition of a patient, such as pre-existing pulmonary and cardiac conditions that can predispose a patient to develop hypoxaemia. Prescience data support clinical observations that the effect of ASA status on the risk of hypoxaemia more than doubles when the ASA status increases from I to V. Advancing age also predicted intraoperative hypoxaemia, probably representing the presence of co-morbidities[27]. These data show that BMI > 30, which meets the clinical definition of obesity[29], is associated with intraoperative hypoxaemia, suggesting impaired pulmonary mechanics. These findings confirm clinical observations and suspicions of the relationship between these patient factors and adverse anaesthesiology outcomes. They also quantify the risk associations, giving a more clinically useful interpretation to anaesthesiologists.
For real-time prediction, measurements from each time series are represented by a set of multiple features. For simplicity, we focus here only on the effect of the shortest time lag exponentially weighted moving average, which essentially represents the most recent reported value in the time series (see Methods for details).

Tidal volume represents the amount of gas exhaled per breath when the patient is either breathing spontaneously or mechanically ventilated during general anaesthesia. As the tidal volume drops below 0.6 l (keeping all other features the same), Prescience risk for hypoxaemia increases. This increase could be due to hypoventilation, in which case anaesthesiologists can take preventative steps to avoid inadequate ventilation.

Figure 6 shows the relationship between end-tidal CO$_2$ and risk of hypoxaemia under general anaesthesia. End-tidal CO$_2$ below 35 mmHg is associated with an increasing risk of intraoperative hypoxaemia. Although we cannot definitively attribute intraoperative hypoxaemia to hypocapnia, this association may represent underlying patient conditions, such as chronic obstructive pulmonary disease, that affect both physiological conditions. Alternately, the low end-tidal CO$_2$ may result from either intentional or unwanted hyperventilation during anaesthesia care.

Examining FiO$_2$ is important because anaesthesiologists can control the amount of oxygen delivered to patients. The current practice is to not provide all patients with 100% FiO$_2$ because not all patients need it, prolonged ventilation with 100% FiO$_2$ is associated with pulmonary atelectasis and delivering oxygen when it is not needed is costly and wasteful. These data show that FiO$_2$ below 40% is independently associated with intraoperative hypoxaemia, irrespective of other features. These findings provide important information regarding the safe provision of FiO$_2$ in patients during general anaesthesia. It is possible that the routine practice of maintaining FiO$_2$ at 30% or close to room air may be harmful to patients and not desirable. Although these effects are adjusted for all other available features, it is important to note that, as with any observational study, some residual confounding with patient risk may still exist. This could explain the increase in hypoxaemia risk we observed for high O$_2$ levels.

These representative features illustrate the ability of our machine-learning-based prediction method to not only provide explained-risk predictions for a complex model, but also quantitative insights into the exact change in risk induced by certain patient or procedure characteristics.

Discussion
Prescience is designed to comprehensively integrate high-fidelity operating room data to predict intraoperative hypoxaemia events before they occur. Based on a comparison with practising anaesthesiologists and existing computational methods applied to other clinical problems, Prescience achieves superior performance when predicting hypoxaemia risk from electronically recorded intraoperative data.

Prescience combines high-accuracy complex models with interpretable explanations. This combination of accuracy and interpretability allows physicians to receive the best possible predictions while also gaining insight into why those predictions were made. To test how Prescience predictions with explanations would impact the ability of an anaesthesiologist to estimate hypoxaemia risk, we compared anaesthesiologist predictions with and without Prescience assistance. We observed a clear increase in prediction accuracy when doctors were assisted by Prescience, demonstrating that anaesthesiologists may make more accurate hypoxaemia risk assessments in the operating room if they had access to Prescience. The augmented-intelligence style approach of Prescience may be particularly helpful for mitigating the unwanted effects of variations in knowledge and/or practice among providers.

