

# Package ‘NMdata’

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**Type** Package

**Title** Preparation, Checking and Post-Processing Data for PK/PD Modeling

**Version** 0.2.0

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**Description** Efficient tools for preparation, checking and post-processing of data in PK/PD (pharmacokinetics/pharmacodynamics) modeling, with focus on use of Nonmem. Attention is paid to ensure consistency, traceability, and Nonmem compatibility of Data. Rigorously checks final Nonmem datasets. Implemented in 'data.table', but easily integrated with 'base' and 'tidyverse'.

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## Contents

addCor	3
addOmegaCorr	4
addTAPD	4
cc	6
cl	7
colLabels	8
compareCols	8
dims	10
dt2mat	11
editCharCols	12
egdt	13
findCovs	14
findVars	15
flagsAssign	16
flagsCount	18
fnAppend	20
fnExtension	21
is.NMdata	22
listMissings	22
mat2dt	23
mergeCheck	24
NMcheckColnames	27
NMcheckData	27
NMdataConf	31
NMdataOperations	34
NMexpandDoses	35
NMextractDataFile	36
NMextractText	37
NMgenText	39
NMinfo	40
NMIsNumeric	41
NMorderColumns	42
NMreadCov	44
NMreadCsv	45
NMreadExt	46
NMreadFilters	47
NMreadInits	48
NMreadParsText	48
NMreadPhi	51
NMreadSection	52
NMreadShk	54
NMreadSizes	54
NMreadTab	55
NMrelate	56
NMreplaceDataFile	57
NMscanData	58

`addCor` 3

NMscanInput . . . . .	61
NMscanMultiple . . . . .	64
NMscanTables . . . . .	65
NMstamp . . . . .	66
NMwriteData . . . . .	67
NMwriteFilters . . . . .	70
NMwriteInits . . . . .	71
NMwriteSection . . . . .	72
NMwriteSizes . . . . .	74
print.summary_NMdata . . . . .	75
renameByContents . . . . .	76
summary.NMdata . . . . .	77
triagSize . . . . .	77
uniquePresent . . . . .	78
unNMdata . . . . .	79

**Index** 80

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<code>addCor</code>	<i>add correlations of off-diagonal OMEGA and SIGMA elements to a parameter table</i>
---------------------	---

---

### Description

add correlations of off-diagonal OMEGA and SIGMA elements to a parameter table

### Usage

```
addCor(pars, by = NULL, as.fun, col.value = "value")
```

### Arguments

<code>pars</code>	A parameter table, like returned by <code>'NMreadExt()'</code> .
<code>by</code>	The name of a column, as a string. Calculate the correlations within a grouping variable? This will often be a column containing the model name.
<code>as.fun</code>	See <code>'?NMdataConf'</code>
<code>col.value</code>	The name of the column from which to take the <code>'OMEGA'</code> values. Default is <code>"value"</code> in alignment with the output from <code>'NMreadExt()'</code> .

### Value

The parameter table with a `'corr'` column added.

---

addOmegaCorr	<i>Deprecated: use addCor. Add correlations to parameter table</i>
--------------	--

---

**Description**

Anything arguments are passed to 'addCor()'. See '?addCor()'.

**Usage**

```
addOmegaCorr(...)
```

**Arguments**

...                    Passed to addCor

**Value**

The parameter table with a 'corr' column added.

---

addTAPD	<i>Add time since previous dose to data, time of previous dose, most recent dose amount, cumulative number of doses, and cumulative dose amount.</i>
---------	--

---

**Description**

For now, doses have to be in data as EVID=1 and/or EVID=4 records. They can be in the format of one row per dose or repeated dosing notation using ADDL and II.

**Usage**

```
addTAPD(
  data,
  col.id,
  col.time,
  col.evid = "EVID",
  col.amt = "AMT",
  col.tpdos = "TPDOS",
  col.tapd = "TAPD",
  col.pdosamt = "PDOSAMT",
  col.doscuma = "DOSCUMA",
  col.doscumn = "DOSCUMN",
  prefix.cols,
  suffix.cols,
  subset.dos,
  subset.is.complete,
```

```

    order.evid = c(3, 0, 2, 4, 1),
    by,
    SDOS = 1,
    quiet,
    as.fun,
    col.ndoses
)

```

## Arguments

<code>data</code>	The data set to add the variables to.
<code>col.id</code>	The name of the column with the subject identifier. All calculations are by default done by subject, so this column name must be provided. Default is controlled by <code>'?NMdataConf()'</code> .
<code>col.time</code>	Name of time column on which calculations of relative times will be based. Default is <code>"TIME"</code> . Default is controlled by <code>'?NMdataConf()'</code> .
<code>col.evid</code>	The name of the event ID column. This must exist in data. Default is <code>EVID</code> .
<code>col.amt</code>	<code>col.evid</code> The name of the dose amount column. This must exist in data. Default is <code>AMT</code> .
<code>col.tpdos</code>	Name of the time of previous dose column (created by <code>addTAPD()</code> ). Default is <code>"TPDOS"</code> . Set to <code>NULL</code> to not create this column.
<code>col.tapd</code>	Name of the time of previous dose column (created by <code>addTAPD()</code> ). Default is <code>"TAPD"</code> . Set to <code>NULL</code> to not create this column.
<code>col.pdosamt</code>	The name of the column to be created holding the previous dose amount. Set to <code>NULL</code> to not create this column.
<code>col.doscuma</code>	The name of the column to be created holding the cumulative dose amount. Set to <code>NULL</code> to not create this column.
<code>col.doscumn</code>	The name of the column (created by <code>addTAPD</code> ) that holds the cumulative number of doses administered to the subject. Set to <code>NULL</code> to not create this column.
<code>prefix.cols</code>	String to be prepended to all generated column names, that is each of <code>col.tpdos</code> , <code>col.tapd</code> , <code>col.ndoses</code> , <code>col.pdosamt</code> , <code>col.doscuma</code> that are not <code>NULL</code> .
<code>suffix.cols</code>	String to be appended to all generated column names, that is each of <code>col.tpdos</code> , <code>col.tapd</code> , <code>col.ndoses</code> , <code>col.pdosamt</code> , <code>col.doscuma</code> that are not <code>NULL</code> .
<code>subset.dos</code>	A string that will be evaluated as a custom expression to identify relevant events. See <code>subset.is.complete</code> as well.
<code>subset.is.complete</code>	Only used in combination with non-missing <code>subset.dos</code> . By default, <code>subset.dos</code> is used in addition to the impact of <code>col.evid</code> (must be 1 or 4) and <code>col.amt</code> (greater than zero). If <code>subset.is.complete=TRUE</code> , <code>subset.dos</code> is used alone, and <code>col.evid</code> and <code>col.amt</code> are completely ignored. This is typically useful if the events are not doses but other events that are not expressed as a typical dose combination of <code>EVID</code> and <code>AMT</code> columns.
<code>order.evid</code>	Order of events. This will only matter if there are simultaneous events of different event types within subjects. Typically if using nominal time, it may be important to specify whether samples at dosing times are pre-dose samples. The default is <code>'c(3,0,4,1,2)'</code> - i.e. samples and simulations are pre-dose. See details.

by	Columns to do calculations within. Default is ID.
SDOS	Scaling value for columns related to dose amount, relative to AMT values. col.pdosamt and col.doscuma are affected and will be derived as AMT/SDOSE.
quiet	Suppress messages? Default can be set using 'NMdataConf()'.
as.fun	The default is to return data as a data.frame. Pass a function (say 'tibble::as_tibble') in as.fun to convert to something else. If data.tables are wanted, use 'as.fun="data.table"'. The default can be configured using NMdataConf.
col.ndoses	Deprecated. Use col.doscolumn instead.

### Details

addTAPD does not require the data to be ordered, and it will not order it. This means you can run addTAPD before ordering data (which may be one of the final steps) in data set preparation. The argument called order.evid is important because of this. If a dosing event and a sample occur at the same time, when which dose was the previous for that sample? Default is to assume the sample is a pre-dose sample, and hence output will be calculated in relation to the dose before. If no dose event is found before, NA's will be assigned.

### Value

A data.frame with additional columns

### See Also

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

---

cc

*Create character vectors without quotation marks*

---

### Description

When creating character vectors with several elements, it becomes a lot of quotes to type. cc provides a simple way to skip the quotes - but only for simple strings.

### Usage

```
cc(...)
```

### Arguments

... The unquoted names that will become character values in the returned vector.

### Details

Don't use cc with any special characters - only alphanumerics and no spaces supported. Also, remember that numerics are converted using as.character. Eg, this means that leading zeros are dropped.

**Value**

A character vector

**See Also**

cl

**Examples**

```
cc(a,b,`a b`)
cc(a,b,"a b")
## be careful with spaces and special characters
cc( d)
cc(" d")
cc()
## Numerics are converted using as.character
cc(001,1,13e3)
```

---

cl	<i>Define a vector with factor levels in the same order as occurring in the vector.</i>
----	---

---

**Description**

This is a shortcut for creating factors with levels as the order of appearance of the specified levels.

**Usage**

```
cl(...)
```

**Arguments**

...                    unique elements or vectors with unique elements

**Value**

A factor (vector)

**See Also**

cc

**Examples**

```
factor("b", "a")
cl("b", "a")
x <- c("b", "a")
factor(x)
cl(x)
```

---

colLabels	<i>Extract column labels as defined in SAS</i>
-----------	--

---

**Description**

Extract column labels as defined in SAS

**Usage**

```
colLabels(...)
```

**Arguments**

... See ‘?compareCols’

**Value**

A data.frame with variable and their labels

**See Also**

compareCols NMinfo

---

compareCols	<i>Compare elements in lists with aim of combining</i>
-------------	--

---

**Description**

Useful interactive tool when merging or binding objects together. It lists the names of elements that differ in presence or class across multiple datasets. Before running rbind, you may want to check the compatibility of the data.

**Usage**

```
compareCols(  
  ...,  
  list.data,  
  keep.names = TRUE,  
  test.equal = FALSE,  
  diff.only = TRUE,  
  cols.wanted,  
  fun.class = base::class,  
  quiet,  
  as.fun,  
  keepNames,  
  testEqual  
)
```



**Arguments**

<code>...</code>	objects which element names to compare
<code>list.data</code>	As alternative to <code>...</code> , you can supply the data sets in a list here.
<code>keep.names</code>	If TRUE, the original dataset names are used in reported table. If not, generic <code>x1, x2,...</code> are used. The latter may be preferred for readability.
<code>test.equal</code>	Do you just want a TRUE/FALSE to whether the names of the two objects are the same? Default is FALSE which means to return an overview for interactive use. You might want to use TRUE in programming. However, notice that this check may be overly rigorous. Many classes are compatible enough (say numeric and integer), and <code>compareCols</code> doesn't take this into account.
<code>diff.only</code>	If TRUE, don't report columns where no difference found. Default is TRUE if number of data sets supplied is greater than one. If only one data set is supplied, the full list of columns is shown by default.
<code>cols.wanted</code>	Columns of special interest. These will always be included in overview and indicated by a prepended <code>*</code> to the column names. This argument is often useful when you start by defining a set of columns that you want to end up with by combining a number of data sets.
<code>fun.class</code>	the function that will be run on each column to check for differences. <code>base::class</code> is default. Notice that the alternative <code>'base::typeof'</code> is different in certain ways. For instance, <code>'typeof'</code> will not report a difference on numeric vs <code>difftime</code> . You could basically submit any function that takes a vector and returns a single value.
<code>quiet</code>	The default is to give some information along the way on what data is found. But consider setting this to TRUE for non-interactive use. Default can be configured using <code>NMdataConf</code> .
<code>as.fun</code>	A function that will be run on the result before returning. If first input data set is a <code>data.table</code> , the default is to return a <code>data.table</code> , if not the default is to return a <code>data.frame</code> . Use whatever to get what fits in with your workflow. Default can be configured with <code>NMdataConf</code> .
<code>keepNames</code>	Deprecated. Use <code>keep.names</code> instead.
<code>testEqual</code>	Deprecated. Use <code>test.equal</code> instead.

**Details**

technically, this function compares classes of elements in lists. However, in relation to `NMdata`, this will most of the time be columns in `data.frames`.

Despite the name of the argument `fun.class`, it can be any function to be evaluated on each element in `'...'`. See examples for how to extract SAS labels on an object read with `'read_sas'` from the `'haven'` package.

**Value**

A `data.frame` with an overview of elements and their classes of objects in `...`. Class as defined by `as.fun`.

**See Also**

Other DataWrangling: [dims\(\)](#), [listMissings\(\)](#)

**Examples**

```
## get SAS labels from objects read with haven::read_sas
## Not run:
compareCols(..., fun.class=function(x)attributes(x)$label)

## End(Not run)
```

---

`dims`

*Get dimensions of multiple objects*

---

**Description**

Get dimensions of multiple objects

**Usage**

```
dims(..., list.data, keep.names = TRUE, as.fun = NULL, keepNames)
```

**Arguments**

<code>...</code>	data sets
<code>list.data</code>	As alternative to <code>...</code> , you can supply the data sets in a list here.
<code>keep.names</code>	If TRUE, the original dataset names are used in reported table. If not, generic <code>x1</code> , <code>x2</code> ,... are used. The latter may be preferred for readability in some cases.
<code>as.fun</code>	A function that will be run on the result before returning. If first input data set is a <code>data.table</code> , the default is to return a <code>data.table</code> , if not the default is to return a <code>data.frame</code> . Use whatever to get what fits in with your workflow. Default can be configured with <code>NMdataConf</code> .
<code>keepNames</code>	Deprecated. Use <code>keep.names</code> instead.

**Value**

A `data.frame` with dimensions of objects in `...` Actual class defined by `as.fun`.

**See Also**

Other DataWrangling: [compareCols\(\)](#), [listMissings\(\)](#)

---

dt2mat	<i>Convert a data.table of parameter estimates to a matrix</i>
--------	--

---

### Description

Often needed when using estimates of Omega or Sigma matrices in further calculations.

### Usage

```
dt2mat(pars, dt.subset = "unique", max.i, fill = 0, col.value)
```

### Arguments

pars	A data.table with parameters. Must contain columns 'i' and 'j' with row and column indexes and 'est' with parameter (matrix) values.
dt.subset	Specifies whether pars contains only a lower or upper triangle of an assumed symmetric matrix (most often the case for variance-covariance matrices), or it contains the full matrix. 'dt.subset="unique"' (default) means that 'pars' only contains either upper or lower diagonal matrix (including diagonal), 'dt.subset="all"' means 'pars' contains both upper and lower triangles. See details.
max.i	By default, the maximum row number is derived as the maximum value in the 'i' column. If more (empty ones) are needed, specify the maximum row number with 'max.i'. This can be necessary in cases where only estimated elements are available but a full matrix including elements related to fixed parameters is needed.
fill	Value to insert for missing elements
col.value	The name of the column from which to take the 'OMEGA' values. Default is "value" in alignment with the output from 'NMreadExt()'.

### Details

If pars does not contain all 'i' values, they will be imputed with zeros. The desired matrix dimension is inferred from 'min(i)' and 'max(i)'. In case 'dt.subset=="unique"' missing 'j' elements will also give imputations of missing elements.

### Value

a matrix

---

editCharCols

*Replace strings in character character columns of a data set*


---

## Description

Replace strings in character character columns of a data set

## Usage

```
editCharCols(data, pattern, replacement, as.fun, ...)
```

## Arguments

data	The data set to edit.
pattern	Pattern to search for in character columns. Passed to 'gsub()'. By default, 'gsub()' works with regular expressions. See ... for how to disable this if you want to replace a specific string.
replacement	pattern or string to replace with. Passed to 'gsub()'.
as.fun	The default is to return data as a data.frame. Pass a function (say tibble::as_tibble) in as.fun to convert to something else. If data.tables are wanted, use as.fun="data.table". The default can be configured using NMdataConf.
...	Additional arguments passed to 'gsub()'. Especially, notice fixed=TRUE will disable interpretation of 'pattern' and 'replace' as regular expressions.

