Step-by-step causal analysis of EHRs to ground decision-making

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Abstract

How to inform clinical decision-making using routine care data? First, individualizing care calls for tailored prediction, eg with machine learning. Second, predictions must be associated with actionable interventions to avoid shortcuts or biases in the data. For a given treatment, a useful model would distinguish responders from non-responders. This requires modeling heterogeneity in causal inference, which faces many pitfalls, particularly so on time-varying data, as in

Causal inference enables machine learning methods to estimate treatment effects of medical interventions from electronic health records (EHRs) or claims. Here, we decompose these pitfalls in a three steps retrospective case-control study: choice of study design, confounding variables, and estimator. Studying. However, researchers should be wary of many pitfalls. Here we explain and study common pitfalls and potential solutions, estimating the effect of albumin on mortality in sepsis in the Medical Information Mart for using an Intensive Care database (MIMIC-IV), we and comparing to outcome data from randomized controlled trials (RCT) as gold-standard. The first step is study design, using the target trial concept and the PICOT framework: Population (patients with sepsis), Intervention (combination of crystalloids and albumin for fluid resuscitation), Control (crystalloids only), Outcome (28-day mortality), Time (intervention start within 24h of admission). We show that a too large treatment-initiation time induces immortal time bias. The second step is selection of the confounding variables based on expert knowledge. Increasingly adding confounders enables to recover the RCT results from observational data (0%, 95% CI -1% to 1%, and 0%, 95% -5% to 5%, respectively). The third step is fitting outcome models, and we assess the influence of multiple models with varying assumptions. A doubly robust estimator (AIPW) with random forests proved to be the most reliable estimator. Results show that these steps are all important to build valid decision-making. In a tutorial spirit, we make all the code and data openly available for valid causal estimates. A valid causal model can then be used to

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individualize decision making: subgroup analyses showed that treatment efficacy of albumin was better for patients >60 years old, males, and patients with septic shock.

Author summary

How to inform clinical decision-making using routine care data? First, individualizing care calls for tailored prediction, eg with machine learning. Second, predictions must be associated with actionable interventions to avoid shortcuts or biases in the data . For a given treatment, a useful model would distinguish responders from non-responders. This requires modeling heterogeneity in causal inference, which faces many—

Rich routine-care data, as EHR or claims, is useful to individualize decision making using machine learning; but guiding interventions requires causal inference. Unlike with an RCT, interventions in routine data do not easily enable an apple-to-apple measure of the effect of an intervention, leading to many analytical pitfalls, particularly so on in time-varying data, as in electronic health records (EHRs) or claims. Here, we decompose these pitfalls in a three steps retrospective case-control study: choice of study design, confounding variables, and estimator. Studying the effect of albumin on mortality in sepsis in the Medical Information Mart for Intensive Care database (MIMIC-IV). We study these in a tutorial spirit, making the code and data openly available. We give 5 analytical steps for data-driven individualized interventions: Step 1) Study design, where common pitfalls are selection bias, with information unequally collected across treatment and control patients, and immortal time bias, where the inclusion-defining event interacts with the intervention time. Step 2) Identification of the causal assumptions and categorization of confounders. Step 3) Estimation of the causal effect of interest by correct aggregation of confounders and selection of an appropriate statistical model. Step 4) Assessing the analysis' robustness to assumptions, and finally Step 5) Individualizing treatment decision, by exploring treatment heterogeneity, eg across subgroups. Studying choice of fluid resuscitation in sepsis, we show that these steps are all important to build valid decision-making. In a tutorial spirit, we make all the code and data openly available common mistakes in steps 1, 2, and 3 equally compromise causal validity.

Introduction: data-driven decisions require causal inference

How can data lead to useful clinical decisions? Prevention strategies using well-validated and calibrated cardiovascular risk models have not reduced the burden of cardiovascular diseases [1], probably because they targeted both responders and non-responders [2]. Addressing treatment heterogeneity is crucial to effective careInforming a care option extends beyond merely predicting the occurrence of an event; it involves estimating the effect of the corresponding treatment effects. Routine-care data comes naturally to mind to guide routine decisions, but they require care to estimate treatment effects as they are observational, unlike Randomized controlled trials (RCTs). This context calls for causal inference statistical frameworks. But merely applying these tools to the data does suffice to ensure the validity of the inferences; numerous considerations must be carefully addressed.

Individualized predictions are not enough Medicine and Machine Learning Challenges Machine learning is central to individualized medicine. Machine learning models recently outperformed plays a pivotal role in individualized medicine

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[3–7]. It demonstrated superior performance over traditional rule-based clinical scores to predict in predicting a patient's readmission risk, mortality, or future comorbidities using Electronic Health Records (EHRs) [3–7]. But there is growing evidence [3–7]. However, mounting evidence suggests that machine-learning models can reproduce and amplify biases inadvertently perpetuate and exacerbate biases present in the data [8], such as including gender or racial biases [9,10], or and the marginalization of under-served populations [11]. The models typically encode these biases These biases are typically encoded by capturing shortcuts:—stereotypical or distorted features in the data [12–14]. For instance, many numerous machine learning algorithms use rely on post-treatment information [15–18]; a paradigmatic example being, exemplified by a diagnostic model for skin cancer relying that depends on surgical marks [13]. A motivating example on Intense For Intensive Care Unit datais provided in, focus of our study, such information markedly improves mortality prediction (Figure S1 5), but cannot inform decisions.

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The Importance of Causal thinking is key to data-driven decision-making Reasoning in Data-Driven Decision-Making [19] While traditional conventional machine learning relies on data resulting from prior interventions retrospective to generate predictions of future effects [20], truly insightful informing decision-making support needs to contrast the needs a comparison of potential outcomes with and without the intervention. This involves estimating a causal effectas in Randomized Control Trials (RCTs) [19]. Yet, RCTs face , mirroring the methodology employed in RCTs [19]. However, RCTs encounter challenges such as selection biases [21,22], failure to recruit minorities, or limited sample size difficulties in recruiting diverse populations, and limited sample sizes for exploring treatment heterogeneity across subgroups. Routinely collected data naturally provides presents a unique opportunity to assess real-life benefit-risk trade-offs associated with a decision [23], with less sampling bias, reduced sampling bias and sufficient data to capture heterogeneity [24]. Estimating Nevertheless, estimating causal effects from this such data is challenging however, as the intervention is confounded due to the confounding of the intervention by indication. Hence Therefore, dedicated statistical techniques are needed imperative to emulate a "target trial" [25] from observational data.

Grounding clinical decisions on data requires considerations scattered across different fields. How lack of individual treatment model limits a cardiovascular risk score The QRISK is a risk score widely used in the UK to assess 10-year and lifetime cardio-vascular risk [26]. It is well calibrated, recommended by the national regulatory agency [27], and the National Institute for Health and Care Excellence (NICE) argues to use such a score to systematically screen cardio-vascular disease (CVD) risks and treat with statins individuals at risk [27], especially if 10-year CVD risk exceeds 10% [28]. A cost-effectiveness analysis guided the choice of the 10% threshold to optimize quality-adjusted life years at the population level [29]. The NICE analysis is based on evidence mostly from three trials, accounting for more than 70% of the effect: LIPID [30] (32% of the effect). PROSPER [31] (22% of the effect), ALLHAT-LLT [32] (23% of the effect). All these trials focused on high-risk patients and secondary prevention. All studies operate under the assumption of a consistent statin effect (risk ratios constant) across different CVD risk groups, irrespective of the baseline CVD risk and LDL-cholesterol levels. This aligns with the outcomes of two meta-analyses which observed broadly similar effectiveness across risk levels [33], but is challenged by another study in primary prevention [34]. Meanwhile the US Preventive Services Task Force concluded that the

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evidence is insufficient to determine the balance of benefits and harms of statin use for the primary prevention of CVD events and mortality in adults 76 years and older [35].

Epidemiologic literature emphasizes the

Critics on the efficiency of systematic screening and treatment heterogeneity Conversely, the World Health Organization (WHO) published a systematic review on screening for CVD risk in 2019 [1] asserting the limited effectiveness of such screenings, aligning with a Cochrane review that also reported poor effectiveness [36] resulting in over-treatment. The surge in statin prescriptions has been linked to a divergence in statin prescriptions at specified risk score levels [37]. This leads to over-prescription for low-risk patients and under-prescription for high-risk patients, casting doubt on the overall efficacy as the constancy assumption of the statin treatment effect might be violated. Accounting for sources of heterogeneity in patient profiles would be instrumental in optimizing treatment allocation.

Moreover, the WHO report critiques suboptimal adoption leading to social biases in the screened populations. In the UK, there are instances of treatment heterogeneity at a given risk level [37] and documented evidence of social inequities in health checks [2]. Evidence suggests that predominantly socially advantaged patients undergo screening, consequently being offered statin treatment. This critique further emphasizes the presence of treatment heterogeneity.

Multiple Perspectives on Evidence-based Decision Making Across different fields, existing literature has emphasized different challenges associated with estimating treatment effects using observational data. While epidemiologic studies underscore the importance of the target trial approach [38-42], but focuses on biases due to confounding variables and less so on those due to poor designs or [38-42], there emphasis primarily lies on biases that arise from temporal effects [25, 43-47] or confounding variables [48-50], with relatively less attention to issues arising from estimator selection. Machine Recent replications of RCTs using observational data did not explore the impact of modern machine learning methods on the robustness of the results [41,51].

