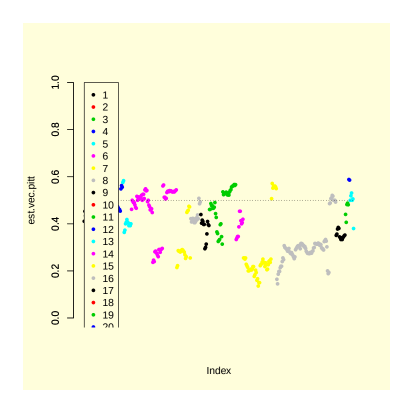


The top CNV component has the followinfg trio-states. Where '1' indicates a deletion and order is $F,\,M,\,O.$

The CNV components with significant p-values (Bonferroni) are given below.

GRanges with 4 ranges and 0 metadata columns:

```
seqnames
                                     ranges strand
             <Rle>
                                  <IRanges>
comp1883
              chr7 [141435189, 141439888]
comp1994
              chr8 [ 39356825,
                                  39370186]
comp2009
              chr8 [ 39476659,
                                  39497557]
             chr11 [ 55204004,
comp2610
                                  55209499]
seqlengths:
           chr1
                   chr1_random
                                          chr2 ...
                                                              {\tt chrY}
                                                                             \mathtt{chrM}
     247249719
                       1663265
                                    242951149 ...
                                                         57772954
                                                                            16571
```



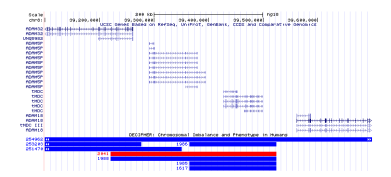


Figure 1: UCSC Genome Browser at chromosome 8 locus.

```
GRanges with 3 ranges and 0 metadata columns:
      seqnames
                               ranges strand
         <Rle>
                            <IRanges> <Rle>
  [1]
          chr7 [141407267, 141441259]
  [2]
          chr8 [ 39341981, 39518214]
  [3]
         chr11 [ 55124465, 55216671]
  seqlengths:
            chr1
                   chr1_random
                                        chr2 ...
                                                           chrY
                                                                         chrM
                                                     57772954
                   1663265
                                   242951149 ...
                                                                        16571
       247249719
> freq.beaty <- colSums(cnv(fe.beaty))/2/nrow(cnv(fe.beaty))</pre>
> freq.pitt <- colSums(cnv(fe.pitt))/2/nrow(cnv(fe.pitt))</pre>
> freq.beaty[index.1.beaty]
comp1883
0.1023152
> freq.pitt[index.1.pitt]
comp1645
0.116408
> freq.beaty[index.2.beaty]
comp1994
0.3278566
> freq.pitt[index.2.pitt]
 comp1793
0.2577605
> freq.beaty[index.3.beaty]
 comp2009
0.3188947
> freq.pitt[index.3.pitt]
 comp1805
0.2588692
> freq.beaty[index.4.beaty]
 comp2610
0.1467513
> freq.pitt[index.4.pitt]
```

```
comp2356
0.154102
> summary(freq.beaty[index.1.full.beaty])
                                 1st Qu.
                                                               Median
                                                                                                 Mean
                                                                                                                 3rd Qu.
0.0003734 \ 0.0067210 \ 0.1070000 \ 0.0690100 \ 0.1131000 \ 0.1135000
> summary(freq.pitt[index.1.full.pitt])
           Min. 1st Qu.
                                                       Median
                                                                                      Mean 3rd Qu.
                                                                                                                                         Max.
0.001109 0.016210 0.121400 0.081830 0.129200 0.129700
> summary(freq.beaty[index.2.full.beaty])
              Min.
                                 1st Qu.
                                                               Median
                                                                                                 Mean
                                                                                                                     3rd Qu.
0.0003734 0.3359000 0.3574000 0.2984000 0.3620000 0.3645000
> summary(freq.pitt[index.2.full.pitt])
              Min.
                                 1st Qu.
                                                               Median
                                                                                                 Mean
                                                                                                                     3rd Qu.
                                                                                                                                                         Max.
0.0005543 0.2633000 0.2716000 0.2352000 0.2752000 0.2788000
> summary(freq.beaty[index.3.full.beaty])
              Min.
                                 1st Qu.
                                                               Median
                                                                                                 Mean
                                                                                                                     3rd Qu.
                                                                                                                                                         Max.
0.0007468\ 0.1822000\ 0.2110000\ 0.1940000\ 0.2255000\ 0.2274000
> summary(freq.pitt[index.3.full.pitt])
                                 1st Qu.
                                                               Median
                                                                                                 Mean
                                                                                                                     3rd Qu.
                                                                                                                                                         Max.
0.0005543 0.1818000 0.1924000 0.1795000 0.1973000 0.2040000
> trioClasses:::trans.tab
function (object)
{
          \label{eq:total_total_total_total} $T <- \sup(\text{object[c("101", "011", "122", "212", "111", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "112", "1
                      "112")], na.rm = TRUE)
          U <- sum(object[c("100", "010", "121", "211", "111", "110",
                      "110")], na.rm = TRUE)
           if (T + U >= 50) {
                      return(binom.test(x = T, n = T + U, p = 0.5, alternative = "greater")p.value)
           }
           else {
                      return(NA)
```