## Value

a data.frame

## Examples

```
### remove commas from character columns
dat <- data.frame(A=1:3, text=cc(a,"a,d","g"))
editCharCols(dat, pattern=",", "")
### factors are not edited but result in an error
## Not run:
dat <- data.frame(A=1:3, text=cc(a,"a,d",g), fac=c1("a", "a,d", "g"))
editCharCols(dat, pattern=",", "")

## End(Not run)
```

---

egdt	<i>Expand grid of data.tables</i>
------	-----------------------------------

---

**Description**

Expand grid of data.tables

**Usage**

```
egdt(dt1, dt2, quiet)
```

**Arguments**

dt1	a data.table.
dt2	another data.table.
quiet	The default is to give some information along the way on what data is found. But consider setting this to TRUE for non-interactive use. Default can be configured using NMdataConf.

**Details**

Merging works mostly similarly for data.table and data.frame. However, for data.table the merge must be done by one or more columns. This means that the convenient way to expand all combinations of all rows in two data.frames is not available for data.tables. This functions provides that functionality. It always returns data.tables.

**Value**

a data.table that expands combinations of rows in dt1 and dt2.

**Examples**

```
df1 <- data.frame(a=1:2,b=3:4)
df2 <- data.frame(c=5:6,d=7:8)
merge(df1,df2)
library(data.table)
## This is not possible
## Not run:
merge(as.data.table(df1),as.data.table(df2),allow.cartesian=TRUE)

## End(Not run)
## Use egdt instead
egdt(as.data.table(df1),as.data.table(df2),quiet=TRUE)
## Dimensions are conveniently listed for interactive use
res <- egdt(as.data.table(df1),as.data.table(df2))
```

---

 findCovs

*Extract columns that vary within values of other columns*


---

### Description

This function provides an automated method to extract covariate-like columns. The user decides which columns these variables cannot vary within. So if you have repeated measures for each ID, this function can find the columns that are constant within ID and their unique values for each ID. Or, you can provide a combination of id.cols, say ID and STUDY, and get variables that do not vary within unique combinations of these.

### Usage

```
findCovs(data, by = NULL, cols.id, as.fun = NULL)
```

### Arguments

data	data.frame in which to look for covariates
by	covariates will be searched for in combinations of values in these columns. Often by will be either empty or ID. But it can also be both say c("ID","DRUG") or c("ID","TRT").
cols.id	Deprecated. Use by instead.
as.fun	The default is to return a data.table if data is a data.table and return a data.frame in all other cases. Pass a function in as.fun to convert to something else. If data is not a data.table, the default can be configured using NMdataConf.

### Value

a data set with one observation per combination of values of variables listed in by.

### See Also

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

### Examples

```
dt1=data.frame(ID=c(1,1,2,2),
               OCC=c(1,2,1,2),
               ## ID level
               eta1=c(1,1,3,3),
               ## occasion level
               eta2=c(1,3,1,5),
               ## not used
               eta3=0
               )
## model level
```

```

findCovs(dt1)
## ID level
findCovs(dt1,"ID")
## actual ID level
findVars(findCovs(dt1,"ID"))
## occasion level
findCovs(findVars(dt1,"ID"),c("ID","OCC"))
## Based on a "real data example"
## Not run:
dat <- NMscanData(system.file("examples/nonmem/xgxr001.lst", package = "NMdata"))
findCovs(dat,by="ID")
### Without an ID column we get non-varying columns
findCovs(dat)

## End(Not run)

```

---

findVars	<i>Extract columns that vary within values of other columns in a data.frame</i>
----------	---

---

### Description

If you want to look at the variability of a number of columns and you want to disregard those that are constant. Like for findCovs, by can be of arbitrary length.

### Usage

```
findVars(data, by = NULL, cols.id, as.fun = NULL)
```

### Arguments

data	data.frame in which to look for covariates
by	optional covariates will be searched for in combinations of values in these columns. Often by will be either empty or ID. But it can also be both say c("ID","DRUG") or c("ID","TRT").
cols.id	Deprecated. Use by instead.
as.fun	The default is to return a data.table if data is a data.table and return a data.frame in all other cases. Pass a function in as.fun to convert to something else. If data is not a data.table, the default can be configured using NMdataConf.

### Details

Use this to exclude columns that are constant within by. If by=ID, this could be to get only time-varying covariates.

### Value

a data set with as many rows as in data.

**See Also**

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

**Examples**

```
dt1 <- data.frame(ID=c(1,1,2,2),
                  OCC=c(1,2,1,2),
                  ## ID level
                  eta1=c(1,1,3,3),
                  ## occasion level
                  eta2=c(1,3,1,5),
                  ## not used
                  eta3=0
                  )
## model level
findCovs(dt1)
## ID level
findCovs(dt1,"ID")
## actual ID level
findVars(findCovs(dt1,"ID"))
## occasion level
findCovs(findVars(dt1,"ID"),c("ID","OCC"))
```

---

 flagsAssign

 Assign exclusion flags to a dataset based on specified table
 

---

**Description**

The aim with this function is to take a (say PK) dataset and a pre-specified table of flags, assign the flags automatically.

**Usage**

```
flagsAssign(
  data,
  tab.flags,
  subset.data,
  col.flagn,
  col.flagc,
  flags.increasing = FALSE,
  grp.incomp = "EVID",
  flagc.0 = "Analysis set",
  as.fun = NULL
)
```



**Arguments**

<code>data</code>	The dataset to assign flags to.
<code>tab.flags</code>	A data.frame containing at least these named columns: FLAG, flag, condition. Condition is disregarded for FLAG==0. FLAG must be numeric and non-negative, flag and condition are characters.
<code>subset.data</code>	An optional string that provides a subset of data to assign flags to. A common example is <code>subset="EVID==0"</code> to only assign to observations. Numerical and character flags will be missing in rows that are not matched by this subset.
<code>col.flagn</code>	The name of the column containing the numerical flag values in <code>tab.flags</code> . This will be added to data. Default value is FLAG and can be configured using <code>NMdataConf</code> .
<code>col.flagc</code>	The name of the column containing the character flag values in <code>tab.flags</code> . This will be added to data. Default value is flag and can be configured using <code>NMdataConf</code> .
<code>flags.increasing</code>	The flags are applied by either decreasing (default) or increasing value of <code>col.flagn</code> . Decreasing order means that conditions associated with higher values of <code>col.flagn</code> will be evaluated first. By using decreasing order, you can easily adjust the Nonmem IGNORE statement from <code>IGNORE(FLAG.NE.0)</code> to say <code>IGNORE(FLAG.GT.10)</code> if BLQ's have FLAG=10, and you decide to include these in the analysis.
<code>grp.incomp</code>	Column(s) that distinct incompatible subsets of data. Default is "EVID" meaning that if different values of EVID are found in data, the function will return an error. This is a safeguard not to mix data unintentionally when counting flags.
<code>flagc.0</code>	The character flag to assign to rows that are not matched by exclusion conditions (numerical flag 0).
<code>as.fun</code>	The default is to return data.tables if input data is a data.table, and return a data.frame for all other input classes. Pass a function in <code>as.fun</code> to convert to something else. If <code>return.all=FALSE</code> , this is applied to data and <code>tab.flags</code> independently.

**Details**

`dt.flags` must contain a column with numerical exclusion flags, one with character exclusion flags, and one with a expressions to evaluate for whether to apply the exclusion flag. The flags are applied sequentially, by increasing value of the numerical exclusion flag.

**Value**

The dataset with flags added. Class as defined by `as.fun`. See parameter `flags.return` as well.

**See Also**

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

**Examples**

```
## Not run:
pk <- readRDS(file=system.file("examples/data/xgxr2.rds",package="NMdata"))
dt.flags <- data.frame(
  flagn=10,
  flagc="Below LLOQ",
  condition=c("BLQ==1")
)
pk <- flagsAssign(pk,dt.flags,subset.data="EVID==0",col.flagn="flagn",col.flagc="flagc")
pk <- flagsAssign(pk,subset.data="EVID==1",flagc.0="Dosing",
  col.flagn="flagn",col.flagc="flagc")
unique(pk[,c("EVID","flagn","flagc","BLQ")])
flagsCount(pk[EVID==0],dt.flags,col.flagn="flagn",col.flagc="flagc")

## End(Not run)
```

---

 flagsCount

---

*Create an overview of number of retained and discarded datapoints.*


---

**Description**

Generate an overview of number of observations disregarded due to different reasons. And how many are left after each exclusion flag.

**Usage**

```
flagsCount(
  data,
  tab.flags,
  file,
  col.id = "ID",
  col.flagn,
  col.flagc,
  by = NULL,
  flags.increasing = FALSE,
  flagc.0 = "Analysis set",
  name.all.data = "All available data",
  grp.incomp = "EVID",
  save = TRUE,
  quiet = FALSE,
  as.fun = NULL
)
```

**Arguments**

data	The dataset including both FLAG and flag columns.
tab.flags	A data.frame containing at least these named columns: FLAG, flag, condition. Condition is disregarded for FLAG==0.

file	A file to write the table of flag counts to. Will probably be removed and put in a separate function.
col.id	The name of the subject ID column. Default is "ID".
col.flagn	The name of the column containing the numerical flag values in tab.flags. This will be added to data. Use the same as when flagsAssign was called (if that was used). Default value is FLAG and can be configured using NMdataConf.
col.flagc	The name of the column containing the character flag values in data and tab.flags. Use the same as when flagsAssign was called (if that was used). Default value is flag and can be configured using NMdataConf.
by	An optional column to group the counting by. This could be "STUDY", "DRUG", "EVID", or a combination of multiple columns.
flags.increasing	The flags are applied by either decreasing (default) or increasing value of col.flagn. By using decreasing order, you can easily adjust the Nonmem IGNORE statement from IGNORE(FLAG.NE.0) to say IGNORE(FLAG.GT.10) if BLQ's have FLAG=10, and you decide to include these in the analysis.
flagc.0	The character flag to assign to rows that are not matched by exclusion conditions (numerical flag 0).
name.all.data	What to call the total set of data before applying exclusion flags. Default is "All available data".
grp.incomp	Column(s) that distinct incompatible subsets of data. Default is "EVID" meaning that if different values of EVID are found in data, the function will return an error. This is a safeguard not to mix data unintentionally when counting flags.
save	Save file? Default is TRUE, meaning that a file will be written if file argument is supplied.
quiet	Suppress non-critical messages? Default is 'FALSE'.
as.fun	The default is to return a data.table if input data is a data.table, and return a data.frame for all other input classes. Pass a function in as.fun to convert to something else. If data is not a data.table, default can be configured using NMdataConf.

## Details

This function is used to count flags as assigned by the flagsAssign function.

Notice that the character flags reported in the output table are taken from tab.flags. The data column named by the value of col.flagc (default is flag) is not used.

In the returned table, N.discarded is the difference in number of subjects since previous step. If two is reported, it can mean that the remaining one observation of these two subjects are discarded due to this flag. The majority of the samples can have been discarded by earlier flags.

## Value

A summary table with number of discarded and retained subjects and observations when applying each condition in the flag table. "discarded" means that the reduction of number of observations and subjects resulting from the flag, "retained" means the numbers that are left after application of the flag. The default is "both" which will report both. Class as defined by as.fun.

**See Also**

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

**Examples**

```
## Not run:
pk <- readRDS(file=system.file("examples/data/xgxr2.rds", package="NMdata"))
dt.flags <- data.frame(
  flagn=10,
  flagc="Below LLOQ",
  condition=c("BLQ==1")
)
pk <- flagsAssign(pk,dt.flags,subset.data="EVID==0",col.flagn="flagn",col.flagc="flagc")
pk <- flagsAssign(pk,subset.data="EVID==1",flagc.0="Dosing",
  col.flagn="flagn",col.flagc="flagc")
unique(pk[,c("EVID","flagn","flagc","BLQ")])
flagsCount(pk[EVID==0],dt.flags,col.flagn="flagn",col.flagc="flagc")

## End(Not run)
```

---

 fnAppend

*paste something before file name extension.*

---

**Description**

Append a file name like file.mod to file\_1.mod or file\_pk.mod. If it's a number, we can pad some zeros if wanted. The separator (default is underscore) can be modified.

**Usage**

```
fnAppend(fn, x, pad0 = 0, sep = "_", allow.noext = FALSE)
```

**Arguments**

fn	The file name or file names to modify.
x	A character string or a numeric to add to the file name
pad0	In case x is numeric, a number of zeros to pad before the appended number. This is useful if you are generating say more than 10 files, and your counter will be 01, 02,..., 10,... and not 1, 2,...,10,...
sep	The separator between the existing file name (until extension) and the addition.
allow.noext	Allow 'fn' to be string(s) without extensions? Default is 'FALSE' in which case an error will be thrown if 'fn' contains strings without extensions. If 'TRUE', 'x' will be appended to fn in these cases.

**Value**

A character (vector)

## Examples

```
fnAppend("plot.png", 1)
fnAppend("plot.png", 1, pad0=2, sep="-")
fnAppend("plot.png", "one")
fnAppend("plot", "one", allow.noext=TRUE)
```

---

fnExtension

*Change file name extension*

---

## Description

Very simple but often applicable function to retrieve or change the file name extension (from say file.lst to file.mod)

## Usage

```
fnExtension(fn, ext)
```

## Arguments

fn	file name. Often ending in an extension after a period but the extension is not needed.
ext	new file name extension. If omitted or NULL, the extension of fn is returned.

## Value

A text string

## Examples

```
fnExtension("file.lst", ".mod")
fnExtension("file.lst", "mod")
fnExtension("file.lst", ".mod")
fnExtension("file.lst", cc(.mod, xml))
fnExtension(cc(file1.lst, file2.lst), cc(.xml))
fnExtension(cc(file1.lst, file2.lst), cc(.xml, .cov))
fnExtension("file.lst", "")
fnExtension("file.lst")
```

---

is.NMdata	<i>Check if an object is 'NMdata'</i>
-----------	---------------------------------------

---

**Description**

Check if an object is 'NMdata'

**Usage**

```
is.NMdata(x)
```

**Arguments**

x	Any object
---	------------

**Value**

logical if x is an 'NMdata' object

---

listMissings	<i>List rows with missing values across multiple columns</i>
--------------	--

---

**Description**

Missing can be NA and for character variables it can be certain strings too. This function is experimental and design may change in future releases.

**Usage**

```
listMissings(data, cols, by, na.strings = c("", "."), quiet = FALSE, as.fun)
```

**Arguments**

data	The data to look into.
cols	The columns to look for missings in.
by	If supplied, we are keeping track of the missings within the values of the by columns. In summary, by is included too.
na.strings	Strings that should be interpreted as missing. All spaces will be removed before we compare to na.strings. The default is c("", ".") so say ". " is a missing by default.
quiet	Keep quiet? Default is not to.
as.fun	A function that will be run on the result before returning. If first input data set is a data.table, the default is to return a data.table, if not the default is to return a data.frame. Use whatever to get what fits in with your workflow. Default can be configured with NMdataConf.

**Value**

Invisibly, a data.frame including all findings

**See Also**

Other DataWrangling: [compareCols\(\)](#), [dims\(\)](#)

---

mat2dt	<i>upper or lower triangle or all values of a matrix as long-format</i>
--------	---

---

**Description**

upper or lower triangle or all values of a matrix as long-format

**Usage**

```
mat2dt(x, triangle = "lower", as.fun)
```

**Arguments**

x	A matrix
triangle	Either "lower" (default) or "upper", or "all" for which triangle to return. "lower" and "upper" are equivalent for covariance or correlation matrices but the returned indexes will differ. "all" will return the full matrix which mostly makes sense if matrix is not a covariance or correlation matrix.
as.fun	See '?NMdataConf'

**Details**

The matrix is assumed ordered and the index numbers for rows and columns will be returned in 'i' and 'j' columns. Row names and column names will be returned in columns 'parameter.i' and 'parameter.j'.

**Value**

A 'data.frame'-like object with indexes 'i' and 'j' for position and matrix element value in 'value' column.

**See Also**

dt2mat

---

mergeCheck

---

*Merge, order, and check resulting rows and columns.*


---

### Description

Stop checking that the number of rows is unchanged after a merge - ‘mergeCheck’ checks what you really want - i.e. x is extended with columns from y while all rows in x are retained, and no new rows are created (plus some more checks). ‘mergeCheck’ is not a merge implementation - it is a useful merge wrapper. The advantage over using much more flexible merge or join function lies in the fully automated checking that the results are consistent with the simple merge described above.

