Practice of Epidemiology

The “Dry-Run” Analysis: A Method for Evaluating Risk Scores for Confounding Control

Richard Wyss*, Ben B. Hansen, Alan R. Ellis, Joshua J. Gagne, Rishi J. Desai, Robert J. Glynn, and Til Stürmer

* Correspondence to Dr. Richard Wyss, Division of Pharmacoepidemiology and Pharmacoeconomics, Department of Medicine, Brigham and Women's Hospital and Harvard Medical School, Boston, 1620 Tremont Street, Suite 3030, Boston, MA 02120 (e-mail: rwyss@partners.org).

Initially submitted August 26, 2015; accepted for publication June 29, 2016.

A propensity score (PS) model’s ability to control confounding can be assessed by evaluating covariate balance across exposure groups after PS adjustment. The optimal strategy for evaluating a disease risk score (DRS) model’s ability to control confounding is less clear. DRS models cannot be evaluated through balance checks within the full population, and they are usually assessed through prediction diagnostics and goodness-of-fit tests. A proposed alternative is the “dry-run” analysis, which divides the unexposed population into “pseudo-exposed” and “pseudo-unexposed” groups so that differences on observed covariates resemble differences between the actual exposed and unexposed populations. With no exposure effect separating the pseudo-exposed and pseudo-unexposed groups, a DRS model is evaluated by its ability to retrieve an unconfounded null estimate after adjustment in this pseudo-population. We used simulations and an empirical example to compare traditional DRS performance metrics with the dry-run validation. In simulations, the dry run often improved assessment of confounding control, compared with the C statistic and goodness-of-fit tests. In the empirical example, PS and DRS matching gave similar results and showed good performance in terms of covariate balance (PS matching) and controlling confounding in the dry-run analysis (DRS matching). The dry-run analysis may prove useful in evaluating confounding control through DRS models.

causal inference; disease risk score; epidemiologic methods; prognostic score; propensity score

Abbreviations: DRS, disease risk score; PS, propensity score.

Summary scores that reduce baseline covariate information to a single dimension have become increasingly popular to control confounding in nonexperimental studies. The propensity score (PS), defined as the conditional probability of exposure given a set of observed covariates, has been the most widely used summary score (1, 2). An alternative to the PS is the prognostic score, often referred to as the disease risk score (DRS). Unlike the PS, which summarizes covariate associations with exposure, the DRS summarizes covariate associations with potential outcomes (3). Both the PS and the DRS control for confounding by acting as balancing scores. Rosenbaum and Rubin (1) showed that, upon conditioning on the PS, covariates are independent of, or balanced across, exposure groups. Hansen (3) showed that the DRS acts as a “prognostic balancing” score in that, conditional on the DRS, covariates are independent of the potential outcome under the control condition. Here, we will refer to the control condition as unexposed.

The DRS has not been as widely used as the PS for confounding control, but it can be advantageous in certain settings. The DRS provides a natural measure to evaluate effect measure modification (4–6). Although the PS allows researchers to detect and account for effect modification, it does not provide the best information for health-care providers in determining what subgroups of the patient population are most likely to benefit from a given exposure or treatment. Further, conditioning on the PS can be more restrictive than conditioning on the DRS (3, 7). A DRS-matched or stratified...
analysis can potentially allow researchers to compare a larger proportion of the population across exposure groups compared with PS analyses (7).

In practice, the PS and DRS are unknown and must be estimated from the available data. While both the estimated PS and DRS are susceptible to model misspecification, the DRS is particularly vulnerable because of the need to extrapolate and make additional assumptions when fitting the risk model (discussed further below) (3). Assessing the validity of fitted DRS models could improve the robustness of DRS analyses; however, although a number of studies have discussed and analyzed methods for evaluating PS models (8–14), there remains little discussion of how DRS models should be evaluated when the goal of the DRS is to control for confounding bias.

In this study we used simulations to compare metrics for evaluating DRS models in their ability to control confounding. We compared traditional metrics for evaluating risk models with an alternative strategy termed the “dry-run” analysis (15). We demonstrate the discussed concepts through an empirical example comparing dabigatran with warfarin for preventing ischemic stroke and all-cause mortality within a population of Medicare beneficiaries.

METHODS

Modeling the DRS

The DRS has typically been estimated either by fitting a regression model within the unexposed population and then extrapolating this model to predict disease risk for the full cohort or by fitting a regression model within the full cohort as a function of baseline covariates and exposure and then assigning risk scores after setting exposure status to zero (16–22). Hansen (3) discussed limitations to these strategies, both of which are examples of “same-sample” estimation. If the exposure effect is misspecified when fitting the risk model to the full cohort (e.g., omitting exposure-covariate interactions), then the estimated scores can carry information about the exposure effect. This noncancillarity in the estimated risk scores can potentially bias effect estimates and generate spurious suggestions of effect modification across levels of disease risk (3, 20). Fitting the DRS only to the unexposed population, however, can lead to overfitting, which can itself cause apparent effect modification and bias overall in effect estimates (3, 23, 24).

