snpStats Bugs

Hin-Tak Leung

November 12, 2011

Contents

1	Introduction	Т
2	Bugs in snpStats GLM estimates 2.1 snpStats::snp.*hs.estimates() returns garbage	3
3	Multiple snpStats issues of data corruption, memory violations and crashes	5
4	Bugs in snpStats GLM score tests 4.1 snpStats::snp.lhs.tests(,robust=TRUE) returns garbage 4.2 Malformed "GlmTests" S4 object from snpStats::snp.lhs.tests() 4.3 Crazy large/negative number of samples from snpStats GLM tests 4.4 snpStats::snp.rhs.tests() returning garbage with or without robust	7 8
5	2-df Bug in snpStats::single.snp.tests()	10
6	64-bit mode for snpStats::single.snp.tests()	13
7	X chromosome conversion	13

1 Introduction

In the Regression and Migration vignette, we discovered that there has been a bug in snpMatrix's single.snp.tests() for many years, which can affect 1% to 2% of SNPs, and fixed it. This vignette uses the testsuite code in snpMatrix to reveal snpStats' bug(s). The quick summary is that most (all?) of the statistical tests were broken to various extent arount October 2008 by the imputation-related changes. That is 3 years of flawed publications. David made an effort with 1.3.7+ (20th October 2011) but did not get very far. This document is usually built against current snpStats HEAD as well as 1.3.6 (17th October 2011), the latter because of the number of flawed results in 3 years.

Despite many routines being of the same names but behaving differently (such as the buggy single.snp.tests() in snpStats vs the correct one in snpMatrix) and a warning about routines shadowing each other, it is possible to use either in the same R session even in alternating statements, as long as either are referenced explicitly in each step. This usually consists of

prefix'ing with explicit namespace references (e.g. snpMatrix::single.snp.tests() instead of single.snp.tests()) or adding package= within.

- > library(snpMatrix)
- > library(snpMatrix124)
- > library(snpStats)
- > sessionInfo()

R version 2.14.0 (2011-10-31)

Platform: i686-redhat-linux-gnu (32-bit)

locale:

[1] LC_CTYPE=en_GB.UTF-8 LC_NUMERIC=C

[3] LC_TIME=en_GB.UTF-8 LC_COLLATE=en_GB.UTF-8
[5] LC_MONETARY=en_GB.UTF-8 LC_MESSAGES=en_GB.UTF-8

[7] LC_PAPER=C LC_NAME=C
[9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=en_GB.UTF-8 LC_IDENTIFICATION=C

attached base packages:

- [1] grDevices datasets splines graphics utils stats methods
- [8] base

other attached packages:

[1] snpStats_1.3.6 snpMatrix124_1.2.4.5 snpMatrix_1.19.0.11 [4] Matrix_1.0-1 lattice_0.20-0 survival_2.36-10

loaded via a namespace (and not attached):

[1] grid_2.14.0 tools_2.14.0

At this point there is a warning:

Attaching package: 'snpStats'

The following object(s) are masked from 'package:snpMatrix':

can.impute, chi.squared, col.summary, deg.freedom, effect.sign, effective.sample.size, filter.rules, Fst, glm.test.control, ibsCount, ibsDist, imputation.maf, imputation.nsnp, imputation.r2, impute.snps, ld, misinherits, mvtests, p.value, plotUncertainty, pool, pool2, pp, qq.chisq, read.beagle, read.impute, read.mach, read.pedfile, read.plink, read.snps.long, row.summary, sample.size, single.snp.tests, snp.cbind, snp.cor, snp.imputation, snp.lhs.estimates, snp.lhs.tests, snp.post.multiply, snp.pre.multiply, snp.rbind, snp.rhs.estimates, snp.rhs.tests, switch.alleles, tdt.snp, test.allele.switch, write.plink, xxt

Loading snpMatrix124 simultaneously can also cause a few warnings of the form, understandably:

```
A specification for class ``X.snp.matrix'' in package 'snpMatrix124' seems equivalent to one from package 'snpMatrix' and is not turning on duplicate class definitions for this class
```

The warning is harmless, if one pays attention to specifying each explicitly, as below.