Empirically derived black box algorithms such as the bispectral index have been used to track the brain states of patients undergoing general anaesthesia by processing real-time electroencephalograms. These algorithms have been criticized because they do not utilize physiological models, do not identify factors associated with risk of events and produce empirically derived metrics to represent neurophysiology of how the anaesthetics affect the brain. The black-box nature of the electroencephalogram algorithms has made it difficult to interpret their output and understand how physiological mechanisms and anaesthetic states determine the algorithm output. A similar danger exists with the application of complex black-box machine-learning models in the operating room, where predictions are difficult to interpret and hence less actionable. Prescience demonstrates a solution that promises to avoid the obscurity traditionally associated with black-box models and instead maintain interpretability even as increasingly complex machine-learning models are applied to operating-room decision support.

It should be clarified that our exercise developing machine-learning methods to predict intraoperative hypoxaemia, although promising, should still be considered an initial attempt. In this first attempt, we did not categorize procedures to assess hypoxaemia predictions in specific types of procedures. For this reason, the clinical interpretation of the results had to be somewhat generic. For enhanced interpretation of risks, future attempts can focus on specific categories of cases and phases of anaesthesia. Another future enhancement would be the integration of additional preoperative data, such as the detailed medical history of a patient, into the prediction models. Higher-fidelity intraoperative data, such as patient monitor waveform data, could enrich machine learning, thus potentially leading to more accurate predictions. Prospective trials of Prescience during live procedures are also needed before deployment to verify the improvements in the performance of anaesthesiologists that we retrospectively observed in pre-recorded procedures.

This paper focuses on hypoxaemia risk during intraoperative anaesthesia care. However, the importance of coupling accurate predictions from complex models with interpretable explanations of why a prediction was made has broad applicability throughout medicine. To support this we have made the explanation tools initially used in Prescience open-source and have continued to improve and extend them. Because Prescience effectively decouples the interpretable explanation from the prediction model, we are also able to continue to refine the core prediction model without changing the user experience for anaesthesiologists.

The global risk profiles learned by Prescience (Figs. 5 and 6) are clinically relevant for a number of reasons. First, they show that in the health system examined, trauma hospital patients may be more critically ill as they have more intraoperative hypoxaemia. When considering the standardization of care to reduce unwanted clinical variation, these data suggest that resources may need to be differentially deployed to address differential rates of adverse events. Second, anaesthesiologists can now quantify risks of intraoperative hypoxaemia adjusted for other factors to the very elderly, those who are overweight and those with more co-morbid conditions. The exact relationships described in Fig. 6 clearly show the patterns and threshold points for the risk. Although low tidal volume is often recommended for patients with acute lung injury, these data suggest that low lung tidal volumes are, in fact, associated with intraoperative hypoxaemia. The relationship between low end-tidal CO$_2$ levels and intraoperative hypoxaemia may reflect underlying critical illness. Despite our inability to fully exclude residual confounding factors, these data shed new light on physiological relationships as well as provide a mechanism to facilitate the provision of anaesthesia care that can mitigate intraoperative hypoxaemia.
As a limitation, we acknowledge that there are several clinical diagnoses that are associated with hypoxaemia, but not directly observable in Prescience. The main clinical diagnoses include main-stem intubation, mucus plug, low FiO2, low tidal volume, tracheal tube balloon leak, and patient factors such as chronic obstructive pulmonary disease from smoking and pulmonary embolus. Among these, only low FiO2 and low tidal volume are directly observable in Prescience as the other data elements are not fully captured in the clinical databases. In these cases, secondary risk indicators will show up in Prescience. The differential diagnosis of hypoxaemia could also have been categorized using ACLS (advanced cardiovascular life support) strategies. However, contrary to a study where factors are a priori identified, Prescience considers all of the available factors and renders an output with associated relative risks. Clinicians must then evaluate the feature relevance on the basis of context and clinical relevance.

The field of medicine is full of data science challenges that have the potential to fundamentally impact the way medicine is practiced. More and more data-driven predictions of patient outcomes are being proposed and used. However, black-box prediction models, which simply provide predictions without explanation, are difficult for physicians to trust and provide little insight into how they should respond. The interpretable explanations used by Prescience represent a technique that can transform any current prediction method from one that provides what the prediction is into one that also explains why.

Methods

Institutional review board statement. The electronic data for this study was retrieved from institutional electronic medical record and data warehouse systems after receiving approval from the Institutional Review Board (University of Washington Human Subjects Division, Approval no. 46889). Protected health information was excluded from the dataset that was used for the machine-learning methods.