To avoid these problems, both Hansen (3) and Glynn et al. (23) have proposed using a historical set of unexposed subjects to fit the DRS model. This strategy can circumvent the problems associated with “same-sample” estimation, but it assumes that the effects of risk factors on the outcome, disease surveillance, and covariate definitions do not change over time. Violation of these assumptions could result in an estimated DRS model that is not generalizable to the study cohort (7, 25, 26).

These challenges in estimating the DRS highlight the importance of evaluating the validity of fitted DRS models as a way to control confounding. While a PS model’s ability to control confounding can be evaluated directly by assessing covariate balance across exposure groups after PS adjustment, the prognostic balance resulting from a DRS model can be evaluated only among unexposed individuals, where the potential outcome under the unexposed condition is observed. Evaluating prognostic balance among only the unexposed can reward models that are overfitted to the unexposed group and does not necessarily indicate how well prognostic balance is achieved within the entire study population (3, 24). Consequently, fitted DRS models have been assessed primarily by using prediction diagnostics and goodness-of-fit tests rather than measures of prognostic balance.

The “dry-run” analysis

Hansen (15) proposed an alternative strategy for evaluating risk scores in their ability to control confounding. Because modeling the PS does not share the same theoretical challenges as modeling the DRS, Hansen (15) explained that researchers can use the estimated PS to divide the unexposed population into “pseudo-exposed” and “pseudo-unexposed” groups in order to create differences on observed covariates that are similar to differences between the actual exposed and unexposed populations. With no exposure effect separating the pseudo-exposed and pseudo-unexposed groups, analysts can perform a dry-run analysis by fitting the DRS model to the pseudo-unexposed group, or a historical set of unexposed subjects, and then evaluating the validity of the risk score setup by its ability to control confounding within the pseudo-population. If subclassification or matching on the estimated DRS results in unconfounded null effect estimates within the pseudo-population, then the modeling procedure should be successful in controlling confounding on the same observed covariates when applied to the original sample. We describe the dry-run analysis in detail below and provide example code in Web Appendix 1 (available at http://aje.oxfordjournals.org/):

1. Estimate the PS within the full study population. This step entails diagnosing the PS model’s validity (e.g., checking positivity violations, calibration, discrimination, covariate balance, etc.).

2. Create a pseudo-exposure group by sampling, without replacement, a subset of unexposed subjects with sampling probabilities arranged so that the odds of selection for pseudo-exposure are proportional to the odds of the estimated PSs from step 1. Sampling should be done so that the proportion of the pseudo-exposed within the full unexposed population is approximately equal to the proportion of the exposed within the full study population. Here, we describe a simple procedure that uses independent Bernoulli sampling to select the pseudo-exposure group. Other sampling procedures could also be employed (see Web Appendix 1).

Let \( \pi_i = \frac{\exp(c + \theta_i)}{1 + \exp(c + \theta_i)} \), where \( \theta_i = \log\left(\frac{\text{PS}_i}{(1 - \text{PS}_i)}\right) \), and \( c \) is a constant such that \( \sum_{i=1}^{n_u} \pi_i = \left[ \frac{n_e}{n_e + n_u} \right] n_u \) (i = 1, . . . , \( n_u \)), where \( n_u \) and \( n_e \) represent the number of unexposed and exposed individuals in the full population). Conduct a single independent Bernoulli trial for each unexposed subject, \( i \), with probability \( \pi_i \), to determine whether subject \( i \) is selected into the pseudo-exposure group. This
sampling results in an expected size of \( \left[ \frac{n_U}{n_U + n_E} \right] n_U \) subjects for the pseudo-exposure group but will vary around this target from sample to sample.

3. Form a pseudo-unexposed group consisting of all individuals in the unexposed population who are not sampled into the pseudo-exposure group.

4. Model the DRS within the pseudo-unexposed group or an external set of unexposed subjects.

5. Estimate the pseudo-exposure effect after stratifying or matching on the estimated DRSs within the pseudo-population, and calculate the pseudo-bias, defined as the difference between the pseudo-effect estimate and the true null effect.

6. Bootstrap (i.e., repeat) steps 2–5 to form a distribution of calculated pseudo-biases whose mean and corresponding confidence limits can be used to evaluate the validity of the fitted DRS model.

