#### 432 Class 10 Slides

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#### Today's Agenda

#### Fitting and evaluating logistic regression models with 1rm

- The framingham example
  - Outcome: chd10 = Developed coronary heart disease in next 10 years?
  - Creating "complete case" data: fram\_cc
  - Single Imputation of Missing Values: fram\_sh
- Use 1rm to predict chd10 using glucose, smoker, sbp and educ
  - on the complete cases (fram\_cc)
  - accounting for missingness via single imputation (fram\_sh)
  - accounting for missingness via multiple imputation
- Consider adding non-linear terms, refit and re-evaluate

## Setup

```
library(magrittr); library(janitor)
library(here); library(knitr)
library(naniar); library(simputation)
library(ROCR)  # one way to draw ROC curves
library(rms)
library(tidyverse)

theme_set(theme_bw())
```

#### Today's Data

```
fram_raw <- read_csv(here("data/framingham.csv")) %>%
    clean_names()
```

See https://www.framinghamheartstudy.org/ for more details. This particular data set has been used by lots of people, in varied settings, with variations all over the net. I don't know who the originators were.

# Managing the Framingham Data

## Data Cleanup

```
fram <- fram raw %>%
    mutate(educ =
               fct recode(factor(education),
                          "Some HS" = "1".
                          "HS grad" = "2",
                          "Some Coll" = "3",
                          "Coll grad" = "4")) %>%
    rename(smoker = "current_smoker",
           cigs = "cigs_per_day",
           stroke = "prevalent_stroke",
           highbp = "prevalent hyp",
           chol = "tot chol",
           sbp = "sys_bp", dbp = "dia_bp",
           hrate = "heart rate",
           chd10 = "ten year chd") %>%
    select(subj_id, chd10, educ, glucose, sbp, smoker,
           everything()) %>% select(-education)
```

# Data Descriptions (Main Variables Today)

The variables describe n=4238 adults examined at baseline, then followed for 10 years to see if they developed incident coronary heart disease.

The main variables we'll use today in developing outcome models are:

Variable	Description
subj_id	identifying code added by Dr. Love
chd10	1=coronary heart disease in next $10$ years
educ	four-level factor: educational attainment
glucose	blood glucose level in mg/dl
sbp	systolic blood pressure (mm Hg)
smoker	$1 = {\sf current} \; {\sf smoker} \; {\sf at} \; {\sf time} \; {\sf of} \; {\sf examination}, \; {\sf else} \; 0$

# Data Descriptions (Other 11 variables)

Here are the other 11 variables in the fram data.

Variable	Description
male	$1={\sf subject}$ is male, else $0$
age	in years (range is 32 to 70)
cigs	number of cigarettes smoked per day
bp_meds	$1 = {\sf using}$ anti-hypertensive medication at time of exam
stroke	1 = history of stroke, else 0
highbp	$1={\sf under}$ treatment for hypertension, else 0
diabetes	1 = history of diabetes, else $0$
chol	total cholesterol (mg/dl)
dbp	diastolic blood pressure (mm Hg)
bmi	body mass index in $kg/m^2$
hrate	heart rate in beats per minute

## Missing Data?

Our outcome chd10 has no missing values.

```
fram %>% tabyl(chd10) %>% adorn_pct_formatting(digits = 1)

chd10    n percent
    0 3594    84.8%
    1 644    15.2%
```

- 3656 (86.3%) of the 4238 subjects in the fram data are complete.
- The remaining 582 observations have something missing.

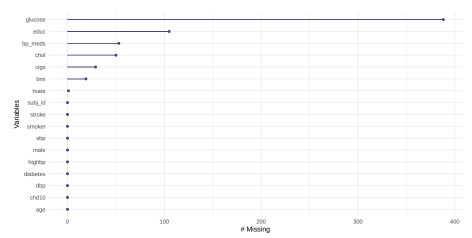
```
n_case_complete(fram); pct_complete_case(fram)
```

- [1] 3656
- [1] 86.26711

## Which variables are missing data?

gg\_miss\_var(fram)

Warning: It is deprecated to specify `guide = FALSE` to remove a guide. Please use `guide = "none"` instead.



## Counts of Missing Data, by Variable

19

1

0.448

0.0236

```
miss_var_summary(fram) %>%
   filter(n miss > 0)
# A tibble: 7 x 3
 variable n_miss pct_miss
 <chr>
          <int>
                   <dbl>
         388 9.16
1 glucose
         105 2.48
2 educ
3 bp_meds
         53 1.25
4 chol
             50 1.18
             29
                  0.684
5 cigs
```

bmi

7 hrate

## Single Imputation

#### We will impute:

- 5 quantitative variables (glucose, bmi, cigs, chol and hrate)
- 1 binary variable (bp\_meds), and
- 1 multi-categorical variable (educ)

```
fram_sh <- bind_shadow(fram)</pre>
fram sh <- fram sh %>%
    data.frame() %>%
    impute_pmm(., bp_meds ~ highbp + sbp + dbp) %>%
    impute_cart(., educ ~ age + smoker + male) %>%
    impute_pmm(., cigs ~ smoker) %>%
    impute rlm(., glucose + chol + hrate + bmi ~
                sbp + diabetes + age + highbp + stroke) %>%
    tibble()
```

## Check multi-categorical single imputation?

80

25

Do the values seem reasonable?

