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RESEARCH

Sequential regression and simulation: a method for estimating causal effects from heterogeneous clinical trials without a common control group

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Abstract

Background The advent of clinical trial data sharing platforms has created opportunities for making new discoveries and answering important questions using already collected data. However, existing methods for meta-analyzing these data require the presence of shared control groups across studies, signifcantly limiting the number of questions that can be confdently addressed. We sought to develop a method for meta-analyzing potentially heterogeneous clinical trials even in the absence of a common control group.

Methods This work was conducted within the context of a broader effort to study comparative efficacy in Crohn's disease. Following a search of clnicaltrials.gov we obtained access to the individual participant data from nine trials of FDA-approved treatments in Crohn's Disease (*N*=3392). We developed a method involving sequences of regression and simulation to separately model the placebo- and drug-attributable efects, and to simulate head-to-head trials against an appropriately normalized background. We validated this method by comparing the outcome of a simulated trial comparing the efficacies of adalimumab and ustekinumab against the recently published results of SEAVUE, an actual head-to-head trial of these drugs. This study was pre-registered on PROSPERO (#157,827) prior to the completion of SEAVUE.

Results Using our method of sequential regression and simulation, we compared the week eight outcomes of two virtual cohorts subject to the same patient selection criteria as SEAVUE and treated with adalimumab or ustekinumab. Our primary analysis replicated the corresponding published results from SEAVUE (*p*=0.9). This fnding proved stable under multiple sensitivity analyses.

Conclusions This new method may help reduce the bias of individual participant data meta-analyses, expand the scope of what can be learned from these already-collected data, and reduce the costs of obtaining high-quality evidence to guide patient care.

Keywords Individual participant data meta-analysis, Randomized clinical trials, Crohn's disease, Comparative effectiveness, Comparative efficacy, Evidence synthesis, Biostatistics

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Background

The individual participant data (IPD) meta-analysis of randomized trials is the gold-standard for clinical research [\[1](#page-11-0), [2](#page-11-1)]. Access to the raw data from trials afords investigators the opportunity to verify published results, ask new questions of these data, and uncover fndings with the potential to impact patient care.

Performing an IPD meta-analyses usually requires multiple trials with negligible heterogeneity across many dimensions: cohort defnition, randomization, blinding, parallel study arms, interventions, and outcomes [\[1](#page-11-0), [2](#page-11-1)]. Although this requirement ensures unbiased estimation, it substantially limits the number of meta-analyses that can be performed due to the rarity of replicate trials.

The method of mixed-effects regression is commonly used to address study heterogeneity when meta-analyzed trials include a shared control group (i.e. placebo). However, there is a paucity of methods for common situations where there is no shared control group across potential studies. The few methods that have been developed include naïve pooling [[3\]](#page-11-2), as well as the Bayesian method of power priors [[4–](#page-11-3)[7](#page-11-4)]. However, these methods fail to address the problem of cohort heterogeneity [[8,](#page-11-5) [9](#page-11-6)]. Another major limitation is the lack of external validation against prospective studies. The result of this methodological gap is the common practice of excluding uncontrolled studies from potential meta-analyses, and ultimately fewer research questions that we are statistically powered to answer using already-collected data.

Here we report a new method for meta-analyzing clinical trials data in the absence of a common control group. We illustrate our method of sequential regression and simulation in the context of a comparative efficacy analysis in Crohn's disease, an immune disorder of the gastrointestinal tract. We use the data from six placebocontrolled trials $(N=3153)$ to develop a model of the placebo efect, then apply this to three placebo-less trials (*N*=239) to normalize and separately model the drugattributable response. Finally, we validate the method by predicting the results of SEAVUE (NCT03464136), a recent head-to-head trial of ustekinumab versus adalimumab [\[10](#page-11-7)].

Methods

This study was approved by the UCSF IRB $(\text{\#18-24,588}).$ It was pre-registered on PROSPERO [[11\]](#page-11-8) (#157,827), YODA [\[12\]](#page-11-9), and Vivli [[13\]](#page-11-10) prior to the initiation of this work and the completion of SEAVUE.

