Introduction to the xps Package: Comparison to Affymetrix Power Tools (APT)

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1 Introduction

The aim of this document is to compare the results obtained with package xps to the results obtained with Affymetrix Power Tools (APT version apt-1.10.2). For this purpose the Exon Array Data Set "Tissue Mixture" will be used, which can be downloaded from the Affymetrix web-site for expression arrays "Human Genome U133 Plus 2.0", "Human Gene 1.0 ST" and "Human Exon 1.0 ST", respectively. First, APT and xps will be compared for each array type seperately, then the results obtained with the different arrays will be compared to each other for both APT and xps. To speed up analysis only two human tissues will be used, namely breast and prostate.
For analyzing expression arrays, including gene and exon arrays, APT contains the command-line program `apt-probeset-summarize`, which supports background correction (RMA, MAS5), normalization (linear scaling, quantile, sketch), and summarization (RMA, MAS5, PLIER) methods.

**Note:** In order to keep this document short, only example code will be presented here, the full source code is available in files "script4xps2apt.R" and "script4bestmatch.R" located in directory "xps/examples".

## 2 Comparison of results for the Human Genome U133 Plus 2.0 Array

### 2.1 Comparison of different implementations of RMA summarization

When doing RMA summarization with `apt-probeset-summarize`, sketch-quantile normalization (using only a subset of data) is applied by default in order to decrease memory consumption. Furthermore, the Affymetrix GUI application "Expression Console" can only use sketch-quantile normalization. For this reason let us first compare RMA using sketch-quantile vs quantile normalization.

The following command will compute RMA using sketch-quantile normalization:

```
apt-probeset-summarize -a rma-bg,quant-norm.sketch=-1.usepm=true.bioc=true, pm-only,med-polish.expon=true -d HG-U133_Plus_2.cdf *.CEL
```

Note that all CEL-files and the corresponding CDF-file must be located in the current directory. The output can be directly imported into R:

```
> apt.rma.sk <- read.delim("rma-bg.quant-norm.pm-only.med-polish.summary.txt", row.names=1, comment.char="\r")
```

To compute RMA using full quantile normalization the following command is used:

```
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, pm-only,med-polish.expon=true -d HG-U133_Plus_2.cdf *.CEL
```

Again, the output can be directly imported into R:

```
> apt.rma <- read.delim("rma-bg.quant-norm.pm-only.med-polish.summary.txt", row.names=1, comment.char="\r")
```

Please note that in both cases the name of the output file is the same.

To compare the results, we will use "MvA-plots" in almost all cases. Thus we can plot:

```
> plot(log2(apt.rma[,1] * apt.rma.sk[,1])/2, log2(apt.rma.sk[,1]/apt.rma[,1]), main="APT: RMA vs RMA-sketch", xlab="A = Log2(SketchRMA*RMA)", ylab="M = Log2(SketchRMA/RMA)",log="",ylim=c(-0.1,0.1))
```

Since we use only the triplicate CEL-files for breast and prostate, respectively, column one of the data.frames `apt.rma.sk` and `apt.rma` show the normalized expression values of tissue "breast_A".

The resulting Figure 1 shows that using sketch-quantile normalization gives a slightly different result, especially at low expression values. Thus, for all following comparisons we will use the full quantile normalization.

**Note:** One advantage of package `xps` compared to APT is, that you can do RMA with full quantile normalization on computers with only 1GB RAM, even for exon arrays.
Now let us compare APT and xps. For this purpose we compute RMA as described in vignette "xps.pdf", and extract data.frame xps.rma from data.rma:

```r
> data.rma <- rma(data.u133p2, "MixU133P2RMA", background="pmonly", normalize=TRUE)
> xps.rma <- validData(data.rma)
```

Now we can compare the results obtained with APT and xps for tissue "breast_A" using an "MvA-plot":

```r
> plot(log2(apt.rma[,1] * xps.rma[,1])/2, log2(xps.rma[,1]/apt.rma[,1]), main="RMA: XPS vs APT", xlab="A = Log2(SketchRMA*RMA)", ylab="M = Log2(XPS/APT)"
```

As shown in Figure 2, the results obtained with APT and xps are identical.
For the sake of completeness, let us compare both results to the results obtained with package affy. For this purpose we compute:

```
> affy.rma <- justRMA()
> affy.rma <- 2^exprs(affy.rma)
```

and plot:

```
> tmp <- cbind(xps.rma[,1],affy.rma[,1],apt.rma[,1])
> colnames(tmp) <- c("xps.rma","affy.rma","apt.rma")
> pairs(log2(tmp), labels=colnames(tmp))
```