Some HS

HS grad

5 NA

NA

#### Data Sets for the rest of our work

- fram\_start includes all 4238 rows and the 6 columns we'll use, including 388 rows missing glucose and 105 missing educ.
- fram\_cc includes only the 3753 complete rows on the 6 columns.
- fram\_sh uses single imputation to get 4238 complete rows, on 8 columns, including the useful missingness indicators.

## **Modeling Plan**

Use 1rm to fit a four-predictor logistic regression model to predict chd10 using glucose, smoker, sbp and educ

- Using the complete cases (fram\_cc)
- Accounting for missingness via single imputation (fram\_sh)
- 3 Accounting for missingness via multiple imputation

Then, we'll consider adding several non-linear terms to the "four-predictor" models, and refit.

# Fitting a Four-Predictor Model using Complete Cases

#### A "Four Predictor" model

First, we'll use the fram\_cc data to perform a complete-case analysis and fix ideas.

This works very nicely when chd10 = 1 (for Yes) or 0 (for No), as it does here. What if your outcome was actually a factor with values Yes and No? Use the following...

#### Main Output for mod\_cc

mod\_cc

```
Logistic Regression Model
 lrm(formula = chd10 ~ glucose + smoker + sbp + educ. data = fram_cc.
    x = TRUE. v = TRUE
                     Model Likelihood
                                      Discrimination
                                                      Rank Discrim.
                          Ratio Test
                                            Indexes
                                                           Indexes
Obs
        3753
                   LR chi2
                             223.29
                                      R2
                                              0.100
                                                             0.682
                                                      С
 0
            3174
                   d.f.
                                      g 0.689
                                                      Dxy 0.363
                                      gr 1.992
             579
                   Pr(> chi2) <0.0001
                                                      gamma 0.364
 max |deriv| 2e-11
                                      gp 0.092
                                                             0.095
                                                      tau-a
                                      Brier 0.122
              Coef S.E. Wald Z Pr(>|Z|)
 Intercept
           -5.5622 0.3217 -17.29 <0.0001
 glucose 0.0081 0.0016 4.93 <0.0001
 smoker 0.3126 0.0955 3.27 0.0011
 sbp
     0.0237 0.0020 12.05 < 0.0001
 educ=HS grad -0.4674 0.1157 -4.04 <0.0001
 educ=Some Coll -0.3924 0.1423 -2.76 0.0058
 educ=Coll grad -0.1356 0.1549 -0.88 0.3815
```

- We'll walk through these summaries in the next few slides.
- Notes Section 12.14 provides additional details.

# Deconstructing the $mod_cc$ summaries (1/5)

```
Obs 3753
0 3174
1 579
max |deriv| 2e-11
```

- 0bs = The number of observations used to fit the model, with <math>0 = the number of zeros and 1 = the number of ones in our outcome, chd10.
- Also specified is the maximum absolute value of the derivative at the point where the maximum likelihood function was estimated.

All you're likely to care about is whether the iterative function-fitting process converged, and R will warn you in other ways if it doesn't.

# Deconstructing the $mod_cc$ summaries (2/5)

```
Model Likelihood
Ratio Test
LR chi2 223.29
d.f. 6
Pr(> chi2) <0.0001
```

- This is a global likelihood ratio test (drop in deviance test.)
- Likelihood Ratio  $\chi^2$  statistic = null deviance residual deviance d.f. = null degrees of freedom residual degrees of freedom
- $\Pr(> \text{chi2})$  is a p value obtained from comparison to a  $\chi^2$  distribution with appropriate d.f.

It's not saying much to suggest that some part of this logistic regression model has some detectable predictive value.

• The null hypothesis here (that the model has no predictive value at all) is rarely interesting in practical work.

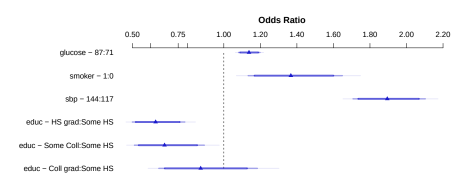
# Deconstructing the $mod_cc$ summaries (3/4)

```
Coef S.E. Wald Z Pr(>|Z|)
Intercept -5.5622 0.3217 -17.29 <0.0001
glucose 0.0081 0.0016 4.93 <0.0001
smoker 0.3126 0.0955 3.27 0.0011
sbp 0.0237 0.0020 12.05 <0.0001
educ=HS grad -0.4674 0.1157 -4.04 <0.0001
educ=Some Coll -0.3924 0.1423 -2.76 0.0058
educ=Coll grad -0.1356 0.1549 -0.88 0.3815
```

- How does each predictor appear to relate to 10-year risk?
  - Which is the baseline educ category?
  - Remember that these estimates are on the logit scale.
  - See the effect size discussion linked in today's README.