Data sources. Our hospital system has installed an AIMS (Merge AIM, Merge Inc.) that automatically captures minute-by-minute hemodynamic and ventilation parameters from the patient monitor and the anæsthesia machine. The system also integrates with other hospital electronic medical record systems to automatically acquire laboratory and patient registration information. The automatic capture of data is supplemented by the manual documentation of medications and anæsthesia interventions to complete the anæsthesia record during a surgical episode. For the current project, we extracted the high-fidelity anæsthesia data from the AIMS database from May 2012 to June 2014. The medical history data of each patient were also extracted from our electronic medical record data warehouse (Cardigem). The high-fidelity anæsthesia record data and the corresponding medical history data from the hospital electronic medical record formed the underlying data for machine learning. The various data elements used for machine learning are outlined in Supplementary Table 1.

SpO2 desaturation labels. We considered SpO2 ≤92% as hypoxaemia, which falls between the intervention level (≤94%) and emergency level (≤90%) recommended by the World Health Organization. Predictions of hypoxaemia were made for a window of five minutes into the future. If the SpO2 was ≤92% at any point during those 5 min, then it was considered a positive label, otherwise it was negative. The machine-learning algorithm was trained using these training labels on all time points where SpO2 was not already ≤92% at that time point.

When evaluating the performance of the machine-learning algorithm by comparing with anæsthesiologists (Fig. 3), we deliberately chose to use hypoxaemia events that were encountered after a period of stable and normal SpO2 (Supplementary Fig. 1). This was done to maximize the separation observed between the different prediction approaches and so minimize the number of time points anæsthesiologists needed to label. For a more generalized prediction of all low SpO2 values, the performance reported on the full test set using training labels should be used (Supplementary Fig. 3). The more stringent testing definition used for Fig. 3 excludes some time points, which leads to a smaller set of anæsthesiologist testing labels. Anæsthesiologist testing labels were positive only if SpO2 was ≥95% for the previous 10 min and then fell below 92% in the following 5 min (Supplementary Fig. 1; left). Anæsthesiologist testing labels were negative only if SpO2 remained ≥95% for the previous 10 min and the following 10 min (Supplementary Fig. 1; right). All other cases do not have anæsthesiologist testing labels. This more restrictive labelling scheme ensures that positive testing labels are clear drops in SpO2 levels that would be hard to predict in advance, whereas negative testing labels are clearly not drops in SpO2 (Supplementary Fig. 1).

An important point to consider when building labels for health-outcome prediction is that anæsthesiologist interventions can affect outcomes. It has been noted that models can learn when a doctor is likely to intervene and hence lower the risk of an otherwise high-risk patient.

This means that patients with low risk (from the model) may still need treatment. To address this, the authors of ref. 35 proposed removing examples from the training set where doctors have intervened. This allows one to train a model that predicts patient outcomes without the effect of doctor's interventions. In contrast, it is not possible to fully identify when and how anæsthesiologist is intervening (and if that intervention prevented hypoxaemia), so we sought to address this issue in two ways:

1. It must be recognized that the model predicts hypoxaemia when following standard procedures, not the occurrence of hypoxaemia if the anæsthesiologist takes no action to influence hypoxaemia. This is a natural assumption in the operating room where interventions that may affect SpO2 levels are performed frequently.

2. By focusing on clear explanations of why a certain risk was predicted, we enable anæsthesiologists to identify when the algorithm may be basing its risk on their actions versus when the risk is based on other factors.

Extracted time-series features. To make a prediction at an arbitrary point in time, a consistent set of extracted features should be computed that capture the information present in all previous time points. All of the data provided about a procedure are associated with a specific data and time. Text data have the time it was provided, minute-by-minute data from the patient monitor have the time at which each measurement was taken and single point measurements have the times at which they were recorded.

