The R package trioClasses for definition of the class FamilyExperiment, an extension of SummarizedExperiment, for use in trio based analyses of genetic data.

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## 1 Packages & Data

```
> library("trioClasses")
> library("CleftCNVAssoc")
> source("~/jhsph/R/packages/CleftCNVAssoc/vignettes/curated/make-data.R")
> num.snp <- 25
> gr.deletion.beaty <- gr.beaty[values(gr.beaty)$numsnp >=
     num.snp & values(gr.beaty)$cn %in% 0:1]
> homos.beaty <- with(values(gr.deletion.beaty), cn ==</pre>
> gr.deletion.pitt <- gr.pitt[values(gr.pitt)$numsnp >=
     num.snp & values(gr.pitt)$cn %in% 0:1]
> homos.pitt <- with(values(gr.deletion.pitt), cn == 0)</pre>
> gr.deletion.beaty.clean <- c(unlist(reduce(split(gr.deletion.beaty,
     values(gr.deletion.beaty)$id))), unlist(reduce(split(gr.deletion.beaty[homos.beaty],
     values(gr.deletion.beaty[homos.beaty])$id))))
> gr.deletion.pitt.clean <- c(unlist(reduce(split(gr.deletion.pitt,
     values(gr.deletion.pitt)$id))), unlist(reduce(split(gr.deletion.pitt[homos.pitt],
     values(gr.deletion.pitt[homos.pitt])$id))))
> system.time(cnv.obj.beaty <- CNVMatrix(gr.deletion.beaty.clean,
     gr = gr.deletion.pitt.clean))
   user system elapsed
948.668
        0.016 951.497
> system.time(cnv.obj.pitt <- CNVMatrix(gr.deletion.pitt.clean,
     gr = gr.deletion.beaty.clean))
   user system elapsed
660.052
        0.000 661.943
> save(cnv.obj.beaty, cnv.obj.pitt, file = "./../data/cnv-25.RData")
```