## Plot of Effects using mod\_cc

#### plot(summary(mod\_cc))

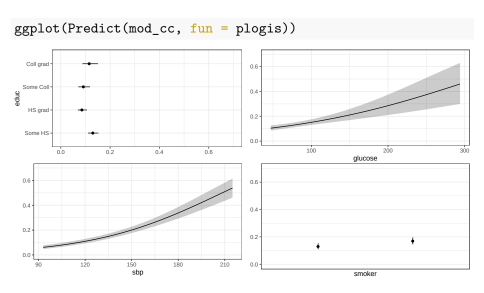


## Effect Size Summary for mod\_cc

#### summary(mod\_cc)

```
Effects
                                 Response : chd10
                        Low High Diff. Effect
                                                 S.E.
                                                          Lower 0.95 Upper 0.95
Factor
alucose
                          71 87
                                         0.12912 0.026171 0.077828
                                                                      0.18041
                                  16
Odds Ratio
                          71 87
                                  16
                                         1.13780
                                                       NA 1.080900
                                                                      1.19770
smoker
                                         0.31259 0.095453
                          0
                                                           0.125510
                                                                      0.49968
Odds Ratio
                          0
                                         1.36700
                                                       NA 1.133700
                                                                      1.64820
sbp
                         117 144
                                  27
                                         0.63907 0.053053
                                                           0.535080
                                                                      0.74305
Odds Ratio
                         117 144
                                  27
                                         1.89470
                                                          1.707600
                                                                      2.10230
educ - HS grad:Some HS
                                  NA
                                        -0.46740 0.115720 -0.694220
                                                                     -0.24059
Odds Ratio
                                                                      0.78616
                                 NA
                                         0.62663
                                                           0.499470
educ - Some Coll:Some HS
                          1 3 NA
                                        -0.39238 0.142310 -0.671310
                                                                     -0.11346
Odds Ratio
                           1 3 NA
                                         0.67544
                                                           0.511040
                                                                      0.89274
educ - Coll grad:Some HS
                                        -0.13556 0.154910 -0.439180
                                                                      0.16806
                                  NA
Odds Ratio
                                  NA
                                         0.87323
                                                           0.644570
                                                                      1.18300
```

## Predict results for mod\_cc



# Deconstructing the $mod_cc$ summaries (4/4)

Discrimination				
	Indexes		Indexes	
R2	0.100	С	0.682	
g	0.689	Dxy	0.363	
gr	1.992	gamma	0.364	
gp	0.092	tau-a	0.095	
Brier	0.122			

The key indexes for our purposes are:

- Nagelkerke R<sup>2</sup>, symbolized R2 here.
- The Brier score, symbolized Brier.
- The area under the ROC curve, or C statistic, shown as C.
- Somers' d statistic, symbolized Dxy here.

Let's walk through each of those, in turn.

# Key Indexes (Nagelkerke $R^2$ )

• In our model, Nagelkerke  $R^2 = 0.100$ 

There are at least three ways to think about  $R^2$  in linear regression, but when you move to a categorical outcome, not all of those ways can be expressed in the same statistic. See our Course Notes Section 10 for details.

#### The Nagelkerke $R^2$ :

- reaches 1 if the fitted model shows as much improvement as possible over the null model (which just predicts the mean response on the 0-1 scale for all subjects).
- is 0 for the null model
- is larger (closer to 1) as the fitted model improves, although it's been criticized for being misleadingly high,
- AND a value of 0.100 no longer means 10% of anything.

A value of 0.100 indicates a model of pretty poor quality.

#### An Alternative: McFadden's R<sup>2</sup>

Consider the McFadden R-square, which can be defined as 1 minus the ratio of (the model deviance over the deviance for the intercept-only model.)

To obtain this for our mod\_cc run with lrm, we can use:

```
1 - (mod_cc$deviance[2] / mod_cc$deviance[1])
```

```
[1] 0.069174
```

This McFadden  $R^2$  corresponds well to the proportionate reduction in error interpretation of an  $R^2$ , but some people don't like it as well.

# Key Indexes (Brier Score = 0.122)

- The lower the Brier score, the better the predictions are calibrated.
- The maximum (worst) score is 1, the best is 0.

From Wikipedia: Suppose you're forecasting the probability P that it will rain on a given day.

- If the forecast is P = 1 (100%) and it rains, the Brier Score is 0.
- If the forecast is P=1 (100%) and it doesn't rain, the Brier Score is 1.
- If the forecast is P = 0.7 and it rains,  $Brier = (0.70 1)^2 = 0.09$ .
- If the forecast is P = 0.3 and it rains, Brier  $= (0.30 1)^2 = 0.49$ .
- If the forecast is P = 0.5, the Brier score is  $(0.50 1)^2 = 0.25$  regardless of whether it rains.

The Brier score can also be decomposed to assess calibration and discrimination separately.

## Receiver Operating Characteristic Curve Analysis

One way to assess the predictive accuracy within the model development sample in a logistic regression is to consider analyses based on the receiver operating characteristic (ROC) curve. ROC curves are commonly used in assessing diagnoses in medical settings, and in signal detection applications.

The accuracy of a test can be evaluated by considering two types of errors: false positives and false negatives.

See Section 12.9 of our Course Notes for more details.