The sampling outlined above results in a pseudo-population in which the odds of selection for pseudo-exposure are proportional to the estimated odds of exposure in the full population. The goal of the dry-run sampling is not to create pseudo-exposed and pseudo-unexposed groups where covariate distributions mirror those of the full exposed and unexposed populations, but rather it is to create differences between the pseudo-exposed and pseudo-unexposed groups that are similar to observed differences in the full population. For example, suppose we have a cohort where the average age among the exposed is 50 years, the average age among the unexposed is 40 years, and there are equal numbers of exposed and unexposed. The goal of the dry-run sampling is to create pseudo-exposed and pseudo-unexposed groups that differ by 10 years on average, matching the difference between the exposed and unexposed. That could happen with a mean age of 45 years within the pseudo-exposed and 35 years within the pseudo-unexposed, resulting in a difference of 10 years while maintaining the overall average of 40 years across unexposed subjects.

Therefore, prior to mounting a full dry-run validation, analysts should check that differences in baseline characteristics between the pseudo-exposed and pseudo-unexposed groups resemble differences between the actual exposed and unexposed populations. If the PS model is well-specified, while positivity assumptions hold also, there should be such a similarity. It will of course be inexact; one is looking for gross departures here.

**Simulation study**

We simulated a dichotomous exposure \( (A) \) and outcome \( (Y) \), 6 binary covariates \( (X_1, X_3, X_5, X_6, X_8, X_{10}) \) and 4 standard-normal covariates \( (X_2, X_4, X_7, X_9) \). We defined the conditional probability of exposure and outcome according to equations 1 and 2.

\[
\logit(E[Y|X_i, A]) = \beta_0 + \beta_1 X_1 + \cdots + \beta_{10} X_{10} + \beta_1 X_2 X_3 + \beta_2 X_4 X_8 + \beta_3 X_5 X_6 + \beta_4 X_7^2 + \beta_5 X_{10}^2.
\]

(1)

The coefficient values for \( \alpha \) and \( \beta \), \( i = 1 \ldots 15 \), were selected by drawing values from separate uniform\((-0.7, 0.7\) distributions. This range of values (i.e., potentially ranging from \(-0.7 \text{ to } 0.7\) ) was chosen to reflect the range for the majority of coefficient values observed in an empirical example comparing dabigatran with warfarin that is described in the next section. We repeated this process 100 times by drawing a separate set of values for \( \alpha \) and \( \beta \), \( i = 1 \ldots 15 \), to consider a total of 100 unique parameter combinations. To avoid issues with the collapsibility of the odds ratio, we held the exposure effect constant at an odds ratio of \( 1 \) (i.e., \( \beta_A = 0 \) (27)). Both \( \alpha_0 \) and \( \beta_0 \) were set so that the baseline prevalence of exposure and baseline incidence of the outcome were 30% (i.e., baseline prevalence and incidence when all covariate values are set to 0).

With each parameter combination we simulated a study cohort and a historical set of unexposed subjects that was similar to the study cohort but with no exposure introduced. We fitted 32 unique DRS models within this historical population using logistic regression with various degrees of model misspecification. Each of the models included main effects for the covariates \( X_i \) through \( X_{10} \) but different sets of higher-order terms. We considered all possible combinations of the higher-order terms shown in equation 2 (32 in total). We evaluated the calibration and discrimination for each DRS model by calculating the Hosmer-Lemeshow \( P \) value and \( C \) statistic within the original cohort. We also evaluated each DRS model by conducting a dry-run validation as described previously.

Because the dry-run analysis relies on using the PS to create the sampling probabilities, we estimated the PS using 4 different logistic models with varying degrees of misspecification: PS model 1 (included all higher-order terms), PS model 2 (excluded 1 interaction term), PS model 3 (excluded 1 interaction term and 1 quadratic term), and PS model 4 (excluded all higher-order terms).

We conducted simulations using sample sizes of 10,000, 5,000, and 2,000. We estimated the exposure effect within the original study population and each of the pseudo-populations after stratifying on quintiles of the estimated DRS and after matching on the estimated DRS. One-to-one caliper matching was done without replacement using a caliper width of 0.2 standard deviations of the respective DRS distribution (28, 29).

For each of the 100 parameter scenarios, we simulated 100 data sets and evaluated the correlation between the mean of each of the described measures and the mean bias in the effect estimates across all simulation runs. Because the \( C \) statistic and Hosmer-Lemeshow \( P \) value do not take the direction of confounding into account, we also evaluated the correlation between the mean of each measure and the mean absolute bias in the effect estimates across all simulation runs (the absolute pseudo-bias was used when comparing with the absolute bias).
Empirical example

We compared the performance of dabigatran versus warfarin in a nonselected population of older US adults using a 20% random sample (n = 67,667) of all patients with fee-for-service Medicare parts A (hospital), B (outpatient), and D (pharmacy) coverage for at least 1 month from October 19, 2010, through December 31, 2012. Details of the study population and cohort creation are provided elsewhere (7).