As shown in Figure 2, the results are identical in all three cases. For example, the differences between xps and APT or affy are less than 4e-6.
2.2 Comparison of different implementations of MAS 5.0 summarization

The command-line to compute MAS5 summarization using APT is:

```
apt-probeset-summarize -a mas5-bg,pm-mm,mas5-signal -d HG-U133_Plus_2.cdf *.CEL
```

The output is imported into R and must be scaled to sc=500 since APT does not allow for signal level normalization across the chip (as mentioned in APT FAQ):

```
> apt.mas5 <- read.delim("mas5-bg.pm-mm.mas5-signal.summary.txt", row.names=1, comment.char="", skip=52)
> apt.mas5 <- apply(apt.mas5, 2, function(x){x*(500/mean(x, trim=0.02))})
```

Now we compute MAS5 with xps as described in vignette "xps.pdf":

```
> data.mas5 <- mas5(data.u133p2,"MixU133P2MAS5All", normalize=TRUE, sc=500, update=TRUE)
> xps.mas5 <- validData(data.mas5)
```

Finally, we compute MAS5 using package affy:

```
> affy <- ReadAffy()
> affy.mas5 <- mas5(affy, normalize=TRUE, sc=500)
> affy.mas5 <- exprs(affy.mas5)
```

Using "MvA-plots" to compare the MAS5 expression levels for sample "Breast_A" we obtain the results shown in Figure 4.

The upper row shows how the implementation of the MAS5 algorithm in packages xps and affy compare to APT. Both packages show clear differences to APT, especially at low expression levels. However, as APT FAQ mentions, "a new implementation of the MAS5 methods was necessary to get this method to work in the APT architecture". Thus, the middle row shows how packages xps and affy compare to the MAS5 results obtained with the Affymetrix "Expression Console", which has implemented MAS5 identical to GCOS. There are still differences seen in the expression levels, with the xps implementation of MAS5 being slightly more close to the GCOS implementation of MAS5. Finally, the lower row shows the differences between APT and GCOS, and between xps and affy, respectively.
2.3 Comparison of different implementations of MAS 5.0 detection calls

Now let us compute MAS5 detection calls. The command-line for APT is:

```
apt-probeset-summarize -a pm-mm,mas5-detect.calls=1.pairs=1 -d HG-U133_Plus_2.cdf -x 10 *.CEL
```

Using package xps MAS5 detection calls are computed as follows:

```
> call.mas5 <- mas5.call(data.u133p2,"MixU133P2Call")
> xps.pval <- validData(call.mas5)
```

while package affy uses the following commands:

```
> affy <- ReadAffy()
> affy.dc5 <- mas5calls(affy)
> affy.pval <- assayData(affy.dc5)[["se.exprs"]]
```
Figure 5 compares the detection p-values obtained with packages \texttt{xps} and \texttt{affy}, as well as Expression Console and APT.

![Comparison of p-values obtained with xps, affy, Expression Console and APT.](image)

While Expression Console and APT result in identical p-values, both \texttt{xps} and \texttt{affy} differ from APT. However, as shown in Figure 6, the detection call p-values obtained with \texttt{xps} and \texttt{affy} are identical.
3 Comparison of results for the Human Gene 1.0 ST Array

3.1 Comparison of the RMA summarization implementations

While the results for expression arrays obtained with xps and APT, respectively, can be directly compared, since the probesets used are identical, comparison of xps and APT with gene/exon arrays requires the definition of common probesets first.

Let us first compute RMA using package xps:

```r
> data.rma.tc <- rma(data.genome,"MixHuGeneRMAcore_tc", background="antigenomic", 
+ normalise=TRUE, option="transcript", exonlevel="core")
> xps.rma.tc <- validData(data.rma.tc)
```

Now we can create a file specifying the probesets to summarize for APT:

```r
> tmp <- as.data.frame(rownames(xps.rma.tc))
> colnames(tmp) <- "probeset_id"
> write.table(tmp, "transcriptList.txt", quote=FALSE, row.names=FALSE)
```

This transcriptList contains now the probesets used by xps and can be added as option when computing RMA using APT:

```bash
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, \
pm-only,med-polish.expon=true \n-p HuGene-1_0-st-v1.r3.pgf -c HuGene-1_0-st-v1.r3.clf \n-s transcriptList.txt *.CEL
```

Figure 7 shows the resulting "MvA-plot" for tissue "breast_A".
As Figure 7 shows, there is clearly a difference of few percent in the expression levels obtained when using xps or APT, respectively, including a non-linearity at low expression levels.

