Create CDISC pilot tables

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Contents

Setup	1
Create from demo data PhUSE scripting program	2
Get the values in the cube	34
Get cube components	36
p	37
p	37
Get observations	37
Get observations with labels	39
Reproduce the metadata for the workbook from cube	41
Session information	42

Setup

This script creates RDF data cubes from .csv files.

```
library(rrdfancillary)
library(rrdfcdisc)
library(rrdfqb)
library(rrdfqbcrnd0)
devtools::load_all(pkg="../..")
```

Loading rrdfqbcrndex

All files are stored in the directory

```
targetDir<- system.file("extdata/sample-rdf", package="rrdfqbcrndex")
(targetDir)</pre>
```

[1] "/home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf"

Create from demo data PhUSE scripting program

The files are generated by a SAS program - extdata/sample-workflow/adsl1.sas. The input files are read directly from .csv files instead of using the workbook, to avoid the extra step with the workbook, as the input files may be updated by running the SAS program.

```
tab1x010bsDataCsvFn<- system.file("extdata/sample-cfg", "TAB1X01.csv", package="rrdfqbcrndex")
# tab1x010bsDataCsvFn<- "TAB1X01.csv"
tab1x010bsDataCsvFn<- "TAB1X01.csv"
tab1x010bsData <- read.csv(tab1x010bsDataCsvFn,stringsAsFactors=FALSE)

##T0D0 add measurefmt; quick hack - affects vignettes/cube-from-workbook.Rmd and
##T0D0 inst/data-raw/create-qb-examples-as-ttl.Rmd
if (!( "measurefmt" %in% names(tab1x010bsData))) {
tab1x010bsData$measurefmt<- "%6.1f"
tab1x010bsData$measurefmt[ tab1x010bsData$procedure %in% c("n", "nmiss", "count") ]<- "%6.0f"
## sprintf( tab1x010bsData$measurefmt, tab1x010bsData$measure)
}
tab1x01MetaDataCsvFn<- system.file("extdata/sample-cfg", "TAB1X01-Components.csv", package="rrdfqbcrnde"
# tab1x01MetaDataCsvFn<- "TAB1X01-Components.csv"
tab1x01MetaData <- read.csv(tab1x01MetaDataCsvFn,stringsAsFactors=FALSE)

tab1x01.cube.fn<- BuildCubeFromDataFrames(tab1x01MetaData, tab1x010bsData)</pre>
```

```
## [1] "prefix rdf: <a href="http://www.w3.org/1999/02/22-rdf-syntax-ns#">http://www.w3.org/2004</a>
##
      DataStructureDefinition
                                              dimension
## 1
               ds:dsd-TAB1X01 crnd-dimension:comp24fl
## 2
               ds:dsd-TAB1X01 crnd-dimension:comp24fl
## 3
               ds:dsd-TAB1X01 crnd-dimension:comp24fl
## 4
               ds:dsd-TAB1X01 crnd-dimension:comp24fl
## 5
               ds:dsd-TAB1X01 crnd-dimension:disconfl
               ds:dsd-TAB1X01 crnd-dimension:disconfl
## 6
## 7
               ds:dsd-TAB1X01 crnd-dimension:disconfl
## 8
               ds:dsd-TAB1X01
                                  crnd-dimension:efffl
## 9
               ds:dsd-TAB1X01
                                  crnd-dimension:efffl
               ds:dsd-TAB1X01
                                  crnd-dimension:efffl
## 10
## 11
               ds:dsd-TAB1X01
                                  crnd-dimension:efffl
               ds:dsd-TAB1X01
                                  crnd-dimension:factor
## 12
## 13
               ds:dsd-TAB1X01
                                 crnd-dimension:factor
               ds:dsd-TAB1X01
                                  crnd-dimension:factor
## 14
## 15
               ds:dsd-TAB1X01
                                 crnd-dimension:factor
## 16
               ds:dsd-TAB1X01
                                  crnd-dimension:ittfl
## 17
               ds:dsd-TAB1X01
                                  crnd-dimension:ittfl
## 18
               ds:dsd-TAB1X01
                                  crnd-dimension:ittfl
## 19
               ds:dsd-TAB1X01 crnd-dimension:procedure
## 20
               ds:dsd-TAB1X01 crnd-dimension:procedure
## 21
               ds:dsd-TAB1X01
                                  crnd-dimension:saffl
## 22
               ds:dsd-TAB1X01
                                  crnd-dimension:saffl
## 23
               ds:dsd-TAB1X01
                                  crnd-dimension:saffl
## 24
               ds:dsd-TAB1X01
                                  crnd-dimension:trt01p
## 25
               ds:dsd-TAB1X01
                                  crnd-dimension:trt01p
## 26
               ds:dsd-TAB1X01
                                  crnd-dimension:trt01p
```

```
## 27
               ds:dsd-TAB1X01
                                   crnd-dimension:trt01p
## 28
                                   crnd-dimension:trt01p
               ds:dsd-TAB1X01