# The C statistic (area under ROC curve) = 0.682

The C statistic and Somers' d (Dxy) are connected:

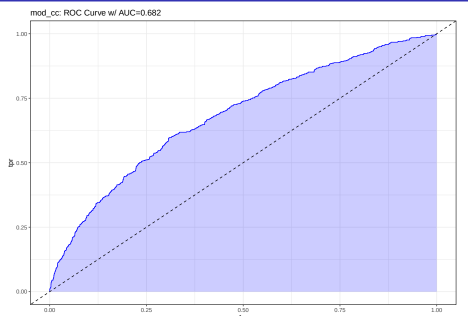
$$C = 0.5 + \frac{d}{2}, d = 2(C - .5)$$

The C statistic ranges from 0 to 1.

- ullet C = 1 indicates a perfect prediction model, one that guesses "yes" for all patients with chd10 = 1 and which guesses "no" for all patients with chd10 = 0.
- Most of the time, the closer to 1, the happier we are:
  - $C \ge 0.8$  usually indicates a moderately strong model (good discrimination)
  - ullet  $C \geq 0.9$  indicates a very strong model (excellent discrimination)

So 0.682 isn't good.

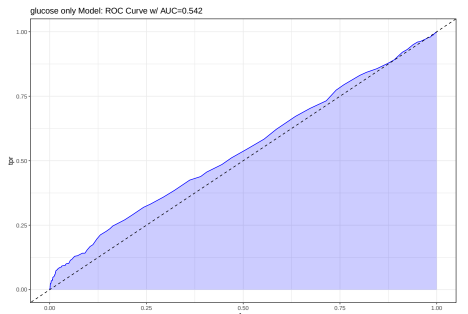
# ROC Curve for our mod\_cc



#### **Code for Previous Slide**

```
## requires ROCR package
prob <- predict(mod_cc, type="fitted")</pre>
pred <- prediction(prob, fram_cc$chd10)</pre>
perf <- performance(pred, measure = "tpr", x.measure = "fpr")</pre>
auc <- performance(pred, measure="auc")</pre>
auc <- round(auc@y.values[[1]],3)</pre>
roc.data <- data.frame(fpr=unlist(perf@x.values),</pre>
                        tpr=unlist(perf@y.values),
                        model="GLM")
ggplot(roc.data, aes(x=fpr, ymin=0, ymax=tpr)) +
    geom ribbon(alpha=0.2, fill = "blue") +
    geom line(aes(y=tpr), col = "blue") +
    geom abline(intercept = 0, slope = 1, lty = "dashed") +
    labs(title = paste0("Model A: ROC Curve w/ AUC=", auc))
```

# ROC Curve for a Simple Model (glucose only)



## Validate Summary Statistics for mod\_cc

 Usual approach (as in ols) to correcting for over-optimism through bootstrap validation, now using 50 bootstrap resamples instead of 40.

```
set.seed(432)
validate(mod_cc, B = 50)
```

```
index.orig training
                              test optimism index.corrected n
Dxy
             0.3634
                     0.3655
                             0.3583
                                     0.0072
                                                    0.3562 50
R2
             0.1001
                     0.1007
                             0.0977
                                     0.0029
                                                    0.0972 50
Intercept
             0.0000
                     0.0000 -0.0196
                                     0.0196
                                                   -0.019650
Slope
            1.0000
                    1.0000
                             0.9873
                                    0.0127
                                                    0.9873 50
                                    0.0064
Emax
            0.0000
                    0.0000 0.0064
                                                    0.0064 50
                    0.0596 0.0578
                                    0.0018
            0.0592
                                                    0.0574 50
                    -0.0005 0.0000
                                    -0.0006
            -0.0005
                                                    0.0000 50
             0.0598
                    0.0601 0.0577
                                     0.0024
                                                    0.0574 50
             0.1216
                    0.1215
                             0.1219
                                    -0.0004
                                                    0.1220 50
            0.6892
                                    0.0105
                    0.6933
                             0.6829
                                                    0.6787 50
             0.0917
                     0.0918
                             0.0907
                                     0.0011
                                                    0.0906 50
```

- Summaries we'll focus on here are Dxy, R2 and B
- Remember that  $C = 0.5 + \frac{Dxy}{2}$ , so our validated C statistic would be 0.5 + (0.3562/2) = 0.6781

#### ANOVA for mod\_cc

Model mod\_cc uses 6 degrees of freedom.

anova(mod\_cc)

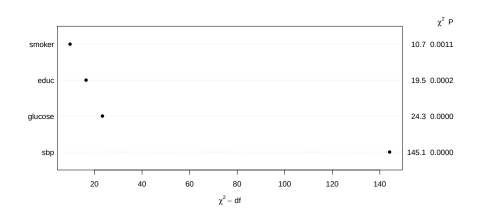
Wald Statistics

Response: chd10

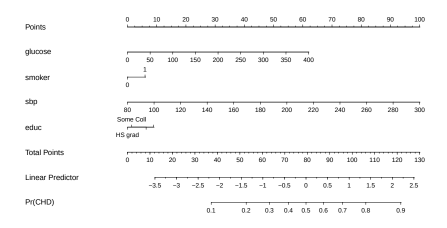
Factor	Chi-Square	d.f.	P
glucose	24.34	1	<.0001
smoker	10.72	1	0.0011
sbp	145.10	1	<.0001
educ	19.45	3	0.0002
TOTAL	208.87	6	<.0001