### Usage

```
mergeCheck(
  x,
  y,
  by,
  by.x,
  by.y,
  common.cols = base::warning,
  ncols.expect,
  track.msg = FALSE,
  quiet,
  df1,
  df2,
  subset.x,
  fun.na.by = base::stop,
  as.fun,
  fun.commoncols,
  ...
)
```

### Arguments

x	A data.frame with the number of rows must should be obtained from the merge. The resulting data.frame will be ordered like x.
y	A data.frame that will be merged onto x.
by	The column(s) to merge by. Character string (vector). by or by.x and by.y must be supplied.
by.x	If the columns to merge by in x and y are named differently. by or by.x and by.y must be supplied.
by.y	If the columns to merge by in x and y are named differently. by or by.x and by.y must be supplied.
common.cols	If common columns are found in x and y, and they are not used in ‘by’, this will by default create columns named like col.x and col.y in result (see ?merge).



Often, this is a mistake, and the default is to throw a warning if this happens. If using 'mergeCheck' in programming, you may want to make sure this is not happening and use `common.cols=stop`. If you want nothing to happen, you can do `common.cols=NULL`. You can also use `'common.cols="drop.x"'` to drop "non-by" columns in 'x' with identical column names in 'y'. Use "drop.y" to drop them in 'y' and avoid the conflicts. The last option is to use `'common.cols="merge.by"'` which means 'by' will automatically be extended to include all common column names.

<code>ncols.expect</code>	If you want to include a check of the number of columns being added to the dimensions of 'x'. So if <code>ncols.expect=1</code> , the resulting data must have exactly one column more than 'x' - if not, an error will be returned.
<code>track.msg</code>	If using 'mergeCheck' inside other functions, it can be useful to use <code>track.msg=TRUE</code> . This will add information to messages/warnings/errors that they came from 'mergeCheck()'. 
<code>quiet</code>	If FALSE, the names of the added columns are reported. Default value controlled by <code>NMdataConf</code> .
<code>df1</code>	Deprecated. Use <code>x</code> .
<code>df2</code>	Deprecated. Use <code>y</code> .
<code>subset.x</code>	Not implemented.
<code>fun.na.by</code>	If NA's are found in (matched) by columns in both x and why, what should we do? This could be OK, but in many cases, it's because something unexpected is happening. Use <code>fun.na.by=NULL</code> if you don't want to be notified and want to go ahead regardless.
<code>as.fun</code>	The default is to return a <code>data.table</code> if x is a <code>data.table</code> and return a <code>data.frame</code> in all other cases. Pass a function in <code>as.fun</code> to convert to something else.
<code>fun.commoncols</code>	Deprecated. Please use 'common.cols'.
<code>...</code>	additional arguments passed to <code>data.table::merge</code> . If all is among them, an error will be returned.

## Details

Besides merging and checking rows, 'mergeCheck' makes sure the order in x is retained in the resulting data (both rows and column order). Also, a warning is given if column names are overlapping, making merge create new column names like `col.x` and `col.y`. Merges and other operations are done using `data.table`. If x is a `data.frame` (and not a `data.table`), it will internally be converted to a `data.table`, and the resulting `data.table` will be converted back to a `data.frame` before returning.

'mergeCheck' is for the kind of merges where we think of x as the data to be enriched with columns from y - rows unchanged. This is even further limited than a left join where you can match rows multiple times. A common example of the use of 'mergeCheck' is for adding covariates to a pk/pd data set. We do not want that to remove or duplicate doses, observations, or simulation records. In those cases, 'mergeCheck' does all needed checks, and you can run full speed without checking dimensions (which is anyway not exactly the right thing to do in the general case) or worry that something might go wrong.

Checks performed:

- x has >0 rows

- by columns are present in x and y
- Merge is not performed on NA values. If by=ID and both x\$ID and y\$ID contain NA's, an error is thrown (see argument fun.na.by).
- Merge is done by all common column names in x and y. A warning is thrown if there are column names that are not being used to merge by. This will result in two columns named like BW.x and BW.y and is often unintended.
- Before merging a row counter is added to x. After the merge, the result is assured to have exactly one occurrence of each of the values of the row counter in x.

Moreover, row and column order from x is retained in the result.

### Value

a data.frame resulting from merging x and y. Class as defined by as.fun.

### See Also

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [tmpcol\(\)](#)

### Examples

```
df1 <- data.frame(x = 1:10,
                  y=letters[1:10],
                  stringsAsFactors=FALSE)
df2 <- data.frame(y=letters[1:11],
                  x2 = 1:11,
                  stringsAsFactors=FALSE)

mc1 <- mergeCheck(x=df1,y=df2,by="y")

## Notice as opposed to most merge/join algorithms, `mergeCheck` by
## default retains both row and column order from x
library(data.table)
merge(as.data.table(df1),as.data.table(df2))
## Here we get a duplicate of a df1 row in the result. If we only
## check dimensions, we make a mistake. `mergeCheck` captures the
## error - and tell us where to find the problem (ID 31 and 180):
## Not run:
pk <- readRDS(file=system.file("examples/data/xgxr2.rds",package="NMdata"))
dt.cov <- pk[,.(ID=unique(ID))]
dt.cov[,COV:=sample(1:5,size=.N,replace=TRUE)]
dt.cov <- dt.cov[c(1,1:(.N-1))]
res.merge <- merge(pk,dt.cov,by="ID")
dims(pk,dt.cov,res.merge)
mergeCheck(pk,dt.cov,by="ID")

## End(Not run)
```

---

NMcheckColnames	<i>Compare \$INPUT in control stream to column names in input data</i>
-----------------	--

---

### Description

Mis-specification of column names in \$DATA is a common source of problems with Nonmem models, and should be one of the first things to check for when seemingly inexplicable things happen. This function lines up input data column names with \$DATA and how NMscanData will interpret \$DATA so you can easily spot if something is off.

### Usage

```
NMcheckColnames(file, as.fun, ...)
```

### Arguments

file	A Nonmem control stream or list file
as.fun	See ?NMdataConf
...	Additional arguments passed to

### Value

An overview of input column names and how they are translated

---

NMcheckData	<i>Check data for Nonmem compatibility or check control stream for data compatibility</i>
-------------	---

---

### Description

Check data in various ways for compatibility with Nonmem. Some findings will be reported even if they will not make Nonmem fail but because they are typical dataset issues.

### Usage

```
NMcheckData(
  data,
  file,
  covs,
  covs.occ,
  cols.num,
  col.id = "ID",
  col.time = "TIME",
  col.dv = "DV",
  col.mdv = "MDV",
```

```

col.cmt = "CMT",
col.amt = "AMT",
col.flagn,
col.row,
col.usubjid,
cols.dup,
type.data = "est",
cols.disable,
na.strings,
return.summary = FALSE,
quiet = FALSE,
as.fun
)

```

### Arguments

<code>data</code>	The data to check. <code>data.frame</code> , <code>data.table</code> , <code>tibble</code> , anything that can be converted to <code>data.table</code> .
<code>file</code>	Alternatively to checking a data object, you can use <code>file</code> to specify a control stream to check. This can either be a (working or non-working) input control stream or an output control stream. In this case, <code>NMdataCheck</code> checks column names in data against control stream (see <code>NMcheckColNames</code> ), reads the data as <code>Nonmem</code> would do, and do the same checks on the data as <code>NMdataCheck</code> would do using the <code>data</code> argument. <code>col.flagn</code> is ignored in this case - instead, <code>ACCEPT/IGNORE</code> statements in control stream are applied. The <code>file</code> argument is useful for debugging a <code>Nonmem</code> model.
<code>covs</code>	columns that contain subject-level covariates. They are expected to be non-missing, numeric and not varying within subjects.
<code>covs.occ</code>	A list specifying columns that contain subject:occasion-level covariates. They are expected to be non-missing, numeric and not varying within combinations of subject and occasion. <code>covs.occ=list(PERIOD=c("FED"))</code> means that <code>FED</code> is the covariate, while <code>PERIOD</code> indicates the occasion.
<code>cols.num</code>	Columns that are expected to be present, numeric and non-NA. If a character vector is given, the columns are expected to be used in all rows. If a column is only used for a subset of rows, use a list and name the elements by subsetting strings. See examples.
<code>col.id</code>	The name of the column that holds the subject identifier. Default is "ID".
<code>col.time</code>	The name of the column holding actual time.
<code>col.dv</code>	The name of the column holding the dependent variable. For now, only one column can be specified, and <code>MDV</code> is assumed to match this column. Default is <code>DV</code> .
<code>col.mdv</code>	The name of the column holding the binary indicator of the dependent variable missing. Default is <code>MDV</code> .
<code>col.cmt</code>	The name(s) of the compartment column(s). These will be checked to be positive integers for all rows. They are also used in checks for row duplicates.
<code>col.amt</code>	The name of the dose amount column.

<code>col.flagn</code>	Optionally, the name of the column holding numeric exclusion flags. Default value is <code>FLAG</code> and can be configured using <code>NMdataConf</code> . Even though <code>FLAG</code> is the default value, no finding will be returned if the column is missing unless explicitly defined as <code>col.flagn="FLAG"</code> . This is because this way of using exclusion flags is only one of many ways you could choose to handle exclusions. Disable completely by using <code>col.flagn=FALSE</code> .
<code>col.row</code>	A column with a unique value for each row. Such a column is recommended to use if possible. Default (" <code>ROW</code> ") can be modified using <code>NMdataConf</code> .
<code>col.usubjid</code>	Optional unique subject identifier. It is recommended to keep a unique subject identifier (typically a character string including an abbreviated study name and the subject id) from the clinical datasets in the analysis set. If you supply the name of the column holding this identifier, <code>NMcheckData</code> will check that it is non-missing, that it is unique within values of <code>col.id</code> (i.e. that the analysis subject ID's are unique across actual subjects), and that <code>col.id</code> is unique within the unique subject ID (a violation of the latter is less likely).
<code>cols.dup</code>	Additional column names to consider in search of duplicate events. <code>col.id</code> , <code>col.cmt</code> , <code>col.evid</code> , and <code>col.time</code> are always considered if found in data, and <code>cols.dup</code> is added to this list if provided.
<code>type.data</code>	" <code>est</code> " for estimation data (default), and " <code>sim</code> " for simulation data. Differences are that <code>col.row</code> is not expected for simulation data, and subjects will be checked to have <code>EVID==0</code> rows for estimation data and <code>EVID==2</code> rows for simulation data.
<code>cols.disable</code>	Columns to not check. This is particularly useful when checking data sets that do not include i.e. ' <code>CMT</code> ', ' <code>EVID</code> ', and others. To skip checking specific columns, provide their names like ' <code>cols.disable=c("CMT","EVID")</code> '.
<code>na.strings</code>	Strings to be accepted when trying to convert characters to numerics. This will typically be a string that represents missing values. Default is " <code>.</code> ". Notice, actual NA, i.e. not a string, is allowed independently of <code>na.strings</code> . See <code>?NMisNumeric</code> .
<code>return.summary</code>	If <code>TRUE</code> (not default), the table summary that is printed if <code>quiet=FALSE</code> is returned as well. In that case, a list is returned, and the findings are in an element called <code>findings</code> .
<code>quiet</code>	Keep quiet? Default is not to.
<code>as.fun</code>	The default is to return data as a <code>data.frame</code> . Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If <code>data.tables</code> are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .

## Details

The following checks are performed. The term "numeric" does not refer to a numeric representation in R, but compatibility with `Nonmem`. The character string "`2`" is in this sense a valid numeric, "`id2`" is not.

- Column names must be unique and not contain special characters
- If an exclusion flag is used (for `ACCEPT/IGNORE` in `Nonmem`), elements must be non-missing and integers. Notice, if an exclusion flag is found, the rest of the checks are performed on rows where that flag equals 0 (zero) only.

- If a unique row identifier is found, it has to be non-missing, increasing integers.
- col.time (TIME), EVID, col.id (ID), col.cmt (CMT), and col.mdv (MDV): If present, elements must be non-missing and numeric.
- col.time (TIME) must be non-negative
- EVID must be in {0,1,2,3,4}.
- CMT must be positive integers. However, can be missing or zero for EVID==3.
- MDV must be the binary (1/0) representation of is.na(DV) for dosing records (EVID==0).
- AMT must be 0 or NA for EVID 0 and 2
- AMT must be positive for EVID 1 and 4
- DV must be numeric
- DV must be missing for EVID in {1,4}.
- If found, RATE must be a numeric, equaling -2 or non-negative for dosing events.
- If found, SS must be a numeric, equaling 0 or 1 for dosing records.
- If found, ADDL must be a non-negative integer for dosing records. II must be present.
- If found, II must be a non-negative integer for dosing records. ADDL must be present.
- ID must be positive and values cannot be disjoint (all records for each ID must be following each other. This is technically not a requirement in Nonmem but most often an error. Use a second ID column if you deliberately want to soften this check)
- TIME cannot be decreasing within ID, unless EVID in {3,4}.
- all ID's must have doses (EVID in {1,4})
- all ID's must have observations (EVID==0)
- ID's should not have leading zeros since these will be lost when Nonmem read, then write the data.
- If a unique row identifier is used, this must be non-missing, increasing, integer
- Character values must not contain commas (they will mess up writing/reading csv)
- Columns specified in covs argument must be non-missing, numeric and not varying within subjects.
- Columns specified in covs.occ must be non-missing, numeric and not varying within combinations of subject and occasion.
- Columns specified in cols.num must be present, numeric and non-NA.
- If a unique subject identifier column (col.usubjid) is provided, 'col.id' must be unique within values of col.usubjid and vice versa.
- Events should not be duplicated. For all rows, the combination of col.id, col.cmt, col.evid, col.time plus the optional columns specified in cols.dup must be unique. In other words, if a subject (col.id) that has say observations (col.evid) at the same time (col.time), this is considered a duplicate. The exception is if there is a reset event (col.evid is 3 or 4) in between the two rows. cols.dup can be used to add columns to this analysis. This is useful for different assays run on the same compartment (say a DVID column) or maybe stacked datasets. If col.cmt is of length>1, this search is repeated for each cmt column.

**Value**

A table with findings

**Examples**

```
## Not run:
dat <- readRDS(system.file("examples/data/xgxr2.rds", package="NMdata"))
NMcheckData(dat)
dat[EVID==0,LLOQ:=3.5]
## expecting LLOQ only for samples
NMcheckData(dat,cols.num=list(c("STUDY"),"EVID==0"=c("LLOQ")))

## End(Not run)
```

---

 NMdataConf

---

*Configure default behavior of NMdata functions*


---

**Description**

Configure default behavior across the functions in NMdata rather than typing the arguments in all function calls. Configure for your file organization, data set column names, and other NMdata behavior. Also, you can control what data class NMdata functions return (say data.tables or tibbles if you prefer one of those over data.frames).

**Usage**

```
NMdataConf(..., allow.unknown = FALSE)
```

**Arguments**

- ...
- NMdata options to modify. These are named arguments, like for base::options. Normally, multiple arguments can be used. The exception is if reset=TRUE is used which means all options are restored to default values. If NULL is passed to an argument, the argument is reset to default. See examples for how to use.
- allow.unknown Allow to store configuration of variables that are not pre-defined in NMdata. This should only be needed in cases where say another package wants to use the NMdata configuration system for variables unknown to NMdata.
- Parameters that can be controlled are:
- args.fread Arguments passed to fread when reading `_input_` data files (fread options for reading Nonmem output tables cannot be configured at this point). If you change this, you are starting from scratch, except from file. This means that existing default argument values are all disregarded.
  - args.fwrite Arguments passed to fwrite when writing csv files (NMwrite-Data). If you use this, you have to supply all arguments you want to use with fwrite, except for x (the data) and file.

