A Novel Strategy to Identify Placebo Responders: Prediction Index of Clinical and Biological Markers in the EMBARC Trial

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Keywords
Placebo responder · Prediction index · EMBARC trial

Abstract
Background: One in three clinical trial patients with major depressive disorder report symptomatic improvement with placebo. Strategies to mitigate the effect of placebo responses have focused on modifying study design with variable success. Identifying and excluding or controlling for individuals with a high likelihood of responding to placebo may improve clinical trial efficiency and avoid unnecessary medication trials. Methods: Participants included those assigned to the placebo arm \( n = 141 \) of the Establishing Modulators and Biosignatures for Antidepressant Response in Clinical Care (EMBARC) trial. The elastic net was used to evaluate 283 baseline clinical, behavioral, imaging, and electrophysiological variables to identify the most robust yet parsimonious features that predicted depression severity at the end of the double-blind 8-week trial. Variables retained in at least 50% of the 100 imputed data sets were used in a Bayesian multiple linear regression model to simultaneously predict the probabilities of response and remission. Results: Lower baseline depression severity, younger age, absence of melancholic features or history of physical abuse, less anxious arousal, less anhedonia, less neuroticism, and higher average theta current density in the rostral anterior cingulate predicted a higher likelihood of improvement with placebo. The Bayesian model predicted remission and response with an actionable degree of accuracy (both AUC > 0.73). An interactive calculator was developed predicting the likelihood of placebo response at the individual level. Conclusion: Easy-

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to measure clinical, behavioral, and electrophysiological assessments can be used to identify placebo responders with a high degree of accuracy. Development of this calculator based on these findings can be used to identify potential placebo responders.

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Introduction

Major depressive disorder (MDD) affects 1 in 6 adults during their lifetime and is estimated to cost the USA over USD 200 billion per year [1, 2]. Yet, only one third of MDD patients achieve remission under any given antidepressant treatment [3], with over one third of the patients not responding to 2 or more antidepressants [4, 5]. Despite substantial financial investments [6, 7], efforts to improve MDD treatment outcomes by developing nonmonoaminergic antidepressants have failed. High placebo response rates have been a common factor in the failure of several novel antidepressant medications in phase 2 and 3 clinical trials [8, 9]. With limited/variable success, attempts to mitigate the increase in placebo response rate over the last three decades [10] have focused mainly on study design-related issues such as: (1) increasing the sample size to account for smaller drug-placebo difference; (2) incorporating a placebo lead-in period; (3) controlling measurement factors by using central raters or standardized interviews; or (4) implementing innovative study designs such as the sequential parallel comparison design [11–14]. A patient-centered approach, which identifies the individual characteristics that define placebo responders, may offer an alternative way to reduce the placebo response rate in clinical trials. These characteristics may be useful, additionally, in clinical practice, where efforts to maximize the placebo response can improve treatment outcomes [15].

The efforts to identify predictors of placebo response have been limited by the focus on subjective disease severity assessment and demographic features. This has frequently resulted in conflicting findings that often did not consider a wide variety of potential predictors from behavioral and biological domains. Among individual factors, low pretreatment symptom severity has been associated with higher likelihood of placebo response [16, 17]. However, other factors such as gender and age, while significant in venlafaxine versus placebo studies [18], did not replicate in a meta-analysis by Holmes et al. [19]. The neurobiological basis of the placebo response is characterized by an increase in the metabolic activity of the frontal and striatal cortical regions [20] and increased endogenous opioid release in the subgenual anterior cingulate cortex, nucleus accumbens, midline thalamus, and amygdala [21, 22]. The placebo response has also been linked to increased baseline resting state functional connectivity of the rostral anterior cingulate cortex (rACC) within the salience network [23] and to increased pretreatment rACC activity in 2 EEG studies [24, 25]. Previous studies of neuroimaging biomarkers of placebo response have been limited by small sample sizes and a lack of comparison with other clinical and biobehavioral markers [21, 23]. Despite extensive research to characterize placebo responders, a set of clinical and objective predictors and tools to filter out this subgroup from clinical trials has yet to be agreed upon and implemented. There is also a potential real-world clinical implication. A portion of treatment-seeking depressed individuals might not need a long-term antidepressant prescription if they are placebo responders. Hence, by identifying such placebo responders in advance, briefer low cost, low side effect interventions may be recommended for these particular patients.