## ANOVA for Model mod\_cc

#### plot(anova(mod\_cc))



## Nomogram for mod\_cc



# Using the Singly Imputed Data to fit the 4-predictor Model

#### Fit mod\_si which is mod\_cc after single imputation

### Model mod\_si with single imputation

#### mod\_si

```
Logistic Regression Model
 lrm(formula = chd10 ~ glucose + smoker + sbp + educ, data = fram_sh,
    x = TRUE, y = TRUE
                      Model Likelihood
                                         Discrimination
                                                          Rank Discrim.
                            Ratio Test
                                                Indexes
                                                               Indexes
                    LR chi2
 Obs
             4238
                                238.36
                                         R2
                                                 0.095
                                                          С
                                                                 0.677
 0
             3594
                    d.f.
                                                 0.673
                                                                 0.354
                                         g
                                                          Dxy
  1
              644
                    Pr(> chi2) <0.0001
                                            1.961
                                                          aamma
                                                                 0.354
                                         ar
 max |deriv| 4e-12
                                                 0.089
                                                                 0.091
                                         qр
                                                          tau-a
                                         Brier
                                                 0.121
               Coef S.E.
                             Wald Z Pr(>|Z|)
              -5.5649 0.3068 -18.14 <0.0001
 Intercept
 glucose
        0.0086 0.0016 5.32 < 0.0001
 smoker
             0.3205 0.0901 3.56 0.0004
             0.0231 0.0019 12.40 < 0.0001
 sbp
 educ=HS grad -0.4707 0.1098 -4.29 <0.0001
 educ=Some Coll -0.3055 0.1336 -2.29 0.0222
 educ=Coll grad -0.0816 0.1470 -0.56 0.5787
```

# **Comparing the Coefficients (exponentiated)**

Comparing the slopes as odds ratios

```
round_half_up(exp(mod_cc$coefficients),3)
    Intercept glucose smoker
                                                sbp
       0.004 1.008
                             1.367
                                               1.024
 educ=HS grad educ=Some Coll educ=Coll grad
       0.627
                    0.675
                                 0.873
round half up(exp(mod si$coefficients),3)
                           smoker
    Intercept glucose
                                                sbp
       0.004
                    1.009
                           1.378
                                               1.023
 educ=HS grad educ=Some Coll educ=Coll grad
       0.625
                    0.737
                                 0.922
```

# **Edited Summaries Comparing The Models**

Summary	mod_si value	mod_cc value
Obs	4238	3753
0	3594	3174
1	644	579
Nagelkerke $R^2$	0.095	0.100
Brier Score	0.121	0.122
C	0.677	0.682
Dxy	0.354	0.363

All of these results came from

mod\_cc
mod\_si

# Validate mod\_si Summary Statistics

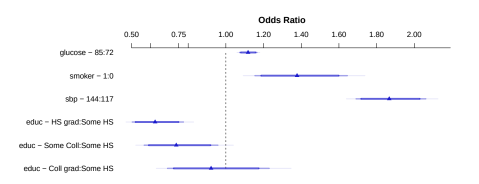
```
set.seed(432)
validate(mod_si, B = 50)
```

```
index.orig training
                             test optimism index.corrected
Dxy
            0.3538
                    0.3555
                           0.3496
                                   0.0058
                                                 0.3480 50
R2
            0.0954
                                   0.0033
                    0.0966
                           0.0933
                                                 0.0921 50
Intercept 0.0000
                    0.0000 -0.0256
                                   0.0256
                                                -0.0256 50
       1.0000 1.0000
Slope
                           0.9860
                                   0.0140
                                                 0.9860 50
Emax
      0.0000 0.0000
                           0.0079
                                   0.0079
                                                0.0079 50
         0.0560
                    0.0568
                           0.0548
                                   0.0021
                                                0.0539 50
D
                                  -0.0005
           -0.0005 -0.0005
                           0.0000
                                                0.0000 50
Q
B
            0.0565
                    0.0573
                           0.0548 0.0026
                                                 0.0539 50
                                  -0.0001
            0.1206
                    0.1207
                           0.1208
                                                 0.1207 50
```

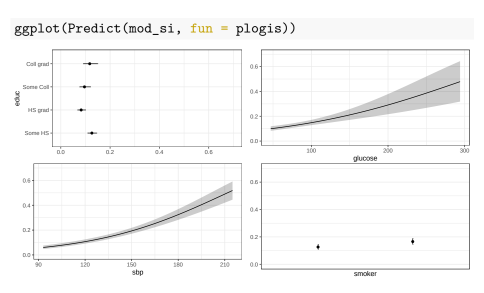
• Again,  $C = 0.5 + \frac{Dxy}{2}$ , so the corrected C statistic estimate will be 0.5 + (0.348/2) = 0.674

# Plot of Effects using mod\_si

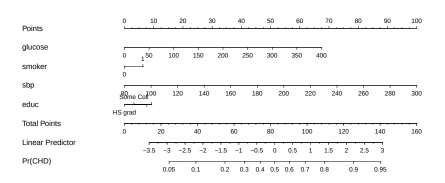
#### plot(summary(mod\_si))



### Predict results for mod\_si



#### Nomogram for mod\_si



# Using Multiple Imputation: The 4-predictor Model

#### Fit the Imputation Model first

We'll use aregImpute here, and create 30 imputed sets.

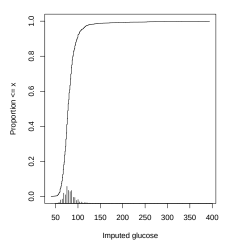
Iteration 1 Iteration 2 Iteration 3 Iteration 4 Iteration 5 It

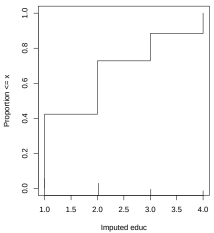
# Imputation Results (abbreviated output)

#### fit\_imp

```
Multiple Imputation using Bootstrap and PMM
aregImpute(formula = ~chd10 + glucose + smoker + sbp + educ,
   data = fram, n.impute = 30, nk = c(0, 3:5), tlinear = FALSE,
   B = 10
n: 4238 p: 5 Imputations: 30 nk: 0
Number of NAs:
 chd10 glucose smoker sbp
                                educ
           388
     0
                                 105
R-squares for Predicting Non-Missing Values for Each Variable
Using Last Imputations of Predictors
glucose educ
  0.046 0.024
```

### Multiply Imputed Values, via plot(fit\_imp)





# What do we need to do our multiple imputation?

Imputation Model

```
fit_imp <-
   aregImpute(~ chd10 + glucose + smoker + sbp + educ,
        nk = c(0, 3:5), tlinear = FALSE, data = fram,
        B = 10, n.impute = 30)</pre>
```

• Outcome Model will be of the following form...

```
lrm(chd10 ~ glucose + smoker + sbp + educ,
    x = TRUE, y = TRUE)
```

# Fitting mod\_mi (mod\_cc with multiple imputation)

- data = fram\_start (which includes NA values)
- xtrans = fit\_imp (results from multiple imputation)
- fitter = lrm (we could actually use glm too)
- pr = FALSE avoids a long printout we don't need

#### Model mod\_mi with multiple imputation

#### mod\_mi

```
Logistic Regression Model
 fit.mult.impute(formula = chd10 ~ glucose + smoker + sbp + educ.
    fitter = 1rm, xtrans = fit imp, data = fram start, pr = FALSE.
    x = TRUE. v = TRUE
                      Model Likelihood
                                        Discrimination
                                                         Rank Discrim.
                           Ratio Test
                                               Indexes
                                                               Indexes
                    LR chi2
Obs
            4238
                               237.84
                                        R2
                                                 0.095
                                                                0.677
                                                         С
            3594
                    d.f.
                                                0.670
                                                         Dxy 0.354
 0
                                        g
                    Pr(> chi2) <0.0001
                                                1.955
                                                         gamma 0.354
             644
                                        ar
 max |deriv| 2e-11
                                        aр
                                               0.088
                                                         tau-a
                                                                0.091
                                        Brier 0.121
              Coef S.E. Wald Z Pr(>|Z|)
 Intercept
              -5.5542 0.3083 -18.02 <0.0001
 alucose 0.0083 0.0016 5.12 < 0.0001
 smoker 0.3188 0.0902 3.54 0.0004
 sbp
        0.0232 0.0019 12.40 < 0.0001
 educ=HS grad -0.4551 0.1120 -4.06 <0.0001
 educ=Some Coll -0.3002 0.1340 -2.24 0.0251
 educ=Coll grad -0.0845 0.1478 -0.57 0.5674
```

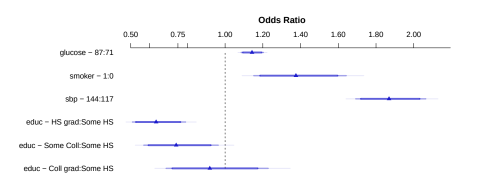
# **Comparing the Coefficients (exponentiated)**

I'll just compare the two models using imputation...

```
round_half_up(exp(mod_mi$coefficients),3)
    Intercept glucose smoker
                                                sbp
       0.004
                    1.008
                             1.376
                                               1.023
 educ=HS grad educ=Some Coll educ=Coll grad
                                 0.919
       0.634
                    0.741
round half up(exp(mod si$coefficients),3)
                           smoker
    Intercept glucose
                                                sbp
       0.004
                    1.009
                            1.378
                                               1.023
 educ=HS grad educ=Some Coll educ=Coll grad
       0.625
                    0.737
                                 0.922
```

# Plot of Effects using mod\_mi

#### plot(summary(mod\_mi))



# **Edited Summaries Comparing Our 3 Models**

Summary	mod_mi value	mod_si value	mod_cc value
Obs	4238	4238	3753
0	3594	3594	3174
1	644	644	579
Nagelkerke $R^2$	0.095	0.095	0.100
Brier Score	0.121	0.121	0.122
C	0.677	0.677	0.682
Dxy	0.354	0.354	0.363

- It's just a coincidence that the mod\_mi and mod\_si values are identical to the level of precision provided in this table.
- What might cause the values to look meaningfully different?

