## Package 'pdInfoBuilder'

October 17, 2024

Title Platform Design Information Package Builder

**Description** Builds platform design information packages.

These consist of a SQLite database containing feature-level data such as x, y position on chip and

featureSet ID. The database also incorporates featureSet-level annotation data. The products of this packages are used by the oligo pkg.

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Collate AllClasses.R AllGenerics.R initialize-methods.R utils.R schema.R initDb.R initDb.snp6.R pmmmBlockToMat.R loaders.R loaders.snp6.R makePdInfoPackage-methods.R chipName-methods.R getGeometry-methods.R pdBuilderV2TiledRegion.R pdBuilderV2ExonTranscription.R pdBuilderV2Gene.R pdBuilderV2HTA2.R pdBuilderV2AffyTiling.R pdBuilderV2NgsExpression.R pdBuilderV2AffyExpressionHT.R pdBuilderV2AffySNP.R pdBuilderV2AffySNPCNV.R pdBuilderV2miRNA.R pdBuilderV3GenericArray.R pdBuilderV2Clariom.R

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## **Description**

PD Info Package Seed for Affymetrix Clariom S Arrays

## **Objects from the Class**

```
Objects can be created by calls of the form new("AffyClariomSPDInfoPkgSeed", pgfFile, clfFile, coreMps, transFile, ...).
```

## **Slots**

```
pgfFile: PGF filename
clfFile: CLF filename
coreMps: MPS filename
transFile: Transcript annotation CSV file
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
```

version: A character vector giving the version number of the package.

license: The license of the package author: Author of the package

email: An email address to use in the Maintainer field

biocViews: Character data for the biocViews field of the DESCRIPTION file

#### Methods

```
chipName chipName
getGeometry initialize
makePdInfoPackage package creator
```

## **Examples**

```
showClass("AffyClariomSPDInfoPkgSeed")
```

AffyExpressionPDInfoPkgSeed-class

Class "AffyExpressionPDInfoPkgSeed"

## **Description**

PD Info Package Seed for Affymetrix Expression Arrays

## **Objects from the Class**

```
Objects can be created by calls of the form new("AffyExpressionPDInfoPkgSeed", cdfFile, csvAnnoFile, tabSeqFile, ...).
```

#### **Slots**

cdfFile: CDF filename celFile: CEL filename

tabSeqFile: TAB sequence file

chipName: Name of the chip or platform manufacturer: chip/platform manufacturer

url: chip URL

genomebuild: The genome build this platform is based upon.

organism: organism for chip. species: species for chip.

version: A character vector giving the version number of the package.

license: The license of the package author: Author of the package

email: An email address to use in the Maintainer field

biocViews: Character data for the biocViews field of the DESCRIPTION file

#### Methods

```
chipName chipName
getGeometry initialize
makePdInfoPackage package creator
```

#### **Examples**

```
showClass("AffyExpressionPDInfoPkgSeed")
```

AffySNPCNVPDInfoPkgSeed-class

Class "AffySNPCNVPDInfoPkgSeed"

## **Description**

This class represents Platform Design (PD) packages for Affymetrix genomewide (SNP 5.0 and SNP 6.0) arrays.

## **Objects from the Class**

Objects can be created by calls of the form new("AffySNPCNVPDInfoPkgSeed", cdfFile, csvAnnoFile, csvSeqFile, csvAnnoFileCnv, csvSeqFileCnv, splineParamFile, crlmmInfoFile, referenceDistFile, ...).