2 Bugs in snpStats GLM estimates

2.1 snpStats::snp.*hs.estimates() returns garbage

snp.*hs.estimates() gives garbage — the way to illustrate this is simply running the corresponding snp.rhs.tests and compare:

```
> data(testdata, package = "snpStats")
> test2 <- snpStats::snp.rhs.estimates(cc ~ region + sex, family = "binomial",
      data = subject.data, snp.data = Autosomes, sets = 1:10)
> test2.t <- snpStats::snp.rhs.tests(cc ~ region + sex, family = "binomial",
      data = subject.data, snp.data = Autosomes, tests = 1:10)
> print(cbind(as(as(test2, "GlmTests"), "data.frame"), as(test2.t,
      "data.frame")))
       Chi.squared Df
                         p.value Chi.squared Df
                                                   p.value
173760 0.002022755 1 0.96412719 0.96172092 1 0.32675367
173761 1.614689875 1 0.20383380 1.61954459 1 0.20315530
173762 2.052781924 1 0.15192836 2.05991420 1 0.15121869
173767 0.777421401 1 0.37793093 0.77858708 1 0.37757361
173769 2.762354385 1 0.09650612 2.92552940 1 0.08718862
               NA NA
173770
                             NΑ
                                          NA O
                             NA 1.02511037 1 0.31130988
173772
               NA NA
173774 0.729732211 1 0.39297000 0.73179195 1 0.39230296
173775 0.952718263 1 0.32902835 0.95584241 1 0.32823661
173776 0.090165195 1 0.76396725 0.09019184 1 0.76393342
  This is the correct result from snpMatrix (1.17.7.11) (Wald test close to the score test):
```

```
> Autosomes <- new("snp.matrix", Autosomes@.Data)
> test2 <- snpMatrix::snp.rhs.estimates(cc ~ region + sex, family = "binomial",
+ data = subject.data, snp.data = Autosomes, sets = 1:10)
> test2.t <- snpMatrix::snp.rhs.tests(cc ~ region + sex, family = "binomial",
+ data = subject.data, snp.data = Autosomes, tests = 1:10)
> print(cbind(as(as(test2, "snp.tests.glm"), "data.frame"), as(test2.t,
+ "data.frame")))
```

```
Chi.squared Df p.value Chi.squared Df p.value 173760 0.68794335 1 0.40686481 0.96171330 1 0.32675559 173761 1.61514861 1 0.20376957 1.61953853 1 0.20315614 173762 2.05337857 1 0.15186885 2.05990584 1 0.15121951 173767 0.77747875 1 0.37791334 0.77858708 1 0.37757361 173769 2.76587572 1 0.09629398 2.92549240 1 0.08719062 173770 NA NA NA NA NA NA NA 173772 0.71127227 1 0.39902177 1.02511036 1 0.31130988 173774 0.72973283 1 0.39296980 0.73179195 1 0.39230296 173775 0.95271902 1 0.32902816 0.95584109 1 0.32823694 173776 0.09016525 1 0.76396718 0.09019184 1 0.76393341
```

See another garbage result from snpStats:

			r			F
173760	NA	NA	NA	NaN	0	NaN
173761	NA	NA	NA	NA	NA	NA
173762	NA	NA	NA	NA	NA	NA
173767	NA	NA	NA	NA	NA	NA
173769	NA	NA	NA	NA	NA	NA
173770	NA	NA	NA	0	0	1
173772	NA	NA	NA	NaN	0	NaN
173774	NA	NA	NA	NA	NA	NA
173775	NA	NA	NA	NA	NA	NA
173776	NA	NA	NA	NA	NA	NA

Chi.squared Df p.value Chi.squared Df p.value

At the time of this writing, snpMatrix (1.17.7.11, unreleased) isn't correct either, but better:

```
2.651885e-01
                                     10.476287 9 0.31331934
173762
        11.155682 9
173767
        17.009456 9 4.856810e-02
                                     15.045105 9 0.08970445
173769 201.767706 9 1.411265e-38
                                     16.267175 9 0.06150760
                                     0.000000 0 1.00000000
173770
               NA NA
173772
         5.449466 9
                      7.935005e-01
                                      1.009439 9 0.99941544
       278.130321 9
173774
                      1.125582e-54
                                     9.411857
                                               9 0.40015665
173775
       337.152659
                      3.352980e-67
                                     13.440413 9 0.14366942
173776 426.044143 9
                      3.773047e-86
                                     16.797680 9 0.05198017
```

3 Multiple snpStats issues of data corruption, memory violations and crashes

There are multiple issues of data corruption, memory violation and crashes in the GLM related code. The best way to demonstrate this is turn on gctorture() and uses the GLM score tests/estimates and see R crash.

4 Bugs in snpStats GLM score tests

4.1 snpStats::snp.lhs.tests(...,robust=TRUE) returns garbage

```
> data(testdata, package = "snpStats")
> snpStats::snp.lhs.tests(Autosomes[, 1:10], ~strata(cc), ~strata(region),
+ data = subject.data, robust = TRUE)
```

	${\tt Chi.squared}$	Df	p.value
173760	NaN	0	NaN
173761	NA	NA	NA
173762	NA	NA	NA
173767	NA	NA	NA
173769	NA	NA	NA
173770	0	0	1
173772	NaN	0	NaN
173774	NA	NA	NA
173775	NA	NA	NA
173776	NA	NA	NA

The correct result should be somewhat close to the non-robust result:

```
> snpStats::snp.lhs.tests(Autosomes[, 1:10], ~strata(cc), ~strata(region),
+ data = subject.data, robust = FALSE)
```

```
Chi.squared Df p.value
173760 NaN 9 NaN
173761 12.272003 9 0.1984058
173762 12.476151 9 0.1877771
173767 14.462676 9 0.1067926
```

```
8.589652
                     9 0.4759812
173769
          0.000000 0 1.0000000
173770
173772
                {\tt NaN}
                     9
                              NaN
                     9 0.7241952
173774
          6.156140
173775
          8.812111
                     9 0.4547958
          7.602749
173776
                     9 0.5746207
```

Now we convert snpStats classes (mixed-cases without ".") to snpMatrix's (lowercases with "."), and re-run the the snpMatrix version of snp.lhs.tests():

```
1:10], "strata(cc), "strata(region), data = subject.data,
      robust = TRUE)
       Chi.squared Df
                         p.value
173760
          1.009134
                    9 0.99941616
173761
         10.263701 9 0.32956204
173762
         10.476287 9 0.31331934
173767
         15.045105 9 0.08970445
         16.267175 9 0.06150760
173769
          0.000000 0 1.00000000
173770
173772
          1.009439 9 0.99941544
173774
          9.411857
                    9 0.40015665
173775
         13.440413
                    9 0.14366942
173776
         16.797680 9 0.05198017
```

> snpMatrix::snp.lhs.tests(new("snp.matrix", Autosomes@.Data)[,

Just to see what snpMatrix does without robust:

```
> snpMatrix::snp.lhs.tests(new("snp.matrix", Autosomes@.Data)[,
      1:10], "strata(cc), "strata(region), data = subject.data,
      robust = FALSE)
       Chi.squared Df
                        p.value
          5.042974 9 0.8305461
173760
173761
         12.272003 9 0.1984058
173762
         12.476151 9 0.1877771
         14.462676 9 0.1067926
173767
173769
          8.589652 9 0.4759812
173770
          0.000000 0 1.0000000
173772
          7.714940
                    9 0.5631090
173774
          6.156140 9 0.7241952
173775
          8.812111 9 0.4547958
173776
          7.602749 9 0.5746207
```

This bug also exist in snpMatrix prior to 1.17.5.10 and was introduced with the imputation-related changes around October 2008.

See also how snpMatrix124 does:

```
> snpMatrix124::snp.lhs.tests(new("snp.matrix", Autosomes@.Data)[,
      1:10], "strata(cc), "strata(region), data = subject.data,
      robust = TRUE)
       Chi.squared Df Df.residual
173760
          1.008943 8
173761
         10.263701 9
                               398
173762
         10.476287
                               396
173767
         15.045105
                               376
         16.006621
173769
                    8
                               394
173770
                NA NA
                               NA
          1.009439
173772
                   8
                               199
173774
          9.411857
                               386
173775
         13.440413
                               397
173776
         16.797680
                               398
> snpMatrix124::snp.lhs.tests(new("snp.matrix", Autosomes@.Data)[,
      1:10], "strata(cc), "strata(region), data = subject.data,
      robust = FALSE)
       Chi.squared Df Df.residual
173760
          5.042974 9
                               198
173761
         12.272003 9
                               398
173762
         12.476151 9
                               396
173767
         14.462676 9
                               376
          8.589652 9
                               394
173769
173770
                NA NA
                               NA
173772
          7.714940
                               199
173774
          6.156140
                    9
                               386
173775
          8.812111
                    9
                               397
173776
          7.602749 9
                               398
```

4.2 Malformed "GlmTests" S4 object from snpStats::snp.lhs.tests()

Upto and including snpStats 1.3.6:

```
> result <- snpStats::snp.lhs.tests(Autosomes[, 1:10], ~strata(cc),
+ ~strata(region), data = subject.data)</pre>
```

Looking at result gives a hard error so I'll just show the message below:

```
> str(result)
Error in FUN(c("snp.names", "var.names", "chisq", "df", "N")[[2L]], ...) :
  no slot of name "var.names" for this object of class "GlmTests"
```

There is no need to show the alternatives as this is clearly broken.

This bug is specific to snpStats and has no equivalent in snpMatrix.

4.3 Crazy large/negative number of samples from snpStats GLM tests

Upto and including snpStats 1.3.6:

> result@N

- [1] 202248192 202248160 202248128 202248096 202248064 202248032 202248000
- [8] 202247968 202247936 202247904

Hundred million samples and negative number of samples?

This bug also exist in snpMatrix prior to 1.17.5.10 and was introduced with the imputation-related changes around October 2008.

4.4 snpStats::snp.rhs.tests() returning garbage with or without robust

```
> snpStats::snp.rhs.tests(cc ~ strata(region, sex), family = "binomial",
+ data = subject.data, snp.data = Autosomes, tests = 1:10)
```

Chi.squared	Df	<pre>p.value</pre>
NaN	1	NaN
NA	0	NA
NaN	1	NaN
	NaN NaN NaN NaN NA NA NaN	NaN 1 NaN 1 NaN 1 NaN 1 NAN 0 NAN 1 NAN 1 NAN 1 NAN 1

The correct result shouldn't be too far from without sex:

```
> snpStats::snp.rhs.tests(cc ~ strata(region), family = "binomial",
+ data = subject.data, snp.data = Autosomes, tests = 1:10)
```

```
Chi.squared Df p.value
173760 1.01538462 1 0.31361630
173761 1.46259571 1 0.22651757
173762 1.92028786 1 0.16582493
173767 0.77609738 1 0.37833736
173769 2.92614948 1 0.08715513
173770 NA 0 NA
173772 1.11008326 1 0.29206385
173774 0.66697270 1 0.41410906
173775 0.96730037 1 0.32535438
173776 0.09831885 1 0.75385649
```

