# *VanillaICE*: Hidden Markov Models for the Assessment of Chromosomal Alterations using High-throughput SNP Arrays

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#### Abstract

This package provides an implementation of a hidden Markov Model for high throughput SNP arrays. Users of this package should already have available locus-level estimates of copy number and B allele frequencies or genotype calls. Copy number estimates can be relative or absolute.

## 1 Overview

This vignette requires that you have

• an absolute estimate of the *total* copy number organized such that rows correspond to loci and columns correspond to samples

and / or

• a matrix of B allele frequencies or genotype calls (1=AA, 2 = AB, 3= BB): rows correspond to loci and columns correspond to samples

If Affymetrix CEL files or Illumina Idat files were processed with the R package *crlmm*, see the vignette crlmmDownstream included with this package.

Other HMM implementations for the joint analysis of copy number and genotype include QuantiSNP [1] and PennCNV [3].

**Data considerations.** The HMM implemented in this package is most relevant for heritable diseases for which integer copy numbers are expected. For somatic cell diseases such as cancer, we suggest circular binary segmentation, as implemented in the R package DNAcopy [2].

**Citing this software.** Robert B Scharpf, Giovanni Parmigiani, Jonathan Pevsner, and Ingo Ruczinski. Hidden Markov models for the assessment of chromosomal alterations using high-throughput SNP arrays. *Annals of Applied Statistics*, 2(2):687–713, 2008.

### 2 Organizing the locus-level data

This package includes simulated genotype and copy number data for 9165 SNPs.

```
> library(oligoClasses)
```

- > library(VanillaICE)
- > library2(SNPchip)
- > library2(Biobase)
- > library2(IRanges)
- > data(locusLevelData)

The copy number estimates in the locusLevelData object were multiplied by 100 and saved as an integer. An object of class oligoSnpSet is instantiated from the simulated data in the next code chunk. Note that the assay data elements for the oligoSnpSet object should be integers. The integerMatrix scales the copy number matrix by a factor of 100 (by default) and returns a matrix of integers. When available, B allele frequencies should be scaled by a factor of 1000. For example,

integerMatrix(bAlleleFreq, scale=1000) \verb

```
> cn <- log2(locusLevelData[["copynumber"]]/100)
> oligoSet <- new("oligoSnpSet",
+ copyNumber=integerMatrix(cn),
+ call=locusLevelData[["genotypes"]],
+ callProbability=locusLevelData[["crlmmConfidence"]],
+ annotation=locusLevelData[["platform"]])
> oligoSet <- oligoSet[!is.na(chromosome(oligoSet)), ]
> oligoSet <- oligoSet[chromosome(oligoSet) < 3, ]</pre>
```

## 3 Fitting the HMM

When jointly modeling the copy number and genotypes/allele frequencies, we assume that the genotype and copy number estimates are independent conditional on the underlying hidden state. The method hmm is defined for several classes, including objects of class oligoSnpSet and BeadStudioSet.

The viterbi algorithm is used to obtain the most likely sequence of hidden states given the observed data. The value returned is an object of class RangedDataHMM with genomic coordinates of the normal and altered regions. We also return the log-likelihood ratio (LLR) of the predicted sequence in an interval versus the likelihood of diploid copy number. For intervals inferred to have diploid copy number, the LLR is zero.

```
> oligoSet <- oligoSet[chromosome(oligoSet) <= 2, ]
> fit.van <- hmm(oligoSet)</pre>
```

Figure 1 plots the data for chromosome 1 and the predictions from the hidden markov model:

To find which markers are included in each genomic interval returned by the hmm method, one can use the findOverlaps method in the *oligoClasses*. For example, the second interval in the RangedDataHMM object fit.van contains 102 markers.

```
> fit.van[2, ]
```

RangedDataHMM with 1 row and 7 value columns across 1 space chrom num.mark space ranges | id <factor> <IRanges> | <integer> <integer> <character> 1 [49825223, 54637167] | NA06993 1 1 102 start.index end.index state LLR <integer> <integer> <integer> <numeric> 1 997 1098 4 14.84152

To find the names of the 102 markers that are included in this genomic interval, one could do the following

```
> mm <- as.matrix(findOverlaps(fit.van, oligoSet))
> markersInRange <- featureNames(oligoSet)[mm[,1]==2]</pre>
```

Multipanel displays can be useful for visualizing the low-level data for copy number alterations. We extend the xyplot method in the R package lattice for two common use-cases: by-locus and by-sample.



Figure 1: Plot of artificial data for one chromosome.

**By locus.** To plot the genomic data for a set of ranges at a given locus, we provide an unevaluated code chunk below (our example contains only a single sample):

> xyplot(cn ~ x | id, oligoSet, range=RangedDataObject, ylim=c(-0.5,4), panel=xypanel)

Note that the default in the above command is to use a common x- and y-scale for each panel. To allow the x-axes to change for each panel, one could set the x-scales to 'free':

> xyplot(cn ~ x | id, oligoSet, range=RangedDataObject, ylim=c(-0.5,4), scales=list(x="free"), + panel=xypanel)

The function **xypanel** provides default colors for annotating the plotting symbols by genotype and by whether the markers are polymorphic. The **RangedDataHMM** must be passed by the name *range* to the **xyplot** method.

By sample. To plot the low-level data for multiple alterations occurring in a single sample, one can again pass a RangedDataHMM object with name *range* to the xyplot. For example, see Figure 2. The code for producing Figure 2 is in the following code chunk. Note that the formula in the example below conditions on *range* instead of *id*. The conditioning variable for displaying multiple panels of a single sample must be called 'range'. We plot a 2 Mb window surrounding each of the alterations in the simulated data by specifying frame=2e6.

```
> ranges.altered <- fit.van[!state(fit.van) %in% c(3,4) & coverage2(fit.van) > 5, ]
> xy.example <- xyplot(cn ~ x | range, oligoSet, range=ranges.altered, frame=2e6,</pre>
```

```
+ scales=list(x="free"), ylim=c(-1,3),
+ panel=xypanel, cex.pch=0.4, pch=21, border="grey",
+ ylab=expression(log[2]("copy number")))
```

See ?xyplot for additional details.

Note also that scales must be set to *free* in the above call to xyplot.

#### 3.1 ICE HMM

This will likely be phased out in favor of using B allele frequencies instead of genotype calls / call probabilities.

To compute emission probabilities that incorporate the *crlmm* genotype confidence scores, we set ICE to TRUE. This option is limited too a few platforms and the Affy 100k platforms is not one of them.

```
> VanillaICE:::icePlatforms()
```

```
[1] "pd.genomewidesnp.6"
```

```
[2] "genomewidesnp6"
```

```
[3] "genomewidesnp6Crlmm"
```

```
[4] "pd.mapping250k.nsp"
```

```
[5] "pd.mapping250k.sty"
```

```
[6] "pd.mapping250k.nsp, pd.mapping250k.sty"
```

For illustration, we assign 'genomewidesnp6' to the annotation slot since this platform is supported.

```
> ann <- annotation(oligoSet)
> annotation(oligoSet) <- "genomewidesnp6"
> fit.ice <- hmm(oligoSet, ICE=TRUE)
> fit.ice
```

RangedDataHMM with 17 rows and 7 value columns across 1 space

	~							
	space			ranges	Ι	chrom	num.mark	id
	<factor></factor>			<iranges></iranges>	Ι	<integer></integer>	<integer></integer>	<character></character>
1	1	[	846864,	49772452]	Ι	1	996	NA06993
2	1	[ 4	49825223,	54637167]	Ι	1	102	NA06993
3	1	[ {	54726362,	70065480]	Ι	1	902	NA06993
4	1	[ ]	70081878,	73401801]	Ι	1	204	NA06993
5	1	[ ]	73804712,	121226982]	Ι	1	2497	NA06993
6	1	[14	44908589,	176547669]	Ι	1	1294	NA06993
7	1	[1]	76548473,	178437444]	Ι	1	98	NA06993
8	1	[1]	78533776,	218173294]	Ι	1	1901	NA06993
9	1	[2:	18219379,	219806187]	Ι	1	99	NA06993
10	1	[2:	19825554,	248794371]	Ι	1	1065	NA06993
11	1	[	170616,	8409113]	Ι	2	4	NA06993
12	1	[ :	15224340,	68466618]	Ι	2	29	NA06993
13	1	[ ]	75654059,	79296379]	Ι	2	3	NA06993
14	1	[ 8	31048974,	88467934]	Ι	2	4	NA06993
15	1	[1(	02998279,	105880487]	Ι	2	2	NA06993
16	1	[1:	14471782,	115837414]	Ι	2	2	NA06993
17	1	[12	23126479,	240767758]	Ι	2	56	NA06993
	start.ind	lex	end.index	x state		LLR		
	<pre><integer> <integer> <integer></integer></integer></integer></pre>					<numeric></numeric>		
1		1	996	3 3		0.0000000		

2	997	1098	4	14.8415228
3	1099	2000	3	0.000000
4	2001	2204	5	195.1615962
5	2205	4701	3	0.000000
6	1	1294	3	0.000000
7	1295	1392	2	166.9786379
8	1393	3293	3	0.000000
9	3294	3392	2	205.2171294
10	3393	4457	3	0.000000
11	1	4	5	-0.5191900
12	5	33	3	0.000000
13	34	36	5	-2.4122910
14	37	40	2	-0.4449246
15	41	42	5	1.3314298
16	43	44	4	0.4059383
17	1	56	3	0.000000

> annotation(oligoSet) <- ann</pre>

#### 3.2 Other options

**Regions of homozygosity.** A HMM for genotype-only data can be used to find long stretches of homozygosity. Note that hemizygous deletions are also identified as 'ROH' when copy number is ignored (as the biallelic genotypte call in a hemizygous deletions tends to be all homozygous calls).

```
> snpSet <- as(oligoSet, "SnpSet")
> fit.gt <- hmm(snpSet, prGtHom=c(0.7, 0.99), normalIndex=1L, S=2L)</pre>
```

A suggested visualization:



## 4 Session Information

The version number of R and packages loaded for generating the vignette were:

- R version 2.15.0 beta (2012-03-22 r58799), x86\_64-unknown-linux-gnu
- Locale: LC\_CTYPE=en\_US.iso885915, LC\_NUMERIC=C, LC\_TIME=en\_US.iso885915, LC\_COLLATE=en\_US.iso885915, LC\_MONETARY=en\_US.iso885915, LC\_MESSAGES=en\_US.iso885915, LC\_PAPER=C, LC\_NAME=C, LC\_ADDRESS=C, LC\_TELEPHONE=C, LC\_MEASUREMENT=en\_US.iso885915, LC\_IDENTIFICATION=C
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: Biobase 2.15.4, BiocGenerics 0.1.14, BiocInstaller 1.3.8, codetools 0.2-8, crlmm 1.13.13, DBI 0.2-5, foreach 1.3.5, IRanges 1.13.32, iterators 1.0.5, oligo 1.19.18, oligoClasses 1.17.36, pd.mapping50k.hind240 1.4.0, pd.mapping50k.xba240 1.4.0, RColorBrewer 1.0-5, RSQLite 0.11.1, SNPchip 2.0.4, VanillaICE 1.17.21
- Loaded via a namespace (and not attached): affxparser 1.27.5, affyio 1.23.2, annotate 1.33.8, AnnotationDbi 1.17.27, Biostrings 2.23.6, bit 1.1-8, compiler 2.15.0, ellipse 0.3-7, ff 2.2-5, genefilter 1.37.1, grid 2.15.0, lattice 0.20-6, msm 1.1, mvtnorm 0.9-9992, preprocessCore 1.17.7, splines 2.15.0, stats4 2.15.0, survival 2.36-12, tools 2.15.0, xtable 1.7-0, zlibbioc 1.1.1

#### References

- [1] Stefano Colella, Christopher Yau, Jennifer M Taylor, Ghazala Mirza, Helen Butler, Penny Clouston, Anne S Bassett, Anneke Seller, Christopher C Holmes, and Jiannis Ragoussis. QuantiSNP: an Objective Bayes Hidden-Markov Model to detect and accurately map copy number variation using SNP genotyping data. Nucleic Acids Res, 35(6):2013–2025, 2007.
- [2] Adam B Olshen, E. S. Venkatraman, Robert Lucito, and Michael Wigler. Circular binary segmentation for the analysis of array-based DNA copy number data. *Biostatistics*, 5(4):557–72, Oct 2004.
- [3] Kai Wang, Mingyao Li, Dexter Hadley, Rui Liu, Joseph Glessner, Struan F A Grant, Hakon Hakonarson, and Maja Bucan. Pennenv: an integrated hidden markov model designed for high-resolution copy number variation detection in whole-genome snp genotyping data. *Genome Res*, 17(11):1665– 1674, Nov 2007.



Figure 2: The method xyplot is used to create a multi-panel display of alterations in a single sample. Each panel displays a single copy number alteration detected by the HMM that is boxed by a rectangle. The alteration is framed by specifying the number of basepairs to plot upstream and downstream of the alteration. Here, we used a frame of 2 Mb. Homozygous SNPs with diallelic geotypes 'AA' and 'BB' are shaded blue; SNPs with diallelic genotype call 'AB' are shaded in red.