The goal of this report is to identify a parsimonious set of markers among assessments across units of analyses (clinical, demographic, neuroimaging, electrophysiological, behavioral, and cognitive assessments) that most strongly predict the likelihood of placebo response and can be implemented in research settings and clinical practice. Traditional approaches to handling a statistical problem like this – such as a stepwise regression or factor analysis – are not well equipped to handle a rich database with (1) a large number of predictors relative to the number of subjects or (2) missing data. To attain this goal of identifying the parsimonious set of markers, we utilized data from the Establishing Moderators and Biosignatures for Antidepressant Response in Clinical Care (EMBARC) trial to systematically explore 283 variables using an advanced variable selection method to identify the variables that most strongly predict the likelihood of improvement with placebo. Next, we used a novel Bayesian method to simultaneously predict the degree of symptom change, as well as the probability of remission and response using these variables. Such a method allows flexibility in the definition of “placebo response” and can be implemented using a simple web-based tool.

Methods

Participants

Participants for this report were recruited as part of the EMBARC trial, which is a 2-stage, multisite, double-blind randomized controlled study designed to evaluate possible modera-
Prediction Index in the EMBARC Trial

Tors and mediators of antidepressant treatment response in patients with MDD. The study design has been described in detail by Trivedi et al. [26]. Data for this report were obtained from participants (n = 141) who were assigned to the placebo arm during stage 1 of the EMBARC trial and received at least 1 placebo dose. Stage 1 of the EMBARC trial included an 8-week double-blind placebo-controlled trial of sertraline that enrolled 309 participants. Participants were 18–65 years old with MDD diagnosed by the Structured Clinical Interview for DSM-IV Axis I Disorders (SCID). A complete list of inclusion/exclusion criteria and their justification has been previously published [26]. Briefly, participants had to have a Quick Inventory of Depressive Symptomatology Self-Report (QIDS-SR) score ≥14 at both screening and randomization visits. Only patients whose first major depressive episode began before the age of 30, with either a chronic current episode (duration ≥2 years) or recurrent MDD (at least 2 lifetime episodes) were eligible. Participants were recruited from 4 clinical sites (UT Southwestern, Massachusetts General Hospital, Columbia University, and University of Michigan) after approval from the institutional review boards at each site and after obtaining written informed consent from each participant. In the EMBARC trial, 141 participants were assigned to placebo and received at least 1 dose and constitute the sample for this report.

Outcome

The 17-item Hamilton Rating Scale for Depression (HAMD17): the 17 items of this clinician-rated scale to assess depression severity have 3–5 choices which are scored either from 0 to 2 or 0 to 4 [27]. The individual items are summed to measure depression severity [none (<6), mild (6–13), moderate (14–18), severe (19–23), and very severe (>23)] [27]. In the EMBARC trial, the HAMD17 was administered at each study visit (baseline and weeks 1, 2, 3, 4, 6, and 8 of stage 1). For the purpose of this report, the primary outcome was the last observed HAMD17 score, which also contributed to the definitions of outcome of remission (defined as last observed HAMD17 ≤7) and response (a decrease in HAMD17 of ≥50% at the last observation as compared to the baseline). In total, 126 subjects completed 8 weeks of treatment, 5 had their last observed value at week 6, 4 each at weeks 3 and 4, and 1 each at weeks 1 and 2.

Baseline Assessments

Clinical and Demographic. 42 clinical and demographic variables were identified a priori. In addition to standard demographic characteristics (e.g., sex, age), clinical factors previously reported to have predictive power for treatment outcome were assessed for moderating effects [28]. Clinical variables were derived from the HAMD17, Altman Self-Rating Mania Scale, Anger Attacks Questionnaire, Childhood Trauma Questionnaire, Concise Associated Symptoms Tracking, Concise Health Risk Tracking, Edinburgh Handedness Inventory, Family History Screen, Mood and Anxiety Symptom Questionnaire (MASQ-30), NEO Five-Factor Inventory, QIDS-SR, Self-Administered Comorbidity Questionnaire, Snaith-Hamilton Pleasure Scale, Social Adjustment Scale Short Form, SCID, and Wechsler Abbreviated Scale of Intelligence. Sixteen of the variables had at least some missing data, with the percentage ranging from 0.7 to 11.3%.