We modeled the 1-year risk of combined ischemic stroke and all-cause mortality within a historical population of new warfarin users (30) with an index date prior to the introduction of dabigatran (from January 1, 2008, through October 18, 2010). This model was then used to predict the disease risk for all individuals within the study cohort. We fitted PS and DRS models that included main effects for 37 a priori selected covariates and an additional 200 empirically selected covariates that were identified within Medicare files containing medication claims, inpatient and outpatient diagnostic codes, and procedural codes. The estimated scores were implemented using 1-to-1 matching without replacement. Details of covariate selection and definitions are provided elsewhere (7).

To evaluate the validity of the fitted DRS model, we created a pseudo-exposure group by sampling new warfarin users within the original study cohort (i.e., index date after October 18, 2010), using the sampling described previously. We created a pseudo-unexposed group consisting of new warfarin users who were not selected into the pseudo-exposure group. We then evaluated the validity of the historically fitted DRS model by observing how well matching on

**Figure 1.** Box plots of the Spearman rank correlation coefficients when stratifying on the estimated disease risk score. Each box plot contains 100 correlation coefficients (1 for each of the 100 parameter combinations considered in the simulation), showing results when considering sample sizes of 10,000 ((A) and (B)); 5,000 ((C) and (D)); and 2,000 ((E) and (F)). The box plots show correlation coefficients when comparing each metric with the absolute bias ((A), (C), and (E)) and bias ((B), (D), and (F)) in the effect estimate. Pseudo1 and abs.Pseudo1 represent the pseudo-bias and absolute pseudo-bias when PS model 1 (including all higher-order terms) was used to create the pseudo-population; Pseudo2 and abs.Pseudo2 correspond to PS model 2 (excluding 1 interaction term); Pseudo3 and abs.Pseudo3 correspond to PS model 3 (excluding 1 interaction and 1 quadratic term); and Pseudo4 and abs.Pseudo4 correspond to PS model 4 (excluding all higher-order terms).
the estimated scores controlled for confounding within the pseudo-population. We bootstrapped this process 1,000 times and took the mean of the pseudo-bias across all bootstrapped runs as the measure for model fit. For comparison, we also evaluated the performance of the estimated DRSs by assessing the calibration (Hosmer-Lemeshow goodness-of-fit test) and discrimination (C statistic) of the predicted values within the original study cohort.

RESULTS

Simulation results

Figures 1 and 2 show box plots for the Spearman correlation coefficients between each of the described measures and both the absolute bias (Figure 1A, 1C, and 1E) and bias (Figure 1B, 1D, and 1F) in the estimated exposure effect. The absolute pseudo-bias was used when comparing measures with the absolute bias. In Figure 1, exposure effects were estimated through DRS stratification, while in Figure 2 the exposure effects were estimated through DRS matching. Each box plot shows the distribution of 100 correlation coefficients (1 correlation coefficient for each of the 100 parameter combinations).

When the estimated PS was a close approximation to the true PS model (PS models 1 and 2), there was a strong correlation between the pseudo-bias and the actual bias within the original study cohort (Figures 1 and 2). As the misspecification in the PS model increased, the strength of this correlation became less pronounced and the C statistic showed

![Box plots](https://example.com/box_plots.png)

**Figure 2.** Box plots of the Spearman rank correlation coefficients when matching on the estimated disease risk score. Each box plot contains 100 correlation coefficients (1 for each of the 100 parameter combinations considered in the simulation), showing results when considering sample sizes of 10,000 ((A) and (B)); 5,000 ((C) and (D)); and 2,000 ((E) and (F)). The box plots show correlation coefficients when comparing each metric with the absolute bias ((A), (C), and (E)) and bias ((B), (D), and (F)) in the effect estimate. Pseudo1 and abs.Pseudo1 represent the pseudo-bias and absolute pseudo-bias when PS model 1 (including all higher-order terms) was used to create the pseudo-population; Pseudo2 and abs.Pseudo2 correspond to PS model 2 (excluding 1 interaction term); Pseudo3 and abs.Pseudo3 correspond to PS model 3 (excluding 1 interaction and 1 quadratic term); and Pseudo4 and abs.Pseudo4 correspond to PS model 4 (excluding all higher-order terms).
a stronger correlation in predicting bias (Figures 1 and 2). Compared with DRS stratification, matching on the estimated DRSs resulted in a weaker correlation between the pseudo-bias and actual bias in the effect estimate (Figures 1 and 2). The pseudo-bias showed a stronger correlation with bias in the effect estimate compared with the correlation between the absolute pseudo-bias and absolute bias in the effect estimate (Figures 1 and 2). As the sample size decreased, the correlation between the pseudo-bias and bias in the effect estimate was attenuated, but the pseudo-bias was still generally stronger in predicting bias compared with the C statistic and Hosmer-Lemeshow P value.