We summarized these unevenly sampled time-registered data into a fixed-length feature vector at any point in time using several complementary methods:

- Patient data, procedure information and pre-operative notes are represented by a ‘last value’ extracted feature, which is zero before any data is recorded and the value of the data afterwards.
- Time-series data are captured using exponentially decaying weighted average and variance estimates using multiple decay rates. These decay rates specify how much impact each past time point has on the computed mean or variance for the time series. We used 6, 1 min and 5 min half-life times to capture both high and low frequency components of the signal in each time series (Supplementary Fig. 12).
- Drug dose data are captured using both an exponentially decaying sum and a time since the last measurement. Decay rates with half-lives of 5 min and 1 h were used to capture both near-term and longer-term average drug dosing effects.

To ensure that there was enough training data for each extracted feature, we removed extracted features that had fewer than 100 recorded data values for the real-time model and fewer than 50 for the initial model. For a full list of the 3,797 extracted features used by Prescience for initial predictions see Supplementary Table 4. For the 3,905 extracted features used in intraoperative predictions see Supplementary Table 5. Note that more than 2,000 of the initial and intraoperative features represent words from text data sources.

Gradient-boosting machines for prediction. The extracted features we compute from real-time operating room data have a variety of complex nonlinear interactions. Capturing these requires a model with significant flexibility and we chose a non-parametric approach called gradient-boosting machines

We compared the performance of gradient boosting against three baseline methods: Lasso penalized linear logistic regression; a linear SVM autoregressive model that was proposed for predicting hypoxaemia based only on the SpO2 data stream in an earlier report 11 and an unsupervised Parzen window method that was used in an earlier report to predict patient deterioration.15 Gradient-boosting machines significantly outperformed all baseline methods for our primary endpoint, real-time hypoxaemia prediction (Supplementary Fig. 3). For our secondary task of initial prediction, gradient-boosting machines were only slightly superior (Supplementary Fig. 2). The large performance gain of gradient-boosting for intraoperative prediction (Supplementary Fig. 3) is probably because there are 8 million training samples, whereas for preoperative predictions (Supplementary Fig. 2) there are only 42,000 samples and no time-series data. Note that for the initial prediction, the autoregressive support vector machine (SVM) and Parzen window methods were not applicable and hence not evaluated.

Gradient-boosting machines are non-parametric models that draw a parallel between boosting and gradient descent in function space. They additively build up simple models, like boosting, and use a variable selection procedure to find the most useful additional features. A gradient-boosting machine is trained by repeatedly fitting a simple model to the residual error of the previous model. In practice, this method is equivalent to minimizing a loss function with the gradient of the loss at which they were recorded.

A vector machine (SVM) and Parzen window methods were not applicable and these methods: Lasso penalized linear logistic regression; a linear SVM autoregressive model that was proposed for predicting hypoxaemia based only on the SpO2 data stream in an earlier report and an unsupervised Parzen window method that was used in an earlier report to predict patient deterioration. Gradient-boosting machines significantly outperformed all baseline methods for our primary endpoint, real-time hypoxaemia prediction (Supplementary Fig. 3). For our secondary task of initial prediction, gradient-boosting machines were only slightly superior (Supplementary Fig. 2). The large performance gain of gradient-boosting for intraoperative prediction (Supplementary Fig. 3) is probably because there are 8 million training samples, whereas for preoperative predictions (Supplementary Fig. 2) there are only 42,000 samples and no time-series data. Note that for the initial prediction, the autoregressive support vector machine (SVM) and Parzen window methods were not applicable and hence not evaluated.

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Gradient-boosting machines are non-parametric models that draw a parallel between boosting and gradient descent in function space. They additively build up simple models, like boosting, and these models are fit to the gradient of the loss at which they were recorded.
Fitting the trees is computationally challenging for large datasets, so we used XGBoost, a high performance implementation of gradient-boosting machines\(^5\). For the real-time model we used \(\eta=0.2\) and 1,242 trees, whereas for the initial model we chose \(\eta=0.1\) and 4,000 trees. Using a smaller \(\eta\) value means that more trees are required for fitting, which requires more time to run but results in a smoother (and generally better) model. For both the initial and real-time models we used bagging, where trees were trained on a random 50% sub-sample of the training data. For the preoperative model, the maximum tree depth was four and the minimum child weight of any branch in the trees was one. For the real-time model, the maximum tree depth was six and the minimum child weight of any branch in the trees was ten.

