EBM Basic Simulation

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Set up

We'll start with the basic cross-sectional model, with a single baseline characteristic, X_i . Here, X_i is binary and about half the population has $X_i = 1$, half has $X_i = 0$. (This could be sex, naturally, but we might consider some other discrete group like race, some comorbidity, sexuality, etc.)

$$X_i \sim_{iid} \text{Bernoulli}(0.5)$$

$$Z_i \sim_{iid} \text{Bernoulli}(0.15)$$

$$S_i | Z_i = 1 \sim_{iid} \text{Uniform}(0, 1)$$

Where $S_i = 0$ whenever $Z_i = 0$.

Additionally let's consider that a person's probability of diagnosis depends on not only the severity of their disease but their baseline characteristics:

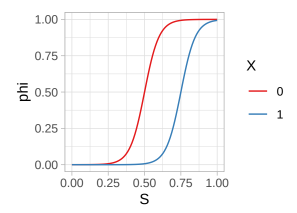
$$D_i \sim_{iid} \text{Bernoulli} (\phi(S_i, X_i))$$

Without loss of generality, let's assume that people with $X_i = 0$ are more likely to be correctly diagnosed than people with $X_i = 1$. Explicitly, let's suppose:

$$\phi(S_i,X_i) = \frac{1}{1 + exp(-(\beta_0 + \beta_1 S_i + \beta_2 X_i))} - c_{x_i},$$

where again, $c_{x_i} = \frac{1}{1 + exp(-(\beta_0 + \beta_2 X_i))}$ is a corrective constant to ensure that $\phi(0, X_i) = 0$. Let Let $\beta_0 = -10$, $\beta_1 = 20$, and $\beta_2 = -5$.

Inessence, this means that people are diagnosed according to a sigmoid function, with people with $X_i = 1$ needing a higher severity to be diagnosed with the same probability as people with $X_i = 0$, shown below:



A Naive Study

Again, we collect a dataset of 5000, recording the baseline characteristic $\{S_i, D_i, X_i\}_{i=1}^n$. Now, our researcher wants to fit a logistic model for predicting disease status from X_i and S_i . However, since the true disease status is unknown, they use diagnosis as a proxy for disease. In regression shorthand they want to model:

$$D_i \sim S_i + X_i$$

The model result of such a study is shown below.

```
##
## Call:
##
   glm(formula = diagnosed ~ x + severity, family = "binomial",
##
       data = df
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
                    -0.0065
##
  -3.0646 -0.0089
                             -0.0006
                                         3.7198
##
##
  Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -10.1429
                            0.9564 -10.605
                                              <2e-16 ***
                                    -8.808
## xTRUE
                -5.4054
                            0.6137
                                              <2e-16 ***
                            1.8502 11.020
## severity
                20.3897
                                              <2e-16 ***
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
##
       Null deviance: 2141.18
                               on 4999
                                        degrees of freedom
## Residual deviance: 220.94
                               on 4997
                                        degrees of freedom
## AIC: 226.94
##
## Number of Fisher Scoring iterations: 11
```

	OR	2.5 %	97.5 %
(Intercept)	0.0000000e+00	0.000	0.000000e+00
xTRUE	4.000000e-03	0.001	1.400000e-02
severity	7.163488e + 08	27331111.273	$3.946674e{+10}$

The researcher records a very high AIC and reports that this model is quite effective at predicting who has the disease. They write a nice paper, talking about how predictive modeling is the gateway to evidence-based personalized medicine. Maybe they suggest that future work might leverage the Awesome Power of Machine Learning. When they interpret their results, they might say that $X_i = 0$ significantly reduces predicted disease risk.

What's wrong with this

The researcher above imagines they are fitting a model for disease risk when in fact they are fitting a model for diagnosis probability. In fact, the model shown above is actually quite a good fit for $\phi(S_i, X_i)$.