Figure 3 shows each measure plotted against the absolute bias (Figure 3A, 3C, and 3E) and bias (Figure 3B, 3D, and 3F) in the effect estimate after combining the values for each measure across all parameter combinations and models. For example, for Figure 3A and 3B we calculated 3,200 pseudo-bias measures (32 DRS models × 100 parameter combinations) when using PS model 1 to create the pseudo-populations. Figure 3A and 3B plot the absolute pseudo-bias (Figure 3A) and pseudo-bias (Figure 3B) under PS model 1 against the corresponding absolute bias (Figure 3A) and bias (Figure 3B) in the exposure effect estimate from the original study cohorts. Similar descriptions can be applied to Figure 3C and 3D for

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Metrics for evaluating DRS models plotted against the absolute bias ((A), (C), and (E)) and bias ((B), (D), and (F)) in the effect estimate. Each point represents a value for the given metric (e.g., C statistic) and the corresponding bias in the treatment effect estimate. Each subplot contains values across all parameter combinations for each of the 32 fitted DRS models. The actual bias in the treatment effect estimate and pseudo-bias were calculated after matching the estimated disease risk scores. Pseudo-bias 1 in (A) corresponds to PS model 1 (including all higher-order terms). The line in each plot is the fitted least squares regression.
the C statistic and Figure 3E and 3F for the Hosmer-Lemeshow P value. Similar to Figures 1 and 2, when the estimated PS was correctly specified, the pseudo-bias showed the strongest correlation with bias and had an intercept from the least-squares regression line of approximately 0 (Figure 3A and 3B). Both the C statistic and the P value from the Hosmer-Lemeshow test showed weaker correlations after combining results across different data-generating models (Figure 3C–3F).

Figures 1 and 2 show that the value of the dry-run pseudo-bias as an indication for bias in the full study cohort is affected by the specification of the PS model and sample size. To examine other factors that may influence the performance of the dry-run analysis in more detail, we conducted a number of sensitivity analyses where we varied the prevalence of exposure, the degree of separation in the PS distributions, the correlation between the PS and DRS, and the exposure effect. A detailed description of these additional simulations is provided in Web Appendix 2. Results showed that the prevalence of exposure, separation in PS distributions, and correlation between the PS and DRS can also influence the performance of the dry-run analysis. When the PS was correctly specified, however, the pseudo-bias generally performed well in predicting bias in the effect estimate compared with the C statistic and Hosmer-Lemeshow P value (Web Figures 1–8).

**Empirical results**

We present results for the empirical study in Figure 4 and Tables 1 and 2. Figure 4 shows good overlap in PS distributions across the dabigatran (exposed) and warfarin (unexposed) exposure groups. Table 1 shows the distribution of 37 a priori selected covariates across exposure groups and across the sampled pseudo-exposed and pseudo-unexposed groups (covariate values for the pseudo-population were averaged over all bootstrapped runs). New users of dabigatran were generally healthier with fewer comorbidities than new users of warfarin (Table 1) (31, 32). In general, differences between the pseudo-exposed and pseudo-unexposed groups paralleled differences between the dabigatran and warfarin groups (Table 1).

Table 2 shows that both PS and DRS matching resulted in similar effect estimates, with hazard ratios of 0.88 (95% confidence interval: 0.81, 0.95) and 0.87 (95% confidence interval: 0.81, 0.94), respectively. The fitted PS model resulted in good predictive performance and model fit in terms of discrimination and calibration, with a C statistic of 0.73 and Hosmer-Lemeshow P value of 0.18 (Table 2). The fitted DRS also resulted in good discrimination, with a C statistic of 0.78, but poor calibration in terms of the Hosmer-Lemeshow goodness-of-fit test, with a P value of <0.01 (Table 2). After matching on the PS, exposure groups were approximately balanced on measured covariates with an average standardized absolute mean difference of <0.01. Matching on the DRS resulted in a pseudo-bias of approximately −0.02 (95% confidence interval: −0.1, 0.06) (Table 2).

**DISCUSSION**

In this study, we used simulations and an empirical example to compare metrics for evaluating DRS models in their ability to reduce bias in effect estimates. We considered 2 traditional measures of model performance: the C statistic and the P value from the Hosmer-Lemeshow goodness-of-fit test. We also considered the dry-run method proposed by Hansen where the PS is used to divide the unexposed population into pseudo-exposed and pseudo-unexposed groups, and the fitted DRS is then evaluated by its ability to control for confounding within this pseudo-population (15).