In contrast, machine learning and causal inference literature focuses on estimators [52–56] predominantly studies estimators [52–56]: propensity score matching [57], inverse probability weighting [58], outcome models [59], doubly robust methods, [53] or deep learning based models [60]. This literature may be opaque for some due to intricate mathematical details and unverifiable assumptions. Guidelines seldom mention address time-related biases, frequent loopholes when data have or covariate aggregation which frequently emerge in datasets with temporal dependencies [43, 45]. Here, putting all steps—choice of study design, confounders, estimator—in a consistent framework (section Recently, the machine learning community shifted its focus from EHR data to simulated data, which may not capture the complexities of real-world data [61–64].

In this work, we bring together epidemiological concepts and principles from statistical and machine learning literature. We adopt an empirical perspective to answer practical needs of applied researchers. A study of choices spread out across the analysis –study design, consideration of confounders, and selection of estimators (refer to Section Step-by-step framework for robust decision-making from EHR data), we show they are equally important to ensure validity of the analysis. We compare the corresponding biasesby studying the effect – highlights their equal importance in ensuring the validity of results. To illustrate and compare biases, we investigate the impact of albumin on sepsis mortality in using data from a publicly available intensive

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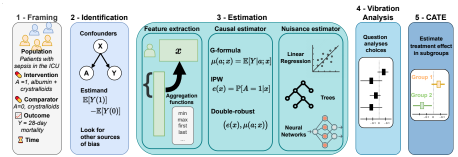


Fig 1. Step-by-step analytic framework – The complete inference pipeline confronts the analyst with many choices, some guided by domain knowledge, others by data insights. Making those choices explicit is necessary to ensure robustness and reproducibility.

care database(, MIMIC-IV)— [65] (section Application: evidence from MIMIC-IV on which resuscitation fluid to use). The main section's focus is on being accessible, while the appendices expand technical details.

The primary focus of the main section is on accessibility, with technical details expanded in the appendices.

Step-by-step framework for robust decision-making from EHR data

Step-by-step framework for robust decision-making from EHR data

Whether or not using machine learning, many pitfalls threaten an analysis' value for decision-making. To avoid these pitfalls, we outline a simple step-by-step analytic framework illustrated in Figure 1 for retrospective case-control studies. We frame the medical question as a target trial [66] to match the design to an RCT giving the gold standard average effect. Then we probe for heterogeneity –predictions on sub-groups-going beyond what RCTs can achieve.

Step 1: study design – Frame the question to avoid biases

Grounding decisions on evidence needs well-framed questions, defined by their PICO(T) components. Population, Intervention, Control, and Outcome [67,68], and in case of EHRs or claims data an additional time component, are necessary to concord with a (hypothetical) target randomized clinical trial [51,69] – Table 1.

Without care in defining these PICO(T) components, non-causal associations between treatment and outcomes can easily be introduced into an analysis [70]. The time-varying nature of EHR calls for checking systematically of the Population and Time components by addressing two commonly encountered types of bias.

PICO component	Description	Notation	Example
Population	What is the target population of interest?	$X \sim \mathbb{P}(X)$, the covariate distribution	Patients with sepsis in the ICU
Intervention	What is the treatment?	$A \sim \mathbb{P}(A=1) = p_A,$ the probability to be treated	Combination of crystalloids and albumin
Control	What is the clinically relevant comparator?	$1 - A \sim 1 - p_A$	Crystalloids only
Outcome	What are the outcomes to compare?	$Y(1), Y(0) \sim \mathbb{P}(Y(1), Y(0)),$ the potential outcomes distribution	28-day mortality
Time	Is the start of follow-up aligned with intervention assignment?	N/A	Intervention within the first day

Table 1. PICO(T) components help to clearly define the medical question of interest.

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Selection Bias: In EHRs, outcomes and treatments are often not directly available and need to be inferred from indirect events. These signals could be missing not-at random, sometimes correlated with the treatment allocation [71]. For example, billing codes can be strongly associated with case-severity and cost. Consider comparing the effectiveness of fluid resuscitation with albumin to crystalloids. As albumin is more costly, this treatment is more likely to have a sepsis billing code. On the contrary, for patients treated with crystalloids, only the most severe cases will have a billing code. Naively comparing patients would overestimate the effect of albumin.

Immortal time bias: Improper alignment of the inclusion defining event and the intervention time is a major source of bias in time-varying data [25, 43, 46]. Immortal time bias (illustrated in Appendix 6) occurs when the follow-up period, i.e. cohort entry, starts before the intervention, e.g. prescription for a second-line treatment. In this case, the treated group will be biased towards patients still alive at the time of assignment and thus overestimating the effect size. Other frequent temporal biases are lead time bias [44, 45] or right censorship [25], and attrition bias [47]. Good practices include explicitly stating the cohort inclusion event [72, Chapter 10:Defining Cohorts] and defining an appropriate grace period between starting time and the intervention assignment [25]. At this step, a population timeline can help.

Step 2: identification – List necessary information to answer the causal question

The identification step builds a causal model to answer the research question. Indeed, the analysis must compensate for differences between treated and non-treated that are not due to the intervention ([73, chapter 1], [40, chapter 1]).

Causal Assumptions Valid causal inference requires assumptions [74]—detailed in S2 Appendix. The analyst should thus review the plausibility of the following: 1) Unconfoundedness: after adjusting for the confounders as ascertained by domain expert insight, treatment allocation should be random; 2) Overlap—also called positivity—the distribution of confounding variables overlaps between the treated and controls—this is the only assumption testable from data [58]—; 3) No interference between units and consistency in the treatment, a reasonable assumption in most clinical questions.

Categorizing covariates Potential predictors –covariates – should be categorized depending on their causal relations with the intervention and the outcome (illustrated in S4 Fig): confounders are common causes of the intervention and the outcome; colliders are caused by both the intervention and the outcome; instrumental variables are a cause of the intervention but not the outcome, mediators are caused by the intervention and is a cause of the outcome. Finally, effect modifiers interact with the treatment, and thus modulate the treatment effect in subpopulations [75].

To capture a valid causal effect, the analysis should only include confounders and possible treatment-effect modifiers to study the resulting heterogeneity. Regressing the outcome on instrumental and post-treatment variables (colliders and mediators) will lead to biased causal estimates [49]. Drawing causal Directed Acyclic Graphs (DAGs) [48], eg with a webtool such as DAGitty [76], helps capturing the relevant variables and defining a suitable estimand or effect measure. The estimand is the final statistical quantity estimated from the data. Depending on the question, different estimands are better suited to contrast the two potential outcomes E[Y(1)] and E[Y(0)] [77, 78]. For continuous outcomes, risk difference is a natural estimand, while for binary outcomes (e.g. events) the choice of estimand depends on the scale. Whereas

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the risk difference is very informative at the population level, e.g. for medico-economic decision-making, the risk ratio and the hazard ratio are more informative at the level of sub-groups or individuals [78].

Step 3: Estimation – Compute the causal effect of interest

Confounder aggregation Confounders captured via measures collected over multiple time points must be aggregated at the patient level. Simple forms of aggregation include taking the first or last value before a time point, or an aggregate such as mean or median over time. More elaborate choices may rely on hourly aggregations providing more detailed information on the disease course such as vital signs. They may reduce confounding bias between rapidly deteriorating and stable patients but also increase the number of confounders making estimation more challenging [79]. The increase of variance occurs either in arbitrarily small propensity scores for treatment models or in hazardous extrapolation from one group to another for outcome model. If multiple choices appear reasonable, one should compare them in a vibration analysis (see Step 4: Vibration analysis – Assess the robustness of the hypotheses).

Beyond tabular data, unstructured clinical text may capture confounding or prognostic information [80,81] which can be added in the causal model [42]. However, high-dimensional confounder space such as text may break the positivity assumption just as hourly aggregation choices for measurements.

Causal estimators or statistical modeling A given estimand can be estimated through different methods. One can model the outcome with regression models also known as G-formula, [59] and use it as a predictive counterfactual model for all possible treatments for a given patient. Alternatively, one can model the propensity of being treated use it for matching or Inverse Propensity Weighting (IPW) [58]. Finally, doubly robust methods model both the outcome and the treatment, benefiting from the convergence of both models [82]. There is a variety of doubly robust models, reviewed in S3 Appendix.

Estimation models of outcome and treatment The causal estimators use models of the outcome or the treatment –called nuisances. There is currently no clear best practice to choose the corresponding statistical model [62,83]. The trade-off lies between simple models risking misspecification of the nuisance parameters versus flexible models risking to overfit the data at small sample sizes. Stacking models of different complexity as in a super-learner is a good solution to navigate the trade-off [84,85].

Step 4: Vibration analysis – Assess the robustness of the hypotheses

Some choices in the pipeline may not be clear cut. Several options should then be explored, to derive conceptual error bars going beyond a single statistical model. When quantifying the bias from unobserved confounders, this process is sometimes called sensitivity analysis [86–88]. Following [89], we use the term vibration analysis to describe the sensitivity of the results to all analytic choices.

Step 5: Treatment heterogeneity – Compute treatment effects on subpopulations

Once the causal design and corresponding estimators are established, they can be used to explore the variation of treatment effects among subgroups. A causally-grounded

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model can be used to predict the effect of the treatment from all the covariates –confounders and effect modifiers– the *Conditional Average Treatment Effect* (CATE) [90]. Practically, CATEs can be estimated by regressing an individual's predictions given by the causal estimator against the sources of heterogeneity (details in S8 Appendix).