Here is how snpMatrix does it:

```
> snpMatrix::snp.rhs.tests(cc ~ strata(region, sex), family = "binomial",
      data = subject.data, snp.data = new("snp.matrix", Autosomes@.Data),
      tests = 1:10)
       Chi.squared Df
                         p.value
173760 1.25356125 1 0.26287339
173761 1.61290542 1 0.20408387
173762 2.04226350 1 0.15298186
173767 0.29671726 1 0.58594776
173769 3.54351327 1 0.05977872
173770
               NA O
                              NA
173772 0.59863946 1 0.43909761
173774 0.82443150 1 0.36388768
173775 0.87744532 1 0.34890234
173776 0.09218633 1 0.76141588
  Just to see that snpMatrix does it without sex:
> snpMatrix::snp.rhs.tests(cc ~ strata(region), family = "binomial",
      data = subject.data, snp.data = new("snp.matrix", Autosomes@.Data),
      tests = 1:10)
       Chi.squared Df
                         p.value
173760 1.01538462 1 0.31361630
173761 1.46259571 1 0.22651757
173762 1.92028786 1 0.16582493
173767 0.77609738 1 0.37833736
173769 2.92614948 1 0.08715513
173770
               NA O
                              NΑ
173772 1.11008326 1 0.29206385
173774 0.66697270 1 0.41410906
173775 0.96730037 1 0.32535438
173776 0.09831885 1 0.75385649
  This bug also exist in snpMatrix prior to 1.17.5.10 and was introduced with the imputation-
related changes around October 2008.
  See how snpMatrix124 does:
> snpMatrix124::snp.rhs.tests(cc ~ strata(region, sex), family = "binomial",
      data = subject.data, snp.data = new("snp.matrix", Autosomes@.Data),
      tests = 1:10)
       Chi.squared Df Df.residual
173760 1.25356125 1
                              373
173761 1.61290542 1
                              376
173762 2.04226350 1
                              374
173767 0.29671726 1
                              350
```

```
173769 3.54958407 1
                              372
173770
               NA NA
                              376
173772 0.59863946 1
                              376
173774 0.82443150 1
                              365
173775 0.87744532 1
                              375
173776 0.09218633 1
                              376
> snpMatrix124::snp.rhs.tests(cc ~ strata(region), family = "binomial",
      data = subject.data, snp.data = new("snp.matrix", Autosomes@.Data),
      tests = 1:10)
       Chi.squared Df Df.residual
173760 1.01538462 1
173761 1.46259571 1
                              390
173762 1.92028786 1
                              388
173767 0.77609738 1
                              368
173769 2.92614948 1
                              386
173770
                              390
                NA NA
173772 1.11008326 1
                              390
173774 0.66697270 1
                              378
173775 0.96730037 1
                              389
173776 0.09831885 1
                              390
    2-df Bug in snpStats::single.snp.tests()
5
> data(for.exercise, package = "snpStats")
> tests.snpStats <- snpStats::single.snp.tests(cc, stratum, data = subject.support,
      snp.data = snps.10)
  Now we convert snpStats classes (mixed-cases without ".") to snpMatrix's (lowercases with "."),
and re-run the the snpMatrix version of single.snp.tests():
> str(snps.10)
Formal class 'SnpMatrix' [package "snpStats"] with 1 slots
  ..@ .Data: raw [1:1000, 1:28501] 01 01 01 01 ...
  ... - attr(*, "dimnames")=List of 2
  .....$: chr [1:1000] "jpt.869" "jpt.862" "jpt.948" "ceu.564" ...
  .....$ : chr [1:28501] "rs7909677" "rs7093061" "rs12773042" "rs7475011" ...
> snps.10 <- new("snp.matrix", snps.100.Data)</pre>
> str(snps.10)
Formal class 'snp.matrix' [package "snpMatrix"] with 1 slots
  ..@ .Data: raw [1:1000, 1:28501] 01 01 01 01 ...
  ...- attr(*, "dimnames")=List of 2
  .....$ : chr [1:1000] "jpt.869" "jpt.862" "jpt.948" "ceu.564" ...
  .....$ : chr [1:28501] "rs7909677" "rs7093061" "rs12773042" "rs7475011" ...
```

```
> tests.snpMatrix <- snpMatrix::single.snp.tests(cc, stratum, data = subject.support,
      snp.data = snps.10)
   Then we use the testsuite code to compare:
> all.equal(tests.snpStats@chisq[, "1 df"], tests.snpMatrix@chisq[,
      "1 df"], tolerance = 0)
[1] TRUE
> all.equal(tests.snpStats@chisq[, "2 df"], tests.snpMatrix@chisq[,
      "2 df"], tolerance = 0)
[1] "'is.NA' value mismatch: 787 in current 807 in target"
> snpMatrix:::.chi2.all.equal(tests.snpStats@chisq[, "2 df"], tests.snpMatrix@chisq[,
      "2 df"])
Max absolute finite difference: 0
Max relative finite difference: 0
Finite in 1st but not in 2nd: 0
Finite in 2nd but not in 1st: 20
    Min. 1st Qu. Median
                              Mean 3rd Qu.
                                                  Max.
0.003281 1.167000 2.272000 2.070000 2.952000 4.258000
Difference
        20
   Or 20 SNP tests failed in snpStats but okay in snpMatrix.
   Now we run the non-stratified tests, but convert in the opposite direction, and compare:
> tests.snpMatrix.crude <- snpMatrix::single.snp.tests(cc, data = subject.support,
      snp.data = snps.10)
> str(snps.10)
Formal class 'snp.matrix' [package "snpMatrix"] with 1 slots
  ..@ .Data: raw [1:1000, 1:28501] 01 01 01 01 ...
  ... - attr(*, "dimnames")=List of 2
  .....$ : chr [1:1000] "jpt.869" "jpt.862" "jpt.948" "ceu.564" ...
  .....$ : chr [1:28501] "rs7909677" "rs7093061" "rs12773042" "rs7475011" ...
> snps.10 <- new("SnpMatrix", snps.10@.Data)
> str(snps.10)
Formal class 'SnpMatrix' [package "snpStats"] with 1 slots
  ..@ .Data: raw [1:1000, 1:28501] 01 01 01 01 ...
  ... - attr(*, "dimnames")=List of 2
  .....$ : chr [1:1000] "jpt.869" "jpt.862" "jpt.948" "ceu.564" ...
  .....$ : chr [1:28501] "rs7909677" "rs7093061" "rs12773042" "rs7475011" ...
```

```
> tests.snpStats.crude <- snpStats::single.snp.tests(cc, data = subject.support,
      snp.data = snps.10)
> all.equal(tests.snpStats.crude@chisq[, "2 df"], tests.snpMatrix.crude@chisq[,
      "2 df"], tolerance = 0)
[1] "'is.NA' value mismatch: 789 in current 807 in target"
> snpMatrix:::.chi2.all.equal(tests.snpStats.crude@chisq[, "2 df"],
      tests.snpMatrix.crude@chisq[, "2 df"])
Max absolute finite difference: 0
Max relative finite difference: 0
Finite in 1st but not in 2nd: 0
Finite in 2nd but not in 1st: 18
  Min. 1st Qu. Median Mean 3rd Qu.
                                           Max.
  1.043
          1.408 2.405 2.323 2.865
                                           3.922
Difference
        18
  Or 18 SNP tests failed in snpStats but okay in snpMatrix.
  This bug also exist in snpMatrix prior to 1.17.4.9 for its entire history (i.e. since pre-1.0), and
differently before 1.5.x also.
  This explains why snpStats and snpMatrix124 differs:
> snps.10 <- new("snp.matrix", snps.10@.Data)
> tests.snpMatrix124 <- snpMatrix124::single.snp.tests(cc, stratum,
      data = subject.support, snp.data = snps.10)
> all.equal(tests.snpStats@chisq[, "1 df"], tests.snpMatrix124$chi2.1df,
      tolerance = 0)
[1] "Mean relative difference: 2.163423e-16"
> all.equal(tests.snpStats@chisq[, "2 df"], tests.snpMatrix124$chi2.2df,
     tolerance = 0)
[1] "'is.NA' value mismatch: 789 in current 807 in target"
> snpMatrix:::.chi2.all.equal(tests.snpMatrix124$chi2.1df, tests.snpMatrix@chisq[,
      "1 df"])
Max absolute finite difference: 7.105427e-15
Max relative finite difference: 8.941748e-13
Finite in 1st but not in 2nd: 0
Finite in 2nd but not in 1st: 0
Difference
```