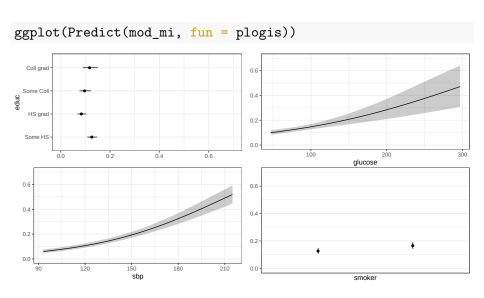
# Validate mod\_mi Summary Statistics

```
set.seed(432)
validate(mod_mi, B = 50)
```

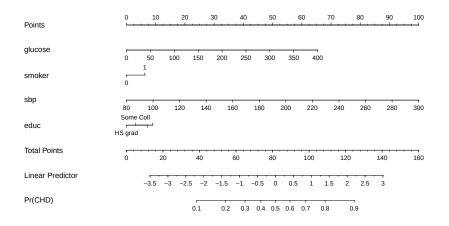
```
index.orig training
                              test optimism index.corrected
DXV
             0.3535
                     0.3551
                            0.3493
                                     0.0058
                                                    0.3477 50
R2
            0.0952
                     0.0958
                            0.0925
                                     0.0033
                                                    0.0919 50
Intercept
            0.0000
                     0.0000 -0.0259
                                     0.0259
                                                   -0.0259 50
Slope
          1.0000 1.0000
                            0.9858
                                     0.0142
                                                    0.9858 50
Emax
          0.0000 0.0000
                            0.0080
                                     0.0080
                                                    0.0080 50
D
            0.0559 0.0564
                            0.0543
                                     0.0021
                                                    0.0538 50
            -0.0005
                    -0.0005
                            0.0000
                                    -0.0005
                                                    0.0000 50
U
             0.0564
                     0.0569
                            0.0543
                                     0.0026
                                                    0.0538 50
Q
                                    -0.0001
             0.1207
                     0.1208
                            0.1209
                                                    0.1208 50
```

• Optimism-corrected C statistic estimate is 0.5 + (0.3477/2) = 0.674

### Predict results for mod\_mi



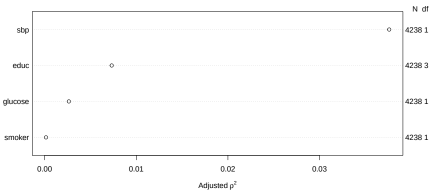
#### Nomogram for mod\_mi



# **Considering Non-Linear Terms**

# **Spearman** $\rho^2$ **Plot**





# Adding some non-linear terms

- We'll add a restricted cubic spline with 5 knots in sbp
- and an interaction between the educ factor and the linear effect of sbp,
- and a quadratic polynomial in glucose

to our main effects model, just to show how to do them...

I'll just show the results including the multiple imputation, since if you
can get those, you should have little difficulty instead applying the
single imputation or the complete case analysis.

#### mod\_big incorporating multiple imputation

Our mod\_big will incorporate several non-linear terms.

#### The mod\_big model with non-linear terms

```
Logistic Regression Model
 fit.mult.impute(formula = chd10 ~ rcs(sbp, 5) + pol(glucose,
        2) + smoker + educ + educ %ia% sbp. fitter = lrm. xtrans = fit imp.
        data = fram_start, pr = FALSE, x = TRUE, y = TRUE)
                                    Model Likelihood
                                                                    Discrimination
                                                                                                  Rank Discrim.
                                              Ratio Test
                                                                                Indexes
                                                                                                           Indexes
 0bs
                    4238
                                 LR chi2
                                                    245.28
                                                                                   0.098
                                                                                                               0.679
   0
                    3594
                                 d.f.
                                                           13
                                                                                   0.710
                                                                                                  Dxv 0.357
                      644
                                 Pr(> chi2) <0.0001
                                                                                   2.034
                                                                                                  gamma 0.357
 max |deriv| 0.02
                                                                     ap
                                                                                   0.092
                                                                                                  tau-a
                                                                                                              0.092
                                                                     Brier
                                                                                   0.120
                                   Coef S.E. Wald Z Pr(>|Z|)
                              -3.2646 2.1123 -1.55 0.1222
 Intercept
                                 0.0034 0.0190 0.18 0.8565
 sbp
 sbp'
                                0.1756 0.1837 0.96 0.3390
                     -0.5056 0.6402 -0.79 0.4296
 sbp''
 sbp'''
                                0.3651 0.6492 0.56 0.5738
                                0.0061 0.0054 1.12 0.2612
 alucose

        glucose
        0.0061
        0.0054
        1.12
        0.2612

        glucose^2
        0.0000
        0.0000
        0.0000
        0.0000

        smoker
        0.3218
        0.0903
        3.56
        0.0004

        educ=HS grad
        -0.4033
        0.6438
        -0.63
        0.5310

        educ=Some Coll
        -1.4405
        0.8055
        -1.79
        0.0737

        educ=Coll grad
        -1.1027
        0.9379
        -1.18
        0.2397

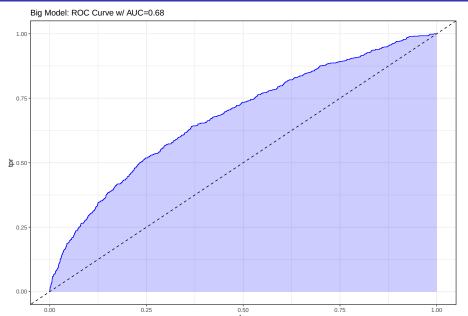
        educ=HS grad * sbp
        -0.0004
        0.0045
        -0.09
        0.9246

 educ=Some Coll * sbp 0.0083 0.0057 1.44 0.1485
 educ=Coll grad * sbp 0.0075 0.0068 1.10 0.2697
```

