## Package "maskBAD"

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#### Abstract

Package "maskBAD" allows to users to identify, inspect and remove probes for which binding affinity differs between two groups of samples. It works for Affymetrix 3' IVT and whole transcript (exon and gene) arrays. When there are SNPs, transcript isoforms or cross hybrydizing transcripts specific to only one of compared groups, probes' binding affinities may differ between groups. Such probes bias estimates of gene expression, introduce false positives in differential gene expression analysis and reduce the power to detect real expression differences. Hence they should be identified and removed from the data at preprocessing stage. For details about impact of BAD (binding affinity difference) probes on differential gene expression estimates, implemented detection method and its power to improve data quality, please consult our papers Dannemann, Lorenc et al.: "The effects of probe binding affinity differences on gene expression measurements and how to deal with them", Bioinformatics, 2009 ([2]) and Dannemann et al.: "maskBAD' - a package to detect and remove Affymetrix probes with binding affinity differences", BMC Bioinformatics 2012 ([1]).

#### **1** Introduction

The "maskBAD" package implements functions to detect and remove probes with different binding affinity (BAD) in Affymetrix array expression data. Identification and removal of BAD probes removes spurious gene expression differences and helps recover true ones. BAD probes are prevalent in comparisons of genetically distinct samples, like belonging to different strains or species, but systematic qualitative differences in transcriptome might introduce them also when samples differ by treatment, health status or tissue type. Masking therefore should be a routine step in data preprocessing. In this package we introduce functions that allow to identify, inspect and remove BAD probes and show how it can be integrated in a standard gene expression analysis pipeline.

### 2 Identification of BAD probes

Detection of BAD (binding affinity difference) probes is done on an AffyBatch object, usually prepared with ReadAffy() from package *affy*. We'll use 100 random probe sets from human and chimpanzee expression dataset ([3]), each with 10 individuals, available with the package.

```
> ## data available by loading data(AffyBatch)
> newAffyBatch
AffyBatch object
size of arrays=640x640 features (22 kb)
cdf=newCdf (100 affyids)
number of samples=20
number of genes=100
annotation=hgu95av2
notes=
> exmask <- mask(newAffyBatch,ind=rep(1:2,each=10),verbose=FALSE,useExpr=F)</pre>
```

The scoring of probes is performed with the function mask and is based on the algorithm presented in Dannemann et. al ([2]). Briefly, for a given probe, its signal is proportional to the amount of RNA in the sample and its binding affinity. When one target is measured with several probes (a probe set), as on Affymetrix arrays, the probes' signals are correlated for each sample. BAD probes correlation differs between groups, hence comparison of probes' pairwise correlations between those groups allows to identify of BAD probes. Function mask() detects probes differing in binding affinity between groups of samples defined by argument ind. The masking algorithm workes on a gene/probe set basis as the expression signal for the same transcript should be correlated among probes. Therefore, to identify BAD probes use probe sets defined at transcript/gene level.

The method works properly only for probe sets with signal above background in both groups. By default (use.expr=TRUE option) only probe sets with mas5calls() "P" in at least 90% of samples from both groups are considered in BAD detection. Another desired set of probe sets might be specified by the parameter exprlist.

#### 2.1 Evaluation of the mask and choice of cutoff

For human-chimpanzee, with assumption of sequence divergence 1%, we'll expect naively  $\sim 22\%$  of probes comprise a SNP. To filter out lower 22% of quality scores, we have to put a cutoff at 0.029.

```
> quantile(exmask[[1]]$quality.score,0.22)
```

22%

0.02922448

Manual inspection of borderline cases with plotProbe() might help to decide about the cutoff. It plots signal intensity of a probe against all other probes from the same probe set.