Application: evidence from MIMIC-IV on which resuscitation fluid to use

We now use the above framework to extract evidence-based decision rules for resuscitation. Ensuring optimal organ perfusion in patients with septic shock requires resuscitation by reestablishing circulatory volume with intravenous fluids. While crystalloids are readily available, inexpensive and safe, a large fraction of the administered volume is not retained in the vasculature. Colloids offer the theoretical benefit of retaining more volume, but might be more costly and have adverse effects [91]. Meta-analyses from multiple pivotal RCTs found no effect of adding albumin to crystalloids [92, 93] on 28-day and 90-day mortality. Given this previous evidence, we thus expect no average effect of albumin on mortality in sepsis patients. However, studies –RCT [94] and observational [95]– have found that septic-shock patients do benefit from albumin.

Emulated trial: Effect of albumin in combination with crystalloids compared to crystalloids alone on 28-day mortality in patients with sepsis Multiple published RCTs can validate the analysis pipeline before investigating sub-population effects for individualized decisions. Using MIMIC-IV [65], we compare the magnitude of biases introduced by reasonable choices in the different analytical steps recalled in Figure 2.

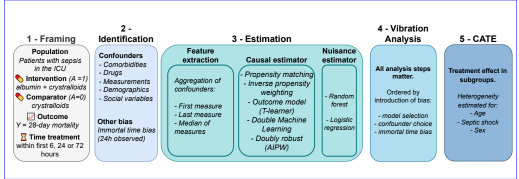


Fig 2. Application of the step-by-step framework on which resuscitation fluid to use.

Study design: effect of crystalloids on mortality in sepsis

- Population: Patients with sepsis in an ICU stay according to the sepsis-3 definition. Other inclusion criteria: sufficient follow-up of at least 24 hours, and age over 18 years described in Appendix 6.
- Intervention: Treatment with a combination of crystalloids and albumin during the first 24 hours of an ICU stay.

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• Control: Treatment with crystalloids only in the first 24 hours of an ICU stay.

- Outcome: 28-day mortality.
- Time: Follow-up begins after the first administration of crystalloids. Thus, we potentially introduce a small immortal time bias by allowing a time gap between follow-up and the start of the albumin treatment –see the full timeline in S3 Fig. Because we are only considering the first 24 hours of an ICU stay, we hypothesize that this gap is insufficient to affect our results. We test this hypothesis in the vibration analysis step.

In MIMIC-IV, these inclusion criteria yield 18,121 patients of which 3,559 were treated with a combination of crystalloids and albumin(. While glycopeptide antibiotic therapy was similar between both groups (51.8% crystalloid vs 51.5% crystalloids + albumin), aminoglycosides, carbapenems, and beta-lactams were more frequent in the crystalloid only group (2.0% vs. 0.7%, 4.3% vs. 2.6%, and 35.5% vs. 13.8%, respectively). The crystalloid only group was more frequently admitted as an emergency (57.3% vs. 30.7%). Vasopressors (80.2% vs 41.7%) and ventilation (96.8% vs 87.0%) were more prevalent in the treated populations, underlying the overall higher severity of patients receiving albumin (mean SOFA at admission 6.9 vs. 5.7). Table 2 details patient characteristics and S5 Fig details the selection flowchart).

	Missing	Overall	Cristalloids only	Cristalloids + Albumin	P-Value
n		18421	14862	3559	
Female, n (%)		7653 (41.5)	6322 (42.5)	1331 (37.4)	
White, n (%)		12366 (67.1)	9808 (66.0)	2558 (71.9)	
Emergency admission, n (%)		9605 (52.1)	8512 (57.3)	1093 (30.7)	
admission_age, mean (SD)	0	66.3 (16.2)	66.1 (16.8)	67.3 (13.1)	< 0.001
SOFA, mean (SD)	0	6.0(3.5)	5.7(3.4)	6.9 (3.6)	< 0.001
lactate, mean (SD)	4616	3.0(2.5)	2.8(2.4)	3.7(2.6)	< 0.001

Table 2. Characteristics of the trial population measured on the first 24 hours of ICU stay. Appendix 6 describes all confounders used in the analysis.

Identification: listing confounders

For confounders selection we use a causal DAG shown in Figure 11. Gray confounders are not controlled for since they are not available in the data. However, resulting confounding biases are captured by proxies such as comorbidity scores (SOFA or SAPS II) or other variables (eg. race, gender, age, weight). S1 Table details confounders summary statistics for treated and controls.

Estimation

Confounder aggregation: We tested multiple aggregations such as the last value before the start of the follow-up period, the first observed value, and both the first and last values as separated features.

Causal estimators: We implemented multiple estimation strategies, including Inverse Propensity Weighting (IPW), outcome modeling (G-formula) with T-Learner, Augmented Inverse Propensity Weighting (AIPW) and Double Machine Learning (DML). We used the python packages dowhy [55] for IPW implementation and EconML [96] for all other estimation strategies. Confidence intervals were estimated by bootstrap (50 repetitions). S3 Appendix and S5 Appendix detail the estimators and the available Python implementations.

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Outcome and treatment estimators: To model the outcome and treatment, we used two common but different estimators: random forests and ridge logistic regression implemented with scikit-learn [97]. We chose the hyperparameters with a random search procedure (S6 Appendix). While logistic regression handles predictors in a linear fashion, random forests bring the benefit of modeling non-linear relations.

Vibration analysis: Comparing sources of systematic errors

Study design — Illustration of immortal time bias: To illustrate the risk of immortal-time bias, we vary the eligibility period of treatment or control in a shorter or longer time window than 24 hours. As explained in section , a longer eligibility period means that patients are more likely to be treated if they survived up to the intervention and hence the study is biased to overestimate the beneficial effect of the intervention. Figure 3a shows that longer eligibility periods lead to albumin being markedly more efficient.

Confounder choice We consider other choice of confounding variables (S8 Appendix). Figure 3b shows that a less thorough choice, neglecting the administrated drugs, makes little to no difference. Major errors, such as omitting the biological measurements or using only socio-demographical variables, lead to sizeable bias.

Estimation choices – Confounder aggregation, causal and nuisance estimators: Figure 3c shows varying confidence intervals (CI) depending on the method. Doubly-robust methods provide the narrowest CIs, whereas the outcome-regression methods have the largest CI. The estimates of the forest models are closer to the consensus across prior studies (no effect) than the logistic regression indicating a better fit of non-linear relationships. We only report the first and last pre-treatment feature aggregation strategy, since detailed analysis showed little differences for other aggregations (S7 Fig). Both methodological studies [98] and consistency with published RCTs suggest to prefer doubly-robust approaches.

Treatment heterogeneity: Which treatment for a sub-population?

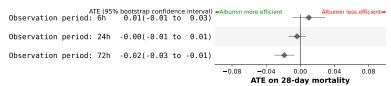
With adequate choice of study design, confounding variables and causal estimator, the average treatment effect matches well published findings: Pooling evidence from high-quality RCTs, no effect of albumin in severe sepsis was demonstrated for both 28-day mortality (odds ratio (OR) 0.93, 95% CI 0.80-1.08) and 90-day mortality (OR 0.88, 95% CI 0.761.01) [92]. Having validated the analytical pipeline, we can use it to inform decision-making. We explore heterogeneity along four binary patient characteristics, displayed in Figure 4. We find that albumin is beneficial with patient with septic shock consistent with one RCT [94]. It is also beneficial for older patients (age >=60) and males.

Discussion and conclusion

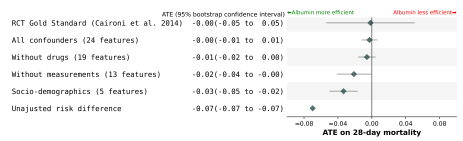
Valid decision-making evidence from EHR data requires a clear causal framework. Indeed, machine-learning algorithms have often extracted non-causal associations between the intervention and the outcome, improper for decision-making [13, 15, 16]. Machine learning studies in medicine often rely on an implicit causal thinking, via a good understanding of the clinical settings. A clear framework helps making sure nothing falls through the cracks.

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(a) Framing - Immortal Time Bias



(b) Identification - confounders choice



(c) Model selection

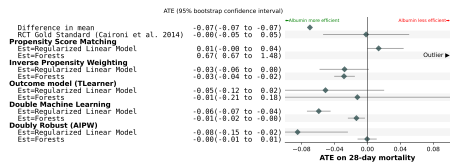


Fig 3. The effect of choices on the three analytical steps – All three analytical steps are equally important for the validity of the analysis. 3a) Framing step: Poor framing introduces time bias: A longer observation period (72h) artificially favors the efficacy of Albumin. 3b) Identification step: Choosing less informed confounders set introduces increasing bias in the results. 3c) Model selection step: Different estimators give different results. Score matching yields unconvincingly high estimates, inconsistent with the published RCT. With other causal approaches, using linear estimators for nuisances suggest a reduced mortality risk for albumin, while using forests for nuisance models points to no effect, which is consistent with the RCT gold standard.

The diamonds depict the mean effect and the bar are the 95% confidence intervals obtained respectively by 30, 30 and 50 bootstrap repetitions. For framing and identification, the estimator is a doubly robust learner (AIPW) with random forests for nuisances. Features are aggregated by taking the first and last measurements for all experiments.

We have separated three steps important for causal validity: the choice of study design, confounders, and estimators. Regarding study design, major caveats arise from the time component, where a poor choice of inclusion time easily brings in significant bias. Regarding choice of prediction variables, forgetting some variables that explains both the treatment allocation and the outcome leads to confounding bias, that however remains small when these variables capture weak links. Regarding choice of causal estimators, preferring flexible models such as random forests reduces the bias, in particular for doubly-robust estimators. We have shown that all these three steps are equally important: paying no attention to one of them leads to invalid estimates of treatment effect, yet imperfect but plausible choices lead to small biases of the same order of magnitude for all steps. For instance, despite the emphasis often put on choice

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Individual Treatment effects: better treatment efficacy for patients older than 60 years, septic shock, and to a lower extent males. The final estimator is ridge regression. The boxes contain the 25th and 75th percentiles of the CATE distributions

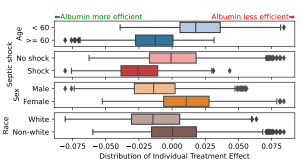
with the median indicated by the

the distribution.

vertical line. The whiskers extend to

1.5 times the inter-quartile range of

Fig 4. Subgroup distributions of



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of confounders, minor deviations from the expert's causal graph did not introduce substantial bias (S8 Appendix), no larger than a too rigid choice of estimator.