#### **Slots**

```
cdfFile: Path to the CDF file for this.
csvAnnoFile: Path to the Affymetrix CSV annotation for the SNP probes.
csvSeqFile: Path to the (SNP) probe sequence file.
csvAnnoFileCnv: Path to the Affymetrix CSV annotation for the CNV probes.
csvSeqFileCnv: Path to the (CNV) probe sequence file.
splineParamFile: Path to the spline parameters file used to compute the predicted accuracy of
     the the genotype calls. Used internally in .predictAccuracy.
crlmmInfoFile: Path to is data file containing regions data used by the crlmm function.
referenceDistFile: Path to a reference distribution file used in the normalization step. This is
     the reference used in snprma.
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
```

version: A character vector giving the version number of the package.

license: The license of the package author: Author of the package

email: An email address to use in the Maintainer field

biocViews: Character data for the biocViews field of the DESCRIPTION file

#### Methods

```
chipName signature(object = "AffySNPCNVPDInfoPkgSeed"): ...
getGeometry signature(object = "AffySNPCNVPDInfoPkgSeed"): ...
makePdInfoPackage signature(object = "AffySNPCNVPDInfoPkgSeed"): ...
```

#### Notes

\*IMPORTANT\* Users are strongly advised to download Affymetrix SNP packages from BioConductor.

The files used for slots splineParamFile, crlmmInfoFile, and referenceDistFile are generated by the Bioconductor project for each chip/platform and are hosted in our svn data repository at <a href="https://hedgehog.fhcrc.org/bioc-data/trunk/annotation/parms\_store">https://hedgehog.fhcrc.org/bioc-data/trunk/annotation/parms\_store</a>. When makePdInfoPackage is run, these files are simply copied to the inst/extdata directory of the generated package.

## Author(s)

Benilton Carvalho

#### **Examples**

```
showClass("AffySNPCNVPDInfoPkgSeed")
```

AffySNPCNVPDInfoPkgSeed2-class

Class "AffySNPCNVPDInfoPkgSeed2"

#### **Description**

A generic annotation package builder for Affymetrix SNP/CNV arrays. This is a simplified version of the annotation package and crlmm will \*NOT\* work for them.

## **Objects from the Class**

```
Objects can be created by calls of the form new("AffySNPCNVPDInfoPkgSeed2", csvAnnoFileCnv, csvSeqFileCnv, ...).
```

#### **Slots**

```
cdfFile: Path to the CDF file for this.
csvAnnoFile: Path to the Affymetrix CSV annotation for the SNP probes.
csvSeqFile: Path to the (SNP) probe sequence file.
csvAnnoFileCnv: Path to the Affymetrix CSV annotation for the CNV probes.
csvSeqFileCnv: Path to the (CNV) probe sequence file.
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
version: A character vector giving the version number of the package.
license: The license of the package
author: Author of the package
email: An email address to use in the Maintainer field
biocViews: Character data for the biocViews field of the DESCRIPTION file
```

#### Methods

```
chipName signature(object = "AffySNPCNVPDInfoPkgSeed2"): ...
makePdInfoPackage signature(object = "AffySNPCNVPDInfoPkgSeed2"): ...
```

## Note

This is a simplified annotation package. CRLMM won't work for these objects.

The user may need to rename the columns or even add column names to the annotation and sequence files. In case problems are found, column names are suggested.

## Author(s)

Benilton Carvalho

```
showClass("AffySNPCNVPDInfoPkgSeed2")
```

```
AffySNPPDInfoPkgSeed-class 
 Class "AffySNPPDInfoPkgSeed"
```

## **Description**

This class represents Platform Design (PD) packages for Affymetrix mapping (SNP chip) arrays.

#### **Objects from the Class**

```
Objects can be created by calls of the form new("AffySNPPDInfoPkgSeed", splineParamFile, crlmmInfoFile, referenceDistFile, ...).
```

## Slots

```
splineParamFile: Spline parameters file used to compute the predicted accuracy of the genotype
crlmmInfoFile: Data file containing regions data used by the crlmm function.
referenceDistFile: Reference distribution file used in the normalization step by snprma.
cdfFile: CDF file for the design.
csvAnnoFile: Affymetrix CSV Annotation file.
csvSeqFile: Affymetrix Probe Sequence file.
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
version: A character vector giving the version number of the package.
license: The license of the package
author: Author of the package
email: An email address to use in the Maintainer field
biocViews: Character data for the biocViews field of the DESCRIPTION file
```