In order to understand this difference let us first compare the "median-polish" summarization step (w/o background correction and quantile normalization).

In package xps this is done as follows:

```r
> expr.mp.tc <- express(data.genome, "MixHuGeneMedPolcore_tc", summarize.method="medianpolish",
+                        summarize.select="pmonly", summarize.option="transcript", summarize.logbase="log2",
+                        summarize.params=c(10, 0.01, 1.0), exonlevel="core")
> xps.mp.tc <- validData(expr.mp.tc)
```

while APT uses the following command:

```
apt-probeset-summarize -a pm-only,med-polish.expon=true \ 
-p HuGene-1_0-st-v1.r3.pgf -c HuGene-1_0-st-v1.r3.clf \ 
-s transcriptList.txt *.CEL
```

The resulting Figure 8 shows that the results obtained with xps and APT are identical (the difference being less than 4e-6, see scale of y-axis).
Since the results of median-polish are identical for xps and APT, the differences seen in Figure 7 are due to background correction and/or quantile normalization, respectively. Package xps uses only those probes to compute background and quantile normalization, which are also used for summarization, as defined by parameter `exonlevel`. The reason for this is that RMA estimates the background from the PM probe intensities only, as described in (Irizarry et al., 2003). Analogously, quantile normalization (Bolstad et al., 2003) is applied only to the probes which are used for summarization.

It seems that APT uses (almost) all probes on the array for background correction and quantile normalization. Since package xps has an (unsupported) option to use (almost) all probes (which include internal control probes and background probes), there is a possibility to use all probes by setting parameter `exonlevel` individually for background correction, quantile normalization and median-polish summarization, respectively. This is done as follows:

```r
> data.rma.tc.bq99 <- rma(data.genome, "HuGeneMixRMAbgqu99mp9", background="antigenomic", 
  normalize=TRUE, exonlevel=c(992316,992316,9216))
> xps.rma.tc.bq99 <- validData(data.rma.tc.bq99)
```

Here, we have set `exonlevel="core+affx+unmapped+exon+intron+antigenomic"` for background correction and quantile normalization, respectively, and `exonlevel="core"` for median-polish summarization (see ?exonLevel). Comparing the results of this setting to APT, we obtain the following "MvA-plot" for tissue "breast_A", see Figure 9.
As Figure 9 shows, the expression levels obtained when using xps or APT, respectively, are now identical, indicating that APT does indeed use all probes on the array for background correction and quantile normalization (excluding only the probes hybridizing to the control oligo).

### 3.2 Comparison of the DABG call implementations

Now let us compute the detected above background (DABG) calls. The command-line for APT is:

```
apt-probeset-summarize -a dabg \
ap -p HuGene-1_0-st-v1.r3.pgf -c HuGene-1_0-st-v1.r3.clf \
b HuGene-1_0-st-v1.r3.bgp -s probesetList.txt -x 8 *.CEL
```

Using package xps DABG calls are computed as follows:

```
> call.dabg.tc <- dabg.call(data.genome, "HuGeneMixDABGcore_tc", 
>   option="transcript", exonlevel="core")
> xps.pval.tc <- validData(call.dabg.tc)
```

Figure 10 shows the difference in the detection p-values obtained with xps and APT, respectively. The detection p-values are identical (the minor apparent difference only due to different numbers of digits exported).
3.3 Comparison of results when using HuGene as exon array

With release r4 of the library files Affymetrix has converted the HuGene-1.0-st-v1 array into an exon array. While the probes for "HuGene-1.0-st-v1.r3.pgf" were grouped according to the transcript_cluster_ids, the probes for "HuGene-1.0-st-v1.r4.pgf" were grouped according to the probeset_ids of the new probeset annotation file.

Thus in order to compare xps and APT at the transcript level it is now necessary to create the transcript probesets for APT using function metaProbesets():

```r
> writeLines(rownames(xps.rma.tc), "core.txt")
> metaProbesets(scheme.genome, "core.txt", "coreList.mps", "core")
```

Now we need to use this 'coreList.mps' file to compute RMA and DABG, respectively, at the transcript level using APT, e.g. we compute RMA as follows:

```r
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, \ pm-only,med-polish.expon=true \ -p HuGene-1.0-st-v1.r4.pgf -c HuGene-1.0-st-v1.r4.clf \ -m coreList.mps *.CEL
```

The results for RMA and DABG are identical to the results obtained with release r3 of the library files shown above.

In addition to the transcript level release r4 allows now to compute both RMA and DABG on the probeset level as well, analogously to the exon arrays. For this purpose we compute first RMA at the probeset level using package xps:

```r
> data.rma.ps <- rma(data.genome, "MixHuGeneRMAcore_ps", background="antigenomic", + normalize=TRUE, option="probeset", exonlevel="core")
> xps.rma.ps <- validData(data.rma.ps)
```
Then we create a file specifying the probesets to summarize for APT:

```r
> tmp <- as.data.frame(rownames(xps.rma.ps))
> colnames(tmp) <- "probeset_id"
> write.table(tmp, "probesetList.txt", quote=FALSE, row.names=FALSE)
```

Finally we use this probesetList to compute RMA using APT:

```r
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, pm-only,med-polish.expon=true \ -p HuGene-1_0-st-v1.r4.pgf -c HuGene-1_0-st-v1.r4.clf -s probesetList.txt *.CEL
```

The resulting figure [11] shows once again a difference of few percent in the expression levels obtained when using xps or APT.

![RMA PS: XPS vs APT](image)

Figure 11: RMA for probesets computed with xps and apt, respectively.

However, when using the same probes for background correction and quantile normalization as APT:

```r
> data.rma.ps.bq99 <- rma(data.genome,"HuGeneMixRMAbgqu99mp9_ps", background="antigenomic", normalize=TRUE, option="probeset", exonlevel=c(992316,992316,9216))
> xps.rma.ps.bq99 <- validData(data.rma.ps.bq99)
```

the results are once again identical to the results obtained with APT, as shown in Figure [12]
4 Comparison of results for the Human Exon 1.0 ST Array

4.1 Comparison of RMA summarization at the probeset level

As already mentioned, comparison of xps and APT with exon arrays requires the definition of common probesets first.

Thus we compute first RMA at the probeset level using package xps:

```r
> data.rma.ps <- rma(data.exon, "MixHuExonRMAcore_ps", background="antigenomic", 
+ normalize=TRUE, option="probeset", exonlevel="core")
> xps.rma.ps <- validData(data.rma.ps)
```

Then we can create a file specifying the probesets to summarize for APT:

```r
> tmp <- as.data.frame(rownames(xps.rma.ps))
> colnames(tmp) <- "probeset_id"
> write.table(tmp, "corePSList.txt", quote=FALSE, row.names=FALSE)
```

Finally we use this probesetList to compute RMA using APT:

```
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, \ 
  pm-only,med-polish.expon=true \ 
  -p HuEx-1_0-st-v2.r2.pgf -c HuEx-1_0-st-v2.r2.clf \ 
  -b HuEx-1_0-st-v2.r2.antigenomic.bgp -s corePSList.txt *.CEL
```
Figure 13: RMA computed with xps and apt, respectively.

As the resulting "MvA-plot" for tissue "breast_A", shown in Figure 13 reveals, there is quite some difference for the probeset levels computed with xps or APT, respectively. Thus let us compare the "median-polish" summarization step only, as done before for the gene array.

In package xps this is done as follows:

```
> expr.mp.ps <- express(data.exon, "HuExonMedPolcorePS", summarize.method="medianpolish",+
    summarize.select="pmonly", summarize.option="probeset", summarize.logbase="log2",+
    summarize.params=c(10, 0.01, 1.0), exonlevel="core")
> xps.mp.ps <- validData(expr.mp.ps)
```

while APT uses the following command:

```
apt-probeset-summarize -a pm-only,med-polish.expon=true \
-p HuEx-1_0-st-v2.r2.pgf -c HuEx-1_0-st-v2.r2.clf \
-s corePSList.txt *.CEL
```

The resulting Figure[14] shows that the results obtained with xps and APT are identical (the difference being less than 6e-6, see scale of y-axis).
As for gene arrays, the differences seen in Figure 13 are due to background correction and quantile normalization, respectively. This time we set parameter `exonlevel` individually to `exonlevel="all+genomic+antigenomic+intron+exon+unmapped"` for background correction and quantile normalization, respectively, and `exonlevel="core"` for median-polish summarization:

```r
> data.rma.ps.bq103 <- rma(data.exon, "HuExonMixRMAbgqu103mp9_ps", background="antigenomic", + normalize=TRUE, option="probeset", exonlevel=c(1032124,1032124,9216))
> xps.rma.ps.bq103 <- validData(data.rma.ps.bq103)
```

Comparing the results of this setting to APT, we obtain the following "MvA-plot" for tissue "breast_A" shown in Figure 15:

![MedPol PS: XPS vs APT](image)

**Figure 14:** Median-polish only, computed with xps and apt, respectively.

As for gene arrays, the differences seen in Figure 13 are due to background correction and quantile normalization, respectively. This time we set parameter `exonlevel` individually to `exonlevel="all+genomic+antigenomic+intron+exon+unmapped"` for background correction and quantile normalization, respectively, and `exonlevel="core"` for median-polish summarization:

```r
> data.rma.ps.bq103 <- rma(data.exon, "HuExonMixRMAbgqu103mp9_ps", background="antigenomic", + normalize=TRUE, option="probeset", exonlevel=c(1032124,1032124,9216))
> xps.rma.ps.bq103 <- validData(data.rma.ps.bq103)
```

Comparing the results of this setting to APT, we obtain the following "MvA-plot" for tissue "breast_A" shown in Figure 15.
As Figure 15 shows, the differences in the expression levels obtained when using xps or APT, respectively, are dramatically reduced. At high expression levels there is no difference, while the difference at low expression levels is reduced to at most 20%. However, in contrast to the HuGene array the difference is still visible. Thus in this case it is not clear to me which probes Affymetrix is using for background correction and quantile normalization.

4.2 Comparison of RMA summarization at the transcript level

Comparing xps and APT with exon arrays at the gene level requires the definition of common probesets for each transcript, i.e. a file containing meta probeset definitions.

First we compute RMA at the transcript level using package xps:

```r
> data.rma.tc <- rma(data.exon, "MixHuExonRMAcore_tc", background="antigenomic", 
+ normalize=TRUE, option="transcript", exonlevel="core")
> xps.rma.tc <- validData(data.rma.tc)
```

Then we can create a meta probeset file using function `metaProbesets()`:

```r
> writeLines(rownames(xps.rma.tc), "core.txt")
> metaProbesets(scheme.exon, "core.txt", "coreList.mps", "core")
```

Now we use this 'coreList.mps' file to compute RMA using APT:

```bash
apt-probeset-summarize -a rma-bg,quant-norm.sketch=0.usepm=true.bioc=true, \
pm-only,med-polish.expon=true \n-p HuEx-1_0-st-v2.r2.pgf -c HuEx-1_0-st-v2.r2.clf \n-m coreList.mps *.CEL
```
As expected, the resulting "Mva-plot" for tissue "breast_A", shown in Figure 16, is similar to Figure 13. Again, we compare the "median-polish" summarization step only.

In package xps this is done as follows:

```r
> expr.mp.tc <- express(data.exon, "HuExonMedPolcore", summarize.method="medianpolish",
+ summarize.select="pmonly", summarize.option="transcript", summarize.logbase="log2",
+ summarize.params=c(10, 0.01, 1.0), exonlevel="core")
> xps.mp.tc <- validData(expr.mp.tc)
```

while APT uses the following command:

```bash
apt-probeset-summarize -a pm-only,med-polish.expon=true \
-p HuEx-1_0-st-v2.r2.pgf -c HuEx-1_0-st-v2.r2.clf \
-m coreList.mps *.CEL
```

As for the probesets, the resulting Figure 17 shows that the results obtained with xps and APT are identical (the difference being less than 6e-6, see scale of y-axis).
When we set parameter \texttt{exonlevel} individually to \texttt{exonlevel="all+genomic+antigenomic+intron+exon+unmapped"} for background correction and quantile normalization, respectively, and \texttt{exonlevel="core"} for median-polish summarization:

\begin{verbatim}
> data.rma.tc.bq103 <- rma(data.exon, "HuExonMixRMAbgqu103mp9_tc", background="antigenomic", normalize=TRUE, option="transcript", exonlevel=c(1032124,1032124,9216))
> xps.rma.tc.bq103 <- validData(data.rma.tc.bq103)
\end{verbatim}

and compare the results to APT, we obtain the "MvA-plot" for tissue "breast_A" shown in Figure 18.
As Figure 18 shows, the differences in the expression levels obtained when using xps or APT, respectively, are also dramatically reduced. At high expression levels there is almost no difference, while the difference at low expression levels is reduced to less than 20%.

4.3 Comparison of the DABG call implementations

Now let us compare the detected above background (DABG) calls at the probeset level.

The command-line for APT is:

```
apt-probeset-summarize -a dabg \
   -p HuEx-1_0-st-v2.r2.pgf -c HuEx-1_0-st-v2.r2.clf \
   -b HuEx-1_0-st-v2.r2.antigenomic.bg� -s corePSList.txt -x 12 *.CEL
```

while package xps computes DABG calls as follows:

```
> call.dabg.ps <- dabg.call(data.exon, "HuExonDABGcorePS", option="probeset", exonlevel="core")
> xps.pval.ps <- validData(call.dabg.ps)
```

Figure 19 shows the difference in the detection p-values obtained with xps and APT, respectively. The detection p-values are identical (the minor apparent difference only due to different numbers of digits exported).
Figure 19: Comparison of p-values obtained with xps and APT.

Note: Another advantage of package xps compared to APT is, that you can compute DABG calls not only at the probeset level but also at the transcript level. This is simply done as follows:

```r
> call.dabg.tc <- dabg.call(data.exon, "HuExonDABGcoreTC", option="transcript", exonlevel="core")
> xps.pval.tc <- pvalData(call.dabg.tc)
```

5 Comparison of Affymetrix gene expression arrays

In this section we want to compare the gene expression levels obtained with arrays HG-U133 Plus_2, HuGene-1_0-st-v1 and HuEx-1_0-st-v2, respectively, for tissues breast and prostate. For this purpose we need to select first a subset of transcripts common to all three arrays. Then we want to find differentially expressed transcripts and compare the results for the normalized expression levels obtained with xps or APT, respectively.

5.1 Comparison of RMA summarization for common transcript probesets

In order to create a subset of transcripts common to all three arrays, we import as a first step the "Array Comparison Spreadsheets", which can be downloaded from the Affymetrix web-site:

```r
> hx2hg <- read.delim("HuExVsHuGene_BestMatch.txt", row.names=3)
> up2hx <- read.delim("U133PlusVsHuEx_BestMatch.txt", row.names=3)
> up2hg <- read.delim("U133PlusVsHuGene_BestMatch.txt", row.names=3)
```

Since we want to compare the RMA normalized expression levels xps.urma (HG-U133_Plus_2), xps.grma (HuGene-1_0-st-v1) and xps.xrma (HuEx-1_0-st-v2), with xps.grma and xps.xrma using only the "core" transcript probesets, we obtain the common subset core, containing unique exon and genome transcripts identical to U133P2 as follows:
> hx2hg <- uniqueframe(hx2hg)
> up2hx <- uniqueframe(up2hx)
> up2hg <- uniqueframe(up2hg)
> u2x <- intersectframes(up2hx, up2hg)
> u2g <- intersectframes(up2hg, up2hx)
> tmp <- intersectrows(u2gx, xps.grma, 1, NULL)
> core <- intersectrows(tmp, xps.xrma, 2, NULL)

The resulting subset core consists of 11033 transcripts common to all arrays.

Note: The full source code and the functions uniqueframe, intersectframes and intersectrows are described in file "script4bestmatch.R" located in directory "examples".

Now we obtain the following subsets of normalized expression data:

> xpsu <- intersectrows(xps.urma, core, NULL, NULL)
> xpsg <- intersectrows(xps.grma, core, NULL, 1, -1)
> xpsx <- intersectrows(xps.xrma, core, NULL, 2, -1)
> aptg <- intersectrows(apt.grma, core, NULL, 1, -1)
> aptx <- intersectrows(apt.xrma, core, NULL, 2, -1)

where aptg and aptx are the corresponding data for APT (w/o aptu since it is identical to xpsu).

Figure 20: Scatter plot of mean expression summaries for breast.

A scatter plot of the averaged gene-level summaries for breast tissue is shown in Figure 20 for the data obtained with package xps. The corresponding figure for the data obtained with APT is almost identical. Furthermore, Figure 20 is very similar to the supplementary figure (page 10) published by Robinson & Speed in Robinson and Speed (2007), however, using the averaged gene-level summaries for one particular mixture, and Ensembl gene-centric probesets.
5.2 Differentially expressed transcripts in breast vs prostate

Now we consider the task of finding differentially expressed genes between the breast and prostate samples. Following Robinson & Speed (Robinson and Speed, 2007) we will use the moderated t-statistics from package limma to compute raw p-values, which are then adjusted by a Benjamini–Hochberg correction (the default setting).

