Big Match - Large Demo

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This rmarkdown is meant for running a larger demo of the big_match functionality.

Import Data

This sample cohort data was provided by Justin Lee at the Quantitative Sciences Unit. It contains $\sim 900,000$ observations of 112 variables.

```
dat <- read_sas("../sample_data/justincohort_june2017.sas7bdat")
# dimensions: ~900,000 x 112
dim(dat)</pre>
```

```
## [1] 893498 112
```

We include only hospitalizations with totalct > 1 and arteryCt < 3. A patient is considered to have recieved treatment if arteryCt is greater than 1. The outcome of this analysis is mortality.

```
# filter and add treatment column
dat <- filter(dat, totalct > 1 & arteryCt < 3) %>%
  mutate(treat = ifelse(arteryCt > 1, 1, 0))
# dimensions: ~900,000 x 112
dim(dat)
```

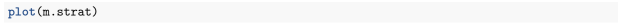
```
## [1] 833657 113
```

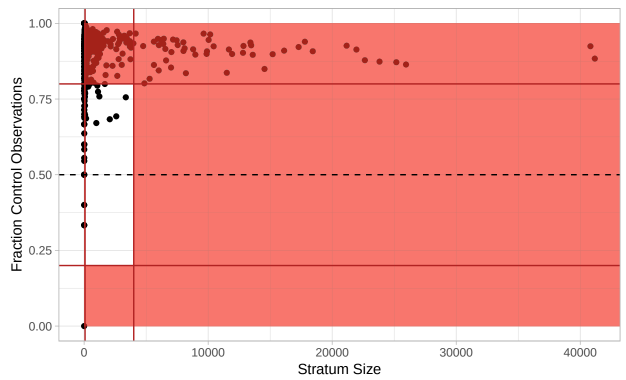
Manual Statify

As a simple example, we can choose to stratify manually by race, sex, and hospital state. This is done as shown below and gives the following strata sizes:

```
## manual_strata object from package big_match.
##
## Function call:
## manual_stratify(data = dat, treat = "treat", covariates = c("Male",
## "race", "hosp_state"))
##
## Analysis set dimensions: 833657 X 114
##
## Number of strata: 704
##
## Min size: 1 Max size: 41175
```

The minimum size is 1 (far too small) and the maximum is 41175 (too large to be computationally feasible). The diagnostic plot below shows all strata based on their treat/control balance and number of observations. The un-tinted rectangle is the target for strata with good balance and appropriate size.





If desired, one can use trial and error to select a covariate set for stratification which gives more favorable strata sizes and balances.

Auto Stratify by Prognostic Score

Another option is to stratify based on prognostic score. There are a few ways to do this. The simplest method is to supply a list of covariates to auto_stratify which details the features we would like to include in a prognostic model. The auto_stratify function will then automatically split the data randomly into a model_set and an analysis_set. By default, the model set contains 1/10 of the controls from the input dataset. A logistic model for prognostic score is then fit on the model_set using the covariates specified. Once the prognostic model is fit, the model set should be kept separate from the data used for the remainder of the analysis.

Using our new prognostic model, we can assign prognostic scores to each observation in the analysis set. Finally, using these scores, the analysis set can be divided into strata of relatively equal size, where observations which appear in the same strata are characterized by similar prognostic scores.

Common Problem: Non-Representative Model Set

The code to automatically stratify the dataset might look like this:

In this example dataset, we see a common problem. When we try to fit a prognostic model which includes the hospital state as a variable, the following error is showm:

Error applying prognostic model: Some categorical variable value(s) in the analysis set do not appear in the modeling set.

This occurs because some value(s) of one or more categorical variables appear in the analysis set which were not seen in the model set. This means that when we try to obtain prognostic scores for our analysis set, we run into some new value that our prognostic model was not prepared to handle. For example, suppose all individuals in the model set had hosp_state equal to some state in the continental US, and the analysis contained one example from with hosp_state equal to "Virgin Islands". We don't know what prognostic score to assign this individual because we did not see any patients in the Virgin Islands when we built our prognostic model.

We have the following options:

- 1. **Group by this variable in auto_stratify:** add the offending categorical variable to group_vars in the call to auto_stratify. This will ensure that the model set contains the same proportions of individuals from each category as in the original dataset.
- 2. **Rejection Sampling:** run auto_stratify again and again with different random seeds until this error does not occur
- 3. Remove the offending examples from the entire data set
- 4. Remove this variable from the prognostic model

In this case, there are only two hospitalizations from the Virgin Islands:

```
filter(dat, hosp_state == "Virgin Islands") %>% nrow()
```

```
## [1] 2
```

For simplicity, we will remove these observations for now.

```
dat <- filter(dat, hosp_state != "Virgin Islands")</pre>
```