Data access

In June 2019 we queried clinicaltrials.gov to identify studies for meta-analysis (Supplementary Fig. 1, Additional File [2,](#page-10-0) Table 1). Our inclusion criteria were completed, phase 2–4, randomized, double-blinded, interventional trials of FDA-approved treatments for Crohn's disease that capture the Crohn's Disease Activity Index (CDAI) at week eight relative to treatment initiation. Our initial search led to 90 candidate studies. Following manual review, we confirmed 9 studies as meeting these criteria (*N* =3392). We requested and were granted access to all IPD from these studies. We manually reviewed their major inclusion and exclusion criteria to ensure comparability, and confirmed that they were at low risk of bias using the Cochrane ROB2 tool.

Study design

We designed this study to emulate a hypothetical headto-head, parallel-design, efficacy trial randomizing participants to two treatment arms (Fig. [1](#page-3-0)). Although a typical meta-analytic study design would have involved pooling cohorts from several internally controlled, parallel-arm trials, this was not possible in our case. Many of the included studies involved open-label induction followed by a randomization event to continue or discontinue the treatment (Fig. [2](#page-3-1)). We noted that three of the nine trials did not include a parallel arm placebo cohort randomized at week 0 and followed for eight weeks. Thus, for our study, they were considered uncontrolled. As a frst step towards developing a method for handling this heterogeneity, we restricted our initial analyses to just the placebo-controlled trials (six trials; *N*=3153; Table [1](#page-4-0)). Subsequent analyses used a second set of placebo-less trials of adalimumab, one of the drugs compared in SEAVUE.

Quality control, harmonization

We performed extensive tests of data quality [\(Supple](#page-10-1)[mentary Methods,](#page-10-1) Additional file [1](#page-10-1)). These included reproducing published results from each trial cohort (Supplementary Figs. 3, 4, Additional File [2\)](#page-10-0). We used domain knowledge to select nine variables that were universally available across trials for subsequent modeling: Age, Sex, body mass index (BMI), baseline CDAI, c-reactive protein (CRP), history of tumor necrosis factor-alpha inhibitor (TNFi) use, steroid use, immunomodulator use, and ileal involvement.

3% of the participants had at least one missing covariate at baseline. Continuous variables were addressed by median imputation, and participants with missing categorical variables were dropped $(N=86)$. 11% of participants had a missing outcome at week eight. We used last-observation-carried-forward to impute these. This is the typical practice for the analysis of these

Fig. 1 Study overview. **A** Clinical trials were found using clinicaltrials.gov and sought for retrieval on the YODA and Vivli platforms. Individual participant data (IPD) from trials that collected CDAI scores at week 8 visits were then aggregated and harmonized. **B** Two linear mixed efect models—placebo-attributable and ADA-attributable—were developed from the harmonized data to partition the CDAI reduction based on baseline covariates (age, sex, BMI, etc.). Disease activity reduction was partitioned into placebo attributable (square) and drug-attributable (circle) efects. IPD (solid lines) were used to predict or simulate data (dashed lines). **C** Using the adalimumab (ADA) attributable model, we simulated the outcomes of the placebo group from the ustekinumab (UST) trials under a counterfactual scenario where they had instead been assigned to receive ADA. **D** Results from a simulated head-to-head trial were compared against a recently completed head-to-head trial, SEAVUE, to externally validate the proposed method

Fig. 2 Summary of randomized controlled trial study designs. Data harmonization required careful understanding of the study designs. All treatment arms that involved 8 weeks of consistent exposure to either placebo or (blue) or active treatment at the FDA-approved doses (red) were included. R=randomized and blinded; O=open label

trials in regulatory submissions and was the prespecifed approach in the protocols of the included trials.

Some important variables could not be included in this study. Ethnicity was not collected in most trials. Race was missing in some trials, but when it was captured, it refected signifcant imbalance (88% of participants were white). Other variables like disease behavior and duration were not uniformly captured across studies.

Table 1 Characterization of baseline covariates of included studies and simulated head-to-head trial. Placebo arms from the CLASSIC, EXTEND, and NCT02499783 studies were **Table 1** Characterization of baseline covariates of included studies and simulated head-to-head trial. Placebo arms from the CLASSIC, EXTEND, and NCT02499783 studies were
not included due to the absence of an 8-week paral =tumor necrosis factor inhibitor =c-reactive protein, TNFi

Modeling assumptions

We incorporated several assumptions when developing and interpreting candidate models. We assumed that the observed week eight reduction in CDAI refected a combination of two distinct efects: a drug-independent (i.e., placebo) efect and drug-attributable efect. These effects were separately modeled as a function of the above predictors and study year. The justification for this is briefy summarized below and additionally presented graphically (Fig. [3\)](#page-7-0).