All method parameters were tuned (and methods were chosen) using a validation set of operating room procedures separate from the final test set used for all final performance results. To ensure that there was no bias towards the final test set, the test data were initially compressed and left uncompressed until after method development was completed.

**Computing feature-importance estimates.** Understanding why a statistical model has made a specific prediction is a key challenge in machine learning. It engenders appropriate trust in predictions and provides insight into how a model may be improved. However, many complex models with excellent accuracy, such as gradient boosting, make predictions that even experts struggle to interpret. This forces a trade-off between accuracy and interpretability. In response to this we chose to use a model agnostic measure of feature importance, where the impact of each feature on the model is represented using Shapley values\(^9,10\), which have been shown to be the only way to assign feature importance while maintaining two important properties, local accuracy and consistency (defined below)\(^9\). The application of these values in Prescience uses fast estimation methods that we have developed to compute the Shapley values (that is, the estimated importance of features for a particular prediction) in real-time\(^9,10\).

Shapley values are from the game theory literature and provide a theoretically justified method for allocation of a coalition’s output among the members of the coalition (see equation 1). In Prescience, the coalition is a set of interpretable model input feature values and the output of the coalition is the value of the prediction made by the model when given those input feature values. Feature local impact is defined as the change in the expected value of the model’s output when a feature is observed versus unknown. Some feature values have a large impact on the prediction, whereas others have a small impact. The Shapley values \(\phi_i(f, x)\), explaining a prediction \(f(x)\), are an allocation of credit among the various features in \(x\) (such as age, weight, time-series features and so on) and are the only such allocation that obeys a set of desirable properties. Note that \(\phi_i(f, x)\) is a single numerical value representing the impact of feature \(i\) on the prediction of the model \(f\) when given the input \(x\). For Prescience \(f\) is a gradient-boosting model and \(x\) is the set of all input features from a time point. We provide a brief summary of these properties below and refer the reader to ref. 10 for a full discussion and for connections with several other recent methods in complex model interpretability. In the properties below \(f_S(f)\) is \(f([x]_S)\) for any subset of features with \(|S| \\leq m\), where \(x_S\) is the subset of the input vector with only the features in the set \(S\) present.

**Local accuracy.** The local accuracy property (also known as completeness or additivity) is given by the following equality

\[
\phi_i(f, x) = \phi_i(f, x_S) + \sum_{S \subseteq [m]} \phi_i(f, x_S) - \phi_i(f, x)
\]

where \(\phi_i(f, x_S) = \mathbb{E}[f([x]_S)]\) (the expected value of the model over the training dataset) and \(M\) is the number of ‘interparable’ inputs, each which correspond to a group of original input features (such as those shown in Fig. 4). The local accuracy assumption forces the attribution values to correctly capture the difference between the expected model output and the output for the current prediction. For Prescience, the input feature groups are the sets of extracted features associated with each time series. For instance, the 6 s, 1 min and 5 min moving average extracted features and the 5 min moving variance extracted feature from the SpO\(_2\) time series are all considered as a single group. This manual grouping process is not strictly necessary but can help improve the interpretation of partially redundant features.

**Consistency.** For any two models \(f\) and \(f'\), if

\[
f'(S) \leq f(S) \leq f(S | \{i\}) + f(S | \{i\}) - f(S| \{i\})
\]

for all \(S \subseteq [2|/l|\) where \(Z\) is the set of all \(M\) input features, then \(\phi_i(f, x) \geq \phi_i(f, x)\). This states that if a feature is more important in one model than another, no matter what other features are also present, then the importance attributed to that feature should also be higher. Note that consistency is known as monotonicity in game theory literature.

Only one allocation of credit satisfies these two properties (and also trivial assumptions about unused model inputs) and that allocation is the one given by the Shapley values\(^9,10\).

Given a specific prediction \(f(x)\), we can compute the Shapley values using a weighted sum that represents the impact of each feature being added to the model averaged over all possible orders of features being introduced:

\[
\phi_i(f, x) = \sum_{S \subseteq [m]} \frac{|S|!(M-|S|-1)!}{M!} \left( f(S) - \left( f(S \cup \{i\}) - f(S) \right) \right)
\]

\[
= \sum_{S \subseteq [m]} \frac{1}{M \text{ choose } |S| \{i\}} \left( f(S) - \left( f(S \cup \{i\}) - f(S) \right) \right)
\]

(1)

In practice, there are far too many terms to evaluate this sum completely, so we can instead approximate it by a sampling procedure\(^9,10\). We have released an open implementation of this explanation approach which also includes additional improvements for tree models (developed after Prescience) at: http://github.com/slundberg/shap.