For this purpose we create the design matrix:

```r
> tissue <- c(rep("Breast", 3), rep("Prostate", 3))
> design <- model.matrix(~factor(tissue))
> colnames(design) <- c("Breast", "BreastvsProstate")
```

which we use to compute the moderated t-statistics for each platform, e.g. for data xpsu:

```r
> tmp <- as.matrix(log2(xpsu))
> fit <- lmFit(tmp, design)
> fit <- eBayes(fit)
> xpsu.lm <- topTable(fit, coef=2, n=length(rownames(tmp)), adjust="BH")
```

Now we can first compare the fold-change values for the different platforms.

Figure 21: Scatter plot of fold-change values.

Figure 21 shows a scatter plot of the fold-change values for the data obtained with package xps. The corresponding figure for the data obtained with APT is (almost) identical.
Regarding the computed p-values, let us first compare the p-values for the HuGene-1_0-st-v1 data xpsg and aptg, obtained with xps or APT, respectively.

![Scatter plot of p-values obtained with xpsg, aptg, and bq99g.](image)

Figure 22: Scatter plot of p-values obtained with xpsg, aptg, and bq99g.

The upper rows of Figure 22 compare the p-values obtained with xpsg and aptg, respectively, whereby the right scatter-plot compares the lowest p-values only, i.e. the most relevant p-values. The lower rows of Figure 22 compare the p-values obtained with bq99g and aptg, respectively, where bq99g is identical to aptg, as shown in Figure 9.

Comparing the upper and lower scatter-plots reveals that the differences between xps and APT, seen in Figure 7 may not have a large effect when computing moderated t-statistics. The same is true when comparing the adjusted p-values. However, the differences between xpsg and aptg are only a few percent, as seen in Figure 7, thus it is not unexpected that the (adjusted) p-values are very similar.

In contrast, the differences between the expression levels obtained with array HuEx-1_0-st-v2, i.e. xpsx and aptx, respectively, are quite pronounced, as demonstrated earlier (see Figure 16). Thus it can be expected that these differences are also reflected in differences in p-values, when computing the moderated t-statistics for xpsx or aptx, respectively. As shown in Figure 23, this is indeed the case.
In this case, the upper rows of Figure 23 compare the BH-adjusted p-values obtained with `xpsx` and `aptx`, respectively, whereby the right scatter-plot compares once again the lowest p-values only, i.e. the most relevant p-values.

This time, the (adjusted) p-values computed for `xpsx` and `aptx`, respectively, show clearly larger variance. Of special interest is the upper right part of the figure, which compares only the the most relevant p-values. It is evident, that the relevant adjusted p-values computed for `xpsx` are smaller than the corresponding p-values computed for `aptx` (as the slope of an imaginary fitted line would show). This indicates that the expression levels `xpsx` obtained with package `xps` result in a better separation between tissues breast and prostate for the most relevant genes than the expression levels `aptx` obtained with APT.

As described earlier, we have also adjusted background correction and quantile normalization to resemble the APT expression levels (see Figure 18), resulting in subset `bq103x`. Comparing the adjusted p-values obtained with these expression levels (`bq103x`), with the p-values obtained with subset `aptx` results in the scatter plot shown in the lower part of Figure 23. This part of the figure looks similar to Figure 22 and shows that the adjusted p-values for `bq103x` and `aptx` are very similar, although there is still a difference of about 20% at low expression levels (Figure 18). However, in both cases the relevant p-values are larger than the p-values obtained with subset `xpsx`. In my opinion, this is a clear indication that only those probes should be used for background correction and quantile normalization, which are also used for median-polish summarization.

6 Summary

Comparison of packages `xps` and `affy` with APT for expression arrays shows that the different implementations of the RMA algorithm result in identical expression levels, as shown in Figure 3. (The negligible difference of less than 5e-6 is due to the fact that package `xps` does all computations with
double-precision (64bit), but saves the results as floats (32bit)).

Interestingly, all three implementations of the MAS5 algorithm (xps, affy, APT) differ from the original GCOS implementation, with xps probably being slightly more close to GCOS than affy (Figure 4), but I may be biased.

The implementations of MAS5 detection calls are identical for xps and affy, but differ slightly from the APT implementation which is identical to the GCOS implementation (Figure 5).