Behavioral and Cognitive Performance. 14 behavioral and cognitive performance measures were assessed for moderating effects. Reaction time, psychomotor slowing, cognitive control, working memory, and reward learning were derived from the following tasks: Choice Reaction Time; Word Fluency Test; “A not B” Working Memory task, Flanker, and Probabilistic Reward Task [29]. All variables had at least some missing data, ranging from 0.7 to 17.7%.

Electrophysiological. 15 EEG measures, derived from a pretreatment EEG recording, were preselected for moderator analysis. These included resting (task-free) measures of power in the alpha (recorded during eyes-closed and eyes-open) and theta (eyes-closed) bands, loudness dependence of auditory evoked potentials, and theta current density extracted from the rACC [25]. For details regarding paradigms and methods, see Tenke et al. [30]. All variables had at least some missing data, ranging from 5.0 to 20.6%.

Neuroimaging. 212 structural or functional imaging variables were analyzed for moderating effects [26]. Structural imaging variables included volumetric magnetic resonance imaging (MRI) assessments using FreeSurfer and diffusion tensor imaging [31, 32]. Functional imaging variables were collected from MRI during resting state (2 separate blocks collected on the same day; pre- and post-task), the Reward Processing Task, and the Emotional Recognition Task. All variables had at least some missing data, ranging from 28.4 to 34.8%.

Statistical Analysis Plan

To accomplish the goals of the analysis, three statistical problems had to be addressed: performing variable selection, accounting for missing data, and estimating the magnitude and direction of the regression coefficients for the variables with the most predictive power after dealing with the first two listed problems. Online supplementary Figure 1 (for all online suppl. material, see www.karger.com/doi/10.1159/000491093) displays the progression of the analysis. A high-level summary of the analysis plan is given below; for a more detailed description (with more technical aspects addressed), see the online supplementary file.

Variable Selection Method

With the goal to identify a parsimonious set of variables that predict response to placebo, the elastic net [33] – a penalized regression technique that is becoming increasingly popular in the statistical literature [34] with respect to variable selection – was used. This procedure introduces bias to the regression estimates in the form of shrinkage – that is, it pulls the estimates towards 0 – while simultaneously reducing the variance of the estimates in order to increase the overall predictive power. The elastic net was selected over other methods such as random forests, support vector machines, gradient boosting machines, and multivariate adaptive regression splines, to ensure the analysis would result in an easily interpretable linear model. The aforementioned alternatives – while popular as tools for prediction – often require large amounts of data [35] and can be difficult to interpret due to the inclusion of complex, nonlinear interactions [36]. The elastic net was implemented via the glmnet package [37] in R 3.3.3 [38].

Approach for Missing Data

Due to missing values of predictor baseline variables, multivariate imputation by chained equations was used [39, 40] under the assumption that the data were missing at random [41] to generate a total of 100 imputed data sets. No imputation was used for the outcome variable, (as described above in the “Outcome” subsection) the last observed HAMD17 score. As there is no universally accepted approach to apply variable selection methods in the
Table 1. Baseline sociodemographic and clinical features of EMBARC trial participants randomized to placebo who received at least 1 dose (n = 141)

<table>
<thead>
<tr>
<th>Categorical variables, n (%)</th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>52 (36.9)</td>
</tr>
<tr>
<td>Female</td>
<td>89 (63.1)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>98 (69.5)</td>
</tr>
<tr>
<td>Black</td>
<td>23 (16.3)</td>
</tr>
<tr>
<td>Other</td>
<td>20 (14.2)</td>
</tr>
<tr>
<td>Employment status</td>
<td></td>
</tr>
<tr>
<td>Employed</td>
<td>78 (55.3)</td>
</tr>
<tr>
<td>Unemployed</td>
<td>58 (41.1)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Continuous variables (mean ± SD)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>37.4±12.9</td>
</tr>
<tr>
<td>Age of onset, years</td>
<td>16.4±5.6</td>
</tr>
<tr>
<td>Years of education</td>
<td>15.2±2.7</td>
</tr>
<tr>
<td>Number of MDE</td>
<td>4±2</td>
</tr>
<tr>
<td>Duration of current episode, months</td>
<td>41.5±75.8</td>
</tr>
<tr>
<td>QIDS-SR</td>
<td>17.8±2.7</td>
</tr>
<tr>
<td>HAMD_{17}</td>
<td>18.6±4.3</td>
</tr>
</tbody>
</table>