The placebo effect was modeled as a function of the nine covariates as well as predictors of trial-specifc heterogeneity. We assumed that much of the spontaneous improvement seen in placebo-assigned participants was related to regression to the mean, as study participation was limited to patients with currently active Crohn's disease. Conversely, we assumed that failure to spontaneously improve was likely to refect chronic and cumulative disease burden with relative stability in symptoms. Thus, variables corresponding to concomitant and prior treatments were treated as proxies of chronic disease burden and included as predictors. Lastly, we considered other infuences on overall heterogeneity, including diferences in cohorts, data capture, outcome ascertainment, and study personnel. To account for these sources of variation, we included study year as well as trial identifer as additional covariates. In mixed-efect models, trial was included as a random efect. Other covariates were fxed efects.

The drug-attributable effect was separately modeled as a function of these same covariates, refecting drug-specifc (interaction) efects on the outcome. Many of these covariates are well-established as modifers of treatment response, such as a history of TNFi and immunomodulators use [\[4](#page-11-3)]. Others (CRP, baseline CDAI) are proxies of bowel infammation, the target of these medications. These variables were included to maximize the explained variation in the outcome.

Development and assessment of a placebo model

We ft a linear mixed efects model utilizing all nine predictors as well as study year as predictors of the placebo efect. To minimize the risks of residual bias due to model misspecifcation (e.g., non-linearities, unmodeled interactions), we compared the predictive performance of this model against other statistical and machine learning models. We further evaluated this model from the perspective of being used to impute unmeasured placebo efects, and thus normalize different trials to the same background. We performed a leave-one-trial-out analysis and inspected the trialaveraged residuals.

Estimation of the drug-attributable efect

To normalize the responses of drug-assigned cohorts that lacked a within-study, parallel-arm control group, we used the fnalized placebo model to simulate and subsequently partition their overall response into drugindependent and drug-dependent (i.e., drug attributable) components (Fig. [1](#page-3-0)b). We performed this using the data from the adalimumab trials $(N=239)$ because they were all lacking an 8-week continuous placebo group and thus required normalization. From the outcomes of these patients, we subtracted the conditional mean outcomes associated with the placebo efect and used the residuals as the new outcome variable of a second mixed-efects model to estimate the adalimumab-attributable efect.

Validation, sensitivity analyses

Using the covariates associated with the placebo recipients of the ustekinumab trials, we used the adalimumab-attributable regression model to simulate their counterfactual week 8 outcomes had they received adalimumab instead (Fig. [1c](#page-3-0)). We identifed the subset of these virtual patients who were naïve to TNFi (an additional inclusion criterion from SEAVUE) and did the same with the ustekinumab recipients. We compared their week 8 outcomes using the same defnition of clinical remission as used in SEAVUE (CDAI < 150) and performed a Fisher's exact test to compare our results with SEAVUE's. We tested the robustness of our result using three sensitivity analyses: 1) removing ENACT and ENCORE from the dataset due to>10% missingness of outcome data, 2) removing participants with missing outcomes, and 3) removing the ustekinumab trials from the placebo model training data, to address potential information leakage. Lastly, we compared our results with what we might have found had we not used our method to normalize cohorts.

Results

See Fig. [1](#page-3-0) for an overview. This method was originally developed in the context of an existing effort to study comparative efficacy in Crohn's disease by reanalyzing the IPD of corresponding clinical trials. As the frst step towards this goal, we sought to address the problem of meta-analyzing data from several potentially heterogeneous trials lacking a common control group.