## Methods

```
chipName signature(object = "AffySNPPDInfoPkgSeed"): ...
getGeometry signature(object = "AffySNPPDInfoPkgSeed"): ...
makePdInfoPackage signature(object = "AffySNPPDInfoPkgSeed"): ...
```

#### Note

\*IMPORTANT\* The user is strongly advised to download Affymetrix SNP packages from BioConductor

The files used for slots splineParamFile, crlmmInfoFile, and referenceDistFile are generated by the Bioconductor project for each chip/platform and are hosted in our svn data repository at <a href="https://hedgehog.fhcrc.org/bioc-data/trunk/annotation/parms\_store">https://hedgehog.fhcrc.org/bioc-data/trunk/annotation/parms\_store</a>. When makePdInfoPackage is run, these files are simply copied to the inst/extdata directory of the generated package.

## **Examples**

```
showClass("AffySNPPDInfoPkgSeed")
cdfFile <- "Mapping250K_Nsp.cdf"
csvAnno <- "Mapping250K_Nsp_annot.csv"</pre>
csvSeq <- "Mapping250K_Nsp_probe_tab"</pre>
spline <- "pd.mapping250k.nsp.spline.params.rda"</pre>
refd <- "pd.mapping250k.nspRef.rda"</pre>
crlmmInf <- "pd.mapping250k.nspCrlmmInfo.rda"</pre>
pkg <- new("AffySNPPDInfoPkgSeed",</pre>
            version="0.1.5",
            author="A. U. Thor", email="au@thor.net",
            biocViews="AnnotationData",
            genomebuild="NCBI Build 35, May 2004",
            cdfFile=cdfFile, csvAnnoFile=csvAnno, csvSeqFile=csvSeq,
            splineParamFile=spline, crlmmInfoFile=crlmmInf,
            referenceDistFile=refd)
showMethods(classes=class(pkg))
```

#### **Description**

A generic annotation package builder for Affymetrix SNP arrays. This is a simplified version of the annotation package and crlmm will \*not\* work for them.

## **Objects from the Class**

```
Objects can be created by calls of the form new("AffySNPPDInfoPkgSeed2", cdfFile, csvAnnoFile, csvSeqFile, ...).
```

#### **Slots**

```
axiom: Logical flag for experimental build of annotation packages for Axiom arrays.

cdfFile: CDF file for the design.

csvAnnoFile: Affymetrix CSV Annotation file.

csvSeqFile: Affymetrix Probe Sequence file.

chipName: Name of the chip or platform

manufacturer: chip/platform manufacturer

url: chip URL

genomebuild: The genome build this platform is based upon.

organism: organism for chip.

species: species for chip.

version: A character vector giving the version number of the package.

license: The license of the package

author: Author of the package

email: An email address to use in the Maintainer field

biocViews: Character data for the biocViews field of the DESCRIPTION file
```

#### Methods

```
chipName signature(object = "AffySNPPDInfoPkgSeed2"): ...
```

#### Note

This is a simplified annotation package. CRLMM won't work for these objects.

The user may need to rename the columns or even add column names to the annotation and sequence files. In case problems are found, column names are suggested.

## **Examples**

```
showClass("AffySNPPDInfoPkgSeed2")
```

```
AffySTPDInfoPkgSeed-class
```

Class "AffySTPDInfoPkgSeed" for the Sense Target gene-level array

#### **Description**

container for parameters related to pdmapping package construction for ST type arrays

#### **Objects from the Class**

```
Objects can be created by calls of the form new("AffySTPDInfoPkgSeed", pgfFile, clfFile, probeFile, transFile, ...).
```