Basic Auto Strification

Now, we can run the code as before:

```
set.seed(123)
a.strat <- auto_stratify(data = dat, treat = "treat",
                         outcome = "dead",
                          covariates = c("totalct", "hosp_state", "AMI_7", "COPD_7",
                                         "ISCHEMICHEART_7", "STROKE_TIA_7", "ATRIAL_FIB_7",
                                         "CHRONICKIDNEY_7", "DIABETES_7", "ALZH_DEMEN_7",
                                         "Male", "race"))
## [1] "Constructing a model set via subsampling."
## [1] "Fitting prognostic model: dead~totalct+hosp_state+AMI_7+COPD_7+ISCHEMICHEART_7+STROKE_TIA_7+ATR
## [1] "Generating strata assignments based on prognostic score."
## [1] "Completing strata diagnostics."
print(a.strat)
## auto_strata object from package big_match.
##
## Function call:
  auto_stratify(data = dat, treat = "treat", outcome = "dead",
##
       covariates = c("totalct", "hosp_state", "AMI_7", "COPD_7",
           "ISCHEMICHEART_7", "STROKE_TIA_7", "ATRIAL_FIB_7", "CHRONICKIDNEY_7",
##
           "DIABETES_7", "ALZH_DEMEN_7", "Male", "race"))
##
## Analysis set dimensions: 758213 X 114
##
## Model set dimensions: 75442 X 113
##
## Prognostic Score Model:
##
## Call: glm(formula = formula(formula_str), family = "binomial", data = model_set)
##
## Coefficients:
##
                       (Intercept)
                                                            totalct
##
                         -9.98541
                                                            0.03078
##
                hosp_stateAlabama
                                                  hosp_stateAlaska
##
                          8.38816
                                                            7.18931
##
                hosp_stateArizona
                                                hosp_stateArkansas
##
                           8.32274
                                                            8.11693
                                                hosp_stateColorado
##
             hosp_stateCalifornia
##
                           8.33747
                                                            8.35222
            hosp_stateConnecticut
##
                                                hosp_stateDelaware
##
                          8.50978
                                                            7.94440
## hosp_stateDistrict of Columbia
                                                 hosp_stateFlorida
##
                           8.63687
                                                            8.30760
##
                hosp_stateGeorgia
                                                  hosp_stateHawaii
##
                           8.30898
                                                            8.76670
##
                  hosp_stateIdaho
                                                hosp_stateIllinois
##
                          7.97234
                                                            8.40512
```

##	hosp_stateIndiana	hosp_stateIowa
## ##	8.31327 hosp_stateKansas	8.46555 hosp_stateKentucky
##	nosp_statekansas 8.14804	nosp_statekentucky 8.26063
##	hosp_stateLouisiana	hosp_stateMaine
##	8.32117	8.29869
##	hosp_stateMaryland	hosp_stateMassachusetts
##	8.34472	8.31328
##	hosp_stateMichigan	hosp_stateMinnesota
##	8.36176	8.32598
##	hosp_stateMississippi	${\tt hosp_stateMissouri}$
##	8.23361	8.30468
##	hosp_stateMontana	hosp_stateNebraska
##	8.57506	8.35863
##	hosp_stateNevada	hosp_stateNew Hampshire
##	8.61286	8.12860
##	hosp_stateNew Jersey	hosp_stateNew Mexico
## ##	8.24553 hosp_stateNew York	8.12606 hosp_stateNorth Carolina
##	nosp_stateNew fork 8.28634	8.25697
##	hosp_stateNorth Dakota	hosp_stateOhio
##	8.46294	8.42554
##	hosp_stateOklahoma	hosp_stateOregon
##	8.28569	8.15285
##	hosp_statePennsylvania	hosp_statePuerto Rico
##	8.34529	8.39532
##	hosp_stateRhode Island	hosp_stateSouth Carolina
##	8.57200	8.07109
##	hosp_stateSouth Dakota	${\tt hosp_stateTennessee}$
##	7.90457	8.23020
##	hosp_stateTexas	hosp_stateUtah
##	8.31432	8.29113
##	hosp_stateVermont	hosp_stateVirginia
##	8.23726	8.15050
##	hosp_stateWashington 8.27598	hosp_stateWest Virginia
## ##	*	8.32826 hosp_stateWyoming
##	hosp_stateWisconsin 8.32709	nosp_statewyoming 7.80117
##	AMI_7	COPD_7
##	0.17583	0.46509
##	ISCHEMICHEART_7	STROKE_TIA_7
##	0.02598	0.21875
##	ATRIAL_FIB_7	CHRONICKIDNEY_7
##	0.61505	0.42109
##	DIABETES_7	ALZH_DEMEN_7
##	0.22020	0.43783
##	Male	raceBlack
##	-0.04238	0.35006
##	raceHispanic	raceNorth American Native
##	0.19132	-0.07869
##	raceOther	raceUnknown
##	-0.05135	0.13405
##	raceWhite	
##	0.17457	

```
## Degrees of Freedom: 75441 Total (i.e. Null); 75373 Residual
## Null Deviance:
## Residual Deviance: 84630
                                     AIC: 84770
## Number of strata: 302
   Min size: 661
                       Max size: 4466
plot(a.strat)
    1.00
 Fraction Control Observations
    0.75
    0.50
    0.25
    0.00
                              1000
                                                 2000
                                                                                       4000
                                                                    3000
                                                 Stratum Size
```

Matching

Since we're matching on a single computer, we'll start with a very small subset of the original data to demonstrate the optimal matching functionality.