# mod\_big vs. mod\_mi comparison

Summary	mod_big	mod_mi
Obs	4238	4238
0	3594	3594
1	644	644
Nagelkerke $R^2$	0.098	0.095
Brier Score	0.120	0.121
C	0.679	0.677
Dxy	0.357	0.354

# ROC Curve for mod\_big

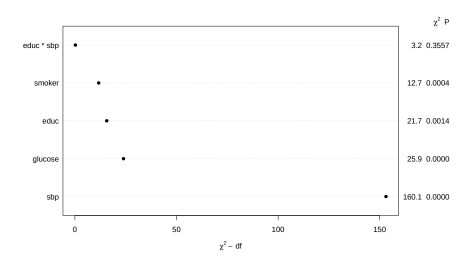


# What does ANOVA suggest about the fit?

#### anova(mod\_big)

Wald Statistics	Response: ch	nd10	
Factor	Chi-Square	d.f.	Р
sbp (Factor+Higher Order Factors)	160.07	7	<.0001
All Interactions	3.24	3	0.3557
Nonlinear	3.03	3	0.3869
glucose	25.92	2	<.0001
Nonlinear	0.21	1	0.6495
smoker	12.71	1	0.0004
educ (Factor+Higher Order Factors)	21.68	6	0.0014
All Interactions	3.24	3	0.3557
educ * sbp (Factor+Higher Order Factors)	3.24	3	0.3557
TOTAL NONLINEAR	3.18	4	0.5280
TOTAL NONLINEAR + INTERACTION	7.14	7	0.4145
TOTAL	222.84	13	<.0001

### plot(anova(mod\_big)) (model includes 13 df)



# Validate mod\_big Summary Statistics

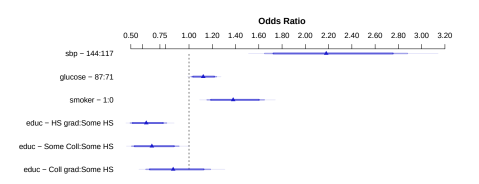
```
set.seed(432)
validate(mod_big, B = 50)
```

```
index.orig training
                              test optimism index.corrected
            0.3577
                     0.3650
                            0.3507
Dxv
                                    0.0143
                                                   0.3434 50
R2
            0.0980
                     0.1022
                            0.0922
                                    0.0100
                                                   0.0880 50
Intercept 0.0000
                     0.0000 -0.0911
                                    0.0911
                                                  -0.0911 50
Slope
        1.0000
                    1.0000
                                                   0.9456 50
                            0.9456
                                    0.0544
Emax
            0.0000
                     0.0000
                            0.0296
                                    0.0296
                                                   0.0296 50
D
            0.0576
                     0.0603
                            0.0541
                                    0.0062
                                                   0.0515 50
U
            -0.0005
                    -0.0005
                            0.0003
                                   -0.0007
                                                   0.0003 50
Q
            0.0581
                     0.0607
                            0.0538
                                    0.0069
                                                   0.0512 50
            0.1204
                     0.1202
                            0.1209
                                   -0.0007
                                                   0.1211 50
```

• Optimism-Corrected C = 0.5 + (.3434/2) = .672

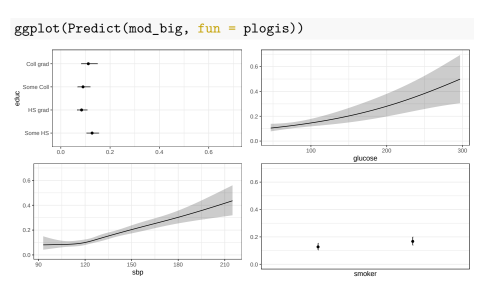
# Plot of Effects using mod\_big

#### plot(summary(mod\_big))



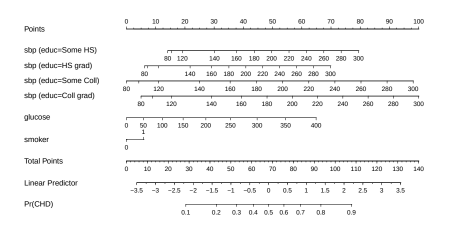
Adjusted to:sbp=128 educ=Some HS

# Predict results for mod\_big



### Nomogram for mod\_big

plot(nomogram(mod\_big, fun = plogis, funlabel = "Pr(CHD)"))



#### What's Next?

 Using the tidymodels framework to fit linear and logistic regression models