To assert the validity of the analysis, we argue to relate as much as possible the average effect to a reference target trial, even when the goal is to capture the heterogeneity of the effect to individualize decisions. EHRs complement RCTs: RCTs cannot address all the subpopulations and local practices [21,99]. EHRs often cover many individuals, with the diversity needed to model treatment heterogeneity. The corresponding model can then inform better decision-making [19]: a sub-population analysis (as in Figure 4) can distill rules on which groups of patients should receive a treatment. Beyond a sub-group perspective, patient-specific estimates facilitate a personalized approach to clinical decision-making [100].

Even without considering a specific intervention, anchoring machine-learning models on causal mechanisms can make them more robust to distributional shift [101], thus safer and fairer for clinical use [20,102]. Yet it is important to keep in mind that better prediction is not per se a goal in healthcare. Establishing strong predictors might be less important than identifying moderately strong but modifiable risk factors as established on in the Framingham cohort [103]. Cardio-vascular risk scores, while others such as the QRISK score are used to optimize population-wide cost effectiveness, but do not inform which patient benefits most from a given treatment ()cost-effectiveness, but not individual treatment effect.

No sophisticated data-processing tool can safeguard against invalid study design or a major missing confounder, loopholes that can undermine decision-making systems. Our framework helps the investigator ensure causal validity by outlining the important steps and relating average effects to RCTs. Causal grounding of individual predictions should reduce the social disparities that they reinforce [8,104,105], as these are driven by historical decisions and not biological mechanisms. At the population level, it leads to better public health decisions. For instance, going back to cardio-vascular diseases, the stakes are to go beyond risk scores and also account for responder status when prescribing prevention drugs.

Availability of data and materials

The datasets are available on PhysioNet (https://doi.org/10.13026/6mm1-ek67). We used MIMIC-IV.v2.2 The code for data preprocessing and analyses are available on github https://github.com/soda-inria/causal_ehr_mimic/.

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Authors contributions

MD and TS designed the study, MD performed the analysis and wrote the manuscript. TS, JA, CM, LAC, GV reviewed and edited the manuscript.

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Supporting information

S1 Fig. Motivating example: Failure of predictive models to predict mortality from pretreatment variables. To illustrate how machine learning frameworks can fail to inform decision making, we present a motivating example from MIMIC-IV. Using the same population and covariates as in the main analysis (described in Table 6), we train a predictive model for 28-day mortality. We split the data into a training set (80%) and a test set (20%). The training set uses the last measurements from the first 24 hours, whereas the validation set only uses the last measurements before the administration of crystalloids. We split the train set into a train and a validation set. We fit a HistGradientBoosting classifier ¹ on the train set and evaluate the performance on the validation set and on the test set. We see good area under the Precision-recall curve (PR AUC) on the validation set, but a deterioration of 10 points on the test set (Figure 5a). The same is seen in Figure 5b when measuring performances with Area Under the Curve of the Receiving Operator Characteristic (ROC AUC). On the contrary, a model trained on pre-treatment features yields competitive performances. This failure illustrates well the shortcuts on which predictive models could rely to make predictions. A clinically useful predictive model should support decision-making -in this case, addition of albumin to crystalloids- rather than maximizing predictive performance. In this example, causal thinking would have helped to identify the bias introduced by post-treatment features. In fact, these features should not be included in a causal analysis since they are post-treatment colliders.

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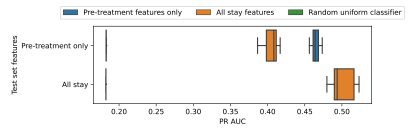
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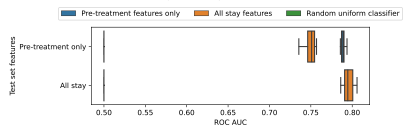
This kind of error might sound naive to a clinical expert but relying on shortcuts -some of them being post-treatment variables- is a common error. Here, we detail some real use cases where machine learning fail in providing useful predictions for decision-making. [15] use deep learning to predict hip fracture using confounding patient and healthcare variables. An example of such covariates shown by the authors is the triage of patients before imaging that results in the model trying to predict the image acquisition machine and rely on it to predict hip fracture. [16] describe the use of algorithm in US extra-care programs. By equating care needs with previous care costs (in a pure predictive fashion), the algorithm falsely conclude that Black patients are healthier than equally white patients, since they do less money is spent on them for a given level of need. Beyond Machine Learning, we also spotted the inclusion of post-treatment variables in the development of the recent SCORE2 cardio-vascular risk score [106]: Our risk models might have underestimated CVD risk be- cause data used to estimate multipliers were likely to include some people already on CVD prevention therapies (e.g. statins or anti-hypertensive medication. This score might be used to inform on the initiation of statins for primary prevention. But, relying on post-treatment, it might under-discover patients who would benefit from statins at screening time.

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¹https://scikit-learn.org/stable/modules/ensemble.html#histogram-based-gradient-boosting



(a) Area under the Precision-Recall curve (PR_AUC)



(b) Area under the Receiving Operator Characteristic (ROC $_AUC$)

Fig 5. Failure to predict 28-day mortality from a model fitted on pre-treatment variables. The model is trained on the last features from the whole stay and tested on two validation sets: one with all stay features and one with last features before crystalloids administration (Pre-treatment only). The all-stay model performance markedly decreases in the pre-treatment only dataset.

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S2 Fig. Immortal time bias illustration.

Figure 6 illustrates the immortal time bias. This time bias is a major pitfall in the retrospective evaluation of screening programs [107].

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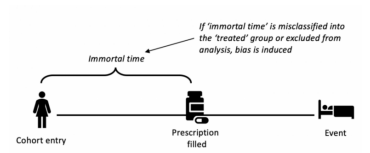


Fig 6. Poor experimental design can introduce Immortal time bias, which leads to a treated group with falsely longer longevity [108].

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S3 Fig. Graphical timeline. Drawing a graphical timeline as the one in Figure 7 during the study design helps to detect and prevent time-related biases.

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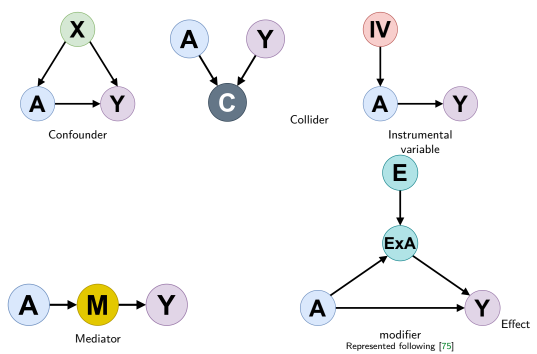
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Fig 7. Defining the inclusion event, the starting time T0 for follow-up, the intervention's assignment time and the observation window for confounders is crucial to avoid time and selection biases. In our study, the gap between the intervention and the inclusion is small compared to the occurrence of the outcome to limit immortal time bias: 6.7 hours vs 40 days for mortality.

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S4 Fig. Types of causal variables.

Figure 8 illustrates the different types of causal variables.



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Fig 8. The five categories of causal variables needed for our framework: A: Treatment, X: Confounder, IV: Instrumental variable, M: mediator, Y: Outcome, C: Collider, E: Effect modifier.

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S1 Appendix. Estimation of Treatment effect with MIMIC data.

We searched for causal inference studies in MIMIC using PubMed and Google scholar with the following search terms ((MIMIC-III OR MIMIC-IV) AND (causal inference OR treatment effect)). We retained eleven treatment effect studies clearly following the PICO framework:

- [109] studied the effect of High-flow nasal cannula oxygen (HFNC) against noninvasive mechanical ventilation on 801 patients with hypoxemia during ventilator weaning on 28-day mortality. They used propensity score matching, and found non-negative effects as previous RCTs reported –though those were focused on reintubation as the main outcome [110,111].
- [112] studied the effect of lower hypoxemia vs higher hypoxemia thresholds for the initiation of invasive ventilation (defined with saturation-to-inspired oxygen ratio (SF)) for 3,357 patients from MIMIC receiving inspired oxygen fraction greater than 0.4 on 28-day mortality. Using bayesian G-computation (time-varying treatment model with gaussian process and outcome-model with BART, taking the treatment model as entry), they found protective effects for initialization at low hypoxemia. However, when externally validation their findings in the AmsterdamUMCdb dataset, they found the highest mortality probability for patients with low hypoxemia. Authors concluded that their model was heavily dependent on clinical context and baseline characteristics. There might be some starting-time bias in this study since it is really close
- [113] studied the effect of indwelling arterial catheters (IACs) vs non-IAC for 1,776 patients who are mechanically ventilated and did not require vasopressor support on 28-day mortality. They used propensity score matching and found no effect. A notebook based on google cloud access to MIMIC-IV replicating the study is available here.
- [114] studied the effect of transthoracic echocardiography vs no intervention for 6,361 patients with sepsis on 28-day mortality. They used IPW, PSM, g-formula and a doubly robust estimation. The propensity score was modeled with boosting and the outcome model with a logistic regression. They found a significant positive reduction of mortality (odd ratio 0.78, 95% CI 0.68-0.90). Study code is open source.
- [115] studied the effect of liberal –target SpO2 greater than 96%– vs conservative oxygenation –target SpO2 between 88-95%– in 4,062 mechanically ventilated patients on 90-day mortality. They found an advantage of the liberal strategy over liberal (ATE=0.13) by adjusting on age and apsii. This is not consistent with previous RCTs where no effects have been reported [116, 117].
- [118] studied the effect of fluid-limiting treatment –caped between 6 and 10 L– vs no cap on fluid administration strategies for 1,639 sepsis patients on 30 day-mortality. Using a dynamic Marginal Structural Model with IPW, they found a protective effect of fluid-limitation on ATE -0.01 (95%CI -0.016, -0.03). This is somehow concordant with the RIFTS RCT that found no effect of fluid limitation [119] and two previous meta-analyses [120, 121].
- [122] studied the effect of statin use prior to ICU admission vs absence of pre-ICU prescription for 8,200 patients with sepsis on 30-day mortality. Using AIPW (no estimator reported) and PSM (logistic regression), they found a decrease on mortality (ATE -0.039, 95%CI -0.084, -0.026). This partly supports previous findings in Propensity Matching bases observational studies [123, 124]. But all RCTs [125, 126] found no improvement for sepsis (not pre-admission administration though).

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The [127] meta-analysis concludes that there is lack of evidence for the use of statins in sepsis with inconsistent results between RCTs (no effect) and observational studies (protective effect).

- [128] studied the effect of higher vs lower positive end-expiratory pressures (PEEP) in 1,411 patients with Acute Respiratory Distress Syndrome (ARDS) syndrome on 30 day mortality. Very few details on the methods were reported, but they found a protective effect for higher PEEP consistent results from a target trial [129].
- [128] also studied the effect of early use of a neuromuscular blocking agent vs placebo in 752 patients moderate-severe ARDS on 30 day mortality. Very few details on the methods were reported, but they found a protective effect for the use of a neuromuscular blocking agent, consistent with the results from a target trial [130].
- [95] studied the administration of a combination of albumin within the first 24-h after crystalloids vs crystalloids alone for 6,641 patients with sepsis on 28-day mortality. Using PSM, they found protective effect of combination on mortality, but insist on the importance of initialization timing. This is consistent with [92], who found a non-significant trend in favor of albumin used for severe sepsis patients and a significant reduction for septic shock patients, both on 90-day mortality. These results are aligned with [94] that found no effect for severe sepsis patient but positive effect for septic shock patients.
- [131] studied early enteral nutrition (EN) -<=53 ICU admission hours- vs delayed EN for 2,364 patients with sepsis and EN on acute kidney injury. With PSM, IPW and g-formula (logistic estimator each time), they found a protective effect (OR 0.319, 95%CI 0.245, 0.413) of EEN.

These eleven studies mainly used propensity score matching (6) and IPW (4), two of them used doubly robust methods, and only one included a non-linear estimator in either the outcome or the treatment model. None of them performed a vibration analysis on confounder selection or feature transformations. They have a strong focus on patients with sepsis. Only four of them found concordant results with previous RCTs [109, 118, 128].

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S2 Appendix. Assumptions: what is needed for causal inference from observational studies.

The following four assumptions, referred as strong ignorability, are needed to assure identifiability of the causal estimands with observational data with most causal-inference methods [74], in particular these we use:

Assumption 1 (Unconfoundedness)

$$\{Y(0), Y(1)\} \perp \!\!\!\perp A|X \tag{1}$$

This condition –also called ignorability– is equivalent to the conditional independence on the propensity score $e(X) = \mathbb{P}(A = 1|X)$ [132]: $\{Y(0), Y(1)\} \perp \!\!\!\perp A|e(X)$.

Assumption 2 (Overlap, also known as Positivity)

$$\eta < e(x) < 1 - \eta \quad \forall x \in \mathcal{X} \text{ and some } \eta > 0$$
(2)

The treatment is not perfectly predictable. Or in other words, every patient has a chance to be treated and not to be treated. For a given set of covariates, we need examples of both to recover the ATE.

As noted by [79], the choice of covariates X can be viewed as a trade-off between these two central assumptions. A bigger covariate set generally reinforces the ignorability assumption. In the contrary, overlap can be weakened by large \mathcal{X} because of the potential inclusion of instrumental variables: variables only linked to the treatment which could lead to arbitrarily small propensity scores.

Assumption 3 (Consistency) The observed outcome is the potential outcome of the assigned treatment:

$$Y = AY(1) + (1 - A)Y(0)$$
(3)

Here, we assume that the intervention A has been well defined. This assumption focuses on the design of the experiment. It clearly states the link between the observed outcome and the potential outcomes through the intervention [40].

Assumption 4 (Generalization) The training data on which we build the estimator and the test data on which we make the estimation are drawn from the same distribution, also known as the "no covariate shift" assumption [133].

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S3 Appendix. Major causal-inference methods: When to use which estimator?

G-formula also called conditional mean regression [83], g-computation [59], or Q-model [134]. This approach is directly modeling the outcome, also referred to as the response surface: $\mu_{(a)}(x) = \mathbb{E}(Y \mid A = a, \mathbf{X} = x)$

Using an outcome estimator to learn a model for the response surface $\hat{\mu}$ (eg. a linear model), the ATE estimator is an average over the n samples:

$$\hat{\tau}_G(f) = \frac{1}{n} \sum_{i=1}^n \hat{\mu}(x_i, 1) - \hat{\mu}(x_i, 0) = \frac{1}{n} \sum_{i=1}^n \hat{\mu}_{(1)}(x_i) - \hat{\mu}_{(0)}(x_i)$$
(4)

This estimator is unbiased if the model of the conditional response surface $\hat{\mu}_{(a)}$ is well-specified. This approach assumes than $Y(a) = \mu_a(X) + \epsilon_a$ with $\mathbb{E}[\epsilon|X] = 0$. The main drawback is the extrapolation of the learned outcome estimator from samples with similar covariates X but different intervention A.

Propensity Score Matching (PSM) To avoid confounding bias, the ignorability assumption 1) requires to contrast treated and control outcomes only between comparable patients with respect to treatment allocation probabilities. A simple way to do this is to group patients into bins, or subgroups, of similar confounders and contrast the two population's outcomes by matching patients inside these bins [57]. However, the number of confounder bins grows exponentially with the number of variables. [132] proved that matching patients on the individual probabilities to receive treatment—propensity scores— is sufficient to verify ignorability. PSM is a conceptually simple method, but has delicate parameters to tune such as choosing a model for the propensity score, deciding what is the maximum distance between two potential matches (the caliper width), the number of matches by sample, and matching with or without replacement. It also prunes data not meeting the caliper width criteria, and suffers from high estimation variance in highly-dimensional data where extreme propensity weights are common. Finally, the simple bootstrap confidence intervals are not theoretically grounded [135] making PSM more difficult to use for applied practitioners.

Inverse Propensity Weighting (IPW)

A simple alternative to propensity score matching is to weight the outcome by the inverse of the propensity score [58]. It relies on a similar idea as matching but automatically builds a balanced population by reweighting the outcomes with the propensity score model \hat{e} to estimate the ATE:

$$\hat{\tau}_{IPW}(\hat{e}) = \frac{1}{n} \sum_{i=1}^{N} \frac{A_i Y_i}{\hat{e}(X_i)} - \frac{(1 - A_i) Y_i}{(1 - \hat{e}(X_i))}$$
 (5)

This estimate is unbiased if \hat{e} is well-specified. IPW suffers from high variance if some weights are too close to 0 or 1. In high dimensional cases where poor overlap between treated and control is common, one can clip extreme weights to limit estimation instability.

Doubly Robust Learning, DRL also called Augmented Inverse Probability Weighting (AIPW) [136].

The underlying idea of DRL is to combine the G-formula and IPW estimators to protect against a mis-specification of one of them. It first requires to estimate the two nuisance parameters: a model for the intervention \hat{e} and a model for the outcome f. If one of the two nuisance is unbiased, the following ATE estimator is as well:

$$\widehat{\tau}_{AIPW} = \frac{1}{n} \sum_{i=1}^{n} \left(\widehat{\mu}_{(1)}(x_i) - \widehat{\mu}_{(0)}(x_i) + a_i \frac{y_i - \widehat{\mu}_{(1)}(x_i)}{\widehat{e}(x_i)} - (1 - a_i) \frac{y_i - \widehat{\mu}_{(0)}(x_i)}{1 - \widehat{e}(x_i)} \right)$$

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Moreover, despite the need to estimate two models, this estimator is more efficient in the sense that it converges quicker than single model estimators [82]. For this propriety to hold, one need to fit and apply the two nuisance models in a cross-fitting manner. This means that we split the data into K folds. Then for each fold, we fit the nuisance models on the K-1 complementary folds, and predict on the remaining fold.

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To recover Conditional Treatment Effects from the AIPW estimator, [137] suggested to regress the Individual Treatment Effect estimates from AIPW on potential sources of heterogeneity X^{cate} : $\hat{t}au = \arg\min_{\tau \in \Theta}(\hat{\tau}_{AIPW}(X) - \tau(X^{cate}))$ for Θ some class of model (eg. linear model).

Double Machine Learning [53] also known as the R-learner [138]. It is based on the R-decomposition, [139], and the modeling of the conditional mean outcome, $m(x) = \mathbb{E}[Y|X=x]$ and the propensity score, $e(x) = \mathbb{E}[A=1|X=x]$:

$$y_i - m(x_i) = (a_i - e(x_i)) \tau(x_i) + \varepsilon_i \text{ with } \varepsilon_i = y_i - \varepsilon[4_i \mid x_i, a_i]$$
 (6)

Note that we can impose that the conditional treatment effect $\tau(x)$ only relies on a subset of the features, x^{cate} on which we want to study treatment heterogeneity.