In simulations, the pseudo-bias from the dry run had the strongest correlation with bias in the effect estimate when the functional form of the PS was a close approximation to the true PS model. When there was moderate to severe misspecification in the PS, this correlation was attenuated and the C statistic showed a stronger correlation with bias in these settings. The C statistic performed well when comparing the relative performance of different models fitted to the same data set. In practice, however, researchers will often fit a single model and want to assess its validity in terms of confounding control. In simulations, the C statistic showed little correlation with bias after combining results across all simulation scenarios,
<table>
<thead>
<tr>
<th>Demographics</th>
<th>Warfarin (Unexposed) (n = 56,260)</th>
<th>Dabigatran (Exposed) (n = 11,407)</th>
<th>Pseudo-Unexposed* (n = 46,774)</th>
<th>Pseudo-Exposed* (n = 9,486)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of Participants</td>
<td>%</td>
<td>No. of Participants</td>
<td>%</td>
</tr>
<tr>
<td><strong>Mean age, years</strong></td>
<td>78.9</td>
<td>76.8</td>
<td>79.3</td>
<td>77.1</td>
</tr>
<tr>
<td>White, %</td>
<td>50,185</td>
<td>89.2</td>
<td>10,462</td>
<td>91.7</td>
</tr>
<tr>
<td>Female sex, %</td>
<td>23,726</td>
<td>42.2</td>
<td>5,584</td>
<td>49.0</td>
</tr>
<tr>
<td><strong>Diagnoses</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chest pain</td>
<td>21,608</td>
<td>38.4</td>
<td>3,998</td>
<td>35.0</td>
</tr>
<tr>
<td>Heart disease</td>
<td>41,945</td>
<td>74.6</td>
<td>7,599</td>
<td>66.6</td>
</tr>
<tr>
<td>Heart failure</td>
<td>17,297</td>
<td>30.7</td>
<td>2,194</td>
<td>19.2</td>
</tr>
<tr>
<td>Hypertension</td>
<td>36,616</td>
<td>65.1</td>
<td>7,221</td>
<td>63.3</td>
</tr>
<tr>
<td>Hyperlipidemia</td>
<td>19,807</td>
<td>35.2</td>
<td>4,687</td>
<td>41.1</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>1,965</td>
<td>3.5</td>
<td>216</td>
<td>1.9</td>
</tr>
<tr>
<td>Cerebrovascular disease</td>
<td>11,979</td>
<td>21.3</td>
<td>1,982</td>
<td>17.4</td>
</tr>
<tr>
<td>Stroke</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ischemic</td>
<td>3,427</td>
<td>6.1</td>
<td>492</td>
<td>4.3</td>
</tr>
<tr>
<td>Hemorrhagic</td>
<td>189</td>
<td>0.3</td>
<td>18</td>
<td>0.2</td>
</tr>
<tr>
<td>TIA</td>
<td>3,882</td>
<td>6.9</td>
<td>723</td>
<td>6.3</td>
</tr>
<tr>
<td>VTE</td>
<td>5,829</td>
<td>10.4</td>
<td>191</td>
<td>1.7</td>
</tr>
<tr>
<td>Diabetes</td>
<td>19,744</td>
<td>35.1</td>
<td>3,424</td>
<td>30.0</td>
</tr>
<tr>
<td>Kidney disease</td>
<td>7,076</td>
<td>12.6</td>
<td>541</td>
<td>4.7</td>
</tr>
<tr>
<td>Renal failure</td>
<td>9,053</td>
<td>16.1</td>
<td>656</td>
<td>5.8</td>
</tr>
<tr>
<td>Bleeding</td>
<td>1,058</td>
<td>1.9</td>
<td>78</td>
<td>0.7</td>
</tr>
<tr>
<td>Anemia</td>
<td>8,792</td>
<td>15.6</td>
<td>1,135</td>
<td>10.0</td>
</tr>
<tr>
<td>Baseline medications</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Antidepressant</td>
<td>15,902</td>
<td>28.3</td>
<td>2,611</td>
<td>22.9</td>
</tr>
<tr>
<td>Antihypertensive</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACE/ARB</td>
<td>29,377</td>
<td>52.2</td>
<td>5,730</td>
<td>50.2</td>
</tr>
<tr>
<td>Loop diuretic</td>
<td>23,018</td>
<td>40.9</td>
<td>3,274</td>
<td>28.7</td>
</tr>
<tr>
<td>Nonloop diuretic</td>
<td>29,565</td>
<td>52.6</td>
<td>4,788</td>
<td>42.0</td>
</tr>
<tr>
<td>Hypolipidemic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Statin</td>
<td>27,795</td>
<td>49.4</td>
<td>5,983</td>
<td>52.5</td>
</tr>
<tr>
<td>Fibrate</td>
<td>2,827</td>
<td>5.0</td>
<td>568</td>
<td>5.0</td>
</tr>
<tr>
<td>Rate control therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beta blocker</td>
<td>39,850</td>
<td>70.8</td>
<td>8,212</td>
<td>72.0</td>
</tr>
<tr>
<td>CCB</td>
<td>24,739</td>
<td>44.0</td>
<td>4,768</td>
<td>41.8</td>
</tr>
<tr>
<td>Glycoside</td>
<td>10,401</td>
<td>18.5</td>
<td>1,951</td>
<td>17.1</td>
</tr>
<tr>
<td>Rhythm control therapy</td>
<td>10,745</td>
<td>19.1</td>
<td>2,648</td>
<td>23.2</td>
</tr>
</tbody>
</table>

Table continues

* Evaluating Risk Scores for Confounding Control 849

**Am J Epidemiol. 2017;185(9):842–852**
illustrating that the actual value of a $C$ statistic does not provide the best information for testing the validity of a single fitted DRS model.