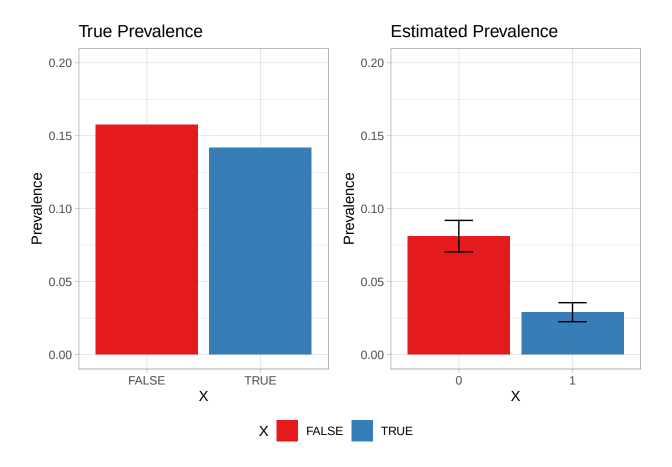
If we actually ran a logistic regression on disease status rather than diagnosis, we would find that X is not at all related to who gets the disease and who does not.

```
##
## Call:
## glm(formula = disease ~ x + severity, family = "binomial", data = df)
##
## Deviance Residuals:
##
          Min
                                                 3Q
                                                             Max
                        1Q
                                Median
                            -0.0000139
##
   -0.0002781
               -0.0002781
                                         -0.0000139
                                                      0.0039523
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
                   -17.068
                               66.529
                                        -0.257
                                                  0.798
##
   (Intercept)
                             1329.697
  xTRUE
                    -5.987
                                        -0.005
                                                  0.996
##
## severity
                 27842.458 219432.652
                                         0.127
                                                  0.899
##
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 4.0605e+03
                                   on 4999
                                             degrees of freedom
## Residual deviance: 1.8230e-04
                                   on 4997
                                             degrees of freedom
##
  AIC: 6.0002
##
## Number of Fisher Scoring iterations: 25
                                         OR
                                               2.5 \%
                                                            97.5 %
                            (Intercept)
                                       0.000
                                                  0
                                                      0.000000e+00
```

(Intercept) 0.000 0 0.000000e+00 xTRUE 0.003 0 3.053422e+17 severity Inf Inf Inf

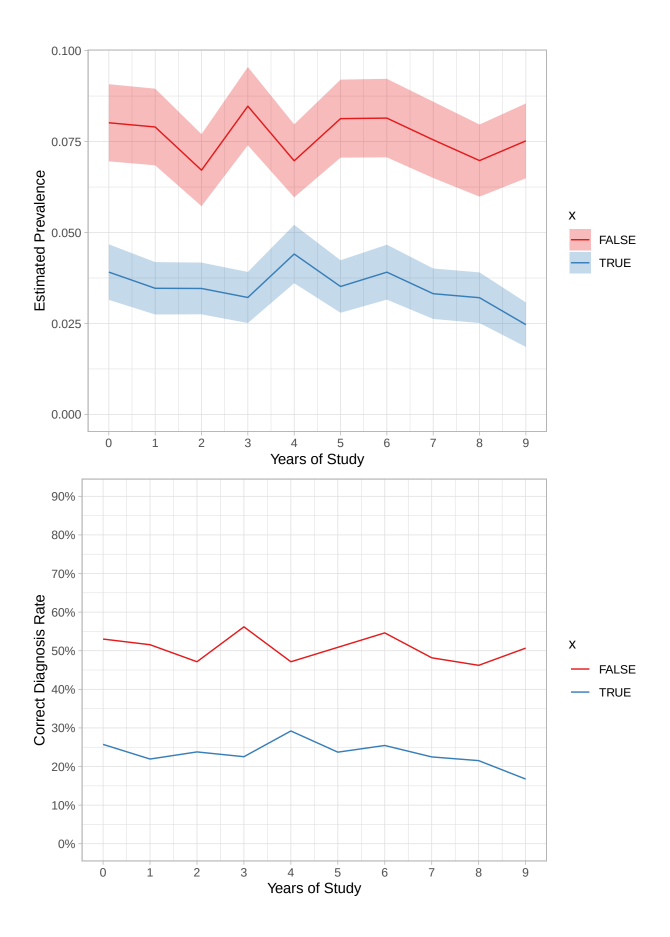
Deployment

Doctors then read about this model and deploy it. In reality, we know that doctors usually do not do exact numeric calculations to decide what diagnosis to give, but let's suppose for a moment that they perfectly follow this published model. In reality, some doctors will be less faithful and some more faithful.



What happens when this process is iterated

Now we imagine that this is part of a learning healthcare system. Every year, a new batch of patients comes to the hospital, and is diagnosed (or isn't). Every year, the researchers retrain their model based on the previous year's diagnoses. The plots below suggest what happens: neither quality of care nor our understanding of the disease improves. Our estimates of disease prevalence are just as incorrect as ever, and the misdiagnosis rates do not improve for either group.



The multi-center case

Now let's suppose there are multiple hospitals using a data-driven approach. Each one trains a "diagnostic model" on it's own set of data in year 0 and each subsequent year. The diagnostic patterns of each hospital appear as somewhat of a random walk. A hospital which has a particularly accurrate or inaccurate diagnostic year by chance will build a particularly accurate or inaccurate model, respectively, in the following year. Unsurprisingly, if the hospitals are very small, their diagnostic rates will vary more wildly from year to year.

