how to make an smlSet from hapmap data

VJ Carey

April 21, 2009

1. Raw data acquisition: Obtain the hapmap files from the bulk data download. A typical filename is

```
genotypes_chrY_YRI_r23_nr.b36_fwd.txt.gz
```

- 2. use snpMatrix read.HapMap.data to obtain the associated snp.matrix and support data frame. We do this for the 24 main chromosome files. We save the snp.matrix for chromosome n to C[nn].rda. Be careful with the ordering of filenames should match desired ordering of chromosomes.
- 3. Create a list of snp.matrix of genotype data:

```
> ofi = dir(patt = "C.*rda")
> allsm = list()
> cn = rep(NA, 24)
> for (i in 1:24) {
+    cat(i)
+    load(ofi[i])
+    fn = gsub(".rda", "", ofi[i])
+    allsm[[i]] = get(fn)[[1]]
+    cn[i] = as.character(get(fn)[[2]][1, "Chromosome"])
+    print(fn)
+    rm(fn)
+    gc()
+ }
```

Don't forget to give names 1:22, X, Y to the list elements.

4. Create an environment and assign the list created above to symbol smList in that environment. This environment is a valid value for the smlEnv slot of a smlSet instance.

- 5. The chromInds slot gives numerical indices indicating which chromosomes are included; see hmceuB36.2021 in GGtools for an example.
- $6.\ {\rm remaining\ slots}$ are as in ExpressionSet