Genotyping with the crlmm Package

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March, 2009

1 Quick intro to crlmm

The crlmm package contains a new implementation for the CRLMM algorithm (Carvalho et. al. 2007). Our focus is on efficient genotyping of SNP 5.0 and 6.0 Affymetrix arrays, although extensions of the method are under development for similar platforms.

This implementation, compared to the previous one (in oligo), offers improved confidence scores, quality scores for SNP's and batches, higher accuracy on different datasets and better performance.

Additionally, this package does not use the pd.genomewidesnp packages created via pdInfoBuilder for oligo. Instead, it uses different annotation packages (genomewidesnp.5 and genomewidesnp.6), which use simple R objects to store only the information needed for genotyping. This allowed us to improve the speed of the method, as SQL queries are no longer performed here.

It is also our priority to make the package simple to use. Below we demonstrate how to get genotype calls with the 'new' CRLMM. We use 3 samples on SNP 5.0 made available via the hapmapsnp5 package.

```
R> require(oligoClasses)
R> library(crlmm)
R> library(hapmapsnp6)
R> path <- system.file("celFiles", package="hapmapsnp6")
R> celFiles <- list.celfiles(path, full.names=TRUE)
R> system.time(crlmmResult <- crlmm(celFiles, verbose=FALSE))
user system elapsed</pre>
```

103.608 0.835 105.056

The crlmmResult is a *SnpSet* (see Biobase) object.

- calls: genotype calls (1 AA; 2 AB; 3 BB);
- confs: confidence scores, which can be translated to probabilities by using:

 $1 - 2^{-}(\text{confs}/1000),$

although we prefer this representation as it saves a significant amount of memory;

- SNPQC: SNP quality score;
- SNR: Signal-to-noise ratio.

R> calls(crlmmResult)[1:10,]

	NA06985_GW6_C.CEL	NA06991_GW6_C.CEL		
SNP_A-2131660	2	2		
SNP_A-1967418	3	3		
SNP_A-1969580	3	3		
SNP_A-4263484	2	1		
SNP_A-1978185	1	1		
SNP_A-4264431	1	1		
SNP_A-1980898	3	3		
SNP_A-1983139	1	1		
SNP_A-4265735	2	2		
SNP_A-1995832	2	3		
NA06993_GW6_C.CEL				
SNP_A-2131660	3			
SNP_A-1967418	3			
SNP_A-1969580	3			
SNP_A-4263484	1			
SNP_A-1978185	1			
SNP_A-4264431	1			
SNP_A-1980898	3			
SNP_A-1983139	1			
SNP_A-4265735	1			
SNP_A-1995832	3			

R> confs(crlmmResult)[1:10,]

	NA06985_GW6_C.CEL	NA06991_GW6_C.CEL		
SNP_A-2131660	0.9999964	0.9999996		
SNP_A-1967418	0.9999969	0.9999997		
SNP_A-1969580	0.9995187	0.9995139		
SNP_A-4263484	0.9999999	1.0000000		
SNP_A-1978185	1.0000000	1.0000000		
SNP_A-4264431	1.0000000	1.0000000		
SNP_A-1980898	0.9995192	0.9995206		
SNP_A-1983139	1.0000000	0.9999877		
SNP_A-4265735	0.9999822	0.9999862		
SNP_A-1995832	0.9999758	1.0000000		
NA06993_GW6_C.CEL				
SNP_A-2131660	0.9999998			
SNP_A-1967418	0.9999969			
SNP_A-1969580	0.9995124			
SNP_A-4263484	1.0000000			

1.0000000
1.0000000
0.9995134
1.0000000
0.9999998
0.9999999

```
R> crlmmResult[["SNR"]]
```

[1] 8.578197 8.368455 7.265255

2 Details

This document was written using:

```
R> sessionInfo()
R version 3.2.3 (2015-12-10)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Ubuntu 14.04.3 LTS
locale:
 [1] LC_CTYPE=en_US.UTF-8
                              LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
                                LC_COLLATE=C
 [5] LC_MONETARY=en_US.UTF-8
                               LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                               LC_NAME=C
 [9] LC_ADDRESS=C
                                LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] stats
              graphics grDevices utils
                                            datasets
[6] methods
              base
other attached packages:
[1] genomewidesnp6Crlmm_1.0.7 hapmapsnp6_1.12.0
[3] crlmm_1.28.2
                              preprocessCore_1.32.0
[5] oligoClasses_1.32.0
loaded via a namespace (and not attached):
 [1] Rcpp_0.12.3
                                XVector_0.10.0
 [3] splines_3.2.3
                                GenomicRanges_1.22.3
 [5] BiocGenerics_0.16.1
                                zlibbioc_1.16.0
 [7] IRanges_2.4.6
                                bit_1.1-12
 [9] ellipse_0.3-8
                                lattice_0.20-33
[11] foreach_1.4.3
                                GenomeInfoDb_1.6.1
[13] tools_3.2.3
                                base64_1.1
```

[15]	grid_3.2.3	SummarizedExperiment_1.0.2
[17]	parallel_3.2.3	Biobase_2.30.0
[19]	ff_2.2-13	DBI_0.3.1
[21]	matrixStats_0.50.1	iterators_1.0.8
[23]	RcppEigen_0.3.2.5.1	affyio_1.40.0
[25]	Matrix_1.2-3	S4Vectors_0.8.8
[27]	codetools_0.2-14	VGAM_1.0-0
[29]	RSQLite_1.0.0	BiocInstaller_1.20.1
[31]	Biostrings_2.38.3	stats4_3.2.3
[33]	mvtnorm_1.0-3	illuminaio_0.12.0