#### Slots

```
pgfFile: Object of class "ScalarCharacter" path to pgf
    clfFile: Object of class "ScalarCharacter" path to clf
    probeFile: Object of class "ScalarCharacter", path to probe sequence file (Optional)
    transFile: Object of class "ScalarCharacter", path to trans file (Optional)
    chipName: Name of the chip or platform
    manufacturer: chip/platform manufacturer
    url: chip URL
    genomebuild: The genome build this platform is based upon.
    organism: organism for chip.
    species: species for chip.
    version: A character vector giving the version number of the package.
    license: The license of the package
    author: Author of the package
    email: An email address to use in the Maintainer field
    biocViews: Character data for the biocViews field of the DESCRIPTION file
Methods
    chipName signature(object = "AffySTPDInfoPkgSeed"): ...
    getGeometry signature(object = "AffySTPDInfoPkgSeed"): ...
    makePdInfoPackage signature(object = "AffySTPDInfoPkgSeed"): ...
Author(s)
    B. Carvalho
Examples
    showClass("AffySTPDInfoPkgSeed")
  AffyTilingPDInfoPkgSeed-class
                            Class "AffyTilingPDInfoPkgSeed"
```

## **Description**

PD Info Package Seed for Affymetrix Tiling Arrays

## **Objects from the Class**

Objects can be created by calls of the form new("AffyTilingPDInfoPkgSeed", ...).

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#### **Slots**

```
bpmapFile: BPMAP File - provided by Affymetrix
celFile: CEL File - provided by Affymetrix
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
version: A character vector giving the version number of the package.
license: The license of the package
author: Author of the package
email: An email address to use in the Maintainer field
biocViews: Character data for the biocViews field of the DESCRIPTION file
```

#### Methods

```
makePdInfoPackage signature(object = "AffyTilingPDInfoPkgSeed"): ...
chipName signature(object = "AffyTilingPDInfoPkgSeed"): ...
```

#### **Examples**

```
showClass("AffyTilingPDInfoPkgSeed")
```

cdf2table

Helper functions to assist the creation of an annotation package for a generic array

## **Description**

Helper functions to assist the creation of an annotation package for a generic array. This includes converting CDF files into flat tables and parsing probe sequence files.

## Usage

```
cdf2table(cdfFile)
sequenceParser(seqFile)
```

## Arguments

cdfFile	name of the CDF file to be used
segFile	name of the probe sequence file

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## **Details**

cdf2table will convert a CDF to a flat table.

seqFile will extract a flat table containing physical location and probe sequences.

chipName

Return an Official Chip/Platform Name

## Description

This generic function returns an official or standard chip/platform name.

## Usage

```
chipName(object)
```

## **Arguments**

object

See showMethods("chipName"), but generally object will be a subclass of PkgSeed.

## **Details**

The idea is that the input files can be used to determine a standard name for each platform. For example, the method for AffySNPPDInfoPkgSeed objects reads the header of the CDF file to extract a name.

## Value

A character vector of length one giving a standard name for the platform.

## Author(s)

Seth Falcon

getGeometry 13

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Return the Chip/Platform geometry

## Description

This generic function returns the geometry for a chip/platform.

#### Usage

```
getGeometry(object)
```

#### **Arguments**

object

See showMethods("getGeometry"), but generally object will be a subclass of PkgSeed.

#### **Details**

The idea is that the input files can be used to determine the geometry for each platform. For example, the method for AffySNPPDInfoPkgSeed objects reads the header of the CDF file to extract the geometry.

## Value

A list with two elements nrows and ncols

## Author(s)

Matt Settles

makePdInfoPackage

Create a Platform Design Info Package

## **Description**

This generic function create a platform design info package based on the parameters contained in object which will generally be an instance of a subclass of PkgSeed. The result is a new directory on the filesystem containing the source for the generated pdInfo package.

## Usage

```
makePdInfoPackage(object, destDir, batch_size = 10000, quiet = FALSE, unlink = FALSE)
```

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## **Arguments**

object See showMethods("makePdInfoPackage") to see available methods.

destDir Path where the resulting pdInfo package source directory will be written.