<environment: namespace:trioClasses>

Methods

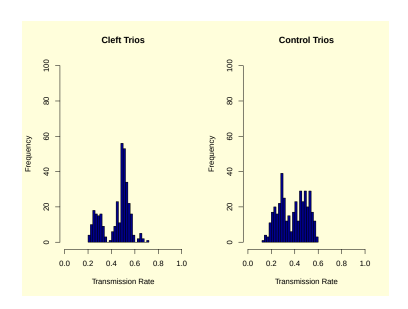
Cleft Data Description

- Performed 336 tests.
- \bullet PennCNV joint HMM
- european, MAD < 0.3, non-WGA, aux $\neq 1$
- coverage > 10
- 14857 hemi/homozygous deletions identified in 445 trios
- 4288 CNV components

```
Common (> 0.01): 660
```

Rare: 3628

- Construct trio-states for all CNV components
 - 0,1,2 variable for normal, hemizygous and homozygous, respectively
- T + U > 50
- count transmissions and perform binom.test (See "trans.tab")



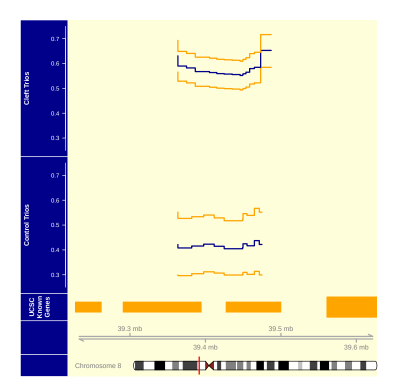


Figure 2:

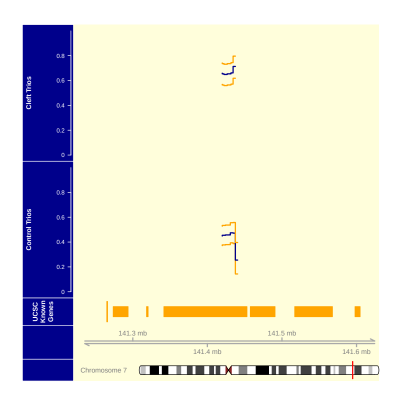


Figure 3:

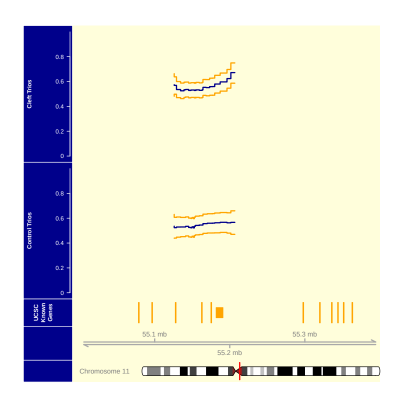


Figure 4: