

# Package ‘pumadata’

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**Type** Package

**Title** Various data sets for use with the puma package

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**Depends** R (>= 3.2.0), affy (>= 1.46.0), Biobase (>= 2.5.5), puma  
, oligo(>= 1.32.0)

**Description** This is a simple data package including various data sets derived from the estrogen data for use with the puma (Propagating Uncertainty in Microarray Analysis) package.

**License** LGPL

**biocViews** ExperimentData, MicroarrayData, SNPData

**URL** <http://umber.sbs.man.ac.uk/resources/puma>

**NeedsCompilation** no

## R topics documented:

affybatch.estrogen . . . . .	2
eset_estrogen_comb . . . . .	3
eset_estrogen_mmgmos . . . . .	3
eset_estrogen_pmmgmos . . . . .	4
eset_estrogen_rma . . . . .	5
HTA_Location . . . . .	6
HTA_probes_transcripts . . . . .	6
HTA_transcript_name . . . . .	7
HTA_transcript_NO . . . . .	7
Human_Location . . . . .	8
Human_probes_transcripts . . . . .	8
Human_transcript_name . . . . .	9
Human_transcript_NO . . . . .	9
Mouse_Location . . . . .	10
Mouse_probes_transcripts . . . . .	10

Mouse_transcript_name . . . . .	11
Mouse_transcript_NO . . . . .	11
oligo.estrogen . . . . .	12
pumapca_estrogen . . . . .	12
Rat_Location . . . . .	13
Rat_probes_transcripts . . . . .	14
Rat_transcript_name . . . . .	14
Rat_transcript_NO . . . . .	15

<b>Index</b>	<b>16</b>
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<b>affybatch.estrogen</b>	<i>The data from the estrogen package as an AffyBatch object</i>
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## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code:

```
library(estrogen)
datadir <- file.path(.find.package("estrogen"), "extdata")
estrogenFilenames <- c("low10-1.cel", "low10-2.cel", "high10-1.cel", "high10-2.cel",
, "low48-1.cel", "low48-2.cel", "high48-1.cel", "high48-2.cel")
affybatch.estrogen <- ReadAffy(
filenames=estrogenFilenames
,celfile.path=datadir
)
pData(affybatch.estrogen) <- data.frame(
"estrogen"=c("absent", "absent", "present", "present"
,"absent", "absent", "present", "present")
,"time.h"=c("10", "10", "10", "10", "48", "48", "48", "48")
, row.names=rownames(pData(affybatch.estrogen)))
)
```

## Usage

```
data(affybatch.estrogen)
```

## Format

An **AffyBatch** object containing 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

---

eset\_estrogen\_comb     *The data from the **estrogen** package processed using the multi-mgMOS and PUMAcomb algorithms*

---

## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code (note this could take a long time to run):

```
data(eset_estrogen_mmgmos)
eset_estrogen_mmgmos_normd <- PUMAnormalize(eset_estrogen_mmgmos, "median")
eset_estrogen_comb <- PUMAcomb(eset_estrogen_mmgmos_normd)
```

## Usage

```
data(eset_estrogen_comb)
```

## Format

An [ExpressionSet](#) object containing the expression levels and standard errors from combining the replicates for each combination of levels of factors from 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

## See Also

[eset\\_estrogen\\_mmgmos](#)

## Examples

```
data(eset_estrogen_comb)
exprs(eset_estrogen_comb)[1:3,1:3]
assayDataElement(eset_estrogen_comb, "se.exprs")[1:3,1:3]
```

---

eset\_estrogen\_mmgmos     *The data from the **estrogen** package processed using the multi-mgMOS algorithm*

---

## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code (note this could take a long time to run):

```
data(oligo.estrogen)
eset_estrogen_mmgmos <- mmgmos(oligo.estrogen)
```

**Usage**

```
data(eset_estrogen_mmgmos)
```

**Format**

An `exprReslt` object containing expression levels and standard errors for 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

**See Also**

[oligo.estrogen](#) [eset\\_estrogen\\_rma](#)

**Examples**

```
data(eset_estrogen_mmgmos)
show(eset_estrogen_mmgmos)
exprs(eset_estrogen_mmgmos)[1:3,1:3]
assayDataElement(eset_estrogen_mmgmos, "se.exprs")[1:3,1:3]
```

**eset\_estrogen\_pmmmgmos**

*The data from the `estrogen` package processed using the multi-mgMOS  
use PM intensities only*

**Description**

This data is taken from the `estrogen` package. It was created to be used in the vignette for the `puma` pacakge. It can be produced using the following code (note this could take a long time to run):

```
data(oligo.estrogen)
eset_estrogen_pmmmgmos <- pmmmgmos(oligo.estrogen)
```

**Usage**

```
data(eset_estrogen_pmmmgmos)
```

**Format**

An `exprReslt` object containing expression levels and standard errors for 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

**See Also**

[oligo.estrogen](#) [eset\\_estrogen\\_rma](#)

## Examples

```
data(eset_estrogen_pmmmgmos)
show(eset_estrogen_pmmmgmos)
exprs(eset_estrogen_pmmmgmos)[1:3,1:3]
assayDataElement(eset_estrogen_pmmmgmos, "se.exprs")[1:3,1:3]
```

---

eset\_estrogen\_rma

*The data from the estrogen package processed using the RMA algorithm*

---

## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code:

```
data(oligo.estrogen)
eset_estrogen_mmgmos <- rma(oligo.estrogen)
```

## Usage

```
data(eset_estrogen_rma)
```

## Format

An **ExpressionSet** object taining expression levels for 8 HG\U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

## See Also

[oligo.estrogen](#) [eset\\_estrogen\\_mmgmos](#)

## Examples

```
data(eset_estrogen_rma)
show(eset_estrogen_rma)
exprs(eset_estrogen_rma)[1:3,1:3]
assayDataElement(eset_estrogen_rma, "se.exprs")[1:3,1:3]
```

HTA_Location	<i>The coordinates of probes and the mapped PM probes for hta2.0 chips</i>
--------------	--

**Description**

This data include the probes location for hta2.0 chips.

**Usage**

```
data(HTA_Location)
```

**Format**

A 1\*5118823 matrix including the location for unique probes in HTA\_transcript\_NO.

**Source**

Danielle Thierry-Mieg ,Jean Thierry-Mieg. Aceview: a comprehensive cDNA-supported gene and transcripts annotation. *Genome Biology*.2006,7(Suppl 1):S12

Manhong Dai, Pinglang Wang, Andrew D. Boyd. (2005) Evolving gene/transcript definitions significantly alter the interpretation of GeneChip data, *Nucleic Acid Research* 33(20):e175.

HTA_probes_transcripts	<i>The number of probes and the number of transcripts mapped to each gene for hta2.0 chips</i>
------------------------	--

**Description**

This data is the number of probes and the number of transcripts mapped to each gene for hta2.0 chips.

**Usage**

```
data(HTA_probes_transcripts)
```

**Format**

A 33394\*2 matrix including the number of probes and the number of transcripts mapped to each of 33394 genes for hta20 chips.

**Source**

Danielle Thierry-Mieg ,Jean Thierry-Mieg. Aceview: a comprehensive cDNA-supported gene and transcripts annotation *Genome Biology*.2006,7(Suppl 1):S12

Manhong Dai, Pinglang Wang, Andrew D. Boyd. (2005) Evolving gene/transcript definitions significantly alter the interpretation of GeneChip data, *Nucleic Acid Research* 33(20):e175.

---

HTA\_transcript\_name      *The names of transcripts mapped to each gene for hta2.0 chips*

---

**Description**

This data include the names of transcripts mapped to each gene for hta2.0 chips.

**Usage**

```
data(HTA_transcript_name)
```

**Format**

A 225456\*1 matrix including 225456 transcript names mapped to genes for hta2.0 chips.

**Source**

Danielle Thierry-Mieg ,Jean Thierry-Mieg. Aceview: a comprehensive cDNA-supported gene and transcripts annotation Genome Biology.2006,7(Suppl 1):S12

Manhong Dai, Pinglang Wang, Andrew D. Boyd. (2005) Evolving gene/transcript definitions significantly alter the interpretation of GeneChip data,Nucleic Acid Research 33(20):e175.

---

HTA\_transcript\_NO      *The coordinates of probes and the mapped transcripts for hta2.0 chips*

---

**Description**

This data include the coordinates of probes and the mapped transcripts for hta2.0 chips.

**Usage**

```
data(HTA_transcript_NO)
```

**Format**

A 20626078\*3 matrix including pos\_x,pos\_y and transcript\_no. pos\_x and pos\_y are respectively X and Y coordinates of probes for hta2.0 chips. Transcript\_no is the mapped transcripts for each probe.

**Source**

Danielle Thierry-Mieg ,Jean Thierry-Mieg. Aceview: a comprehensive cDNA-supported gene and transcripts annotation Genome Biology.2006,7(Suppl 1):S12

Manhong Dai, Pinglang Wang, Andrew D. Boyd. (2005) Evolving gene/transcript definitions significantly alter the interpretation of GeneChip data,Nucleic Acid Research 33(20):e175.

<i>Human_Location</i>	<i>The coordinates of probes and the mapped PM for human exon chips</i>
-----------------------	---

**Description**

This data include the probes location for human exon chips.

**Usage**

```
data(Human_Location)
```

**Format**

A 1\*1565476 matrix including the location for unique probes in Human\_transcript\_NO.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

<i>Human_probes_transcripts</i>
---------------------------------

<i>The number of probes and the number of transcripts mapped to each gene for human exon chips</i>
--

**Description**

This data is the number of probes and the number of transcripts mapped to each gene for human exon chips.

**Usage**

```
data(Human_probes_transcripts)
```

**Format**

A 40174\*2 matrix including the number of probes and the number of transcripts mapped to each of 40174 genes for human exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

---

*Human\_transcript\_name*    *The names of transcripts mapped to each gene for human exon chips*

---

**Description**

This data include the names of transcripts mapped to each gene for human exon chips.

**Usage**

```
data(Human_transcript_name)
```

**Format**

A 121741\*1 matrix including 121741 transcript names mapped to genes for human exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

---

*Human\_transcript\_NO*    *The coordinates of probes and the mapped transcripts for human exon chips*

---

**Description**

This data include the coordinates of probes and the mapped transcripts for human exon chips.

**Usage**

```
data(Human_transcript_NO)
```

**Format**

A 4598850\*3 matrix including pos\_x,pos\_y and transcript\_no. pos\_x and pos\_y are respectively X and Y coordinates of probes for human exon chips. Transcript\_no is the mapped transcripts for each probe.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

Mouse_Location	<i>The coordinates of probes and the mapped PM for Mouse exon chips</i>
----------------	---

**Description**

This data include the probes location for Mouse exon chips.

**Usage**

```
data(Mouse_Location)
```

**Format**

A 1\*1278936 matrix including the location for unique probes in Mouse\_transcript\_NO.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

Mouse_probes_transcripts	
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<i>The number of probes and the number of transcripts mapped to each gene for mouse exon chips</i>
--

**Description**

This data include the number of probes and the number of transcripts mapped to each gene for mouse exon chips.

**Usage**

```
data(Mouse_probes_transcripts)
```

**Format**

A 27719\*2 matrix including the number of probes and the number of transcripts mapped to each of 27719 genes for mouse exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

---

Mouse\_transcript\_name    *The names of transcripts mapped to each gene for mouse exon chips*

---

**Description**

This data include the names of transcripts mapped to each gene for mouse exon chips

**Usage**

```
data(Mouse_transcript_name)
```

**Format**

A 75751\*1 matrix including 75751 transcript names mapped to genes for mouse exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

---

Mouse\_transcript\_NO    *The coordinates of probes and the mapped transcripts mapped for mouse exon chips*

---

**Description**

This data include the coordinates of probes and the mapped transcripts for mouse exon chips.

**Usage**

```
data(Mouse_transcript_NO)
```

**Format**

A 2928848\*3 matrix including pos\_x,pos\_y and transcript\_no. pos\_x and pos\_y are respectively X and Y coordinates of probes for mouse exon chips. Transcript\_no data is the mapped transcripts for each probe.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

oligo.estrogen

*The data from the estrogen package as an ExpressionFeatureSet object*

## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code:

```
library(estrogen)
datadir <- file.path(find.package("estrogen"), "extdata")
estrogenFilenames <- c("low10-1.cel", "low10-2.cel", "high10-1.cel", "high10-2.cel",
                      "low48-1.cel", "low48-2.cel", "high48-1.cel", "high48-2.cel")
setwd(datadir)
oligo.estrogen <- read.celfiles(
  filenames=estrogenFilenames
)
pData(oligo.estrogen) <- data.frame(
  "estrogen"=c("absent", "absent", "present", "present",
              "absent", "absent", "present", "present"),
  "time.h"=c("10", "10", "10", "10", "48", "48", "48", "48"),
  row.names=rownames(pData(oligo.estrogen))
)
```

## Usage

```
data(oligo.estrogen)
```

## Format

An **ExpressionFeatureSet** object containing 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

pumapca\_estrogen

*The data from the estrogen package processed using the pumaPCA algorithm*

## Description

This data is taken from the **estrogen** package. It was created to be used in the vignette for the **puma** pacakge. It can be produced using the following code (note this could take a long time to run):

```
data(eset_estrogen_mmgmos)
pumapca_estrogen <- pumaPCA(eset_estrogen_mmgmos)
```

**Usage**

```
data(pumapca_estrogen)
```

**Format**

An [pumaPCARes](#) object containing principal components (created using pumaPCA) of 8 HG\_U95Av2 arrays, in a 2 x 2 factorial design, with 2 replicates for each combination of factors. The factors are estrogen (absent or present) and time.h (10 or 48).

**See Also**

[eset\\_estrogen\\_mmgmos](#)

**Examples**

```
data(pumapca_estrogen)
plot(pumapca_estrogen,legend1pos="right",legend2pos="top")
```

---

Rat\_Location

*The coordinates of probes and the mapped PM for Rat exon chips*

---

**Description**

This data include the probes location for Rat exon chips.

**Usage**

```
data(Rat_Location)
```

**Format**

A 1\*931210 matrix including the location for unique probes in Rat\_transcript\_NO.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

**Rat\_probes\_transcripts**

*The number of probes and the number of transcripts mapped to each gene for rat exon chips*

**Description**

This data is the number of probes and the number of transcripts mapped to each gene for rat exon chips.

**Usage**

```
data(Rat_probes_transcripts)
```

**Format**

A 23585\*2 matrix including the number of probes and the number of transcripts mapped to each of 23585 genes for rat exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

**Rat\_transcript\_name**

*The names of transcripts mapped to each gene for rat exon chips*

**Description**

This data is the names of transcripts mapped to each gene for rat exon chips

**Usage**

```
data(Rat_transcript_name)
```

**Format**

A 334851\*1 matrix including 334851 transcript names mapped to each gene for rat exon chips.

**Source**

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

---

Rat_transcript_NO	<i>The coordinates of probes and the mapped transcripts for rat exon chips</i>
-------------------	--

---

### Description

This data include the coordinates of probes and the mapped transcripts for rat exon chips.

### Usage

```
data(Rat_transcript_NO)
```

### Format

A 1491570\*3 matrix including pos\_x, pos\_y and transcript\_no. pos\_x and pos\_y are respectively X and Y coordinates of probes for rat exon chips. Transcript\_no is the mapped transcripts for each probe.

### Source

Risueno A, Fontanillo C, Dinger ME, De Las Rivas J. GATEExplorer: genomic and transcriptomic explorer; mapping expression probes to gene loci, transcripts, exons and ncRNAs. BMC Bioinformatics.2010 Apr 29;11:221.

# Index

## \*Topic datasets

affybatch.estrogen, 2  
eset\_estrogen\_comb, 3  
eset\_estrogen\_mmgmos, 3  
eset\_estrogen\_pmmgmos, 4  
eset\_estrogen\_rma, 5  
HTA\_Location, 6  
HTA\_probes\_transcripts, 6  
HTA\_transcript\_name, 7  
HTA\_transcript\_N0, 7  
Human\_Location, 8  
Human\_probes\_transcripts, 8  
Human\_transcript\_name, 9  
Human\_transcript\_N0, 9  
Mouse\_Location, 10  
Mouse\_probes\_transcripts, 10  
Mouse\_transcript\_name, 11  
Mouse\_transcript\_N0, 11  
oligo.estrogen, 4, 5, 12  
pumapca\_estrogen, 12  
pumaPCARes, 13  
Rat\_Location, 13  
Rat\_probes\_transcripts, 14  
Rat\_transcript\_name, 14  
Rat\_transcript\_N0, 15  
  
AffyBatch, 2  
affybatch.estrogen, 2  
  
eset\_estrogen\_comb, 3  
eset\_estrogen\_mmgmos, 3, 3, 5, 13  
eset\_estrogen\_pmmgmos, 4  
eset\_estrogen\_rma, 4, 5  
ExpressionFeatureSet, 12  
ExpressionSet, 3, 5  
exprReslt, 4  
  
HTA\_Location, 6  
HTA\_probes\_transcripts, 6  
HTA\_transcript\_name, 7  
HTA\_transcript\_N0, 7  
  
Human\_Location, 8  
Human\_probes\_transcripts, 8  
Human\_transcript\_name, 9  
Human\_transcript\_N0, 9  
  
Mouse\_Location, 10  
Mouse\_probes\_transcripts, 10  
Mouse\_transcript\_name, 11  
Mouse\_transcript\_N0, 11  
  
oligo.estrogen, 4, 5, 12  
  
pumapca\_estrogen, 12  
pumaPCARes, 13  
  
Rat\_Location, 13  
Rat\_probes\_transcripts, 14  
Rat\_transcript\_name, 14  
Rat\_transcript\_N0, 15