EMBARC, Establishing Moderators and Biosignatures for Antidepressant Response in Clinical Care; SD, standard deviation; MDE, major depressive episode; QIDS-SR, Quick Inventory of Depressive Symptomatology Self-Report version; HAMD_{17}, 17-item Hamilton Rating Scale for Depression. * Median and interquartile range reported due to outlying values.

To estimate the magnitude and direction of the regression coefficients for the variables selected by the elastic net, we employed a Bayesian linear regression model that kept the same outcome variable; this is beneficial for several reasons. First, we can mitigate some of the concerns that would arise were we to instead apply ordinary least squares regression after choosing variables via the elastic net [45, 46] by assigning prior distributions to the regression parameters that mimic the shrinkage that occurs when applying the elastic net to all variables. For more details, see the supplementary file or Makalic and Schmidt [47]. Next, we can simultaneously account for the multiply imputed data [48] by running separate analyses and then mixing posterior draws from each run [49, 50] to create a more comprehensive, singular posterior distribution. Finally, we can sample from the posterior distribution of the regression parameters to estimate the predicted HAMD_{17} score at the end of stage 1 (i.e., the outcome variable) and construct a posterior distribution of these predicted scores. This distribution can then be used to estimate the probability of remission (by calculating the proportion of posterior distribution scores ≤7 in the distribution) and response (by calculating the proportion of posterior distribution scores ≤50% of the baseline HAMD_{17} value) for each subject.

To correct for some of the biases that exist when training and testing a model on the same data, each imputed data set was bootstrapped 20 times (resulting in 2,000 imputed/bootstrapped data sets: 100 data sets × 20 bootstraps), and the Bayesian model was subsequently fit on each bootstrapped data set to better estimate the variability of the selected predictors; the posterior draws were then mixed as described earlier. While 20 bootstrap repetitions are smaller than one might typically see, given that there were 100 imputed data sets and 2,000 draws from the posterior distributions for each imputed/bootstrapped pair, we ended with a total of 4 million data points – making more than 20 bootstrap replications nearly computationally intractable. The Bayesian regression model was implemented using the bayesreg package in R [47].

To understand the individual contributions, predictions were made with each variable at a time (univariate area under the curve, AUC) and the decrement to overall model-fit was estimated after removal of variables one at a time.

Sensitivity Analysis
In addition to the a priori cutoff of retention in at least 50% of the data sets in the elastic net model for selection of variables in the previously described Bayesian model, the percentage of time each variable was retained was plotted to visually identify dropoffs in clustering of candidate variables. Visual inspection of this ordered index suggested two additional thresholds: variables retained in 100% of elastic net run and those retained in 30% of the runs; see supplementary Figure 2 for the ordered index. AUC curves are presented along with the comparison of these three thresholds (100, 50, and 30%) using the DeLong test [51], implemented in the pROC package in R [52].

Visualization of Study Findings
To demonstrate the practical application of the Bayesian linear regression model in clinical practice, an interactive calculator was developed using the Shiny package in R [53].

Results
Of the 141 participants randomized to the placebo arm of the EMBARC trial, the mean (standard deviation) HAMD_{17} at baseline was 18.6 (4.3) and at study exit it was 12.0 (7.5). At study exit, 47/141 (33.3%) attained remission, and 55/141 (39.0%) attained response. Baseline clinical and sociodemographic features are presented in Table 1.