An integer controlling the size of batches processed when reading the flatfiles and loading the DB. In general, larger values of batch\_size will use more memory and less time (unless you exceed physical memory, in which case more time will be used as well).

A logical value. When TRUE, diagnostic and status messages are not printed.

unlink A logical value. If 'TRUE', and 'destDir' already contains a file or directory

with the name 'pkgname', try to unlink (remove) it.

#### **Details**

In general, creating the SQLite database will be a time and memory intensive task.

## Value

This function is called for its side-effect of producing a pdInfo source package directory.

#### Author(s)

Seth Falcon

```
Ngs {\tt ExpressionPDInfoPkgSeed-class} \\ {\tt Class~"NgsExpressionPDInfoPkgSeed"}
```

## **Description**

PDInfo package Seed for NimbleGen Expression arrays

#### **Objects from the Class**

```
Objects can be created by calls of the form new("NgsExpressionPDInfoPkgSeed", ndfFile, pairFile, xysFile, ngdFile...).
```

#### Slots

```
ndfFile: NDF (NimbleGen Design) file

xysFile: XYS File - used as template
chipName: Name of the chip or platform
manufacturer: chip/platform manufacturer
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
version: A character vector giving the version number of the package.
license: The license of the package
author: Author of the package
email: An email address to use in the Maintainer field
biocViews: Character data for the biocViews field of the DESCRIPTION file
```

#### Methods

```
makePdInfoPackage signature(.Object = "NgsExpressionPDInfoPkgSeed"): ...
chipName signature(object = "NimbleGenPDInfoPkgSeed"): ...
getGeometry signature(.Object = "NimbleGenPDInfoPkgSeed"): ...
```

```
showClass("NgsExpressionPDInfoPkgSeed")
```

## **Description**

PDInfo package Seed for NimbleGen Tiling arrays

## **Objects from the Class**

```
Objects can be created by calls of the form new("NgsTilingPDInfoPkgSeed", ndfFile, xysFile, pairFile, posFile...).
```

#### **Slots**

```
ndfFile: NDF (NimbleGen Design) file

xysFile: XYS File - used as template

posFile: POS (Positions) file

chipName: Name of the chip or platform

manufacturer: chip/platform manufacturer

url: chip URL

genomebuild: The genome build this platform is based upon.

organism: organism for chip.

species: species for chip.

version: A character vector giving the version number of the package.

license: The license of the package

author: Author of the package

email: An email address to use in the Maintainer field

biocViews: Character data for the biocViews field of the DESCRIPTION file
```

#### Methods

```
makePdInfoPackage signature(.Object = "NgsTilingPDInfoPkgSeed"): ...
chipName signature(object = "NimbleGenPDInfoPkgSeed"): ...
getGeometry signature(object = "NimbleGenPDInfoPkgSeed"): ...
```

```
showClass("NgsTilingPDInfoPkgSeed")
```

NimbleGenPDInfoPkgSeed-class

 ${\it Class~"Nimble GenPD Info Pkg Seed"}$ 

#### **Description**

PDInfo package Seed for all NimbleGen arrays

## **Objects from the Class**

Objects can be created by calls of the form new("NimbleGenPDInfoPkgSeed", ...).

#### **Slots**

```
manufacturer: Manufacturer = NimbleGen
chipName: Name of the chip or platform
url: chip URL
genomebuild: The genome build this platform is based upon.
organism: organism for chip.
species: species for chip.
version: A character vector giving the version number of the package.
license: The license of the package
author: Author of the package
email: An email address to use in the Maintainer field
biocViews: Character data for the biocViews field of the DESCRIPTION file
```

## Methods

```
chipName signature(object = "NimbleGenPDInfoPkgSeed"): ...
getGeometry signature(object = "NimbleGenPDInfoPkgSeed"): ...
```

```
showClass("NimbleGenPDInfoPkgSeed")
```

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