To compute the Shapley values of each prediction, we need to estimate the predictions of the model when specific input features are missing (those not in the set \(S\)). Given that the model was not trained to support missing values, we approximate what the model would predict (if retrained on that subset of input features) by sampling from the training dataset and replacing the missing features with the values they would have had in that sample. By repeating this process on each sample, we can estimate the expected value of \(f(S)\) only using evaluations of \(f(S\_\{S\})\) where no features are missing.

The approach above requires nested sampling, once to estimate the Shapley value and then from each sample we again sample to estimate \(f(S)\) and \(f(S \cup \{i\})\). To reduce the number of samples in the inner step, we used 4\(^1\)medians to generate 20 medians of the entire dataset and then performed a weighted evaluation for only these 20 summary inputs as an approximation for the entire dataset. This removes the need for nested sampling.

In Prescience we also used a nonlinear link function \(h\) such that:

\[
h(x) = \sum_{x \in x_S} \phi_i(f, x)
\]

As Prescience uses logistic regression, the use of a \(h = \logit\) link function transforms the output space from probabilities to log odds. Assuming the importance of features is additive in the log-odds space is much more natural than assuming they are additive in the space of probabilities (which must fall between 0 and 1). The same reasoning also drives the use of the logit link function during standard logistic regression.

We were able to get stable feature-importance estimates for thousands of features in less than \(5\) s on our server (in large part because these inputs typically had fewer than \(100\) non-zero entries). We compared these theoretically grounded explanations with a simple estimation of feature importance to verify that they showed reasonable agreement. The simple method we chose was to replace a single feature group with random values from other samples in the dataset and determine the average model output over different possible samplings. We then subtracted this mean value from the original model prediction to get a difference from a prediction with a typical value of that feature versus the current value. This method is not very scalable and does not account for interactions with other features, yet it is useful to compare with the Prescience explanations to ensure that the Prescience estimates of feature effects are consistent with an intuition of how much a feature’s change from its typical value affects the current risk of hypoxaemia (Supplementary Fig. 4).

**Physician evaluation.** The potential benefit Prescience provides to physicians was evaluated using previously recorded procedures. Both before a procedure begins and at several time points during the operation, all of the available electronically recorded data were shown to the anaesthesiologist and they were asked to predict if a desaturation (as defined above) will occur in the next 5 min (Supplementary Figs. 5–7). For half of the procedures, anaesthesiologists are given Prescience explained risks (Supplementary Figs. 5 and 6) and for the other half they are given the same data but without any Prescience assistance (Supplementary Fig. 7).

In both cases anaesthesiologists are asked to provide a fold change in the risk that desaturation will occur.

The test procedures were divided into two equal sized groups, replicate 1 and replicate 2. Anaesthesiologists were also divided into two groups, A and B. Group A was given Prescience assistance on replicate 1 but not on replicate 2, whereas group B was given Prescience assistance on replicate 2 but not replicate 1. After randomly assigning anaesthesiologists to the groups, three anaesthesiologists from group A and two anaesthesiologists from group B completed the evaluation. We pooled the results within each group and between groups, and the results of this evaluation are shown in Fig. 3. The order in which anaesthesiologists were presented with cases was random across both replicate sets.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Code availability.** The model-explanation code originally used, and subsequently improved, for Prescience is available open-source at https://github.com/slundberg/shap.
shap. Modelling, processing and web-interface codes specific to Prescience are available for reference purposes at https://gitlab.cs.washington.edu/prescience.

Data availability
Owing to patient-privacy considerations, the operating-room datasets from participating hospitals are not publicly available. The raw data from the anaesthesiologist comparisons in Fig. 3 are available in Supplementary Tables 6 and 7, and data from Fig. 5 are available in Supplementary Tables 8 and 9.