#### 2.2 Including external data

Mask might be calibrated with a subset of probes with known BAD status (called later on external mask), for example probes with target sequences known in both compared groups. External mask contains x and y coordinates and probe set assignement for the probes, along with their BAD status coded as 0/1 (BAD probe/not BAD probe). For a given cutoff, compare BAD status according to expression-based mask and given by external mask. Here we use the object sequenceMask - the information whether the probe sequence is affected by a SNP between the groups (0) or not (1).

> head(exmask\$probes)

x y quality.score probeset 399.559 399 559 0.139886115 1000\_at 544.185 544 185 0.485398831 1000\_at 530.505 530 505 0.704649407 1000\_at

```
617.349 617 349
                  0.008627547
                                1000_at
459.489 459 489
                                1000_at
                  0.072944350
408.545 408 545
                  0.342924723
                               1000_at
> head(sequenceMask)
               affy id match
       х
           у
    291 575 1195_s_at
340
                           0
788 617 349
               1000_at
                            0
1134
      6 591
               1024_at
                           0
                           0
1624 15 427 1015_s_at
2305 26 515
                           0
               1270_at
2563 31 541 1015_s_at
                            0
> rownames(sequenceMask) <- paste(sequenceMask$x,sequenceMask$y,sep="_")</pre>
> rownames(exmask$probes)<- paste(exmask$probes$x,exmask$probes$y,sep="_")</pre>
> cutoff=0.029
> table((exmask$probes[rownames(sequenceMask),"quality.score"]>cutoff)+0,
+
        sequenceMask$match)
      0
          1
    93 151
 0
```

1 79 859

Here, 952 probes are concordant between masks, whereas 79 probes with known SNP are not detected by expression-based mask.

With overlapExprExtMasks() expression mask may be compared with any other designation of BAD probes along a range of cutoffs. Type 1 (fraction of external mask not BAD probes, identified as BAD by expression mask) and type 2 (fraction of external mask BAD probes, not detected as BAD in expression mask) errors are plotted with their binomial confidence intervals.

```
> head(exmask$probes[,1:3])
```

	x	у	quality.score					
399_559	399	559	0.139886115					
544_185	544	185	0.485398831					
530_505	530	505	0.704649407					
617_349	617	349	0.008627547					
459_489	459	489	0.072944350					
408_545	408	545	0.342924723					
<pre>&gt; head(sequenceMask[,c(1,2,4)])</pre>								
	v	v	match					
291 575			match 0					
291_575 617_349	291	575						
617_349	291	575 349	0					
617_349	291 617 6	575 349 591	0 0					
617_349 6_591	291 617 6 15	575 349 591 427	0 0 0 0					
617_349 6_591 15_427	291 617 6 15 26	575 349 591 427 515	0 0 0 0					

The function overlapExprExtMasks() applied on our example data, produces a plot like shown in Figure 1.

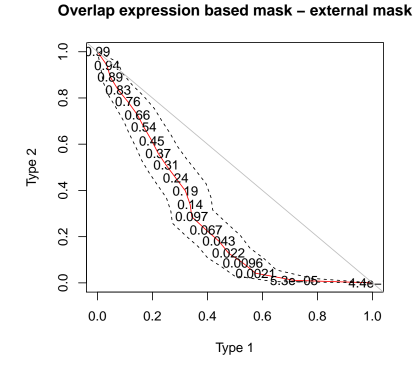


Figure 1: Overlap of expression-based mask and external mask (red line), with binomial confidence intervals (dashed lines). Several cutoff levels are marked on the plot.

The cutoff depends on the goal of masking. SFPs detection needs tighter control of type 2 than type 1 error - low cutoff values. Removal of differential expression bias needs higher cutoff. The distribution of quality scores for probes declared in external mask as BAD and not BAD may be compared with Wilcoxon rank sum test and Kolomogornov-Smirnow test for several cutoff levels. When those distributions do not differ (high p-values on 2), cutoff is too high.

```
> overlapTests <- overlapExprExtMasks(exmask$probes[,1:3],verbose=FALSE,
+ sequenceMask[,c(1,2,4)],
+ wilcox.ks=T,sample=100)</pre>
```

When no information about sequence divergence between the groups is available, a cutoff might be suggested by the distribution of quality scores for the mask. On Figure 3, an excess of probes with low quality scores is obvious. Assuming a uniform distribution of quality scores for probes not different between groups, a cutoff should be set so that the remaining distribution is uniform - like.

