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")
> data("cnv", package = "trioClasses")
> data("pedigrees", package = "CleftCNVAssoc")
See vignette "CNVMatrix."
```

2 SummarizedExperiment

3 Pedigree

```
> beaty.trios <- MinimumDistance:::trios(beaty.pedigree)
> beaty.ped <- DataFrame(famid = do.call("rbind", strsplit(beaty.trios$0,
        "_"))[, 1], id = beaty.trios$0, fid = beaty.trios$F,
        mid = beaty.trios$M, sex = NA, dx = NA)
> ped <- PedClass(beaty.ped)</pre>
```

4 FamilyExperiment

```
> (ste <- FamilyExperiment(se, pedigree = ped))

class: FamilyExperiment
dim: 12915 1341
exptData(0):
assays(1): cnv
rownames(12915): comp1 comp2 ... comp12914 comp12915
rowData metadata column names(0):
colnames(1341): 11005_03@1008472481 11005_02@1008472482 ...
18117_02@0070298660 18117_01@0070298681
colData names(1): id
pedigree(2082): famid id fid mid sex dx
complete trios(447):</pre>
```