While the dry-run validation can assess whether a fitted DRS model is insufficient in terms of confounding control, it does not provide information on whether the lack of confounding control is due to model misspecification or model extrapolation. Extrapolation being done by the risk score, however, can potentially be mitigated by the process of fitting and checking the validity of the PS (e.g., identifying positivity violations). Although the presence of a few high-propensity unexposed subjects could lead to especially problematic extrapolation, this situation would reduce the pseudo-variance relative to the pseudo-bias, increasing the likelihood of rejecting the risk score setup. In other words, the dry-run validation penalizes a risk score for extrapolation.

A few limitations of the dry-run validation deserve attention. Because the dry run uses the PS to create sampling probabilities, the method’s application is constrained by the assumptions of the PS, even though analyses using the DRS do not inherit these stronger assumptions. In particular, the PS requires that there be no covariate patterns at which exposure is received with certainty (i.e., positivity) (33). The DRS requires a weaker condition, that there be no levels of disease risk at which exposure is certain (i.e., risk positivity) (3). While analyses using the DRS may include some individuals for whom positivity is violated, the dry run’s application is limited to the population where positivity holds. If the analyst determines that positivity is violated when assessing the appropriateness of the fitted PS model, then the analyst must decide whether to respecify the PS and try again, restrict the analysis to a subgroup for which positivity does appear to hold, or seek other modes of validation for the risk score model.

The dry-run analysis also requires accurate estimation of the PS. In this case, one could simply use the PS for confounding control. The DRS, however, can be valuable even when a

<table>
<thead>
<tr>
<th>Health-care utilization, average no. of claims</th>
<th>Warfarin (Unexposed) $n=56,260$</th>
<th>Dabigatran (Exposed) $n=11,407$</th>
<th>Pseudo-Unexposed$^a$ $n=46,774$</th>
<th>Pseudo-Exposed$^a$ $n=9,486$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of Participants</td>
<td>%</td>
<td>No. of Participants</td>
<td>%</td>
</tr>
<tr>
<td>ECG</td>
<td>3.74</td>
<td>3.80</td>
<td>3.73</td>
<td>3.78</td>
</tr>
<tr>
<td>PSA</td>
<td>0.36</td>
<td>0.49</td>
<td>0.34</td>
<td>0.46</td>
</tr>
<tr>
<td>Fecal occult blood testing</td>
<td>0.12</td>
<td>0.13</td>
<td>0.11</td>
<td>0.13</td>
</tr>
<tr>
<td>Colonoscopy</td>
<td>0.14</td>
<td>0.14</td>
<td>0.14</td>
<td>0.14</td>
</tr>
<tr>
<td>Flu shot</td>
<td>0.76</td>
<td>0.79</td>
<td>0.75</td>
<td>0.79</td>
</tr>
<tr>
<td>Lipid assessment</td>
<td>1.52</td>
<td>1.72</td>
<td>1.48</td>
<td>1.68</td>
</tr>
<tr>
<td>Mammography</td>
<td>0.25</td>
<td>0.29</td>
<td>0.24</td>
<td>0.28</td>
</tr>
<tr>
<td>Pap smear</td>
<td>0.05</td>
<td>0.07</td>
<td>0.05</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Abbreviations: ACE/ARB, angiotensin-converting enzyme/angiotensin II receptor blocker; CCB, calcium channel blockers; ECG, electrocardiography; PSA, prostate-specific antigen; TIA, transient ischemic attack; VTE, venous thromboembolism.

$^a$ Mean values for each covariate averaged over 1,000 sampled pseudo-populations. The total $n$ and numbers of each covariate in the pseudo-exposed and pseudo-unexposed groups were rounded to the nearest whole number.

Table 2. A Comparison of Effect Estimates on the Basis of Propensity Score or Disease Risk Score Matching in New Users of Dabigatran and Warfarin in a Population of Medicare Beneficiaries ($n=67,667$), United States, 2010–2012

<table>
<thead>
<tr>
<th>Method$^a$</th>
<th>HR</th>
<th>95% CI</th>
<th>Pseudo-Bias$^b$</th>
<th>95% CI</th>
<th>ASAMD$^c$</th>
<th>C Statistic</th>
<th>HL P Value$^d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unadjusted</td>
<td>0.48</td>
<td>0.46, 0.50</td>
<td>−0.57</td>
<td>−0.65, −0.50</td>
<td>0.14</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PS matching</td>
<td>0.88</td>
<td>0.81, 0.95</td>
<td>&lt;0.01</td>
<td>0.73</td>
<td>0.18</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DRS matching</td>
<td>0.87</td>
<td>0.81, 0.94</td>
<td>0.02</td>
<td>0.02</td>
<td>0.78</td>
<td>&lt;0.01</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ASAMD, average standardized absolute mean difference; CI, confidence interval; DRS, disease risk score; HL, Hosmer-Lemeshow; HR, hazard ratio; PS, propensity score.

$^a$ PS and DRS models included 200 empirically selected covariates and 37 covariates selected a priori.

$^b$ Bias in the pseudo–effect estimate on the log scale (averaged across 1,000 bootstrapped samples). The unadjusted pseudo-bias was calculated by taking the log of the unadjusted hazard ratio within the pseudo-population. The DRS-matched pseudo-bias was calculated by taking the log of the hazard ratio obtained after matching pseudo-exposed to pseudo-unexposed subjects.

$^c$ ASAMD of covariates across exposure groups.

$^d$ P value from the Hosmer-Lemeshow goodness-of-fit test.
correctly specified PS is available. As discussed in the Introduction, risk scores are often preferred for evaluating effect measure modification. The use of risk scores for evaluating effect modification, however, is challenging because misspecified or overfitted DRS models can produce spurious suggestions of effect modification across levels of disease risk (3, 15, 34). In theory, the dry-run validation could be used to detect false signals of effect modification (15). If the modeling procedure produces risk scores that result in the appearance of effect modification within the pseudo-population, this would bring into question the value of the risk score setup for evaluating effect modification when applied to the full study population.

With a correctly specified PS, one may wish to evaluate a DRS model by comparing results from DRS and PS analyses. The dry-run strategy, however, maintains objectivity in study design (35). By restricting to the unexposed population when evaluating risk models, the dry-run analysis does not allow information about the exposure-outcome association to contribute to decisions on model selection. Researchers can evaluate and modify fitted risk models within the sampled pseudo-population without the risk of degrading inference (3, 15).

Finally, the optimal strategy for sampling from the unexposed population when performing a dry-run analysis remains unclear. The without-replacement sampling outlined in this study performed well for the scenarios considered. For smaller samples, other without-replacement sampling techniques may be more appropriate. These could include maximum entropy sampling, which allows the analyst to explicitly specify the number to be sampled (36), or a rejection sampling scheme that throws out a particular pseudo-exposure group selection if its size falls outside of a predetermined window. Regardless of the sampling technique used, we emphasize that the dry-run analysis does not provide information about bias caused by unmeasured confounding. Subject-matter expertise is a necessary component when performing PS or DRS analyses (37, 38).

We conclude that accurately modeling the DRS within the study cohort or within a historical set of unexposed subjects presents unique challenges that are not shared by the PS. Measures of predictive performance and goodness-of-fit tests do not necessarily describe the ability of a DRS model to control confounding. If the PS can be accurately modeled, evaluating the ability of the DRS model to control confounding within a dry-run analysis can provide insight into the validity of fitted DRS models.

ACKNOWLEDGMENTS

Author affiliations: Department of Epidemiology, Gillings School of Global Public Health, University of North Carolina, Chapel Hill, North Carolina (Richard Wyss, Til Stürmer); Division of Pharmacoepidemiology and Pharmacoeconomics, Department of Medicine, Brigham and Women’s Hospital and Harvard Medical School, Boston, Massachusetts (Richard Wyss, Joshua J. Gagne, Rishi J. Desai, Robert J. Glynn); Department of Statistics, College of Literature, Science, and the Arts, University of Michigan, Ann Arbor, Michigan (Ben B. Hansen); and Department of Social Work, College of Humanities and Social Sciences, North Carolina State University, Raleigh, North Carolina (Alan R. Ellis).

This work was funded by the National Institute on Aging (grant R01 AG023178). The database infrastructure used for this project was funded by the Pharmacoepidemiology Gillings Innovation Lab (PEGIL) for the Population-Based Evaluation of Drug Benefits and Harms in Older US Adults (grant GIL200811.0010); the Center for Pharmacoepidemiology, Department of Epidemiology, UNC Gillings School of Global Public Health; the CER Strategic Initiative of UNC’s Clinical Translational Science Award (1 ULI RR025747); the Cecil G. Sheps Center for Health Services Research, UNC; and the UNC School of Medicine.

Conflict of interest: none declared.

REFERENCES

13. Stuart EA, Lee BK, Leacy FP. Prognostic score-based balance measures can be a useful diagnostic for propensity score