- `as.fun` A function that will be applied to data returned by various data reading functions (NMscanData, NMreadTab, NMreadCsv, NMscanInput, NMscanTables). Also, data processing functions like mergeCheck, findCovs, findVars, flagsAssign, flagsCount take this into account, but slightly differently. For these functions that take data as arguments, the `as.fun` configuration is only taken into account if the data passed to the functions are not of class `data.table`. The argument `as.fun` to these functions is always adhered to. Pass an actual function, say `as.fun=tibble::as_tibble`. If you want `data.table`, use `as.fun="data.table"` (not a function).
- `check.time` Logical, applies to NMscanData only. NMscanData by default checks if output control stream is newer than input control stream and input data. Set this to FALSE if you are in an environment where time stamps cannot be relied on.
- `col.flagc` The name of the column containing the character flag values for data row omission. Default value is `flag`. Used by flagsAssign, flagsCount.
- `col.flagn` The name of the column containing numerical flag values for data row omission. Default value is `FLAG`. Used by flagsAssign, flagsCount, NMcheckData.
- `col.model` The name of the column that will hold the name of the model. See `modelname` too (which defines the values that the column will hold).
- `col.nmout` A column of this name will be a logical representing whether row was in output table or not.
- `col.nomtime` The name of the column holding nominal time. This is only used for sorting columns by NMorderColumns.
- `col.row` The name of the column containing a unique row identifier. This is used by NMscanData when `merge.by.row=TRUE`, and by NMorderColumns (row counter will be first column in data).
- `col.id` The name of the column holding the numeric subject ID. As of ‘NMdata’ 0.1.5 this is only used for sorting columns by NMorderColumns.
- `col.time` The name of the column holding actual time. As of ‘NMdata’ 0.1.5 this is only used for sorting columns by NMorderColumns.
- `dir.psn` The directory in which to find psn executables like ‘execute’ and ‘update\_inits’. Default is "" meaning that executables must be in the system search path. Not used by NMdata.
- `dir.res` Directory in which ‘NMsim’ will store simulation results files. Not used by NMdata. See `dir.sims` too.
- `dir.sims` Directory in which ‘NMsim’ will store Nonmem simulations. Not used by NMdata. See `dir.res` too.
- `file.cov` A function that will derive the path to the covariance (.cov) output file stream based on the path to the output control stream. Technically, it can be a string too, but when using NMdataConf, this would make little sense because it would direct all output control streams to the same input control streams.
- `file.ext` A function that will derive the path to the parameter (.ext) output file stream based on the path to the output control stream. Technically, it can be a string too, but when using NMdataConf, this would make little sense



because it would direct all output control streams to the same input control streams.

- `file.mod` A function that will derive the path to the input control stream based on the path to the output control stream. Technically, it can be a string too, but when using `NMdataConf`, this would make little sense because it would direct all output control streams to the same input control streams.
- `file.phi` A function that will derive the path to the Nonmem output (`.phi`) file containing individual ETA, ETC, and/or PHI values stream based on the path to the output control stream. Technically, it can be a string too, but when using `NMdataConf`, this would make little sense because it would direct all output control streams to the same input control streams.
- `file.data` A function that will derive the path to the input data based on the path to the output control stream. Technically, it can be a string too, but when using `NMdataConf`, this would make little sense because it would direct all output control streams to the same input control streams.
- `formats.read` Prioritized input data file formats to look for and use if found. Default is `c("rds","csv")` which means `rds` will be used if found, and `csv` if not. `fst` is possible too.
- `formats.write` character vector of formats.write. Default is `c("csv","rds")`. `fst` is possible too.
- `merge.by.row` Adjust the default combine method in `NMscanData`.
- `modelname` A function that will translate the output control stream path to a model name. Default is to strip `.lst`, so `/path/to/run1.lst` will become `run1`. Technically, it can be a string too, but when using `NMdataConf`, this would make little sense because it would translate all output control streams model name.
- `path.nonmem` Path (a character string) to a nonmem executable. Not used by `NMdata`. Default is `NULL`.
- `quiet` For non-interactive scripts, you can switch off the chatty behavior once and for all using this setting.
- `recover.rows` In `NMscanData`, Include rows from input data files that do not exist in output tables? This will be added to the `$row` dataset only, and `$run`, `$id`, and `$occ` datasets are created before this is taken into account. A column called `nmout` will be `TRUE` when the row was found in output tables, and `FALSE` when not. Default is `FALSE`.
- `use.input` In `NMscanData`, merge with columns in input data? Using this, you don't have to worry about remembering including all relevant variables in the output tables. Default is `TRUE`.
- `use.rds` Deprecated, use `formats.read` and `formats.write` instead. Affects `NMscanData()`, `NMscanInput()`, `NMwriteData()`.

## Details

Recommendation: Use this function transparently in the code and not in a configuration file hidden from other users.

**Value**

If no arguments given, a list of active settings. If arguments given and no issues found, TRUE invisibly.

**Examples**

```
## get current defaults
NMdataConf()
## change a parameter
NMdataConf(check.time=FALSE)
## reset one parameter to default value
NMdataConf(modelname=NULL)
## reset all parameters to defaults
NMdataConf(reset=TRUE)
```

---

NMdataOperations

*Basic arithmetic on NMdata objects*

---

**Description**

Basic arithmetic on NMdata objects

**Usage**

```
## S3 method for class 'NMdata'
merge(x, ...)

## S3 method for class 'NMdata'
t(x, ...)

## S3 method for class 'NMdata'
dimnames(x, ...)

## S3 method for class 'NMdata'
rbind(x, ...)

## S3 method for class 'NMdata'
cbind(x, ...)
```

**Arguments**

x                    an NMdata object  
...                   arguments passed to other methods.

**Details**

When 'dimnames', 'merge', 'cbind', 'rbind', or 't' is called on an 'NMdata' object, the 'NMdata' class is dropped, and then the operation is performed. So if an 'NMdata' object inherits from 'data.frame' and no other classes (which is default), these operations will be performed using the 'data.frame' methods. But for example, if you use 'as.fun' to get a 'data.table' or 'tbl', their respective methods are used instead.

**Value**

An object that is not of class 'NMdata'.

---

NMexpandDoses	<i>Transform repeated dosing events (ADDL/II) to individual dosing events</i>
---------------	---

---

**Description**

Replaces single row repeated dosing events by multiple lines, then reorders rows with respect to ID and TIME. If the row order is different, you have to reorder the output manually.

**Usage**

```

NMexpandDoses(
  data,
  col.time = "TIME",
  col.id = "ID",
  col.evid = "EVID",
  track.expand = FALSE,
  subset.dos,
  quiet = FALSE,
  as.fun
)

```

**Arguments**

data	The data set to expand
col.time	The name of the column holding the time on which time since previous dose will be based. This is typically actual or nominal time since first dose.
col.id	The subject identifier. All new columns will be derived within unique values of this column.
col.evid	The name of the event ID column. This must exist in data. Default is EVID.
track.expand	Keep track of what rows were in data originally and which ones are added by NMexpandDoses by including a column called nmexpand? nmexpand will be TRUE if the row is "generated" by NMexpandDoses.
subset.dos	A string that will be evaluated as a custom expression to identify relevant events.

quiet	Suppress messages back to user (default is FALSE)
as.fun	The default is to return data as a data.frame. Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If <code>data.tables</code> are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .

**Value**

A data set with at least as many rows as `data`. If doses are found to expand, these will be added.

---

NMextractDataFile      *Extract the data file used in a control stream*

---

**Description**

A function that identifies the input data file based on a control stream. The default is to look at the `$DATA` section of of the output control stream (or input control stream if `file.mod` argument is used). This can be partly or fully overruled by using the `dir.data` or `file.data` arguments.

**Usage**

```
NMextractDataFile(file, dir.data = NULL, file.mod, file.data = NULL)
```

**Arguments**

file	The input control stream or the list file.
dir.data	See <code>NMscanInput</code> . If used, only the file name mentioned in <code>\$DATA</code> is used. <code>dir.data</code> will be used as the path, and the existence of the file in that directory is not checked.
file.mod	The input control stream. Default is to look for <code>"file\"</code> with extension changed to <code>'mod'</code> (PSN style). You can also supply the path to the file, or you can provide a function that translates the output file path to the input file path. The default behavior can be configured using <code>NMdataConf</code> . See <code>dir.data</code> too.
file.data	Specification of the data file path. When this is used, the control streams are not used at all.

**Value**

The path to the input data file.

---

NMextractText	<i>Versatile text extractor from Nonmem (input or output) control streams</i>
---------------	---

---

### Description

If you want to extract input sections like \$PROBLEM, \$DATA etc, see NMreadSection. This function is more general and can be used to extract eg result sections.

### Usage

```

NMextractText(
  file,
  lines,
  text,
  section,
  char.section,
  char.end = char.section,
  return = "text",
  keep.empty = FALSE,
  keep.name = TRUE,
  keep.comments = TRUE,
  as.one = TRUE,
  clean.spaces = FALSE,
  simplify = TRUE,
  match.exactly = TRUE,
  type = "mod",
  linesep = "\n",
  keepEmpty,
  keepName,
  keepComments,
  asOne
)

```

### Arguments

file	A file path to read from. Normally a .mod or .lst. See lines and text as well.
lines	Text lines to process. This is an alternative to using the file and text arguments.
text	Use this argument if the text to process is one long character string, and indicate the line separator with the linesep argument. Use only one of file, lines, and text.
section	The name of section to extract. Examples: "INPUT", "PK", "TABLE", etc. It can also be result sections like "MINIMIZATION".
char.section	The section denoted as a string compatible with regular expressions. "\$" (remember to escape properly) for sections in .mod files, "0" for results in .lst files.
char.end	A regular expression to capture the end of the section. The default is to look for the next occurrence of char.section.

return	If "text", plain text lines are returned. If "idx", matching line numbers are returned. "text" is default.
keep.empty	Keep empty lines in output? Default is FALSE. Notice, comments are removed before empty lines are handled if 'keep.comments=TRUE'.
keep.name	Keep the section name in output (say, "\$PROBLEM") Default is TRUE. It can only be FALSE, if return="text".
keep.comments	Default is to keep comments. If FALSE, they will be removed.
as.one	If multiple hits, concatenate into one. This will most often be relevant with name="TABLE". If FALSE, a list will be returned, each element representing a table. Default is TRUE. So if you want to process the tables separately, you probably want FALSE here.
clean.spaces	If TRUE, leading and trailing are removed, and multiplied succeeding white spaces are reduced to single white spaces.
simplify	If asOne=FALSE, do you want the result to be simplified if only one table is found? Default is TRUE which is desirable for interactive analysis. For programming, you probably want FALSE.
match.exactly	Default is to search for exact matches of 'section'. If FALSE, only the first three characters are matched. E.G., this allows "ESTIMATION" to match "ESTIMATION" or "EST".
type	Either mod, res or NULL. mod is for information that is given in .mod (.lst file can be used but results section is disregarded). If NULL, NA or empty string, everything is considered.
linesep	If using the text argument, use linesep to indicate how lines should be separated.
keepEmpty	Deprecated. See keep.empty.
keepName	Deprecated. See keep.name.
keepComments	Deprecated. See keep.comments.
asOne	Deprecated. See as.one.

### Details

This function is planned to get a more general name and then be called by NMreadSection.

### Value

character vector with extracted lines.

### See Also

Other Nonmem: [NMapplyFilters\(\)](#), [NMgenText\(\)](#), [NMreadSection\(\)](#), [NMreplaceDataFile\(\)](#), [NMwriteSection\(\)](#)

### Examples

```
NMreadSection(system.file("examples/nonmem/xgxr001.lst", package = "NMdata"), section="DATA")
```

---

NMgenText	<i>Generate text for INPUT and possibly DATA sections of NONMEM control streams.</i>
-----------	--

---

### Description

The user is provided with text to use in Nonmem. NMwriteSection can use the results to update the control streams. INPUT lists names of the data columns while DATA provides a path to data and ACCEPT/IGNORE statements. Once a column is reached that Nonmem will not be able to read as a numeric and column is not in nm.drop, the list is stopped. Only exception is TIME which is not tested for whether character or not.

### Usage

```

NMgenText(
  data,
  drop,
  col.flagn,
  rename,
  copy,
  file,
  dir.data,
  capitalize = FALSE,
  until,
  allow.char.TIME = TRUE,
  width,
  quiet
)

```

### Arguments

data	The data that NONMEM will read. Either as a 'data.frame', or if a path to an rds or a delimited text file, the data will automatically be read first.
drop	Only used for generation of proposed text for INPUT section. Columns to drop in Nonmem \$INPUT. This has two implications. One is that the proposed \$INPUT indicates =DROP after the given column names. The other that in case it is a non-numeric column, succeeding columns will still be included in \$INPUT and can be read by NONMEM.
col.flagn	Name of a numeric column with zero value for rows to include in Nonmem run, non-zero for rows to skip. The argument is only used for generating the proposed \$DATA text to paste into the Nonmem control stream. Default is defined by 'NMdataConf()'. To skip this feature, use 'col.flagn=FALSE'.
rename	For the \$INPUT text proposal only. If you want to rename columns in NONMEM \$DATA, NMwriteData can adjust the suggested \$DATA text. If you plan to use BBW instead of BWBASE in Nonmem, consider rename=c(BBW="BWBASE"). The result will include BBW and not BWBASE.

copy	For the \$INPUT text proposal only. If you plan to use additional names for columns in Nonmem \$INPUT, NMwriteData can adjust the suggested \$INPUT text. Say you plan to use CONC as DV in Nonmem, use copy=c(DV="CONC"), i.e. copy=c(newname="existing"). INPUT suggestion will in this case contain DV=CONC.
file	The file name NONMEM will read the data from (for the \$DATA section). It can be a full path.
dir.data	For the \$DATA text proposal only. The path to the input datafile to be used in the Nonmem \$DATA section. Often, a relative path to the actual Nonmem run is wanted here. If this is used, only the file name and not the path from the file argument is used.
capitalize	For the \$INPUT text proposal only. If TRUE, all column names in \$INPUT text will be converted to capital letters.
until	Use this to truncate the columns in \$INPUT. until can either be a character (column name) or a numeric (column number). If a character is given, it is matched against the resulting column name representation in \$INPUT, i.e. this could be "DV=CONC" if you are using in this case the copy argument. In case until is of length>1, the maximum will be used (probably only interesting if character values are supplied).
allow.char.TIME	For the \$INPUT text proposal only. Assume Nonmem can read TIME and DATE even if it can't be translated to numeric. This is necessary if using the 00:00 format. Default is TRUE.
width	If positive, will be passed to strwrap for the \$INPUT text. If missing or NULL, strwrap will be called with default value. If negative or zero, strwrap will not be called.
quiet	Hold messages back? Default is defined by NMdataConf.

**Value**

Text for inclusion in Nonmem control stream, invisibly. A list with elements 'DATA' and 'INPUT'.

**See Also**

Other Nonmem: [NMapplyFilters\(\)](#), [NMextractText\(\)](#), [NMreadSection\(\)](#), [NMreplaceDataFile\(\)](#), [NMwriteSection\(\)](#)

---

 NMinfo

---

*Get metadata from an NMdata object*


---

**Description**

Extract metadata such as info on tables, columns and further details in your favorite class



**Usage**

```
NMinfo(data, info, as.fun)
```

**Arguments**

<code>data</code>	An object of class <code>NMdata</code> (a result of <code>'NMscanData()'</code> )
<code>info</code>	If not passed, all the metadata is returned. You can use "details", "tables", or "columns" to get only these subsets. If <code>info</code> is "tables" or "columns"
<code>as.fun</code>	The default is to return data as a <code>'data.frame'</code> . Pass a function (say <code>'tibble::as_tibble'</code> ) in <code>as.fun</code> to convert to something else. If <code>'data.table'</code> 's are wanted, use <code>'as.fun="data.table"'</code> . The default can be configured using <code>'NMdataConf()'</code> .

**Value**

A table of class as defined by `as.fun` in case `info` is "columns" or "tables". A list if `info` missing or equal to "details".

---

<code>NMisNumeric</code>	<i>Test if a variable can be interpreted by Nonmem</i>
--------------------------	--

---

**Description**

`Nonmem` can only interpret numeric data. However, a factor or a character variable may very well be interpretable by `Nonmem` (e.g. "33"). This function tells whether `Nonmem` will be able to read it.

**Usage**

```
NMisNumeric(x, na.strings = ".", each = FALSE)
```

**Arguments**

<code>x</code>	The vector to check Don't export
<code>na.strings</code>	Tolerated strings that do not translate to numerics. Default is to accept "." because it's common to write missing values that way to <code>Nonmem</code> (even if <code>Nonmem</code> will handle them as zeros rather than missing). Notice actual NA's are accepted so you may want to use <code>na.strings=NULL</code> if you don't code missings as "." and just do this when writing the data set to a delimited file (like <code>NMwriteData</code> will do for you).
<code>each</code>	Use <code>each=TRUE</code> to evaluate each element in a vector individually. The default ( <code>each=FALSE</code> ) is to return a single-length logical for a vector <code>x</code> summarizing whether all the elements are numeric-compatible.

**Value**

TRUE or FALSE

---

NMorderColumns      *Standardize column order in Nonmem input data*

---

### Description

Order data columns for easy export to Nonmem. No data values are edited. The order is configurable through multiple arguments. See details.

### Usage

```
NMorderColumns(
  data,
  first,
  last,
  lower.last = FALSE,
  chars.last = TRUE,
  alpha = TRUE,
  col.id,
  col.nomtime,
  col.time,
  col.row,
  col.flagn,
  col.dv = "DV",
  allow.char.TIME = TRUE,
  as.fun = NULL,
  quiet
)
```

### Arguments

data	The dataset which columns to reorder.
first	Columns that should come almost first. See details.
last	Columns to move to back of dataset. If you work with a large dataset, and some columns are irrelevant for the Nonmem runs, you can use this argument.
lower.last	Should columns which names contain lowercase characters be moved towards the back? Some people use a standard of lowercase variables (say "race") being character representations ("Asian", "Caucasian", etc.) variables and the uppercase (1,2,...) being the numeric representation for Nonmem.
chars.last	Should columns which cannot be converted to numeric be put towards the end? A column can be a character or a factor in R, but still be valid in Nonmem (often the case for ID which can only contain numeric digits but really is a character or factor). So rather than only looking at the column class, the columns are attempted converted to numeric. Notice, it will attempted to be converted to numeric to test whether Nonmem will be able to make sense of it, but the values in the resulting dataset will be untouched. No values will be edited. If TRUE, logicals will always be put last. NA's must be NA or ".".

alpha	Sort columns alphabetically. Notice, this is the last order priority applied.
col.id	Name of the (numeric) unique subject ID. Can be controlled with 'NMdataConf()'.
col.nomtime	The name of the column containing nominal time. If given, it will put the column quite far left, just after row counter and 'col.id'. Default value is NOMTIME and can be configured with 'NMdataConf()'.
col.time	The name of the column containing actual time. If given, it will put the column quite far left, just after row counter, subject ID, and nominal time. Default value is 'TIME'. Can be controlled with 'NMdataConf()'.
col.row	A row counter column. This will be the first column in the dataset. Technically, you can use it for whatever column you want first. Default value is 'ROW' and can be configured with 'NMdataConf()'.
col.flagn	The name of the column containing numerical flag values for data row omission. Default value is FLAG and can be configured with 'NMdataConf()'.
col.dv	a vector of column names to put early to represent dependent variable(s). Default is DV.
allow.char.TIME	For the \$INPUT text proposal only. Assume Nonmem can read TIME and DATE even if it can't be translated to numeric. This is necessary if using the 00:00 format. Default is TRUE.
as.fun	The default is to return a data.table if data is a data.table and return a data.frame in all other cases. Pass a function in as.fun to convert to something else. The default can be configured using 'NMdataConf()'. However, if data is a data.table, settings via 'NMdataConf()' are ignored.
quiet	If true, no warning will be given about missing standard Nonmem columns.

### Details

This function will change the order of columns but it will never edit values in any columns. The ordering is by the following steps, each step depending on corresponding argument.

**"col.row** - " Row id if argument row is non-NULL

**"not editable** - " ID (if a column is called ID)

**"col.nomtime** - " Nominal time.

**"col.time** - " Actual time.

**"first** - " user-specified first columns

**"Only col.dv editable** - " Standard Nonmem columns: EVID, CMT, AMT, RATE, col.dv, MDV

**"last** - " user-specified last columns

**"chars.last** - " numeric, or interpretable as numeric

**"not editable** - " less often used Nonmem names: col.flagn, OCC, ROUTE, GRP, TRIAL, DRUG, STUDY

**"lower.last** - " lower case in name

**"alpha** - " Alphabetic/numeric sorting

**Value**

data with modified column order.

**See Also**

Other DataCreate: [NMstamp\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

---

NMreadCov

*Read in data file*

---

**Description**

Read in data file

**Usage**

```
NMreadCov(file, auto.ext, ...)
```

**Arguments**

file	The .cov covariance Nonmem matrix file to read
auto.ext	If 'TRUE' (default) the extension will automatically be modified using 'NMdataConf()\$file.cov'. This means 'file' can be the path to an input or output control stream, and 'NMreadCov()' will still read the '.cov' file.
...	Passed to fread

**Details**

This function is taken from nonmem2rx::nmcov which was based on NMdata::NMreadTab.

**Value**

A matrix with covariance step from NONMEM

**Author(s)**

Philip Delff and Matthew L. Fidler

---

NMreadCsv	<i>Read input data formatted for Nonmem</i>
-----------	---

---

### Description

This function is especially useful if the csv file was written using NMwriteData.

### Usage

```
NMreadCsv(file, args.fread, as.fun = NULL, format, args.fst)
```

### Arguments

file	The file to read. Must be pure text.
args.fread	List of arguments passed to fread. Notice that except for "file", you need to supply all arguments to fread if you use this argument. Default values can be configured using NMdataConf.
as.fun	The default is to return data as a data.frame. Pass a function (say tibble::as_tibble) in as.fun to convert to something else. If data.tables are wanted, use as.fun="data.table". The default can be configured using NMdataConf.
format	Format of file to read. Can be of length>1 in which case the first format found will be used (i.e. format is a prioritized vector). If not one of "rds" or "fst", it is assumed to be a delimited text file. Default is to determine this from the file name extension. Notice, if a delimited format is used, the extension can very well be different from "csv" (say file name is "input.tab"). This will work for any delimited format supported by fread.
args.fst	Optional arguments to pass to read_fst if format="fst" is used.

### Details

This is almost just a shortcut to fread so you don't have to remember how to read the data that was exported for Nonmem. The only added feature is that meta data as written by NMwriteData is read and attached as NMdata metadata before data is returned.

### Value

A data set of class as defined by as.fun.

### See Also

NMwriteData

Other DataRead: [NMreadTab\(\)](#), [NMscanData\(\)](#), [NMscanInput\(\)](#), [NMscanTables\(\)](#)

---

 NMreadExt

*Read information from Nonmem ext files*


---

### Description

Read information from Nonmem ext files

### Usage

```

NMreadExt(
  file,
  return,
  as.fun,
  modelname,
  col.model,
  auto.ext,
  tableno = "max",
  file.ext
)

```

### Arguments

file	Path to the ext file
return	The .ext file contains both final parameter estimates and iterations of the estimates. If return="pars" (default) the final estimates are returned in addition to what other parameter-level information is found, like FIX, sd etc. as columns. If return="iterations", the iterations are returned (including objective function value). If return="obj" objective function value at final estimate is returned. If return="all", all are returned, though in separate data.frames compiled in a list.
as.fun	The default is to return data as a data.frame. Pass a function (say 'tibble::as_tibble') in as.fun to convert to something else. If 'data.table's are wanted, use 'as.fun="data.table"'. The default can be configured using 'NMdataConf()'.
modelname	See '?NMscanData'
col.model	See '?NMscanData'
auto.ext	If 'TRUE' (default) the extension will automatically be modified using 'NMdataConf()\$file.ext'. This means 'file' can be the path to an input or output control stream, and 'NMreadExt' will still read the '.ext' file.
tableno	In case the ext file contains multiple tables, this argument controls which one to choose. The options are <ul style="list-style-type: none"> <li>• "max" (default) Pick the table with the highest table number. This typically means the results from the last '\$ESTIMATION' step are used.</li> <li>• "min" Pick results from the first table available.</li> <li>• "all" Keep all results. The tables can be distinguished by the 'tableno' column.</li> </ul>

- an integer greater than 0, in which case the table with this table number will be picked.

file.ext      Deprecated. Please use file instead.

### Details

The parameter table returned if return="pars" or return="all" will contain columns based on the Nonmem 7.5 manual. It defines codes for different parameter-level values. They are:

-1e+09: se -1000000002: eigCor -1000000003: cond -1000000004: stdDevCor -1000000005: seStdDevCor -1000000006: FIX -1000000007: termStat -1000000008: partLik

The parameter name is in the parameter column. The "parameter type", like "THETA", "OMEGA", "SIGMA" are available in the par.type column. Counters are available in i and j columns. j will be NA for par.type=="THETA"

The objective function value is included as a parameter.

Notice that in case multiple tables are available in the 'ext' file, the column names are taken from the first table. E.g., in case of SAEM/IMP estimation, the objective function values will be in the 'SAEMOBJ' column, even for the IMP step. This may change in the future.

### Value

If return="all", a list with a final parameter table and a table of the iterations. If return="pars", only the parameter table, and if return="iterations" only the iterations table. If you need both, it may be more efficient to only read the file once and use return="all". Often, only one of the two are needed, and it more convenient to just extract one.

---

NMreadFilters      *Read data filters from a NONMEM model*

---

### Description

Read data filters from a NONMEM model

### Usage

```
NMreadFilters(file, lines, filters.only = TRUE, as.fun)
```

### Arguments

file	Control stream path
lines	Control stream lines if already read from file
filters.only	Return the filters only or also return the remaining text in a separate object? If 'FALSE', a list with the two objects is returned.
as.fun	Function to run on the tables with filters.

### Value

A 'data.frame' with filters

---

NMreadInits                      *Tabulate information from parameter sections in control streams*

---

### Description

Tabulate information from parameter sections in control streams

### Usage

```
NMreadInits(file, lines, section, return = "pars", as.fun)
```

### Arguments

file	Path to a control stream. See 'lines' too.
lines	A control stream as text lines. Use this or 'file'.
section	The section to read. Typically, "theta", "omega", or "sigma". Default is those three.
return	By default (when return="pars", a parameter table with initial values, FIX, lower and upper bounds etc. In most cases, that is what is needed to derive information about parameter definitions. If return="all", two additional tables are returned which can be used if the aim is to modify and write the resulting parameters to a control stream.
as.fun	See ?NMscanData

### Value

A 'data.frame' with parameter values. If 'return="all"', a list of three tables.

---

NMreadParsText                      *Read comments to parameter definitions in Nonmem control streams*

---

### Description

When interpreting parameter estimates, it is often needed to recover information about the meaning of the different parameters from control stream. 'NMreadParsText' provides a flexible way to organize the comments in the parameter sections into a 'data.frame'. This can subsequently easily be merged with parameter values as obtained with 'NMreadExt'.



**Usage**

```

NMreadParsText(
  file,
  lines,
  format,
  format.omega = format,
  format.sigma = format.omega,
  spaces.split = FALSE,
  unique.matches = TRUE,
  field.idx = "idx",
  use.idx = FALSE,
  modelname,
  col.model,
  as.fun,
  use.theta.idx,
  fields,
  fields.omega = fields,
  fields.sigma = fields.omega
)

```

**Arguments**

<code>file</code>	Path to the control stream to read.
<code>lines</code>	As an alternative to ‘file’, the control stream or selected lines of the control stream can be provided as a vector of lines.
<code>format</code>	Defines naming and splitting of contents of lines in parameter sections. Default is <code>"%init;%idx;%symbol;%label;%unit"</code> . Be careful to remember percentage symbols in front of any variable names.
<code>format.omega</code>	Like ‘format’, applied to ‘\$OMEGA’ section. Default is to reuse ‘format’.
<code>format.sigma</code>	Like ‘format’, applied to ‘\$SIGMA’ section. Default is to reuse ‘format.omega’.
<code>spaces.split</code>	Is a blank in ‘fields’ to be treated as a field separator? Default is not to (i.e. neglect spaces in ‘fields’).
<code>unique.matches</code>	If TRUE, each line in the control stream is assigned to one parameter, at most. This means, if two parameters are listed in one line, the comments will only be used for one of the parameters, and only that parameter will be kept in output. Where this will typically happen is in ‘\$OMEGA’ and ‘\$SIGMA’ sections where off-diagonal may be put on the same line as diagonal elements. Since the off-diagonal elements are covariances of variables that have already been identified by the diagonals, the off-diagonal elements can be automatically described. For example, if ‘OMEGA(1,1)’ is between-subject variability (BSV) on CL and ‘OMEGA(2,2)’ is BSV on V, then we know that ‘OMEGA(2,1)’ is covariance of (BSV on) CL and V.
<code>field.idx</code>	If an index field is manually provided in the control stream comments, define the name of that field in ‘format’ and tell ‘NMreadParsTab()’ to use this idx to organize especially OMEGA and SIGMA elements by pointing to it with ‘field.idx’. The default is to look for a variable called ‘idx’. If the index has

	values like 1-2 on an OMEGA or SIGMA row, the row is interpreted as the covariance between OMEGA/SIGMA 1 and 2.
use.idx	The default method is to automatically identify element numbering ('i' for THETAs, 'i' and 'j' for OMEGAs and SIGMAs). The automated method is based on identification of 'BLOCK()' structures and numbers of initial values. Should this fail, or should you want to control this manually, you can include a parameter counter in the comments and have 'NMreadParsText()' use that to assign the numbering. 'use.idx=FALSE' is default and means all blocks are handled automatically, 'use.idx=TRUE' assumes you have a counter in all sections, and a character vector like 'use.idx="omega"' can be used to denote which sections use such a counter from the control stream. When using a counter on OMEGA and SIGMA, off-diagonal elements MUST be denoted by 'i-j', like '2-1' for OMEGA(2,1). See 'field.idx' too.
modelName	See ?NMscanData
col.model	See ?NMscanData
as.fun	See ?NMscanData
use.theta.idx	If an index field in comments should be used to number thetas. The index field is used to organize '\$OMEGA's and '\$SIGMA's because they are matrices but I do not see where this is advantageous to do for '\$THETA's. Default 'use.theta.idx=FALSE' which means '\$THETA's are simply counted.
fields	Deprecated. Use 'format'.
fields.omega	Deprecated. Use 'format.omega'.
fields.sigma	Deprecated. Use 'format.sigma'.

### Details

Off-diagonal omega and sigma elements will only be correctly treated if their num field specifies say 1-2 to specify it is covariance between 1 and 2.

SAME elements in \$OMEGA will be skipped altogether.

### Value

data.frame with parameter names and fields read from comments

### Examples

```
## setDTthreads() is only needed for CRAN. Users should not do this.
data.table::setDTthreads(1)
## end setDTthreads() for CRAN

## notice, examples on explicitly stated lines. Most often in
## practice, one would use the file argument to automatically
## extract the $THETA, $OMEGA and $SIGMA sections from a control
## stream.

text <- c("

```

```

$THETA (.1) ;[1]; LTVKA (mL/h)
$OMEGA BLOCK(3)
0.126303 ; IIV.CL ; 1 ;IIV ;Between-subject variability on CL;-
0.024 ; IIV.CL.V2.cov ; 1-2 ;IIV ;Covariance of BSV on CL and V2;-
0.127 ; IIV.V2 ; 2 ;IIV ;Between-subject variability on V2;-
0.2 ; IIV.CL.V3.cov ; 1-3 ;IIV ;Covariance of BSV on CL and V3;-
0.2 ; IIV.V2.V3.cov ; 2-3 ;IIV ;Covariance of BSV on V2 and V3;-
0.38 ; IIV.V3 ; 3 ;IIV ;Between-subject variability on V3;-
$OMEGA 0 FIX ; IIV.KA ; 4 ;IIV ;Between-subject variability on KA;-
$SIGMA 1
")
lines <- strsplit(text,split="\n")[[1]]

res <- NMreadParsText(lines=lines,
format="%init;[%num];%symbol",
format.omega="%init; %symbol ; %num ; %type ; %label ; %unit",
field.idx="num")

## BLOCK() SAME are skipped
text <- c("
$THETA
(0,0.1) ; THE1 - 1) 1st theta
(0,4.2) ; THE2 - 2) 2nd theta
$OMEGA 0.08 ; IIV.TH1 ; 1 ;IIV
$OMEGA BLOCK(1)
0.547465 ; IOV.TH1 ; 2 ;IOV
$OMEGA BLOCK(1) SAME
$OMEGA BLOCK(1) SAME")
lines <- strsplit(text,split="\n")[[1]]
res <- NMreadParsText(lines=lines,
format="%init;%symbol - %idx) %label",
format.omega="%init; %symbol ; %idx ; %label "
)

```

---

NMreadPhi

*Read information from Nonmem phi files*


---

## Description

Read information from Nonmem phi files

## Usage

```
NMreadPhi(file, as.fun, modelname, col.model, auto.ext, file.phi)
```

## Arguments

file Path to the phi file. See 'auto.ext' too.

as.fun	The default is to return data as a data.frame. Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If data.tables are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .
modelName	See <code>?NMscanData</code>
col.model	See <code>?NMscanData</code>
auto.ext	If <code>'auto.ext=TRUE'</code> , the file name extension will automatically be changed using the setting in <code>'NMdataConf()\$file.phi'</code> - this by default means that the <code>'phi'</code> extension will be used no matter what extension the provided file name has.
file.phi	Deprecated. Use <code>'file'</code> .