Data access

We queried clinicaltrials.gov and performed manual review to confrm 16 trials as meeting these criteria: completed, phase 2–4, randomized, double-blinded, interventional trials of FDA-approved treatments for Crohn's disease as of June 2019 (Fig. [1](#page-3-0)a, Supplementary Fig. 1, Additional fle [2\)](#page-10-0). Included trials had common inclusion/ exclusion criteria or had participant-level data available

Fig. 3 A directed acyclic graph of the modeling strategy. A A directed acyclic graph (DAG) of the drug attributable effect. In addition to the active treatment itself, patient demographics (age, sex, BMI), baseline Crohn's disease activity (baseline CDAI, CRP, location), and treatment history (prior use of TNFis, current use of oral corticosteroids and immunomodulators) are all modelled as contributing to the drug attributable efect. The non-drug covariates are efect modifers and are implicitly modeled as two-way interaction terms with the active drug. **B** A DAG of the drug independent efect (i.e., placebo efect). The same covariates except for the treatment term are modeled as efect modifers and are implicitly represented as two-way interactions with the placebo efect. **C** Drug attributable and drug independent efects have additive efects on the overall clinical remission at week 8 (CDAI<150), with any individual trial refecting a noisy measurement of the true efect due to unmodeled heterogeneity in study design and execution (random efect)

to control for this heterogeneity (Supplementary Table 1, Additional File 2). They all measured the same endpoint (CDAI) at week eight and were at low risk of bias (Supplementary Fig. 2, Additional File 2). We obtained access to the IPD for 15 studies (*N*=5703), corresponding to trials of all six FDA-approved biologics as of 2019.

Development and assessment of a placebo model

We fit a linear mixed effects model utilizing nine clinical features and study year as predictors of the placebo efect (Fig. [1](#page-3-0)b, Table [2](#page-7-1)). To minimize the risk of residual bias due to model misspecifcation, we compared the predictive performance of this model against other

machine learning models (Supplementary Table 2, Additional file [2](#page-10-0)). We found no significant differences in the root-mean-squared-error. Thus, we selected the mixed-efects model for downstream analyses.

We evaluated this model from the perspective of being used to impute unmeasured placebo efects, and thus normalize diferent trials to the same background placebo response. A leave-one-trial-out analysis suggested that the model predictions were robust and unbiased (Supplementary Figs. 4, 5, Additional fle [2](#page-10-0)). The trial-averaged residuals were consistent with normality ($p = 0.4$; Shapiro–Wilk test).

Table 2 Mixed effect linear regression outputs for the placebo attributable (*n*=1310) and ADA attributable (*n*=239) models. For training, Year was centered by subtracting 2000, Baseline CDAI was centered by subtracting 300, Age was centered by subtracting 35, BMI was centered by subtracting 20, and CRP (mg/L) was centered by subtracting 10. A. The placebo attributable model (intraclass correlation coefficient 0.02) trial random intercepts were found to be -12.808 (PRECISE1), -7.975 (UNITI1), -6.328 (CERTIFI), 6.077 (ENACT), 8.669 (ENCORE), and 12.366 (UNITI2). B. ADA attributable model (intraclass correlation coefcient 0.05) trial random intercepts were found to be -20.215 (CLASSIC), 9.439 (EXTEND), and 10.775 (NCT02499783)

We noted that the unmodeled variation in the placebo efect was relatively large and was independent of the choice of model (Supplementary Table 2, Additional file 2). These results explain the large placebo effects that have been seen in Crohn's disease randomized trials (regression to the mean) and suggest that more work will be needed to improve the measurement of Crohn's disease activity.

To study the placebo efect and identify potential opportunities to improve trial efficiency, we reviewed all signifcant predictors. A history of TNFi was associated with a 38-point reduction in the placebo effect. We interpreted this as refecting a greater cumulative disease burden in patients who failed to improve with TNFis, with disease complications (e.g., minor intestinal strictures) that are unlikely to spontaneously regress over 8 weeks. Similarly, CRP was a negative predictor, suggesting that untreated acute infammation is unlikely to improve over short time periods. The baseline CDAI was a positive predictor, likely refecting regression to the mean efects. Age, sex, BMI, concomitant medications, and ileal involvement were not found signifcant, potentially due to multicollinearity.

Estimation of the drug-attributable efect

We sought to normalize the responses of drug-assigned cohorts that lacked a within-study, parallel-arm control group. Our strategy was to use the fnalized placebo model to partition the overall response into drug-independent and drug-attributable components (Fig. [1b](#page-3-0)). We applied this approach to the data from three study cohorts assigned to receive adalimumab at the FDAapproved dose for treatment induction (*N*=239; Table [1](#page-4-0)). We selected this medication because it is one of the two treatments that were compared against each other in SEAVUE, the target of our emulation and validation efforts.

