## Combining Matches Within Subgroups

Josh Errickson, Mark Fredrickson, Josh Buckner, Peter Solenberger and Ben B. Hansen, with embedded

2022-03-14

When utilizing full matching, a user may want to introduce restrictions to the potential match sets. There are two main reasons to do this.

- There may be certain variables which matches should either always or never agree upon. For example,
  if there is a binary gender variable, you may wish to only match treatment members to control members
  of the same gender.
- When matching on a medium-to-large data set, speed concerns can become paramount. By splitting
  the matching problem into a series of smaller subproblems, we can realize substantial performance
  improvements.

## Combining matches

In optmatch 0.9-11 and above, optmatch objects can be easily combined to facilitate breaking a problem into smaller sub-problems and reconstituting a matched structure on the entire data set. To demonstrate this, let's consider the infert data set.

```
> data(infert)
> head(infert)
     education age parity induced case spontaneous stratum pooled.stratum
## 1
         0-5yrs
                           6
                                         1
                                                       2
                                                                1
                 26
                                    1
                                                                                 3
## 2
                                                       0
                                                                2
         0-5yrs
                  42
                           1
                                    1
                                         1
                                                                                 1
## 3
         0-5yrs
                  39
                           6
                                    2
                                         1
                                                       0
                                                                3
                                                                                 4
         0-5yrs
                  34
                                    2
                                         1
                                                       0
                                                                4
                                                                                 2
                                                                5
                                                                                32
## 5
        6-11yrs
                           3
                                    1
                                         1
                                                       1
                  35
       6-11yrs
```

The "case" variable indicates treatment (1) versus control (0) status. We'll want to match upon "age".

```
> table(infert$case)
##
## 0 1
## 165 83
> table(infert$education, infert$case)
##
## 0 0 1
## 0-5yrs 8 4
## 6-11yrs 80 40
## 12+ yrs 77 39
```

Due to the sample size, if we were to compute matches on the entire data set, the fullmatch call would generate a distance matrix of size  $165 \times 83 = 13,695$ . However, if we were instead to compute a match within each level of the "education" variable, we'd compute three different distance matrices, of total size  $8 \times 4 + 80 \times 40 + 77 \times 39 = 6,235$ , a reduction of 55%.

We'll do this by splitting the data within each match.

```
> f1 <- fullmatch(case ~ age, data = infert[infert$education == "0-5yrs", ])</pre>
> f2 <- fullmatch(case ~ age, data = infert[infert$education == "6-11yrs", ])</pre>
> f3 <- fullmatch(case ~ age, data = infert[infert$education == "12+ yrs", ])</pre>
> summary(f1)
## Structure of matched sets:
## 1:2
## Effective Sample Size: 5.3
## (equivalent number of matched pairs).
> summary(f2)
## Structure of matched sets:
## 1:1 1:2 1:3 1:4 1:5+
   20
        8 6 4
## Effective Sample Size: 49.4
## (equivalent number of matched pairs).
> summary(f3)
## Structure of matched sets:
## 1:1 1:2 1:3 1:4 1:5+
## 23
        5
             6 2
## Effective Sample Size: 47
## (equivalent number of matched pairs).
```

Some of the matched sets are quite large (1:5+) so let's put some restrictions.

```
> f2 <- fullmatch(case ~ age, data = infert[infert$education == "6-11yrs", ],</pre>
                 max.controls = 4)
> f3 <- fullmatch(case ~ age, data = infert[infert$education == "12+ yrs", ],
                 \max.controls = 4)
> summary(f2)
## Structure of matched sets:
## 1:1 1:2 1:3 1:4
## 18 10 6 6
## Effective Sample Size: 49.9
## (equivalent number of matched pairs).
> summary(f3)
## Structure of matched sets:
## 1:1 1:2 1:3 1:4
## 20 6 7
## Effective Sample Size: 48.1
## (equivalent number of matched pairs).
```

Now we simply combine the three matches.

```
> fcombine <- c(f1, f2, f3)
> summary(fcombine)
## Structure of matched sets:
## 1:1 1:2 1:3 1:4
## 38 20 13 12
## Effective Sample Size: 103.4
## (equivalent number of matched pairs).
> infert$match <- fcombine</pre>
```

## Using the within argument

An alternative approach would be using the within argument and the exactMatch function to define subproblems.

Observe that we obtain equivalent matched structure. A few notes comparing the two approaches:

1. When using the within argument, restrictions must be the same across subproblems. That is, max.controls, min.controls and omit.fraction will be equivalent. By running the subproblems separately, you can set different restrictions per subproblem. E.g.,

```
> f1 <- fullmatch(z ~ x, data = d[d$group == 1, ], max.controls = 2)
> f2 <- fullmatch(z ~ x, data = d[d$group == 2, ], min.controls = 1/3)
> c(f1, f2)
```

2. While the matched structures will be equivalent between these two approaches (if the restrictions are the same across subproblems), the actual matched sets themselves may differ if you have observations of equal distance. In general this should not be considered a problem.