Prediction of Placebo Outcomes
After applying the elastic net to 100 imputed data sets, 8 out of 283 variables were retained in at least 50% of the runs. These variables, listed in Table 2, included baseline HAMD_{17}, age, anhedonia, and anxious arousal as measured by MASQ, neuroticism, the presence of melancholic-
ic features, history of physical abuse, and average theta current density in the rACC. These variables were then entered in a Bayesian linear regression model to predict final HAMD$_{17}$ scores at the end of stage 1 of the EMBARC trial. Table 2 gives the posterior median values for the chosen set of predictors. Higher posterior median values predicted higher HAMD$_{17}$ scores at the end of stage 1 of the EMBARC trial, thus signifying worse outcomes. For example, a 1-unit increase in baseline HAMD$_{17}$ was associated with a final HAMD$_{17}$ score that was (on average) 0.52 points higher, holding all other predictors in the model constant. Thus, while higher baseline depression severity, age, neuroticism, anxiety, and anhedonia severity, as well as the presence of melancholic features and history of physical abuse, predicted worse outcomes with placebo, larger values of pretreatment theta current density localized to the rACC predicted better outcomes with placebo.

Taking advantage of the posterior distribution of the predicted HAMD$_{17}$ scores in the Bayesian framework, the probability of remission (HAMD$_{17} \leq 7$) and response (reduction of HAMD$_{17}$ by $\geq 50\%$) at the end of stage 1 of the EMBARC trial were estimated simultaneously and compared with the observed values at individual participant level; the receiver operating curve for both remission and response are shown in Figure 1. The AUC values (0.758 for remission and 0.728 for response) indicate a moderate fit, supporting the validity of using a Bayesian linear regression model to derive these values.

In univariate analyses to predict remission, baseline depression severity had the highest AUC and the greatest decrement in AUC to the model, as shown in Table 2. Two other variables with significant univariate AUC (i.e., their bootstrapped confidence interval did not include 0.50) were age and history of physical abuse. Notably, the highest univariate AUC (0.635 for baseline depression severity) was substantially lower than the AUC including all

<table>
<thead>
<tr>
<th>Variable</th>
<th>Posterior median</th>
<th>95% CI</th>
<th>Univariate AUC for remission</th>
<th>Decrement in AUC for remission upon removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline HAMD$_{17}$ score</td>
<td>0.52</td>
<td>0.186, 0.877</td>
<td>0.635 (0.541 to 0.729)</td>
<td>0.092 (–0.003 to 0.187)</td>
</tr>
<tr>
<td>Age</td>
<td>0.12</td>
<td>0.016, 0.222</td>
<td>0.611 (0.517 to 0.705)</td>
<td>0.058 (–0.032 to 0.148)</td>
</tr>
<tr>
<td>Melancholic depression indicator$^a$</td>
<td>2.06</td>
<td>–0.764, 4.959</td>
<td>0.388 (0.309 to 0.468)</td>
<td>0.003 (–0.077 to 0.083)</td>
</tr>
<tr>
<td>Anhedonia$^b$</td>
<td>0.13</td>
<td>–0.123, 0.367</td>
<td>0.537 (0.454 to 0.619)</td>
<td>0.017 (–0.067 to 0.100)</td>
</tr>
<tr>
<td>Anxious arousal$^c$</td>
<td>0.22</td>
<td>–0.085, 0.537</td>
<td>0.545 (0.444 to 0.646)</td>
<td>0.014 (–0.072 to 0.100)</td>
</tr>
<tr>
<td>Neuroticism$^d$</td>
<td>0.12</td>
<td>–0.092, 0.336</td>
<td>0.505 (0.411 to 0.600)</td>
<td>0.004 (–0.077 to 0.084)</td>
</tr>
<tr>
<td>Physical abuse$^e$</td>
<td>0.18</td>
<td>–0.185, 0.543</td>
<td>0.625 (0.533 to 0.718)</td>
<td>–0.010 (–0.090 to 0.070)</td>
</tr>
<tr>
<td>rACC theta current density</td>
<td>–5.93</td>
<td>–13.417, 1.166</td>
<td>0.510 (0.412 to 0.608)</td>
<td>0.019 (–0.067 to 0.105)</td>
</tr>
</tbody>
</table>

HAMD$_{17}$, 17-item Hamilton Rating Scale for Depression; rACC, rostral anterior cingulate. $^a$ Based on specifier questions on the SCID. $^b$ Anhedonic depression scale from the Mood and Anxiety Symptom Questionnaire. $^c$ Anxiety specific scale (anxious arousal) from the Mood and Anxiety Symptom Questionnaire. $^d$ Based on 12 neuroticism items from the NEO Five-Factor Inventory. $^e$ Scale from the Childhood Trauma Questionnaire.
In this large sample of depressed outpatients, a systematic exploration of a broad range of clinical and biological markers identified baseline depression severity, age, neuroticism, anhedonia, anxious arousal, presence of melanc}-
cholic features, history of physical abuse, and theta current density in the rACC as predictors of placebo response in a double-blind randomized clinical trial. These markers can be implemented within a Bayesian framework to simultaneously predict the likelihood of response as well as remission with an easy-to-use calculator. Clinicians or clinical researchers could use this calculator to make more informed decisions about whether to prescribe a treatment or enroll subjects in clinical trials with a placebo arm.