We used the coefficients of the fitted placebo model to predict and remove the placebo-attributable component from the observed outcomes of these participants. The residuals from this process were interpreted as refecting the adalimumab-attributable efect (Fig. [1](#page-3-0)b). Across these patients the mean drug-attributable CDAI reduction was 68 points. We used these residuals to ft a second model for the adalimumab-attributable efect (Table [2\)](#page-7-1).

As an exploratory analysis we reviewed the signifcant predictors of a response to adalimumab and compared these to the corresponding results from the placebo model. Although the sample size was relatively small, we noted a strong signal for age as a negative predictor: additional decades of life were associated with an 18-point reduction in the response to adalimumab. Interestingly, the direction of this efect was the opposite of that seen in the placebo-only model, suggesting that this coefficient might not have been identifed as signifcant had it not been handled as an interaction term as we did.

External validation

To validate our method, we designed an *in-silico* study to emulate SEAVUE, the only head-to-head study of FDA-approved biologics for Crohn's disease to date [\[3](#page-11-2)]. In SEAVUE, biologic-naive patients with active Crohn's disease were randomly to receive either adalimumab or ustekinumab as treatment. The primary endpoint was clinical remission at week 52, defned as a CDAI less than 150. Secondary endpoints included clinical remission at the time of all study visits, including week eight.

We identifed all participants from the three ustekinumab-related trials who were biologic-naive. We identifed 149 subjects who were assigned to ustekinumab and 135 participants assigned to placebo. We noted that the observed responses of the 135 placebo recipients refected a combination of individual-specifc variability and trial-specifc variability (Fig. [3](#page-7-0)). We therefore reasoned that to simulate the efect of treatment assignment, we needed to 'add back' the conditional mean efect associated with adalimumab to the outcomes of the placebo recipients (Fig. $1c$ $1c$). Using the model coefficients identifed in the adalimumab-attributable regression model (Table [2\)](#page-7-1), we computed and added this extra reduction in the CDAI to the observed week eight outcomes of the placebo cohort.

Finally, we computed the proportion of patients who were in clinical remission at week eight, comparing the results of the observed ustekinumab recipients with that of the patients simulated to have received adalimumab and subject to the same background placebo efect (Fig. [1d](#page-3-0)). We found that ustekinumab and adalimumab appeared to be equally efficacious, with 45% and 46% of the cohorts in remission. This result closely matched that of SEAVUE ($p=0.9$), which found 50% and 48% of these corresponding cohorts in remission (Table [3](#page-9-0)). Our simulated trial was similar in sample size to SEAVUE, with 149 and 135 patients receiving ustekinumab and adalimumab in our study, compared to 191 and 195 in SEAVUE.

We tested the robustness of this result using three sensitivity analyses. In the frst we removed two trials (PRECISE1, ENACT) associated with the greatest degree of outcome missing data (Supplementary Fig. 3, Additional File [2\)](#page-10-0). In the second, we performed a complete case analysis (deleted patient data associated with missing outcomes) as an alternative to last-observationcarried-forward imputation. In the third we removed all participant data emanating from an ustekinumab trial from the placebo training data, to address a possibility of information leakage. Our results remained

Table 3 Comparison of clinical remission rates at week 8 for the TNF-naive ustekinumab (UST) cohort and TNF-naive adalimumab (ADA) cohort for the SEAVUE study, our primary analysis (simulation of SEAVUE), sensitivity analyses, and negative control. Because missing week 8 CDAI values were highest for trials PRECISE1 and ENACT, their participant-level data was removed (N=1482) from the frst sensitivity analysis to account for potential bias. In the complete case sensitivity analysis, all participants with missing week 8 CDAI values (N=361) were removed. In the information leakage sensitivity analysis, participants from an ustekinumab study (*N*=1191) were removed from training the placebo-attributable model to avoid potential information leakage when simulating the adalimumab (ADA) arm (Fig. [1](#page-3-0)c). The negative control summarizes the clinical remission rates at week 8 for TNF-naive participants from the adalimumab studies without applying our regression-based correction method. The fnal column corresponds to the results of null hypothesis testing, that of no statistically signifcant diference between each simulated result and the published SEAVUE results

unchanged over all sensitivity analyses (Table [3\)](#page-9-0), supporting the robustness of our primary fndings as well as the validity of our overall methodology.