## 3 Removal of BAD probes and estimation of expression values

#### 3.1 Removal of BAD probes

To remove probes from affybatch, use prepareMaskedAffybatch(). It produces an affybatch object (without the masked probes) and an according CDF environment. When one probe appears in several probe sets (as is a case for some custom annotations) and it is detected as BAD in one probe set only, it will be nevertheless removed from all probe sets it appears. With cdftable argument, prepareMaskedAffybatch() allows also to remove any list of probes.

```
> affyBatchAfterMasking <-</pre>
    prepareMaskedAffybatch(affy=newAffyBatch,exmask=exmask$probes,
+
                           cutoff=quantile(exmask[[1]]$quality.score,0.22))
Inferring mismatch positions from perfect match
Renaming columns: probeset x y as probeset px py
Regular run: more than NA probe per grouping
Made list.
Making env
finished making CDF!
> newAffyBatch
AffyBatch object
size of arrays=640x640 features (22 kb)
cdf=newCdf (100 affyids)
number of samples=20
number of genes=100
annotation=hgu95av2
notes=
> affyBatchAfterMasking
```

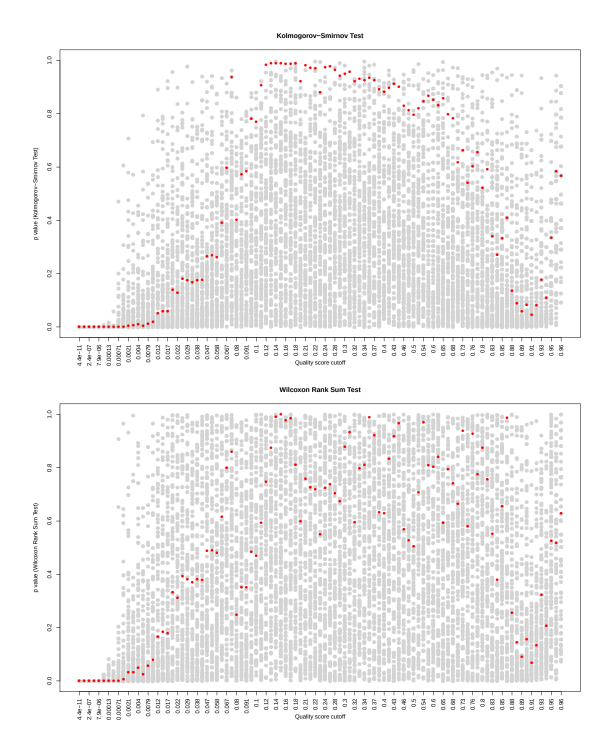


Figure 2: Kolmogorow-Smirnov-Test (upper panel) and Wilcoxon-Rank-Sum-Test (lower panel) comparing distributions of quality scores of BAD and not BAD probes. For low cutoffs, two distributions are different (red - actual tests; gray - tests on bootstraped data, to estimate variance).

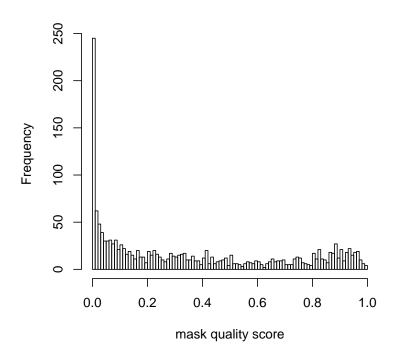


Figure 3: Probes' quality scores.

```
$newAffyBatch
AffyBatch object
size of arrays=640x640 features (22 kb)
cdf=new_cdf (100 affyids)
number of samples=20
number of genes=100
annotation=hgu95av2
notes=
```

# \$newCdf <environment: 0x7cf3b18>

Note changed number of genes (probe sets) in masked affybatch.

#### 3.2 Expression estimation with rma

For the new affybatch expression values might be estimated as usual. New affybatch object has a masked cdf declared, instead of a standard one.

```
> cdfName(affyBatchAfterMasking[[1]])
```

```
[1] "new_cdf"
```

Therefore it is important to save new cdf for further use and make it available in the workspace (with a name matching cdfName slot of masked affybatch) whenever a function to estimate expression is called.

> new\_cdf=affyBatchAfterMasking[[2]] > head(exprs(rma(affyBatchAfterMasking[[1]]))) Background correcting Normalizing Calculating Expression a\_c1a.CEL a\_c1b.CEL a\_c1c.CEL a\_c1d.CEL a\_c1e.CEL a\_c1f.CEL a\_c2a.CEL 8.519381 8.239144 8.698188 8.426627 8.213712 8.631725 8.702705 1000\_at 1001\_at 4.926438 4.926092 4.976691 5.100733 5.071768 4.901918 4.899042 1002\_f\_at 3.994510 3.957382 4.015966 4.146419 4.027846 4.028062 3.938472 1003\_s\_at 6.156051 6.242018 6.284055 6.408501 6.484892 6.239529 5.993723 5.491737 5.487617 5.606666 5.861494 5.653364 5.564533 5.545361 1004\_at 6.606239 6.125861 6.718311 6.343611 6.174270 6.464771 5.276121 1005\_at a\_c2b.CEL a\_c2c.CEL a\_c2d.CEL a\_h1a.CEL a\_h1b.CEL a\_h1c.CEL a\_h1d.CEL 1000\_at 8.147788 8.531981 8.406286 8.617236 8.265444 8.821593 8.607421 5.082391 5.025381 5.166789 4.935324 4.966834 4.984837 5.063387 1001\_at 1002\_f\_at 4.085803 3.981629 3.984972 4.155437 4.396664 4.026082 4.319549 1003\_s\_at 6.178582 6.249877 6.329526 6.465071 6.704101 6.283548 6.401511 1004\_at 5.559185 5.647424 5.666993 5.935005 5.918048 5.894887 5.806115 5.237686 5.070708 5.382761 4.831896 5.298492 5.073494 1005\_at 4.665798 a\_h1e.CEL a\_h1f.CEL a\_h2a.CEL a\_h2b.CEL a\_h2c.CEL a\_h2d.CEL 1000\_at 8.344701 8.542945 8.607421 8.156907 8.882432 8.567411 1001\_at 5.308487 4.971850 4.911928 4.815596 4.822864 5.187153 1002\_f\_at 4.511715 4.240136 3.977734 4.016085 3.964241 4.185346 1003\_s\_at 6.747904 6.560276 6.206139 6.275659 6.210961 6.311143 6.409215 6.004868 5.643354 5.551346 5.601732 5.835371 1004\_at 1005\_at 5.680619 5.317797 5.036664 5.414302 4.848811 4.793108

#### 3.3 Expression estimation with gcrma

Normalization and expression estimates with gcrma() require probe affinities. Those should be computed with standard cdf. Then both affybatch object and probe affinities object should be filtered with the mask.

```
> library(gcrma)
> library(hgu95av2probe)
> affy.affinities=compute.affinities("hgu95av2",verbose=FALSE)
> affy.affinities.m=prepareMaskedAffybatch(affy=affy.affinities,
+ exmask=exmask$probes,cdfName="MaskedAffinitesCdf")
Inferring mismatch positions from perfect match
Renaming columns: probeset x y as probeset px py
Regular run: more than NA probe per grouping
Made list.
Making env
finished making CDF!
```

To calculate expression values, just use both masked objects (make sure that a cdf with name matching cdf slot of masked affyBatch is accessible!).

> head(exprs(gcrma(affyBatchAfterMasking[[1]],affy.affinities.m)))

Adjusting for optical effect.....Done. Normalizing

Calculating Expression									
	a_c1a.CEL	a_c1b.CEL	a_c1c.CEL	a_c1d.CEL	a_c1e.CEL	a_c1f.CEL	a_c2a.CEL		
1000_at	8.634916	8.358381	8.796348	8.549656	8.423818	8.733339	8.771972		
1001_at	5.846012	5.845099	5.895825	5.982687	5.989225	5.821090	5.813759		
1002_f_at	5.249379	5.206399	5.273889	5.380924	5.271458	5.265842	5.193669		
1003_s_at	6.663292	6.720045	6.750878	6.852030	6.921729	6.700775	6.548312		
1004_at	6.181949	6.167532	6.273633	6.458531	6.284642	6.262856	6.222565		
1005_at	6.984931	6.588212	7.071891	6.788985	6.807228	6.902829	6.077073		
	a_c2b.CEL	a_c2c.CEL	a_c2d.CEL	a_h1a.CEL	a_h1b.CEL	a_h1c.CEL	a_h1d.CEL		
1000_at	8.394478	8.649110	8.534103	8.713059	8.384302	8.905719	8.721932		
1001_at	5.982168	5.928388	6.013410	5.856875	5.855905	5.881186	5.956344		
1002_f_at	5.323003	5.253987	5.225674	5.386190	5.636114	5.265202	5.553035		
1003_s_at	6.667764	6.724276	6.773942	6.892581	7.056563	6.739612	6.906102		
1004_at	6.213381	6.283493	6.307053	6.507925	6.546897	6.487002	6.475916		
1005_at	6.046712	5.928464	6.171188	5.813974	6.096748	5.925406	5.679128		
	a_h1e.CEL	$a_h1f.CEL$	a_h2a.CEL	a_h2b.CEL	$a_h2c.CEL$	a_h2d.CEL			
1000_at	8.467408	8.657795	8.721932	8.343586	8.929597	8.677108			
1001_at	6.232725	5.881665	5.825770	5.759558	5.768511	6.070740			
1002_f_at	5.603963	5.516864	5.233100	5.253987	5.213435	5.418177			
1003_s_at	7.192853	6.996384	6.675245	6.745539	6.720009	6.769744			
1004_at	6.895957	6.569566	6.299328	6.119514	6.234432	6.440966			
1005_at	6.509646	6.098351	5.886391	6.187611	5.805249	5.720689			

#### 3.4 Exon and gene arrays

Exon and gene arrays require setting useExpr=FALSE, as those arrays do not contain MM probes and mas5calls() - based definition of expressed genes is not applicable. As computing time increases with number of probe sets, we recommend to use a subset of probe sets (specified with exprlist) for exon arrays.

#### References

- Michael Dannemann, Michael Lachmann, and Anna Lorenc. 'maskBAD' a package to detect and remove affymetrix probes with binding affinity differences. *BMC Bioinformatics*, 13(1):56+, April 2012.
- [2] Michael Dannemann, Anna Lorenc, Ines Hellmann, Philipp Khaitovich, and Michael Lachmann. The effects of probe binding affinity differences on gene expression measurements and how to deal with them. *Bioinformatics (Oxford, England)*, 25(21):2772–2779, November 2009.
- [3] Philipp Khaitovich, Bjoern Muetzel, Xinwei She, Michael Lachmann, Ines Hellmann, Janko Dietzsch, Stephan Steigele, Hong-Hai H. Do, Gunter Weiss, Wolfgang Enard, Florian Heissig, Thomas Arendt, Kay Nieselt-Struwe, Evan E. Eichler, and Svante Pääbo. Regional patterns of gene expression in human and chimpanzee brains. *Genome Research*, 14(8):1462–1473, August 2004.