The current findings are consistent with previous reports that have found that higher baseline depression severity is associated with lower likelihood of improvement in the placebo arm [10, 17]. The most powerful single variable predicting poorer outcome in response to placebo treatment was greater depression severity. This is a very important finding, as enrichment in terms of severity of illness can be easily implemented in the context of a clinical trial. The failure to reduce placebo response by implementing a greater illness severity threshold for inclusion in studies has been attributed to the “grade inflation” when the severity of illness is determined only by clinicians at the site, where the bias towards enrollment can be significant, thereby driving up severity measures [12]. Independent verifications of subject severity at entry by themselves can reduce the placebo response by allowing a greater depression severity enrichment [55].

The finding that higher theta current density in the rACC predicts greater improvement with placebo differs from previous work by Korb et al. [56] which did not find any difference in rACC theta density between placebo responders and nonresponders. This difference may be related to the smaller sample size of the study of Korb et al. Moreover, in a later study, the same group reported that increased rACC theta current density predicted a greater placebo response. Notably, in the International Study to Predict Optimized Treatment in Depression (iSPOT-D)
trial, higher rACC theta was associated with worse outcomes in depressed outpatients treated with either escitalopram, sertraline, or placebo [57]. Theta current density and other objective biomarkers, such as resting-state connectivity, may offer greater predictive value at more cost but may prove over costly drug treatments that are potentially ineffective or harmful to the outcome.

There are several strengths of this report. The a priori defined threshold of 50% retention in elastic net runs has identified clinical and demographic assessments as well as electrophysiological measures, which are relatively inexpensive and easily implemented. The more expensive neuroimaging variables were retained when the threshold for inclusion was more liberal, i.e., included in ≥30% of elastic net runs. Arguably, the most meaningful benefit of fitting the model in the Bayesian framework was to take advantage of the posterior distribution of the predicted HAMD17 scores to simultaneously predict the likelihood of response and remission. In a more traditional analysis, three separate models would have to be run to accomplish this: a multiple linear regression to predict HAMD17 at exit and two logistic regression models, one with remission as the outcome and one with a response as the outcome. Additionally, the interactive calculator informed by our predictive model provides a web-based tool, which could be easily implemented in clinical practice or research settings to predict the likelihood of a placebo response at the individual level.

These findings should be interpreted in the context of certain limitations. Response to placebo is a complex phenomenon and may be related to a multitude of factors such as treatment setting, environmental factors, and illness or patient characteristics [58] that were not assessed in the EMBARC study. Consistent with the recommendations of Fava et al. [58], future clinical trials should comprehensively capture the “multifactorial ingredients of treatment outcome” to better understand the nondrug contributions to improvement with antidepressant treatments. Next, the mean HAMD-17 score in this report is lower than that typically reported for randomized controlled trials of antidepressant medications. Further, while extensive, the variable selection procedure may have missed some additional clinical or biological features that could increase the predictive ability of placebo response. However, when identified, such features may be integrated into future iterations of the calculator. The choice of elastic net was deliberate to maximize the practical application of these findings; nonetheless, this method may have missed more complicated (i.e., nonlinear) interactions between features that might be uncovered with other variable selection methods such as random forest. Lastly, due to the unique combination of variables collected for the EMBARC trial, we cannot validate the model on an external data set. The cross-validation and bootstrap procedures employed add strength to the analysis, but there is no substitute for external validation.

To conclude, a set of 8 clinical and biological markers can predict treatment outcomes with placebo with a fair degree of accuracy. It is also possible to further increase the accuracy of prediction if all 15 variables are included. By integrating these markers in an easy-to-use interactive calculator, the findings of this report can be implemented in research and clinical care.

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