From this decomposition, we can derive an estimation of the ATE τ , where the right hand-side term is the empirical R-Loss:

$$\hat{\tau}(\cdot) = \operatorname{argmin}_{\tau} \left\{ \frac{1}{n} \sum_{i=1}^{n} \left((y_i - m(x_i)) - (a_i - e(x_i)) \tau(x_i^{cate}) \right)^2 \right\}$$
 (7)

The full procedure for R-learning is first to fit the nuisances: \hat{m} and \hat{e} . Then, minimize the estimated R-loss eq.7, where the oracle nuisances (e, m) have been replaced by their estimated counterparts (\hat{e}, \hat{m}) . Minimization can be done by regressing the outcome residuals weighted by the treatment residuals. Finally, get the ATE by averaging conditional treatment effect $\tau(x^{cate})$ over the population.

This estimator has also the doubly robust proprieties described for AIPW. it should have less variance than AIPW since it does not use the propensity score in the denominator.

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	$estimation_method$	$compute_time$	$outcome_model$	event_aggregation
2	LinearDML	1127.977827	Forests	['first', 'last']
3	backdoor.propensity_score_matching	199.765587	Forests	['first', 'last']
4	backdoor.propensity_score_weighting	86.149872	Forests	['first', 'last']
5	TLearner	284.066786	Forests	['first', 'last']
6	LinearDRLearner	2855.403709	Forests	['first', 'last']
7	LinearDML	49.911035	Regularized LR	['first', 'last']
8	backdoor.propensity_score_matching	127.929910	Regularized LR	['first', 'last']
9	backdoor.propensity_score_weighting	6.407206	Regularized LR	['first', 'last']
10	TLearner	6.843931	Regularized LR	['first', 'last']
11	LinearDRLearner	80.747301	Regularized LR	['first', 'last']

Table 3. Compute times for the different estimation methods with 50 bootstrap replicates.

S4 Appendix. Statistical considerations when implementing estimation. Counterfactual prediction lacks off-the-shelf cross-fitting estimators

Doubly robust methods use cross-fit estimation of the nuisance parameters, which is not available off-the-shelf for IPW and T-Learner estimators. For reproducibility purposes, we did not reimplement internal cross-fitting for treatment or outcome estimators. However, when flexible models such as random forests are used, a fairer comparison between single and double robust methods should use cross-fitting for both. This lack in the scikit-learn API [97] reflects different needs between purely predictive machine learning focused on generalization performances and counterfactual prediction aiming at unbiased inference on the input data.

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Good practices for imputation not implemented in EconML

Good practices in machine learning recommend to input distinctly each fold when performing cross-fitting 2 . However, EconML estimators test for missing data at instantiation preventing the use of scikit-learn imputation pipelines. We thus have been forced to transform the full dataset before feeding it to causal estimators. An issue mentioning the problem has been filed, so we can hope that future versions of the package will comply with best practices. 3

Bootstrap may not yield the most efficient confidence intervals

To ensure a fair comparison between causal estimators, we always used bootstrap estimates for confidence intervals. However, closed form confidence intervals are available for some estimators – see [82] for IPW and AIPW (DRLeaner) variance estimations. These formulas exploit the estimator properties, thus tend to have smaller confidence intervals. On the other hand, they usually do not include the variance of the outcome and treatment estimators, which is naturally dealt with in bootstrapped confidence intervals. Closed form confidence intervals are rarely implemented in any of the packages as Dowhy for the IPW estimator, or in EconML for AIPW.

Bootstrap was particularly costly to run for the EconML doubly robust estimators (AIPW and Double ML), especially when combined with random forest nuisance estimators (from 10 to 47 min depending on the aggregation choice and the estimator). See Table 3 for details.

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²https://scikit-learn.org/stable/modules/compose.html#combining-estimators

³https://github.com/py-why/EconML/issues/664

S5 Appendix. Packages for causal estimation in the python ecosystem. We searched for causal inference packages in the python ecosystem. The focus was on the identification methods. Important features were ease of installation, sklearn estimator support, sklearn pipeline support, doubly robust estimators, confidence interval computation, honest splitting (cross-validation), Targeted Maximum Likelihood Estimation. These criteria are summarized in 4. We finally chose EconML despite lacking sklearn. BaseImputer support through the sklearn. Pipeline object as well as a TMLE implementation.

The zEpid package is primarily intended for epidemiologists. It is well documented and provides pedagogical tutorials. It does not support sklearn estimators, pipelines and honest splitting.

EconML [96] implements almost all estimators except propensity score methods. Despite focusing on Conditional Average Treatment Effect, it provides all. One downside is the lack of support for scikit-learn pipelines with missing value imputers. This opens the door to information leakage when imputing data before splitting into train/test folds.

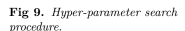
Dowhy [140] focuses on graphical models and relies on EconML for most of the causal inference methods (identifications) and estimators. Despite, being interesting for complex inference –such as mediation analysis or instrumental variables–, we considered that it added an unnecessary layer of complexity for our use case where a backdoor criterion is the most standard adjustment methodology.

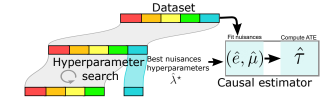
Causalml implements all methods, but has a lot of package dependencies which makes it hard to install.

Packages	Simple	Confidence	sklearn	sklearn	Propensity	Doubly Robust	TMLE	Honest splitting
rackages	installation	Intervals	estimator	pipeline	estimators	estimators	estimator	(cross validation)
dowhy	✓	1	1	1	✓	Х	Х	Х
EconML	1	1	1	Yes except	¥	1	Y	Only for doubly
ECOMVIL	•	•	*	for imputers	<i>^</i>	•	^	robust estimators
zEpid	1	1	Х	Х	1	✓	1	Only for TMLE
causalml	Y	1	1	1	1	1	1	Only for doubly
Causaiiii 🗡	^	^ '	•	· •	*	•	١ ٠	robust estimators

Table 4. Selection criteria for causal python packages

S6 Appendix. Hyper-parameter search for the nuisance models. We followed a two-step procedure to train the nuisance models (eg. $(\hat{e}, \hat{\mu})$ for the AIPW causal estimator), taking inspiration from the computationally cheap procedure from [141, section 3.3]. First, for each nuisance model, we fit a random parameter search with 5-fold cross validation and 10 iterations on the full dataset. Each iteration fit a model with a random combination of parameters in a predefined grid, then evaluate the performance by cross-validation. The best hyper-parameters $\hat{\lambda}^*$ are selected as the ones reaching the minimal score across all iterations. Then, we feed this parameters to the causal estimator. The single robust estimators (matching, IPW and TLearner) refit the corresponding estimator only once on the full dataset, then estimate the ATE. The doubly-robust estimators use a cross-fitting procedure (K=5) to fit the





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	estimator	nuisance	Grid
Estimator type			
Linear	LogisticRegression	treatment	{'C': logspace(-3, 2, 10)}
Linear	Ridge	outcome	{'alpha': logspace(-3, 2, 10)}
Forest	RandomForestClassifier	treatment	{'n_estimators': ['10', '100', '200'], 'max_depth': ['3', '10', '50']}
Forest	RandomForestRegressor	outcome	{'n_estimators': ['10', '100', '200'], 'max_depth': ['3', '10', '50']}

 ${\bf Table~5.}~Hyper-parameter~grid~used~during~random~search~optimization.$

nuisances then estimate the ATE. Figure 9 illustrates the procedure and Table 5 details the hyper-parameters grid for the random search.

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S7	Appendix.	Computing	resources
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The whole project was run on a laptop running Ubuntu 22.04.2 LTS with the following hardware: CPU 12th Gen Intel(R) Core(TM) i7-1270P with 16 threads and 15 GB of RAM.

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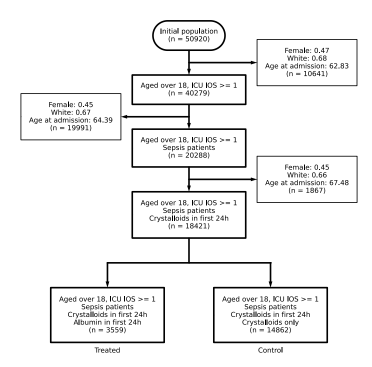


Fig 10. Selection flowchart on MIMIC-IV for the emulated trial.

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S1 Table. Complete description of the confounders for the main analysis.