> snpMatrix:::.chi2.all.equal(tests.snpStats@chisq[, "2 df"], tests.snpMatrix124\$chi2.2df)

```
Max absolute finite difference: 3.907985e-14
Max relative finite difference: 2.035703e-14
Finite in 1st but not in 2nd: 0
Finite in 2nd but not in 1st: 18
  Min. 1st Qu. Median Mean 3rd Qu.
                                           Max.
0.8604 1.2830 2.4020 2.3000 2.9840 4.2580
Difference
        18
> snpMatrix:::.chi2.all.equal(tests.snpMatrix124$chi2.2df, tests.snpMatrix@chisq[,
      "2 df"])
Max absolute finite difference: 3.907985e-14
Max relative finite difference: 3.896745e-14
Finite in 1st but not in 2nd: 0
Finite in 2nd but not in 1st: 2
   Min. 1st Qu.
                   Median
                              Mean 3rd Qu.
0.003281 0.003378 0.003476 0.003476 0.003573 0.003671
Difference
```

So out of 20 of those SNPs that snpStats failed, snpMatrix124 can do 18 and much closer to $snpMatrix\ 1.17.6.10+$.

6 64-bit mode for snpStats::single.snp.tests()

This returns all zeros in 64-bit machine occasionally (corruption?):

```
snp.lhs.tests(Autosomes[,1:10], ~cc, ~region, data=subject.data)
```

7 X chromosome conversion

> data(testdata, package = "snpStats")

The corresponding X-chromosome conversion is as follows:

```
> str(Xchromosome)

Formal class 'XSnpMatrix' [package "snpStats"] with 2 slots
    ..@ .Data : raw [1:400, 1:155] 03 03 03 01 ...
    ... - attr(*, "dimnames")=List of 2
    ... ..$ : chr [1:400] "1987" "436" "762" "1199" ...
    ... ..$ : chr [1:155] "174193" "174196" "174197" "174208" ...
    ... .@ diploid: Named logi [1:400] TRUE FALSE TRUE FALSE FALSE FALSE ...
    ... - attr(*, "names")= chr [1:400] "1987" "436" "762" "1199" ...
```

```
> Xchromosome <- new("X.snp.matrix", Xchromosome@.Data, Female = Xchromosome@diploid)
> str(Xchromosome)

Formal class 'X.snp.matrix' [package "snpMatrix"] with 2 slots
    ..@ .Data : raw [1:400, 1:155] 03 03 03 01 ...
    ... - attr(*, "dimnames")=List of 2
    ... ...$ : chr [1:400] "1987" "436" "762" "1199" ...
    ... ...$ : chr [1:155] "174193" "174196" "174197" "174208" ...
    ... ..@ Female: Named logi [1:400] TRUE FALSE TRUE FALSE FALSE FALSE ...
    ... - attr(*, "names")= chr [1:400] "1987" "436" "762" "1199" ...
```