**Value**

A list with a final parameter table and a table of the iterations

---

NMreadSection	<i>Extract sections of Nonmem control streams</i>
---------------	---

---

**Description**

This is a very commonly used wrapper for the input part of the model file. Look `NMextractText` for more general functionality suitable for the results part too.

**Usage**

```
NMreadSection(
  file = NULL,
  lines = NULL,
  text = NULL,
  section,
  return = "text",
  keep.empty = FALSE,
  keep.name = TRUE,
  keep.comments = TRUE,
  as.one = TRUE,
  clean.spaces = FALSE,
  simplify = TRUE,
  keepEmpty,
  keepName,
  keepComments,
  asOne,
  ...
)
```

**Arguments**

file	A file path to read from. Normally a .mod or .lst. See lines also.
lines	Text lines to process. This is an alternative to using the file argument.
text	Use this argument if the text to process is one long character string, and indicate the line separator with the linesep argument (handled by NMextractText). Use only one of file, lines, and text.
section	The name of section to extract without "\$". Examples: "INPUT", "PK", "TABLE", etc. Not case sensitive.
return	If "text", plain text lines are returned. If "idx", matching line numbers are returned. "text" is default.
keep.empty	Keep empty lines in output? Default is FALSE. Notice, comments are removed before empty lines are handled if 'keep.comments=TRUE'.
keep.name	Keep the section name in output (say, "\$PROBLEM") Default is FALSE. It can only be FALSE, if return="text".
keep.comments	Default is to keep comments. If FALSE, they will be removed. See keep.empty too. Notice, there is no way for NMreadSection to keep comments and also drop lines that only contain comments.
as.one	If multiple hits, concatenate into one. This will most often be relevant with name="TABLE". If FALSE, a list will be returned, each element representing a table. Default is TRUE. So if you want to process the tables separately, you probably want FALSE here.
clean.spaces	If TRUE, leading and trailing are removed, and multiplied succeeding white spaces are reduced to single white spaces.
simplify	If asOne=FALSE, do you want the result to be simplified if only one section is found? Default is TRUE which is desirable for interactive analysis. For programming, you probably want FALSE.
keepEmpty	Deprecated. See keep.empty.
keepName	Deprecated. See keep.name.
keepComments	Deprecated. See keep.comments.
asOne	Deprecated. See as.one.
...	Additional arguments passed to NMextractText

**Value**

character vector with extracted lines.

**See Also**

Other Nonmem: [NMapplyFilters\(\)](#), [NMextractText\(\)](#), [NMgenText\(\)](#), [NMreplaceDataFile\(\)](#), [NMwriteSection\(\)](#)

**Examples**

```
NMreadSection(system.file("examples/nonmem/xgxr001.lst", package="NMdata"), section="DATA")
```

---

NMreadShk                      *Read Shrinkage data reported by Nonmem*

---

### Description

Read Shrinkage data reported by Nonmem

### Usage

```
NMreadShk(file, auto.ext, as.fun)
```

### Arguments

file	A model file. Extension will be replaced by ".shk".
auto.ext	If 'TRUE' (default) the extension will automatically be modified using 'NMdataConf()\$file.shk'. This means 'file' can be the path to an input or output control stream, and 'NMreadShk' will still read the '.shk' file.
as.fun	See ?NMdataConf

### Details

Type 1=etabar Type 2=Etabar SE Type 3=P val Type 4= Type 5= Type 6= Type 7=number of subjects used. Type 8= Type 9= Type 10= Type 11=

### Value

A 'data.frame' with shrinkage values, indexes, and name of related parameter, like 'OMEGA(1,1)'.

---

NMreadSizes                      *Read SIZES info from a control stream*

---

### Description

Read SIZES info from a control stream

### Usage

```
NMreadSizes(file.mod = NULL, lines = NULL)
```

### Arguments

file.mod	Control stream path.
lines	Character vector with control stream file.

### Value

A list with SIZES parameter values

---

NMreadTab	<i>Read an output table file from Nonmem</i>
-----------	--

---

### Description

Read a table generated by a \$TABLE statement in Nonmem. Generally, these files cannot be read by read.table or similar because formatting depends on options in the \$TABLE statement, and because Nonmem sometimes includes extra lines in the output that have to be filtered out. NMreadTab can do this automatically based on the table file alone.

### Usage

```
NMreadTab(
  file,
  col.tableno,
  col.nmrep,
  col.table.name,
  header = TRUE,
  skip,
  quiet = TRUE,
  as.fun,
  ...
)
```

### Arguments

file	path to Nonmem table file
col.tableno	In case of simulations where tables are being repeated, a counter of the repetition number can be useful to include in the output. For now, this will only work if the NOHEADER option is not used. This is because NMreadTab searches for the "TABLE NO..." strings in Nonmem output tables. If col.tableno is TRUE (default), a counter of tables is included as a column called NMREP. Notice, the table numbers in NMREP are cumulatively counting the number of tables reported in the file. NMREP is not the actual table number as given by Nonmem.
col.nmrep	col.nmrep If tables are repeated, include a counter? It does not relate to the order of the \$TABLE statements but to cases where a \$TABLE statement is run repeatedly. E.g., in combination with the SUBPROBLEMS feature in Nonmem, it is useful to keep track of the table (repetition) number. If col.nmrep is TRUE, this will be carried forward and added as a column called NMREP. This is default behavior when more than one \$TABLE repetition is found in data. Set it to a different string to request the column with a different name. The argument is passed to NMscanTables.
col.table.name	The name of a column containing the name or description of the table (generated by Nonmem). The default is "table.name". Use FALSE not to include this column.
header	Use header=FALSE if table was created with NOHEADER option in \$TABLE.

skip	The number of rows to skip. The default is skip=1 if header==TRUE and skip=0 if header==FALSE.
quiet	logical stating whether or not information is printed about what is being done. Default can be configured using NMdataConf.
as.fun	The default is to return data as a data.frame. Pass a function (say tibble::as_tibble) in as.fun to convert to something else. If data.tables are wanted, use as.fun="data.table". The default can be configured using NMdataConf.
...	Arguments passed to data.table::fread().

### Details

The actual reading of data is based on data.table::fread. Generally, the function is fast thanks to data.table.

### Value

The Nonmem table data.

### See Also

Other DataRead: [NMreadCsv\(\)](#), [NMscanData\(\)](#), [NMscanInput\(\)](#), [NMscanTables\(\)](#)

---

NMrelate	<i>Relate parameter names and variables based on control stream code sections.</i>
----------	--

---

### Description

Relate parameter names and variables based on control stream code sections.

### Usage

```
NMrelate(file, lines, modelname, par.type, col.model, sections, as.fun)
```

### Arguments

file	Path to a control stream to process. See 'lines' too.
lines	If the control stream has been read already, the text can be provided here instead of using the 'file' argument. Character vector of text lines.
modelname	Either a model name (like "Base") or a function that derives the model name from the control stream file path. The default is dropping the file name extension on the control stream file name.
par.type	Parameter type(s) to include. Default is all three possible which is c("THETA", "OMEGA", "SIGMA").
col.model	Name of the column containing the model name.
sections	Sections of the control stream to consider. Default is all of c("PRED", "PK", "ERROR").
as.fun	The default is to return data as a data.frame. Pass a function (say tibble::as_tibble) in as.fun to convert to something else. If data.tables are wanted, use as.fun="data.table". The default can be configured using NMdataConf.



**Details**

'NMrelate()' processes \$PRED, \$PK and \$ERROR sections. It does not read ext files or \$THETA, \$OMEGA, \$SIGMA sections to gain information but only extracts what it can from the model code. You can then merge with information from functions such as 'NMreadExt()' and 'NMread-ParText()'.

**Value**

data.frame relating parameters to variable names

---

NMreplaceDataFile	<i>Replace data file used in Nonmem control stream</i>
-------------------	--

---

**Description**

Replace data file used in Nonmem control stream

**Usage**

```
NMreplaceDataFile(files, file.pattern, dir, path.data, newfile = file.mod, ...)
```

**Arguments**

files	Paths to input control streams to modify. See file.pattern and dir too.
file.pattern	A pattern to look for if 'dir' is supplied too (and not 'file.mod'). This is used to modify multiple input control streams at once.
dir	Directory in which to look for 'file.pattern'. Notice, use either just 'file.mod' or both 'dir' and 'file.pattern'.
path.data	Path to input control stream to use in newfile
newfile	A path to a new control stream to write to (and don't edit contents of 'file.mod'). Default is to overwrite 'file.mod'.
...	Additional arguments to pass to NMwriteSection.

**Value**

Lines for a new control stream (invisibly)

**See Also**

Other Nonmem: [NMapplyFilters\(\)](#), [NMextractText\(\)](#), [NMgenText\(\)](#), [NMreadSection\(\)](#), [NMwriteSection\(\)](#)

---

**NMscanData***Automatically find Nonmem input and output tables and organize data*

---

**Description**

This is a very general solution to automatically identifying, reading, and merging all output and input data in a Nonmem model. The most important steps are

- Read and combine output tables,
- If wanted, read input data and restore variables that were not output from the Nonmem model
- If wanted, also restore rows from input data that were disregarded in Nonmem (e.g. observations or subjects that are not part of the analysis)

**Usage**

```
NMscanData(  
  file,  
  col.row,  
  use.input,  
  merge.by.row,  
  recover.rows,  
  file.mod,  
  dir.data,  
  file.data,  
  translate.input = TRUE,  
  quiet,  
  formats.read,  
  args.fread,  
  as.fun,  
  col.id = "ID",  
  modelname,  
  col.model,  
  col.nmout,  
  col.nmrep,  
  order.columns = TRUE,  
  check.time,  
  tz.lst,  
  skip.absent = FALSE,  
  tab.count,  
  use.rds  
)
```

**Arguments**

**file** Path to a Nonmem control stream or output file from Nonmem (.mod or .lst)

<code>col.row</code>	A column with a unique value for each row. Such a column is recommended to use if possible. See <code>merge.by.row</code> and details as well. Default ("ROW") can be modified using <code>NMdataConf</code> .
<code>use.input</code>	Should the input data be added to the output data. Only column names that are not found in output data will be retrieved from the input data. Default is TRUE which can be modified using <code>NMdataConf</code> . See <code>merge.by.row</code> too.
<code>merge.by.row</code>	If <code>use.input=TRUE</code> , this argument determines the method by which the input data is added to output data. The default method ( <code>merge.by.row=FALSE</code> ) is to interpret the Nonmem code to imitate the data filtering (IGNORE and ACCEPT statements), but the recommended method is <code>merge.by.row=TRUE</code> which means that data will be merged by a unique row identifier. The row identifier must be present in input and at least one full length output data table. See argument <code>col.row</code> too.
<code>recover.rows</code>	Include rows from input data files that do not exist in output tables? This will be added to the <code>\$row</code> dataset only, and <code>\$run</code> , <code>\$id</code> , and <code>\$occ</code> datasets are created before this is taken into account. A column called <code>nmout</code> will be TRUE when the row was found in output tables, and FALSE when not. Default is FALSE and can be configured using <code>NMdataConf</code> .
<code>file.mod</code>	The input control stream file path. Default is to look for <code>"file\"</code> with extension changed to <code>.mod</code> (PSN style). You can also supply the path to the file, or you can provide a function that translates the output file path to the input file path. The default behavior can be configured using <code>NMdataConf</code> . See <code>dir.data</code> too.
<code>dir.data</code>	The data directory can only be read from the control stream ( <code>.mod</code> ) and not from the output file ( <code>.lst</code> ). So if you only have the output control stream, use <code>dir.data</code> to tell in which directory to find the data file. If <code>dir.data</code> is provided, the <code>.mod</code> file is not used at all.
<code>file.data</code>	Specification of the data file path. When this is used, the control streams are not used at all.
<code>translate.input</code>	Default is TRUE, meaning that input data column names are translated according to <code>\$INPUT</code> section in Nonmem listing file.
<code>quiet</code>	The default is to give some information along the way on what data is found. But consider setting this to TRUE for non-interactive use. Default can be configured using <code>NMdataConf</code> .
<code>formats.read</code>	Prioritized input data file formats to look for and use if found. Default is <code>c("rds","csv")</code> which means <code>rds</code> will be used if found, and <code>csv</code> if not. <code>fst</code> is possible too. Default can be modified using <code>NMdataConf()</code> .
<code>args.fread</code>	List of arguments passed to when reading <code>_input_</code> data. Notice that except for "input" and "file", you need to supply all arguments to <code>fread</code> if you use this argument. Default values can be configured using <code>NMdataConf</code> .
<code>as.fun</code>	The default is to return data as a <code>data.frame</code> . Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If <code>data.tables</code> are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .
<code>col.id</code>	The name of the subject ID variable, default is "ID".

<code>modelName</code>	The model name to be stored if <code>col.model</code> is not NULL. If not supplied, the name will be taken from the control stream file name by omitting the directory/path and deleting the <code>.lst</code> extension ( <code>path/run001.lst</code> becomes <code>run001</code> ). This can be a character string or a function which is called on the value of <code>file</code> ( <code>file</code> is another argument to <code>NMscanData</code> ). The function must take one character argument and return another character string. As example, see <code>NMdataConf()\$modelName</code> . The default can be configured using <code>NMdataConf</code> .
<code>col.model</code>	A column of this name containing the model name will be included in the returned data. The default is to store this in a column called "model". See argument "modelName" as well. Set to NULL if not wanted. Default can be configured using <code>NMdataConf</code> .
<code>col.nmout</code>	A column of this name will be a logical representing whether row was in output table or not. Default can be modified using <code>NMdataConf</code> .
<code>col.nmrep</code>	If tables are repeated, include a counter? It does not relate to the order of the <code>\$TABLE</code> statements but to cases where a <code>\$TABLE</code> statement is run repeatedly. E.g., in combination with the <code>SUBPROBLEMS</code> feature in <code>Nonmem</code> , it is useful to keep track of the table (repetition) number. If <code>col.nmrep</code> is <code>TRUE</code> , this will be carried forward and added as a column called <code>NMREP</code> . This is default behavior when more than one <code>\$TABLE</code> repetition is found in data. Set it to a different string to request the column with a different name. The argument is passed to <code>NMscanTables</code> .
<code>order.columns</code>	If <code>TRUE</code> (default), <code>NMorderColumns</code> is used to reorder the columns before returning the data. <code>NMorderColumns</code> will be called with <code>alpha=FALSE</code> , so columns are not sorted alphabetically. But standard <code>Nonmem</code> columns like <code>ID</code> , <code>TIME</code> , and other will be first. If <code>col.row</code> is used, this will be passed to <code>NMorderColumns</code> too.
<code>check.time</code>	If <code>TRUE</code> (default) and if input data is used, input control stream and input data are checked to be newer than output control stream and output tables. These are important assumptions for the way information is merged by <code>NMscanData</code> . However, if data has been transferred from another system where <code>Nonmem</code> was run, these checks may not make sense, and you may not want to see these warnings. The default can be configured using <code>NMdataConf</code> . For the output control stream, the time stamp recorded by <code>Nonmem</code> is used if possible, and if the input data is created with <code>NMwriteData</code> , the recorded creation time is used if possible. If not, and for all other files, the file modification times are used.
<code>tz.lst</code>	If supplied, the timezone to be used when reading the time stamp in the output control stream. Please supply something listed in <code>OlsonNames()</code> . Can be configured using <code>NMdataConf()</code> too.
<code>skip.absent</code>	Skip missing output table files with a warning? Default is <code>FALSE</code> in which case an error is thrown.
<code>tab.count</code>	Deprecated. Use <code>col.tableno</code> .
<code>use.rds</code>	Deprecated - use <code>formats.read</code> instead. If provided (though not recommended), this will overwrite <code>formats.read</code> , and only <code>formats.rds</code> and <code>csv</code> can be used.

## Details

This function makes it very easy to collect the data from a `Nonmem` run.

A useful feature of this function is that it can automatically combine "input" data (the data read by Nonmem in \$INPUT or \$INFILE) with "output" data (tables written by Nonmem in \$TABLE). There are two implemented methods for doing so. One (the default but not recommended) relies on interpretation of filter (IGNORE and ACCEPT) statements in \$INPUT. This will work in most cases, and checks for consistency with Nonmem results. However, the recommended method is using a unique row identifier in both input data and at least one output data file (not a FIRSTONLY or LASTONLY table). Supply the name of this column using the col.row argument.