	Missing	Overall	Cristalloids only	Cristalloids + Albumin	P-Value
n		18421	14862	3559	
Glycopeptide, n (%)		9492 (51.5)	7650 (51.5)	1842 (51.8)	
Beta-lactams, n (%)		5761 (31.3)	5271 (35.5)	490 (13.8)	
Carbapenems, n (%)		727 (3.9)	636 (4.3)	91 (2.6)	
Aminoglycosides, n (%)		314 (1.7)	290 (2.0)	24 (0.7)	
suspected_infection_blood, n (%)		170 (0.9)	149 (1.0)	21 (0.6)	
RRT, n (%)		229 (1.2)	205 (1.4)	24 (0.7)	
ventilation, n (%)		16376 (88.9)	12931 (87.0)	3445 (96.8)	
vasopressors, n (%)		9058 (49.2)	6204 (41.7)	2854 (80.2)	
Female, n (%)		7653 (41.5)	6322 (42.5)	1331 (37.4)	
White, n (%)		12366 (67.1)	9808 (66.0)	2558 (71.9)	
Emergency admission, n (%)		9605 (52.1)	8512 (57.3)	1093 (30.7)	
Insurance, Medicare, n (%)		9727 (52.8)	7958 (53.5)	1769 (49.7)	
myocardial_infarct, n (%)		3135 (17.0)	2492 (16.8)	643 (18.1)	
malignant cancer, n (%)		2465 (13.4)	2128 (14.3)	337 (9.5)	
diabetes_with_cc, n (%)		1633 (8.9)	1362 (9.2)	271 (7.6)	
diabetes_without_cc, n (%)		4369 (23.7)	3532 (23.8)	837 (23.5)	
metastatic solid tumor, n (%)		1127 (6.1)	1016 (6.8)	111 (3.1)	
severe liver disease, n (%)		1289 (7.0)	880 (5.9)	409 (11.5)	
renal disease, n (%)		3765 (20.4)	3159 (21.3)	606 (17.0)	
aki_stage_0.0, n (%)		7368 (40.0)	6284 (42.3)	1084 (30.5)	
aki_stage_1.0, n (%)		4019 (21.8)	3222 (21.7)	797 (22.4)	
aki_stage_2.0, n (%)		6087 (33.0)	4605 (31.0)	1482 (41.6)	
aki_stage_3.0, n (%)		947 (5.1)	751 (5.1)	196 (5.5)	
SOFA, mean (SD)	0	6.0 (3.5)	5.7 (3.4)	6.9 (3.6)	< 0.001
SAPSII, mean (SD)	0	40.3 (14.1)	39.8 (14.1)	42.8 (13.6)	< 0.001
Weight, mean (SD)	97	83.3 (23.7)	82.5 (24.2)	86.4 (21.2)	< 0.001
temperature, mean (SD)	966	36.9 (0.6)	36.9 (0.6)	36.8 (0.6)	< 0.001
mbp, mean (SD)	0	75.6 (10.2)	76.3 (10.7)	72.4 (7.2)	< 0.001
resp_rate, mean (SD)	9	19.3 (4.3)	19.6 (4.4)	18.0 (3.8)	< 0.001
heart_rate, mean (SD)	0	86.2 (16.3)	86.2 (16.8)	86.5 (14.3)	0.197
spo2, mean (SD)	4	97.4 (2.2)	97.3 (2.3)	98.0 (2.1)	< 0.001
lactate, mean (SD)	4616	3.0 (2.5)	2.8 (2.4)	3.7 (2.6)	< 0.001
urineoutput, mean (SD)	301	24.0 (52.7)	24.7 (58.2)	21.1 (16.6)	< 0.001
admission_age, mean (SD)	0	66.3 (16.2)	66.1 (16.8)	67.3 (13.1)	< 0.001
delta mortality to inclusion, mean (SD)	11121	316.9 (640.2)	309.6 (628.8)	365.0 (708.9)	0.022
delta intervention to inclusion, mean (SD)	14862	0.3 (0.2)	nan (nan)	0.3 (0.2)	nan
delta inclusion to intime, mean (SD)	0	0.1 (0.2)	0.1 (0.2)	0.1 (0.1)	0.041
delta ICU intime to hospital admission, mean (SD)	0	1.1 (3.7)	1.0 (3.7)	1.6 (3.4)	< 0.001
los_hospital, mean (SD)	0	12.6 (12.5)	12.6 (12.5)	12.9 (12.4)	0.189
los_icu, mean (SD)	0	5.5 (6.7)	5.5 (6.5)	5.5 (7.2)	0.605

Table 6. Characteristics of the trial population measured on the first 24 hours of ICU stay. Risk scores (AKI, SOFA, SAPSII) and lactates have been summarized as the maximum value during the 24 hour period for each stay. Total cumulative urine output has been computed. Other variables have been aggregated by taking mean during the 24 hour period.

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The expert DAG in figure depicts the known causal links between these variables.

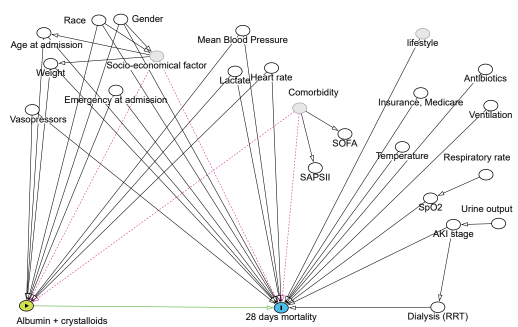


Fig 11. Causal graph for the Albumin vs crystalloids emulated trial – The green arrow indicates the effect studied. Black arrows show causal links known to medical expertise. Dotted red arrows highlight confounders not directly observed. For readability, we draw only the most important edges from an expert point of view. All white nodes correspond to variables included in our study.

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S7 Fig. Complete results for the main analysis.

Compared to Figure 3, we also report in Figure 12 the estimates for Causal forest estimators and other choices of feature aggregation (first and last).



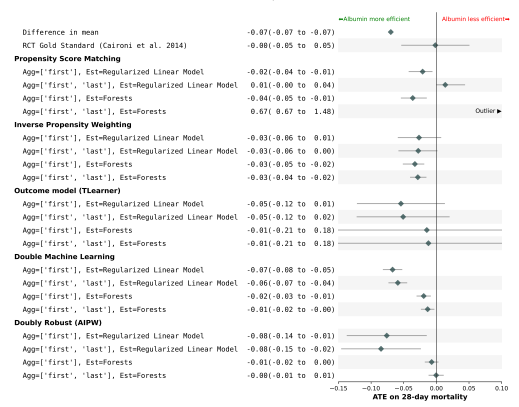


Fig 12. Full sensitivity analysis: The estimators with forest nuisances point to no effect for almost every causal estimator consistently with the RCT gold standard. Only matching with forest yields an unconvincingly high estimate. Linear nuisance used with doubly robust methods suggest a reduced mortality risk for albumin. The choices of aggregation only marginally modify the results expect for propensity score matching. The green diamonds depict the mean effect and the bar are the 95% confidence intervals obtained by 50 bootstrap repetitions.

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S8 Fig. Complete results for the Immortal time bias.

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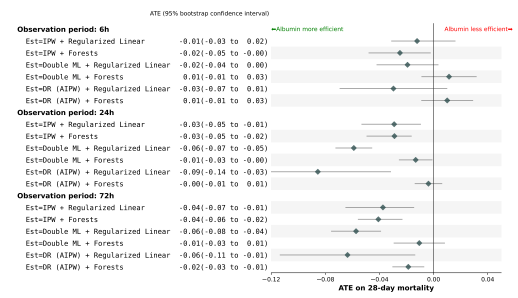


Fig 13. Sensitivity analysis for immortal time bias: Every choice of estimates show an improvement of the albumin treatment when increasing the observation period, thus increasing the blank period between inclusion and administration of albumin. Aggregation was concatenation of first and last features. The green diamonds depict the mean effect and the bar are the 95% confidence intervals obtained by 50 bootstrap repetitions.

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S8 Appendix. Deviating from expert ignorability – Impact of smaller confounders sets.

We conducted a dedicated vibration analysis on the different choices of confounders. We created three confounder subsets in addition to all confounders (24 variables): all confounders without antibiotics (Glycopeptides, Beta-lactams, Carbapenems, Aminoglycosides), all confounders without any measurement (weight, lactate, heart rate, spo2, mbp, urine output, temperature, AKI stage, SAPSII, respiratory rate, SOFA), only socio-demographics (admission age, female, emergency admission, insurance–medicare, race).

Figure 3b shows that small deviation from the ignorability assumptions is tolerable: for example, removing antibiotics does not impact the estimate. However, the larger the deviation from graph 11, the larger the bias compared to the gold-standard. Adjusting only for socio-demographics features is the closest from an unadjusted risk difference, indicating that we lack important confounders on the patient health state. This stability of the treatment effect estimator once sufficient confounders have been included has already been described and suggested as a confounder selection method [50].

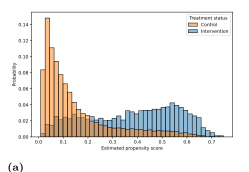
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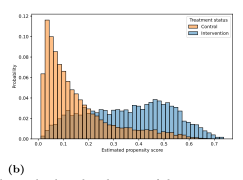
S9 Fig. Vibration analysis for aggregation.

We conducted a dedicated vibration analysis on the different choices of features aggregation, studying the impact on the estimated ATE. We also studied if some choices of aggregation led to substantially poorer overlap.

We assessed overlap with two different methods. As recommended by [58], we did a graphical assessment by plotting the distribution of the estimated. The treatment model hyper-parameters were chosen by random search, then predicted propensity scores were obtained by refitting this estimator with cross-fitting on the full dataset.

As shown in Figure 14, we did not find substantial differences between methods when plotting graphically the distribution of the estimated propensity score.





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Fig 14. Different choices of aggregation yield qualitatively close distributions of the propensity score: Figure 14a)a) shows a concatenation of first, last and median measures whereas Figure 14b)b) shows an aggregation by taking the first measure only. The underlying treatment effect estimator is a random forest.

We also used normalized total variation (NTV) as a summary statistic of the estimated propensity score to measure the distance between treated and control population [85]. This statistic varies between 0 – perfect overlap – and 1 – no overlap at all. Fig 15 shows no marked differences in overlap as measured by NTV between aggregation choices, comforting us in our expert-driven choice of the aggregation: a concatenation of first and last feature observed before inclusion time.