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Author contributions
S.-I.L., S.M.L., B.N. and J.K. initiated the study. S.-I.L. and S.M.L. developed the Prescience algorithms and designed data analyses and experiments. S.M.L. performed data analyses, experiments and data preprocessing. B.N. and S.-F.N. provided the electronic medical record data. J.K. recruited anaesthesiologists and helped design the anaesthesiologist test and survey. M.H., M.J.E., T.A., D.E.L. and D.K.-W.L. performed the web-based anaesthesiologist experiments and provided survey data. M.S.V provided clinical assessment, interpretation of feature importance and connections with anaesthesiologists’ workflow. S.-I.L. and S.M.L. wrote the paper in conjunction with B.N., J.K. and M.S.V. who wrote the sections on clinical interpretation and integration with current practices. M.H. provided manuscript feedback.

Competing interests
B.N. is an advisor for Perimetrics LLC and holds equity in the company. D.K.-W.L. is a Chief Medical Officer for MDmetrix, Inc. The other authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41551-018-0304-0. Reprints and permissions information is available at www.nature.com/reprints. Correspondence and requests for materials should be addressed to S.-I.L.

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Experimental design

1. Sample size

Describe how sample size was determined.

For software training we took all surgical records from two university-affiliated hospitals over the course of two years. For software testing we used 5 anaesthesiologists, chosen because they were willing to participate. Our criterium was to include at least 4 anaesthesiologists, to provide replicate results across two groups.

2. Data exclusions

Describe any data exclusions.

We excluded surgical cases with a heart transplant, lung transplant, tracheostomy, and coronary-artery bypass surgeries in which SpO2 and other hemodynamic parameters can be significantly affected by non-physiological measurements such as during cardiopulmonary bypass.

3. Replication

Describe whether the experimental findings were reliably reproduced.

For human test results we observed consistent patterns across anaesthesiologists, as shown in the manuscript. For learning task evaluations we used a held-out test set with thousands of samples from surgery cases that were not used during training, implying strong statistical significance.

4. Randomization

Describe how samples/organisms/participants were allocated into experimental groups.

Anaesthesiologists were randomized between the two comparison groups. Events used to test anaesthesiologists were randomly selected from the held-out test dataset within criteria described in the manuscript. The training and validation sets were split randomly by surgical procedure.

5. Blinding

Describe whether the investigators were blinded to group allocation during data collection and/or analysis.

Anaesthesiologists were blind to their group assignment. Scott Lundberg and Su-In Lee were aware of the group allocation. All investigators were blind to the test set during method development since the test set was randomly divided when the data was first delivered and left in a compressed ZIP file until after algorithm development was frozen.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.
6. Statistical parameters

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

| n/a | Confirmed |
|-----|-----------|
| ✔   | The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.) |
| ✔   | A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly |
| ✔   | A statement indicating how many times each experiment was replicated |
| ✔   | The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section) |
| ✔   | A description of any assumptions or corrections, such as an adjustment for multiple comparisons |
| ✔   | The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted |
| ✔   | A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range) |
| ✔   | Clearly defined error bars |

See the web collection on statistics for biologists for further resources and guidance.

7. Software

Describe the software used to analyze the data in this study.

The algorithms used in this work used custom data-processing methods written in Python and Julia. XGBoost, Scikit-learn and related packages were used during implementation. Code specific to the paper’s experiments is available at https://gitlab.cs.washington.edu/prescience (primarily for reference, since the original patient data is not publicly available), and the more broadly applicable explanation methods originally developed for Prescience are available at https://github.com/slundberg/shap

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

8. Materials and reagents

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

No unique materials were used.

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

No antibodies were used.

a. State the source of each eukaryotic cell line used.

No cell lines were used.

b. Describe the method of cell line authentication used.

No cell lines were used.

c. Report whether the cell lines were tested for mycoplasma contamination.

No cell lines were used.

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

No cell lines were used.
Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
   Provide details on animals and/or animal-derived materials used in the study.
   No animals were used.

Policy information about studies involving human research participants

12. Description of human research participants
   Describe the covariate-relevant population characteristics of the human research participants.
   Demographic data is provided in Fig. 2 of the paper.