Limitations. A number of Nonmem features are not supported. Most of this can be overcome by using merge.by.row=TRUE. Incomplete list of known limitations:

**character TIME** If Nonmem is used to translate DAY and a character TIME column, TIME has to be available in an output table. NMscanData does not do the translation to numeric.

**RECORDS** The RECORDS option to limit the part of the input data being used is not searched for. Using merge.by.row=TRUE will work unaffectedly.

**NULL** The NULL argument to specify missing value string in input data is not respected. If delimited input data is read (as opposed to rds files), missing values are assumed to be represented by dots (.).

### Value

A data set of class 'NMdata'.

### See Also

Other DataRead: [NMreadCsv\(\)](#), [NMreadTab\(\)](#), [NMscanInput\(\)](#), [NMscanTables\(\)](#)

### Examples

```
## Not run:
res1 <- NMscanData(system.file("examples/nonmem/xgxr001.lst", package="NMdata"))

## End(Not run)
```

---

NMscanInput

*Find and read input data and optionally translate column names according to the \$INPUT section*

---

### Description

This function finds and reads the input data based on a control stream file path. It can align the column names to the definitions in \$INPUT in the control stream, and it can subset the data based on ACCEPT/IGNORE statements in \$DATA. It supports a few other ways to identify the input data file than reading the control stream, and it can also read an rds or fst file instead of the delimited text file used by Nonmem.

**Usage**

```

NMscanInput(
  file,
  formats.read,
  file.mod,
  dir.data = NULL,
  file.data = NULL,
  apply.filters = FALSE,
  translate = TRUE,
  recover.cols = TRUE,
  details = TRUE,
  col.id = "ID",
  col.row,
  quiet,
  args.fread,
  invert = FALSE,
  as.fun,
  applyFilters,
  use.rds
)

```

**Arguments**

<code>file</code>	a .lst (output) or a .mod (input) control stream file. The filename does not need to end in .lst. It is recommended to use the output control stream because it reflects the model as it was run rather than how it is planned for next run. However, see <code>file.mod</code> and <code>dir.data</code> .
<code>formats.read</code>	Prioritized input data file formats to look for and use if found. Default is <code>c("rds", "csv")</code> which means <code>rds</code> will be used if found, and <code>csv</code> if not. <code>fst</code> is possible too. Default can be modified using <code>NMdataConf()</code> .
<code>file.mod</code>	The input control stream file path. Default is to look for <code>"file\"</code> with extension changed to <code>.mod</code> (PSN style). You can also supply the path to the file, or you can provide a function that translates the output file path to the input file path. If <code>dir.data</code> is missing, the input control stream is needed. This is because the .lst does not contain the path to the data file. The .mod file is only used for finding the data file. How to interpret the datafile is read from the .lst file. The default can be configured using <code>NMdataConf</code> . See <code>dir.data</code> too.
<code>dir.data</code>	The data directory can only be read from the control stream (.mod) and not from the output file (.lst). So if you only have the output file, use <code>dir.data</code> to tell in which directory to find the data file. If <code>dir.data</code> is provided, the .mod file is not used at all.
<code>file.data</code>	Specification of the data file path. When this is used, the control streams are not used at all.
<code>apply.filters</code>	If <code>TRUE</code> (default), <code>IGNORE</code> and <code>ACCEPT</code> statements in the Nonmem control streams are applied before returning the data. This affects what rows are returned, not columns.

<code>translate</code>	If TRUE (default), data columns are named as interpreted by Nonmem (in ‘\$INPUT’).
<code>recover.cols</code>	recover columns that were not used in the Nonmem control stream? This means adding column from the input data file that are not used in ‘\$INPUT’. If data file contains more columns than mentioned in ‘\$INPUT’, these will be named as in data file (if data file contains named variables). This affects what columns are returned, not rows.
<code>details</code>	If TRUE, metadata is added to output. In this case, you get a list. Typically, this is mostly useful if programming up functions which behavior must depend on properties of the output. See details.
<code>col.id</code>	The name of the subject ID column. Optional and only used to calculate number of subjects in data. Default is modified by NMdataConf.
<code>col.row</code>	The name of the row counter column. Optional and only used to check whether the row counter is in the data.
<code>quiet</code>	Default is to inform a little, but TRUE is useful for non-interactive stuff.
<code>args.fread</code>	List of arguments passed to fread. Notice that except for "input" and "file", you need to supply all arguments to fread if you use this argument. Default values can be configured using ‘NMdataConf()’.
<code>invert</code>	If TRUE, the data rows that are dismissed by the Nonmem data filters (ACCEPT and IGNORE) and only this will be returned. Only used if ‘apply.filters’ is ‘TRUE’.
<code>as.fun</code>	The default is to return data as a data.frame. Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If data.tables are wanted, use <code>as.fun="data.table"</code> . The default can be configured using NMdataConf.
<code>applyFilters</code>	Deprecated - use <code>apply.filters</code> .
<code>use.rds</code>	Deprecated - use <code>formats.read</code> instead. If provided (though not recommended), this will overwrite <code>formats.read</code> , and only formats rds and csv can be used.

## Details

Columns that are dropped (using ‘DROP’ or ‘SKIP’ in ‘\$INPUT’) in the model will be included in the output.

It may not work if a column is dropped, and a new column is renamed to the same name. Say you have DV and CONC as the only two columns (not possible but illustrative), and in Nonmem you do DV=DROP DV. Not sure it will work in Nonmem, and it probably won’t work in NMscanInput.

## Value

A data set, class defined by ‘`as.fun`’

## See Also

Other DataRead: [NMreadCsv\(\)](#), [NMreadTab\(\)](#), [NMscanData\(\)](#), [NMscanTables\(\)](#)

---

 NMscanMultiple

*Run NMscanData on multiple models and stack results*


---

## Description

Useful function for meta analyses when multiple models are stored in one folder and can be read with NMscanData using the same arguments.

## Usage

```
NMscanMultiple(files, dir, file.pattern, as.fun, ...)
```

## Arguments

files	File paths to the models (control stream) to edit. See file.pattern too.
dir	The directory in which to find the models. Passed to list.files(). See file.pattern argument too.
file.pattern	The pattern used to match the filenames to read with NMscanData. Passed to list.files(). If dir is supplied and files is not (or is NULL), the default is ".*\..lst" which means all files ending in '.lst'. See dir argument too.
as.fun	The default is to return data as a data.frame. Pass a function (say tibble::as_tibble) in as.fun to convert to something else. If data.tables are wanted, use as.fun="data.table". The default can be configured using NMdataConf.
...	Additional arguments passed to NMscanData.

## Value

All results stacked, class as defined by as.fun

## Examples

```
## Not run:
res <- NMscanMultiple(dir=system.file("examples/nonmem", package="NMdata"),
  file.pattern="xgxr01.*\\.lst", as.fun="data.table")
res.mean <- res[,.(meanPRED=exp(mean(log(PRED))), by=. (model, NOMTIME)]
library(ggplot2)
ggplot(res.mean, aes(NOMTIME, meanPRED, colour=model))+geom_line()

## End(Not run)
```



---

 NMscanTables

*Find and read all output data tables in Nonmem run*


---

## Description

Find and read all output data tables in Nonmem run

## Usage

```
NMscanTables(
  file,
  as.fun,
  quiet,
  col.nmrep = TRUE,
  col.tableno = FALSE,
  col.id = "ID",
  col.row,
  details,
  skip.absent = FALSE,
  meta.only = FALSE,
  modelname,
  col.model
)
```

## Arguments

file	the Nonmem file to read (normally .mod or .lst)
as.fun	The default is to return data as a data.frame. Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If data.tables are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .
quiet	The default is to give some information along the way on what data is found. But consider setting this to <code>TRUE</code> for non-interactive use. Default can be configured using <code>NMdataConf</code> .
col.nmrep	<code>col.nmrep</code> If tables are repeated, include a counter? It does not relate to the order of the <code>\$TABLE</code> statements but to cases where a <code>\$TABLE</code> statement is run repeatedly. E.g., in combination with the <code>SUBPROBLEMS</code> feature in Nonmem, it is useful to keep track of the table (repetition) number. If <code>col.nmrep</code> is <code>TRUE</code> , this will be carried forward and added as a column called <code>NMREP</code> . This is default behavior when more than one <code>\$TABLE</code> repetition is found in data. Set it to a different string to request the column with a different name. The argument is passed to <code>NMscanTables</code> .
col.tableno	Nonmem includes a counter of tables in the written data files. These are often not useful. However, if <code>col.tableno</code> is <code>TRUE</code> (not default), this will be carried forward and added as a column called <code>NMREP</code> . Even if <code>NMREP</code> is generated by <code>NMscanTables</code> , it is treated like any other table column in meta ( <code>?NMInfo</code> ) data.

<code>col.id</code>	name of the subject ID column. Used for calculation of the number of subjects in each table.
<code>col.row</code>	The name of the row counter column. Optional and only used to check whether the row counter is in the data.
<code>details</code>	If TRUE, metadata is added to output. In this case, you get a list. Typically, this is mostly useful if programming up functions which behavior must depend on properties of the output.
<code>skip.absent</code>	Skip missing output table files with a warning? Default is FALSE in which case an error is thrown.
<code>meta.only</code>	If TRUE, tables are not read, only a table is returned showing what tables were found and some available meta information. Notice, not all meta information (e.g., dimensions) are available because the tables need to be read to derive that.
<code>modelName</code>	Only affects meta data table. The model name to be stored if <code>col.model</code> is not NULL. If not supplied, the name will be taken from the control stream file name by omitting the directory/path and deleting the <code>.lst</code> extension (path/run001.lst becomes run001). This can be a character string or a function which is called on the value of file (file is another argument to <code>NMscanData</code> ). The function must take one character argument and return another character string. As example, see <code>NMdataConf()\$modelName</code> . The default can be configured using <code>NMdataConf</code> .
<code>col.model</code>	Only affects meta data table. A column of this name containing the model name will be included in the returned data. The default is to store this in a column called "model". See argument "modelName" as well. Set to NULL if not wanted. Default can be configured using <code>NMdataConf</code> .

### Value

A list of all the tables as data.frames. If `details=TRUE`, this is in one element, called `data`, and `meta` is another element. If not, only the data is returned.

### See Also

Other DataRead: [NMreadCsv\(\)](#), [NMreadTab\(\)](#), [NMscanData\(\)](#), [NMscanInput\(\)](#)

### Examples

```
tabs1 <- NMscanTables(system.file("examples/nonmem/xgxr001.lst", package="NMdata"))
```

---

NMstamp

*stamp a dataset or any other object*

---

### Description

Dataset metadata can be valuable, eg. by tracing an archived dataset back to the code that generated it. The metadata added by `NMstamp` can be accessed using the function `NMinfo`.

**Usage**

```
NMstamp(data, script, time = Sys.time(), ...)
```

**Arguments**

data	The dataset to stamp.
script	path to the script where the dataset was generated.
time	the time stamp to attach. Default is to use cpu clock.
...	other named metadata elements to add to the dataset. Example: Description="PK data for phase 1 trials in project".

**Details**

NMstamp modifies the meta data by reference. See example.

**Value**

data with meta data attached. Class unchanged.

**See Also**

NMinfo

Other DataCreate: [NMorderColumns\(\)](#), [NMwriteData\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

**Examples**

```
x=1
NMstamp(x,script="example.R",description="Example data")
NMinfo(x)
```

---

NMwriteData

*Write dataset for use in Nonmem (and R)*

---

**Description**

Instead of trying to remember the arguments to pass to write.csv, use this wrapper. It tells you what to write in \$DATA and \$INPUT in Nonmem, and it (additionally) exports an rds file as well which is highly preferable for use in R. It never edits the data before writing the datafile. The filenames for csv, rds etc. are derived by replacing the extension to the filename given in the file argument.

**Usage**

```

NMwriteData(
  data,
  file,
  formats.write = c("csv", "rds"),
  script,
  args.stamp,
  args.fwrite,
  args.rds,
  args.RData,
  args.write_fst,
  quiet,
  args.NMgenText,
  csv.trunc.as.nm = FALSE,
  genText,
  save = TRUE,
  write.csv,
  write.rds,
  write.RData,
  nm.drop,
  nm.dir.data,
  col.flagn,
  nm.rename,
  nm.copy,
  nm.capitalize,
  allow.char.TIME
)

```

**Arguments**

<code>data</code>	The dataset to write to file for use in Nonmem.
<code>file</code>	The file to write to. The extension (everything after and including last ".") is dropped. csv, rds and other standard file name extensions are added.
<code>formats.write</code>	character vector of formats.write. Default is c("csv","rds"). "fst" is possible too. Default can be modified with <code>NMdataConf()</code> .
<code>script</code>	If provided, the object will be stamped with this script name before saved to rds or RData. See <code>?NMstamp</code> .
<code>args.stamp</code>	A list of arguments to be passed to <code>NMstamp</code> .
<code>args.fwrite</code>	List of arguments passed to <code>fwrite</code> . Notice that except for "x" and "file", you need to supply all arguments to <code>fwrite</code> if you use this argument. Default values can be configured using <code>NMdataConf</code> .
<code>args.rds</code>	A list of arguments to be passed to <code>saveRDS</code> .
<code>args.RData</code>	A list of arguments to be passed to <code>save</code> . Please note that writing RData is deprecated.
<code>args.write_fst</code>	An optional list of arguments to be passed to <code>write_fst</code> .

<code>quiet</code>	The default is to give some information along the way on what data is found. But consider setting this to TRUE for non-interactive use. Default can be configured using <code>NMdataConf</code> .
<code>args.NMgenText</code>	List of arguments to pass to <code>NMgenText</code> - the function that generates text suggestion for INPUT and DATA sections in the Nonmem control stream. You can use these arguments to get a text suggestion you can use directly in Nonmem - and <code>NMwriteSection</code> can even update multiple Nonmem control streams based on the result. This will update your control streams to match your new data file with just one command.
<code>csv.trunc.as.nm</code>	If TRUE, csv file will be truncated horizontally (columns will be dropped) to match the \$INPUT text generated for Nonmem ( <code>genText</code> must be TRUE for this option to be allowed). This can be a great advantage when dealing with large datasets that can create problems in parallelization. Combined with <code>write.rds=TRUE</code> , the full data set will still be written to an rds file, so this can be used when combining output and input data when reading model results. This is done by default by <code>NMscanData</code> . This means writing a lean (narrow) csv file for Nonmem while keeping columns of non-numeric class like character and factor for post-processing.
<code>genText</code>	Run and report results of <code>NMgenText</code> ? Default is 'TRUE' if a csv file is written, otherwise 'FALSE'. You may want to disable this if data set is not for Nonmem.
<code>save</code>	Save defined files? Default is TRUE. If a variable is used to control whether a script generates outputs (say <code>writeOutputs=TRUE/FALSE</code> ), if you use <code>save=writeOutputs</code> to comply with this.
<code>write.csv</code>	Write to csv file? Deprecated, use 'formats.write' instead.
<code>write.rds</code>	write an rds file? Deprecated, use 'formats.write' instead.
<code>write.RData</code>	Deprecated and not recommended - will be removed. RData is not an adequate format for a dataset (but is for environments). Please use <code>write.rds</code> instead.
<code>nm.drop</code>	Deprecated, use <code>args.NMgenText=list(drop=c("column"))</code> instead.
<code>nmdir.data</code>	Deprecated, use <code>args.NMgenText=list(dir.data="your/path")</code> instead.
<code>col.flagn</code>	Deprecated, use <code>args.NMgenText=list(col.flagn="column.name")</code> . Name of a numeric column with zero value for rows to include in Nonmem run, non-zero for rows to skip. The argument is only used for generating the proposed \$DATA text to paste into the Nonmem control stream. To skip this feature, use 'col.flagn=FALSE'.
<code>nm.rename</code>	Deprecated, use <code>args.NMgenText=list(rename=c(newname="existing"))</code> instead.
<code>nm.copy</code>	Deprecated, use <code>args.NMgenText=list(copy=c(newname="existing"))</code> instead.
<code>nm.capitalize</code>	Deprecated, use <code>args.NMgenText=list(capitalize=TRUE)</code> instead.
<code>allow.char.TIME</code>	Deprecated, use <code>args.NMgenText=list(allow.char.TIME=TRUE)</code> instead.