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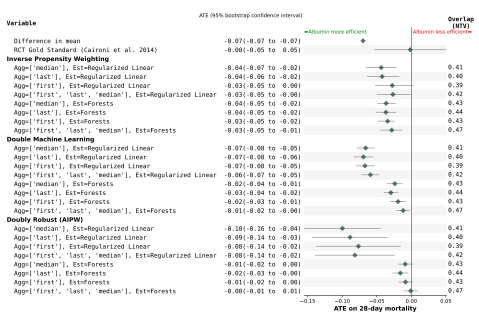


Fig 15. Vibration analysis dedicated to the aggregation choices. The choices of aggregation only marginally modify the results. When assessed with Normalized Total Variation, the overlap assumption is respected for all our choices of aggregation. The green diamonds depict the mean effect and the bar are the 95% confidence intervals obtained by 50 bootstrap repetitions.

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S8 Appendix. Details on treatment heterogeneity analysis. Detailed estimation procedure

The estimation of heterogeneous effect based on Double Machine Learning adds another step after the computation, regressing the residuals of the outcome nuisance $\tilde{Y} - \mu(X)$ against the residuals of the treatment nuisance $\tilde{A} = A - e(X)$ with the heterogeneity features X_{CATE} . Noting the final CATE model θ , Double ML solves:

$$\arg\min_{\theta} \mathbb{E}_n \left[(\tilde{Y} - \tau (X_C A T E) \cdot \tilde{A})^2 \right]$$

Where
$$\tilde{Y} = Y - \hat{m}(X)$$
 and $\tilde{A} = A - \hat{e}(X)$

To avoid the over-fitting of this last regression model, we split the dataset of the main analysis into a train set (size=0.8) where the causal estimator and the final model are learned, and a test set (size=0.2) on which we report the predicted Conditional Average Treatment Effects.

Known heterogeneity of treatment for the emulated trial

[94] observed statistical differences in the post-hoc subgroup analysis between patient with and without septic shock at inclusion. They found increasing treatment effect measured as relative risk for patients with septic shock (RR=0.87; 95% CI, 0.77 to 0.99 vs 1.13;95% CI, 0.92 to 1.39).

[142] conducted a post-hoc subgroup analysis of patients with or without brain injury –defined as Glasgow Coma Scale between 3 to 8-. The initial population was patients with traumatic brain injury (defined as history or evidence on A CT scan of head trauma, and a GCS score <= 13). They found higher mortality rate at 24 months in the albumin group for patients with severe head injuries.

[95] conducted a subgroup analysis on age (<60 vs >60), septic shock and sex. They conclude for increasing treatment effect measured as Restricted Mean Survival Time for Sepsis vs septic shock (3.47 vs. 2.58), for age >=60 (3.75 vs 2.44), for Male (3.4 vs 2.69). None of these differences were statistically significant.

Vibration analysis

The choice of the final model for the CATE estimation should also be informed by statistical and clinical rationals. Figure 16 shows the distribution of the individual effects of a final random forest estimator, yielding CATE estimates that are not consistent with the main ATE analysis. Figure 17 shows that the choice of this final model imposes an inductive bias on the form of the heterogeneity and different sources of noise depending of the nature of the model. A random forest is noisier than a linear model. Figure 17 shows the difference of modelization on the subpopulation of non-white male patients without septic shock. One can see that the decreasing linear trend is reflected by the random forest model only for patients aged between 55 and 80.

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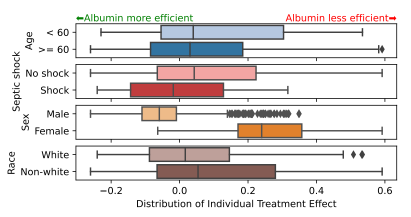


Fig 16. Distribution of Conditional Average Treatment effects on sex, age, race and pre-treatment septic shock estimated with a final forest estimator. The CATE are positive for each subgroups, which is not consistent with the null treatment effect obtained in the main analysis. The boxes contain between the 25th and 75th percentiles of the CATE distributions with the median indicated by a vertical line. The whiskers extends to 1.5 the inter-quartile range of the distribution.

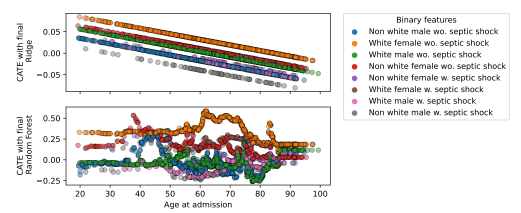


Fig 17. Distribution of Conditional Average Treatment effects on sex, age, race and pre-treatment septic shock plotted for different ages. On the top the final estimator is a linear model; on the bottom, it is a random forest. The forest-based CATE displays more noisy trends than the linear-based CATE. This suggest that the flexibility of the random forest might be underfitting the data.

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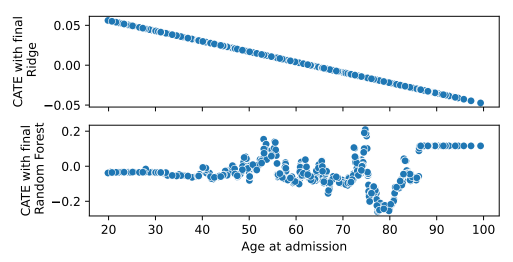


Fig 18. Figure 17 on the subpopulation of white male patients without septic shock. Contrary to the ridge regression (on top) inducing a nicely interpretable trend, using random forests as the final estimator failed to recover CATE on ages: the predicted estimates do not exhibit any trend and display inconsistently large effect sizes, suggesting data underfitting.

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S9 Appendix. Risk assessment tools for prevention.

An example of nationally deployed cardiovascular prevention tool: QRISK

The QRISK is a risk score widely used in the UK to assess 10-year cardio-vascular risk [26]. It is well calibrated and have satisfactory discriminative performances. As such, it is recommended by the national regulatory agency to engage the patient into various preventive interventions such as physical activity, cardio-protective diet, statins initiation [27]. The NICE argues in favor of systematic screening of cardio-vascular risks to increase the number of people receiving statins [27].

The QRISK seems to mostly inform the initiation of statins for patients exceeding 10% 10-year CVD risk (as measured with QRISK) following the 2014 NICE guidelines [28].

They base their evidence on a cost-effectiveness analysis choosing the 10% threshold to optimize quality-adjusted life year at the population level based on the cost of life-long statin treatments [29]. This study assumes constant statins effect among different CVD risk groups: It is assumed that the risk ratios given for treatment with each class of statins are constant regardless of the baseline CV risk—that is, someone with low CV risk will receive the same proportional reduction in that risk as would someone with a high CV risk. This is unproven, but is consistent with the results of meta-analysis carried out by the Cholesterol Treatment Trialists, which found effectiveness to be broadly similar for those at different risk levels. This hypothesis is supported by the meta-analysis of [33] at least for the important risk factors of sex and age [33]. It is also assumed that these risk ratios are constant regardless of baseline LDL-cholesterol levels. More recent work showed that risk ratio stratified by patient baseline CVD risk levels were not significant, raising doubts about the overall efficacy of statins for primary prevention [34, Table 1] [34].

The updated guidelines from the US Preventive Services Task Force found respectively moderate and small net benefice of statins for patients aged 40 to 75 with above 10% CVD risk and from 7.5 to 10% without history of CVD [35]. They conclude that the evidence is insufficient to determine the balance of benefits and harms of statin use for the primary prevention of CVD events and mortality in adults 76 years or older with no history of CVD.

Critics on the efficiency of systematic screening points to treatment heterogeneity

On the other side, the WHO systematic review on systematic screening for cardio-vascular risks from 2019 [1] is categorical on the poor effectiveness of systematic cardiovascular risk screening, consistently with the Cochrane review on poor effectiveness [36]. For the WHO, systematic screening leads to over-diagnosis and over-treatment. They mention in their argumentation no positive effect of statins and even adverse effect for mild hypertension [143]. They also mention critics against poor uptake of the systematic screening in UK or Albany leading to documented social bias in the screened population: for example, in the UK, treatment heterogeneity at a given risk level [37] or documentation of social inequities of health checks [2].

• The first critic shows that the augmentation of statin prescriptions (up to 400% increase for high risk patients), has been associated with heterogeneity in statin prescriptions for given risk score levels [37]. Low risk patients have been over-prescribed and high risk patient under-prescribed, resulting in dubious questionable overall improvement. That suggests that some patients are responders and others are non-responders. In this case, the constant effect hypothesis of statin taken by proposed by the NICE cost-effectiveness study is falsein doubt. Having access to sources of heterogeneity would help to optimize treatment allocation. The risk score might capture interesting heterogeneity

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directions but, we lack a RCT conducted on different risk levels to conclude, but we lack data from RCTs to draw such conclusions.

• The second critic points to a treatment allocation bias for risk assessment. It seems that mostly socially advantaged patients are receiving risk assessment and thus the possibility to be treated with statins. This critic also points to treatment heterogeneity. Suppose that every patient benefits the same from statins, then even a small part of the population –such as the socially advantaged at more than 10% risk—would benefit from it and make the intervention effective at the population level.

NICE evidence for statin efficacy

In the NICE analysis, three trials weight for more than 70% of the effect. All focus on high risk patients and secondary prevention. It is not clear if those effects are used for the cost-effectiveness.

- LIPID [30] (32% of the effect): 9,014 patients with a history of myocardial infraction or hospitalization ie. secondary prevention.
- PROSPER [31] (22% of the effect): an elderly cohort of 5,804 men and women with, or at high risk of developing, cardiovascular disease and stroke.
- ALLHAT-LLT [32] (23% of the effect): 10,355 patients age more than 55 years and stage 1 or 2 hypertension with at least 1 additional CHD risk factor.

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