For the gene and exon arrays I have only compared xps and APT, which both use the new CLF-files and PGF-files as library files, and are able to use the separate probeset groups ‘core’, ‘extended’ and/or ‘full’, which are defined in the Affymetrix annotation files.

Although the xps and APT implementations of the RMA algorithm give identical results for expression arrays, the results for both gene arrays and exon arrays seem to differ, as shown in Figures 6 and 10 respectively. As we have demonstrated earlier, this difference is caused by using different probes for background correction and quantile normalization, respectively, since using identical probes (HuGene) or applying only median-polish summarization gives identical results (Figures 9, 8 and 17).

While package xps uses by default the identical ‘PM’ probes for all three steps of the RMA algorithm, e.g. for exonlevel="core" only probes belonging to the ‘core’ probesets defined in the Affymetrix annotation files are used, APT uses (almost) all probes on the respective array. Thus xps follows the implementation of RMA for expression arrays, while APT differs in the probes usage at the different steps.

In order to determine whether this difference in the probes usage may have an influence on subsequent expression analysis steps, I have applied moderated t-statistics from package limma to find differentially expressed genes between tissues breast and prostate. Computing raw and adjusted p-values for the gene array data results in almost identical p-values, indicating that the small difference seen in Figure 7 does not affect further processing. In contrast, computing raw and adjusted p-values for the exon array data reveals that the xps default usage of identical probes for RMA normalization results in lower (better) p-values in the relevant range to select genes of interest.

The implementation of the DABG call algorithm in xps is identical to APT, however for exon arrays, xps allows to apply DABG at both the probeset level as well as the transcript level.

In summary, package xps offers the following advantages compared to APT:

- full quantile normalization can be done on computers with 1GB RAM only.
- for RMA the same probes are used by default for background correction, quantile normalization and median-polish summarization for all three types of arrays.
- in addition, for gene and exon arrays it is possible to select probes to be used for background correction, quantile normalization and median-polish summarization, respectively, independently.
- MAS5 detection calls can be applied to all three types of arrays.
- DABG calls can be applied to all three types of arrays.
- for exon arrays DABG can be applied at the probeset level and the transcript level, respectively.
A Appendices

A.1 Some notes on PLIER

As an alternative to RMA, Affymetrix has developed a new algorithm, called PLIER: “The PLIER (Probe Logarithmic Error Intensity Estimate) method produces an improved signal by accounting for experimentally observed patterns in feature behavior and handling error at the appropriately at low and high signal values.” (according to the APT manual)

Using the HG-U133_Plus_2 data set, the command-line for APT is:

```
apt-probeset-summarize -a plier-mm -d HG-U133_Plus_2.cdf *.CEL
```

Now we import the results into R for visualization:

```r
> apt.plier <- read.delim("plier-mm.summary.txt", row.names=1, comment.char="", skip=52)
```

There are many methods to visualize the quality of the normalization step, often based on advanced statistical methods. However, often a simple visualization, e.g. a simple scatter-plot is already very helpful. Thus let us create a scatter-plot of the normalized data for the replicated breast tissue:

```r
> plot(apt.plier[,1], apt.plier[,2], main="APT: PLIER", xlab="Breast_A", ylab="Breast_B",log="xy")
```

![APT: PLIER](image)

Figure 24: Scatter-plot.

For comparison purposes, we create also "MvA-plots" for "Breast_A" vs "Breast_B" using either PLIER or RMA for normalization:

```r
> plot(log2(apt.plier[,1] * apt.plier[,2])/2, log2(apt.plier[,1]/apt.plier[,2]), main="APT: PLIER", xlab="A = Log2(BrA*BrB)", ylab="M = Log2(BrA/BrB)", log=""
> plot(log2(apt.rma[,1] * apt.rma[,2])/2, log2(apt.rma[,1]/apt.rma[,2]), main="APT: RMA", xlab="A = Log2(BrA*BrB)", ylab="M = Log2(BrA/BrB)", log="")
```
As Figure 25 shows, the differences between PLIER and RMA are dramatically. Please note that the following is only my personal subjective opinion:

As both Figure 24 and Figure 25 reveal, PLIER produces severe artifacts, even for replicate data, which are supposed to have very similar expression values. Probably other people will not agree with me, however for this reason I never use PLIER, and this is one of the reasons why I have decided not to implement PLIER in package xps.

References