## Details

When writing csv files, the file will be comma-separated. Because Nonmem does not support quoted fields, you must avoid commas in character fields. An error is returned if commas are found in strings.

The user is provided with text to use in Nonmem. This lists names of the data columns. Once a column is reached that Nonmem will not be able to read as a numeric and column is not in nm.drop, the list is stopped. Only exception is TIME which is not tested for whether character or not.

### Value

Text for inclusion in Nonmem control stream, invisibly.

### See Also

Other DataCreate: [NMorderColumns\(\)](#), [NMstamp\(\)](#), [addTAPD\(\)](#), [findCovs\(\)](#), [findVars\(\)](#), [flagsAssign\(\)](#), [flagsCount\(\)](#), [mergeCheck\(\)](#), [tmpcol\(\)](#)

---

NMwriteFilters

*Write IGNORE/ACCEPT filters to NONMEM model*

---

### Description

Write IGNORE/ACCEPT filters to NONMEM model

### Usage

```
NMwriteFilters(file = NULL, lines = NULL, filters, write)
```

### Arguments

file	Path to control stream. Use 'file' or 'lines'.
lines	Control stream as text lines. Use 'file' or 'lines'.
filters	A data frame with filters, like returned by 'NMreadFilters()'.
write	If 'file' is provided, write the results to file? If 'lines' is used, 'write' cannot be used.

### Value

Resulting control stream (lines) as character vector

---

NMwriteInits	<i>Writes a parameter values to a control stream</i>
--------------	--

---

**Description**

Edit parameter values, fix/unfix them, or edit lower/upper bounds.

**Usage**

```

NMwriteInits(
    file.mod,
    update = TRUE,
    file.ext = NULL,
    ext,
    values,
    newfile,
    ...
)

```

**Arguments**

file.mod	Path to control stream.
update	If 'TRUE' (default), the parameter values are updated based on the '.ext' file.
file.ext	Optionally provide the path to an '.ext' file. If not provided, the default is to replace the file name extension on 'file.mod' with '.ext'. This is only used if 'update=TRUE'.
ext	Not implemented.
values	A list of lists. Each list specifies a parameter with named elements. Must be named by the parameter name. 'lower', 'upper' and 'fix' can be supplied to modify the parameter. See examples. Notice, you can use '...' instead. 'values' may be easier for programming but other than that, most users will find '...' more intuitive.
newfile	If provided, the results are written to this file as a new input control stream.
...	Parameter specifications. See examples,

**Details**

Limitations:

- lower, init, and upper must be on same line
- If using something like CL=(.1,4,15), two of those cannot be on the same line

**Value**

a control stream as lines in a character vector.

**Examples**

```

## Requires NMdata 0.1.9
## Not run:
file.mod <- system.file("examples/nonmem/xgxr021.mod", package="NMsim")
NMwriteInits(file.mod,
values=list( "theta(2)"=list(init=1.4),
            "THETA(3)"=list(FIX=1),
            "omega(2,2)"=list(init=0.1))
)
NMwriteInits(file.mod,
            "theta(2)"=list(init=1.4),
            "THETA(3)"=list(FIX=1),
            "omega(2,2)"=list(init=0.1)
)

## End(Not run)

```

---

NMwriteSection

*Replace (\$)sections of a Nonmem control stream*


---

**Description**

Just give the section name, the new lines and the file path, and the "\$section", and the input to Nonmem will be updated.

**Usage**

```

NMwriteSection(
  files,
  file.pattern,
  dir,
  section,
  newlines,
  list.sections,
  location = "replace",
  newfile,
  backup = TRUE,
  blank.append = TRUE,
  data.file,
  write = TRUE,
  quiet,
  simplify = TRUE
)

```

**Arguments**

files                    File paths to the models (control stream) to edit. See file.pattern too.



file.pattern	Alternatively to files, you can supply a regular expression which will be passed to list.files as the pattern argument. If this is used, use 'dir' argument as well. Also see data.file to only process models that use a specific data file.
dir	If file.pattern is used, 'dir' is the directory to search in.
section	The name of the section to update with or without "\$". Example: 'section="EST"' or 'section="\$EST"' to edit the sections starting by '\$EST'. Section specification is not case-sensitive. See '?NMreadSection' too.
newlines	The new text (including "\$SECTION"). Better be broken into lines in a character vector since this is simply past to writeLines().
list.sections	Named list of new sections, each element containing a section. Names must be section names, contents of each element are the new section lines for each section.
location	In combination with 'section', this determines where the new section is inserted. Possible values are "replace" (default), "before", "after", "first", "last".
newfile	path and filename to new run. If missing, the original file (from files or file.pattern) is overwritten (see the backup option below). If NULL, output is returned as a character vector rather than written.
backup	In case you are overwriting the old file, do you want to backup the file (to say, backup_run001.mod)?
blank.append	Append a blank line to output?
data.file	Use this to limit the scope of models to those that use a specific input data file. The string has to exactly match the one in '\$DATA' or '\$INFILE' in Nonmem.
write	Default is to write to file. If write=FALSE, 'NMwriteSection()' returns the resulting input.txt without writing it to disk. Default is 'TRUE'.
quiet	The default is to give some information along the way on what data is found. But consider setting this to TRUE for non-interactive use. Default can be configured using 'NMdataConf()'.
simplify	If TRUE (default) and only one file is edited, the resulting rows are returned directly. If more than one file is edited, the result will always be a list with one element per file.

### Details

The new file will be written with unix-style line endings.

### Value

The new section text is returned. If write=TRUE, this is done invisibly.

### See Also

Other Nonmem: [NMapplyFilters\(\)](#), [NMextractText\(\)](#), [NMgenText\(\)](#), [NMreadSection\(\)](#), [NMreplaceDataFile\(\)](#)

**Examples**

```

newlines <- "$EST POSTHOC INTERACTION METHOD=1 NOABORT PRINT=5 MAXEVAL=9999 SIG=3"
NMwriteSection(files=system.file("examples/nonmem/xgxr001.mod", package = "NMdata"),
section="EST", newlines=newlines,newfile=NULL)
## Not run:
text.nm <- NMwriteData(data)
NMwriteSection(dir="nonmem",
               file.pattern="^run.*\\.mod",
               list.sections=text.nm["INPUT"])

## End(Not run)

```

---

NMwriteSizes

*Create or update \$SIZES in a control stream*


---

**Description**

Update \$SIZES parameters in a control stream. The control stream can be in a file or provided as a character vector (file lines).

**Usage**

```

NMwriteSizes(
  file.mod = NULL,
  newfile,
  lines = NULL,
  wipe = FALSE,
  write = !is.null(newfile),
  ...
)

```

**Arguments**

file.mod	A path to a control stream. See also alternative ‘lines’ argument. Notice, if ‘write’ is ‘TRUE’ (default) and ‘newfile’ is not provided, ‘file.mod’ will be overwritten.
newfile	An optional path to write the resulting control stream to. If nothing is provided, the default is to overwrite ‘file.mod’.
lines	Control stream lines as a character vector. If you already read the control stream - say using ‘NMdata::NMreadSection()’, use this to modify the text lines.
wipe	The default behavior (‘wipe=FALSE’) is to add the ‘\$SIZES’ values to any existing values found. If SIZES parameter names are overlapping with existing, the values will be updated. If ‘wipe=TRUE’, any existing ‘\$SIZES’ section is disregarded.
write	Write results to ‘newfile’?
...	The \$SIZES parameters. Provided anything, like ‘PD=40’ See examples.

**Value**

Character lines with updated control stream

**Examples**

```
## No existing SIZES in control stream
## Not run:
file.mod <- system.file("examples/nonmem/xgxr132.mod", package="NMdata")
newmod <- NMwriteSizes(file.mod, LTV=50, write=FALSE)
head(newmod)

## End(Not run)
## provide control stream as text lines
## Not run:
file.mod <- system.file("examples/nonmem/xgxr032.mod", package="NMdata")
lines <- readLines(file.mod)
newmod <- NMwriteSizes(lines=lines, LTV=50, write=FALSE)
head(newmod)

## End(Not run)
## By default (wipe=FALSE) variabls are added to SIZES
## Not run:
lines.mod <- NMwriteSizes(file.mod, LTV=50, write=FALSE)
newmod <- NMwriteSizes(lines=lines.mod, PD=51, write=FALSE)
head(newmod)

## End(Not run)
```

---

print.summary\_NMdata *print method for NMdata summaries*

---

**Description**

print method for NMdata summaries

**Usage**

```
## S3 method for class 'summary_NMdata'
print(x, ...)
```

**Arguments**

x                   The summary object to be printed. See ?summary.NMdata  
 ...                 Arguments passed to other print methods.

**Value**

NULL (invisibly)

---

renameByContents	<i>Rename columns matching properties of data contents</i>
------------------	--

---

### Description

For instance, lowercase all columns that Nonmem cannot interpret (as numeric).

### Usage

```
renameByContents(data, fun.test, fun.rename, invert.test = FALSE, as.fun)
```

### Arguments

<code>data</code>	data.frame in which to rename columns
<code>fun.test</code>	Function that returns TRUE for columns to be renamed.
<code>fun.rename</code>	Function that takes the existing column name and returns the new one.
<code>invert.test</code>	Rename those where FALSE is returned from <code>fun.test</code> .
<code>as.fun</code>	The default is to return data as a data.frame. Pass a function (say <code>tibble::as_tibble</code> ) in <code>as.fun</code> to convert to something else. If data.tables are wanted, use <code>as.fun="data.table"</code> . The default can be configured using <code>NMdataConf</code> .

### Value

data with (some) new column names. Class as defined by `as.fun`.

### Examples

```
pk <- readRDS(file=system.file("examples/data/xgxr2.rds", package="NMdata"))
pk[,trtact:=NULL]
pk <- renameByContents(data=pk,
                       fun.test = NMisNumeric,
                       fun.rename = tolower,
                       invert.test = TRUE)
## Or append a "C" to the same column names
pk <- readRDS(file=system.file("examples/data/xgxr2.rds", package="NMdata"))
pk[,trtact:=NULL]
pk <- renameByContents(data=pk,
                       fun.test = NMisNumeric,
                       fun.rename = function(x)paste0(x, "C"),
                       invert.test = TRUE)
```

---

summary.NMdata	<i>summary method for NMdata objects</i>
----------------	--

---

**Description**

summary method for NMdata objects

**Usage**

```
## S3 method for class 'NMdata'
summary(object, ...)
```

**Arguments**

object	An NMdata object (from NMscanData).
...	Only passed to the summary generic if object is missing NMdata meta data (this should not happen anyway).

**Details**

The subjects are counted conditioned on the nmout column. If only id-level output tables are present, there are no nmout=TRUE rows. This means that in this case it will report that no IDs are found in output. The correct statement is that records are found for zero subjects in output tables.

**Value**

A list with summary information on the NMdata object.

---

triagSize	<i>Calculate number of elements for matrix specification</i>
-----------	--

---

**Description**

calculate number of elements in the diagonal and lower triangle of a squared matrix, based on the length of the diagonal.

**Usage**

```
triagSize(diagSize)
```

**Arguments**

diagSize	The length of the diagonal. Same as number of rows or columns.
----------	--

**Value**

An integer

**Examples**

```
triagSize(1:5)
```

---

uniquePresent      *Extract unique non-missing value from vector*

---

**Description**

Extract unique non-missing value from vector

**Usage**

```
uniquePresent(x, req.n1 = TRUE, na.pattern)
```

**Arguments**

x	A vector, either numeric or character.
req.n1	Require one unique value? If 'TRUE' (default), an error is thrown if non-unique values found. If 'FALSE', all the unique values are returned.
na.pattern	In addition to NA-elements, what text strings should be considered missing? Default is empty strings and strings only containing white spaces ('na.pattern="'^*\$").

**Details**

This function is particularly useful when combining data sets of which only some contain certain variables. `uniquePresent` with `'req.n1=TRUE'` makes sure the result is a single unique value (e.g., within subjects). A typical use is carrying subject-level covariates from one data set to another in a longitudinal analysis.

**Value**

a vector of same class as 'x'

---

unNMdata	<i>Remove NMdata class and discard NMdata meta data</i>
----------	---

---

**Description**

Remove NMdata class and discard NMdata meta data

**Usage**

```
unNMdata(x)
```

**Arguments**

x                    An 'NMdata' object.

**Value**

x stripped from the 'NMdata' class

# Index

- \* **DataCreate**
  - addTAPD, 4
  - findCovs, 14
  - findVars, 15
  - flagsAssign, 16
  - flagsCount, 18
  - mergeCheck, 24
  - NMorderColumns, 42
  - NMstamp, 66
  - NMwriteData, 67
- \* **DataRead**
  - NMreadCsv, 45
  - NMreadTab, 55
  - NMscanData, 58
  - NMscanInput, 61
  - NMscanTables, 65
- \* **DataWrangling**
  - compareCols, 8
  - dims, 10
  - listMissings, 22
- \* **Nonmem**
  - NMextractText, 37
  - NMgenText, 39
  - NMreadSection, 52
  - NMreplaceDataFile, 57
  - NMwriteSection, 72
- \* **debug**
  - NMcheckColnames, 27
- addCor, 3
- addOmegaCorr, 4
- addTAPD, 4, 14, 16, 17, 20, 26, 44, 67, 70
- cbind.NMdata (NMdataOperations), 34
- cc, 6
- cl, 7
- colLabels, 8
- compareCols, 8, 10, 23
- dimnames.NMdata (NMdataOperations), 34
- dims, 10, 10, 23
- dt2mat, 11
- editCharCols, 12
- egdt, 13
- findCovs, 6, 14, 16, 17, 20, 26, 44, 67, 70
- findVars, 6, 14, 15, 17, 20, 26, 44, 67, 70
- flagsAssign, 6, 14, 16, 16, 20, 26, 44, 67, 70
- flagsCount, 6, 14, 16, 17, 18, 26, 44, 67, 70
- fnAppend, 20
- fnExtension, 21
- is.NMdata, 22
- listMissings, 10, 22
- mat2dt, 23
- merge.NMdata (NMdataOperations), 34
- mergeCheck, 6, 14, 16, 17, 20, 24, 44, 67, 70
- NMapplyFilters, 38, 40, 53, 57, 73
- NMcheckColnames, 27
- NMcheckData, 27
- NMdataConf, 31
- NMdataOperations, 34
- NMexpandDoses, 35
- NMextractDataFile, 36
- NMextractText, 37, 40, 53, 57, 73
- NMgenText, 38, 39, 53, 57, 73
- NMinfo, 40
- NMisNumeric, 41
- NMorderColumns, 6, 14, 16, 17, 20, 26, 42, 67, 70
- NMreadCov, 44
- NMreadCsv, 45, 56, 61, 63, 66
- NMreadExt, 46
- NMreadFilters, 47
- NMreadInits, 48
- NMreadParsText, 48
- NMreadPhi, 51



NMreadSection, [38](#), [40](#), [52](#), [57](#), [73](#)  
NMreadShk, [54](#)  
NMreadSizes, [54](#)  
NMreadTab, [45](#), [55](#), [61](#), [63](#), [66](#)  
NMrelate, [56](#)  
NMreplaceDataFile, [38](#), [40](#), [53](#), [57](#), [73](#)  
NMscanData, [45](#), [56](#), [58](#), [63](#), [66](#)  
NMscanInput, [45](#), [56](#), [61](#), [61](#), [66](#)  
NMscanMultiple, [64](#)  
NMscanTables, [45](#), [56](#), [61](#), [63](#), [65](#)  
NMstamp, [6](#), [14](#), [16](#), [17](#), [20](#), [26](#), [44](#), [66](#), [70](#)  
NMwriteData, [6](#), [14](#), [16](#), [17](#), [20](#), [26](#), [44](#), [67](#), [67](#)  
NMwriteFilters, [70](#)  
NMwriteInits, [71](#)  
NMwriteSection, [38](#), [40](#), [53](#), [57](#), [72](#)  
NMwriteSizes, [74](#)

print.summary\_NMdata, [75](#)

rbind.NMdata (NMdataOperations), [34](#)  
renameByContents, [76](#)

summary.NMdata, [77](#)

t.NMdata (NMdataOperations), [34](#)  
tmpcol, [6](#), [14](#), [16](#), [17](#), [20](#), [26](#), [44](#), [67](#), [70](#)  
triagSize, [77](#)

uniquePresent, [78](#)  
unNMdata, [79](#)