Finally, we sought to evaluate the value of using our modeling approach compared to a simpler approach using published trial results. One barrier we noted to the latter was that the aggregated response of the TNFinaive subcohorts at week eight was only published in one out of the six trials that we included for this comparison of ustekinumab and adalimumab, making it impossible to emulate SEAVUE using this approach. Separate from this, and to specifcally evaluate the value of normalizing disparate cohorts using placebo models, we simulated the potential results of our head-tohead assessment without a normalization step. Under this scenario, the unnormalized adalimumab cohort in clinical remission was 50% (Table [3\)](#page-9-0). While this was not statistically signifcant compared to the observed ustekinumab arm (45%; $p=0.4$), it reflects a trend towards a diference. We interpreted this as refecting a degree of bias that could plausibly result in false positives in other similar studies, but one that is analytically controllable using our method.

Discussion

We developed a new method for meta-analyzing individual participant data (IPD) from heterogenous randomized trials lacking a shared control group. We validated our methodology by successfully reproducing a major endpoint of SEAVUE, a recent head-to-head trial of biologic therapies in Crohn's disease [\[3](#page-11-2)]. Our method involved several steps (Fig. [1\)](#page-3-0):

- Identifying and isolating parallel arm cohorts from the available trials
- Harmonization and quality control
- Separately modeling the placebo effect from drugattributable efects
- Sequentially partitioning and assembling different sources of variation to accurately simulate the outcomes of a suitably normalized comparator group.

After decades of calls for greater data sharing [[14–](#page-12-0)[16](#page-12-1)] we are now seeing many new platforms for accessing clinical trials data. The availability of these data has opened opportunities for researchers to verify published results as well as answer new questions using these data. This has never been more important, with the cost of new phase 3 clinical trials current at \$20 M and climbing [\[17\]](#page-12-2).

Although the growing availability of IPD portends well for the future of research, it has revealed new analytical challenges that require new methods. Existing methods for conducting IPD meta-analyses typically involve including trials with near-identical study designs, including fully parallel-design cohorts and shared placebo comparator arms. When these criteria are not met, problematic trials are often excluded from a given meta-analysis, sometimes in subtle ways. This substantially limits the numbers of questions that might already be answerable using existing clinical data. In some cases, this common practice might even introduce bias.

This work suggests that there may be better ways to handle this heterogeneity and discover new and trustworthy signals from these data. This method as well as extensions therein may substantially increase the numbers of studies that can be done, uncovering new evidence on

comparative efficacy, safety, and ultimately precision medicine. Taking the example of Crohn's disease, a major motivation for conducting the SEAVUE trial is the current level of uncertainty regarding the comparative efficacy of already approved treatments. While the use of causal inference methods to emulate randomized trials from observational data are receiving great attention, our approach leverages already randomized data and is much less susceptible to bias. Thus, methods such as what we propose here can address these gaps, particularly as more therapies are approved and thus the number of potential head-to-head comparisons grows exponentially.

While we have illustrated this methodology in a comparative efficacy analysis, this approach may have signifcant value in other contexts. Models for the placebo efect, such as we demonstrate here, may help improve the design and statistical power of clinical trials across diseases. Moreover, the use of cohort normalization methods may be useful to improve the robustness of external control arm studies. These are studies that typically utilize real-world data to draw indirect inferences against controlled cohorts, typically single-arm intervention studies. However, our analysis suggests that a major driver of the large placebo efects in Crohn's disease is the large unmodeled variation in the CDAI. Future work is needed to improve the measurement of Crohn's disease activity.

We acknowledge several limitations. First, although we undertook extensive eforts to harmonize the data, we could not perfectly reproduce all covariate statistics as published. It is likely that we could have overcome these issues with access to the original analytical code. Nonetheless, the degree of deviations from published results was small, and our primary results remained robust to many sensitivity analyses. Future efforts involving preharmonization to a common data model may improve the reproducibility and feasibility of these IPD meta-analyses. Second, we were unable to include many important covariates like race and ethnicity. Most included studies did not capture ethnicity. Some studies did capture race but showed evidence of signifcant skew towards white participants. This likely reflects the historical underrecognition of the importance of these factors.

Closely related to this point is potentially important role of variable selection and model selection. Our methodology rests on the ability to accurately model and neutralize interstudy heterogeneity with captured covariates. This itself is a function of what covariates were captured, which is subject to clinical knowledge about a given disease at the time that diferent trials were conducted. It is also a function of the model form. Future studies are needed to explore the sensitivity of this approach to unmodeled efect modifers and model misspecifcation.

Lastly, we note that our validation was somewhat underpowered and was performed in the context of just one disease. This is largely a function of the relative rarity of clinical trials (the source of our data and sample size), and especially head-to-head trials like SEAVUE. This underscores the importance of methods for learning more from these small but high-quality data. Future studies are needed to confrm the robustness and generalizability of our methodology to other diseases.

Conclusion

In conclusion, we developed a new method for meta-analyzing data from heterogeneous trials lacking a common control group. We validated this method by reproducing the results of a recent comparative efficacy trial using pre-existing data. We are sharing our code for others to replicate and build upon these methods, and ultimately uncover new insights using the data we already have.

Author's contributors

VAR conceived the study and obtained access to the data. SW, VGR, DVA, and VAR designed the study, analyzed the data, and drafted the manuscript. AM performed the risk of bias assessment. SW performed an independent review of the analytical code. VAR, SW, VGR, DVA, AM, and AJB interpreted the data and critically edited the manuscript.

Abbreviations

- ADA Adalimumab
- BMI Body Mass Index
- CDAI Crohn's Disease Activity Index
- CRP C-Reactive Protein
- CZP Certolizumab Pego
- IPD Individual Participant Data
- NTZ Natalizumab
- TNFi Tumor Necrosis Factor inhibitor
- UST Ustekinumab

Supplementary Information

The online version contains supplementary material available at [https://doi.](https://doi.org/10.1186/s12874-023-02020-5) [org/10.1186/s12874-023-02020-5](https://doi.org/10.1186/s12874-023-02020-5).

Additional fle 1: Supplementary methods.

Additional fle 2: Figure S1. PRISMA-IPD fow diagram. **Figure S2.** Risk of bias. **Figure S3.** Reproducibility of published data. **Figure S4.** Leave-onetrial-out analysis. **Figure S5.** Checking model assumptions. **Table S1.** Major inclusion/exclusion criteria of included studies. **Table S2.** Model selection.

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Authors' contributions

VAR conceived the study and obtained access to the data. SW, VGR, DVA, and VAR designed the study, analyzed the data, and drafted the manuscript. AM performed the risk of bias assessment. SW performed an independent review of the analytical code. VAR, SW, VGR, DVA, AM, and AJB interpreted the data and critically edited the manuscript.

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Availability of data and materials

The data that support the fndings of this study are available from YODA and Vivli, but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of YODA and Vivli. The analytical R code for data processing and modeling can be found in the GitHub repository, [https://github.com/ibd-ipd](https://github.com/ibd-ipd-ma/SequentialRegressionSimulation)[ma/SequentialRegressionSimulation](https://github.com/ibd-ipd-ma/SequentialRegressionSimulation).

Declarations

Ethics approval and consent to participate

This study was approved in accordance with the Declaration of Helsinki and with the oversight of the UCSF IRB (#18–24588). This review board waived the need for participant consent given that the underlying data had already been deidentifed by the owners of the raw data.

Consent for publication

Not applicable.

Competing interests

VAR received grant support from Janssen Inc and Alnylam Inc for unrelated work during this study. DVA is currently an employee at Bristol Myers Squibb. AJB is a co-founder and consultant to Personalis and NuMedii; consultant to Mango Tree Corporation, and in the recent past, Samsung, 10 x Genomics, Helix, Pathway Genomics, and Verinata (Illumina); has served on paid advisory panels or boards for Geisinger Health, Regenstrief Institute, Gerson Lehman Group, AlphaSights, Covance, Novartis, Genentech, and Merck, and Roche; is a shareholder in Personalis and NuMedii; is a minor shareholder in Apple, Meta (Facebook), Alphabet (Google), Microsoft, Amazon, Snap, 10 x Genomics, Illumina, Regeneron, Sanof, Pfzer, Royalty Pharma, Moderna, Sutro, Doximity, BioNtech, Invitae, Pacifc Biosciences, Editas Medicine, Nuna Health, Assay Depot, and Vet24seven, and several other non-health related companies and mutual funds; and has received honoraria and travel reimbursement for

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