## 29
                ds:dsd-TAB1X01
                                   crnd-dimension:trt01p
##
                       cprefLabel
                                                                   c1
##
  1
       Codelist scheme: comp24fl
                                                     code:comp24f1-N
##
  2
       Codelist scheme: comp24fl
                                                     code:comp24fl-Y
       Codelist scheme: comp24fl
                                                 code:comp24fl- ALL
                                            code:comp24fl- NONMISS
## 4
       Codelist scheme: comp24fl
## 5
       Codelist scheme: disconfl
                                                     code:disconfl-Y
       Codelist scheme: disconfl
                                                 code:disconfl-_ALL_
## 6
## 7
       Codelist scheme: disconfl
                                            code:disconfl-_NONMISS_
          Codelist scheme: efffl
## 8
                                                        code:efffl-N
## 9
          Codelist scheme: efffl
                                                        code:efffl-Y
## 10
          Codelist scheme: efffl
                                                    code:efffl-_ALL_
## 11
          Codelist scheme: efffl
                                               code:efffl-_NONMISS_
## 12
         Codelist scheme: factor
                                                   code:factor-_ALL_
## 13
         Codelist scheme: factor
                                              code:factor-_NONMISS_
## 14
         Codelist scheme: factor
                                             code:factor-proportion
## 15
         Codelist scheme: factor
                                               code:factor-quantity
## 16
          Codelist scheme: ittfl
                                                        code:ittfl-Y
## 17
          Codelist scheme: ittfl
                                                    code:ittfl-_ALL_
          Codelist scheme: ittfl
                                               code:ittfl-_NONMISS_
## 19 Codelist scheme: procedure
                                               code:procedure-count
      Codelist scheme: procedure
## 20
                                             code:procedure-percent
## 21
          Codelist scheme: saffl
                                                        code:saffl-Y
## 22
          Codelist scheme: saffl
                                                    code:saffl- ALL
## 23
          Codelist scheme: saffl
                                               code:saffl-_NONMISS_
## 24
         Codelist scheme: trt01p
                                                        code:trt01p-
## 25
         Codelist scheme: trt01p
                                                code:trt01p-Placebo
## 26
         Codelist scheme: trt01p code:trt01p-Xanomeline_High_Dose
## 27
         Codelist scheme: trt01p code:trt01p-Xanomeline_Low_Dose
## 28
         Codelist scheme: trt01p
                                                   code:trt01p-_ALL_
## 29
         Codelist scheme: trt01p
                                              code:trt01p-_NONMISS_
##
               clprefLabel
                                    vn vct
## 1
                              comp24fl DATA rrdfgbcrnd0:ADSL COMP24FL
## 2
                             comp24fl DATA rrdfqbcrnd0:ADSL_COMP24FL
                          Y
## 3
                      ALL
                             comp24fl DATA rrdfqbcrnd0:ADSL COMP24FL
## 4
                  {\tt NONMISS}_{-}
                              comp24fl DATA rrdfqbcrnd0:ADSL_COMP24FL
## 5
                             disconfl DATA rrdfqbcrnd0:ADSL DISCONFL
## 6
                             disconfl DATA rrdfqbcrnd0:ADSL_DISCONFL
                      _ALL_
## 7
                             disconfl DATA rrdfqbcrnd0:ADSL DISCONFL
                  NONMISS
## 8
                          N
                                 efffl DATA
                                               rrdfqbcrnd0:ADSL EFFFL
                          Y
                                 efffl DATA
                                               rrdfqbcrnd0:ADSL EFFFL
## 9
## 10
                                efffl DATA
                                               rrdfqbcrnd0:ADSL_EFFFL
                      _{
m ALL}_{
m }
## 11
                  _NONMISS_
                                efffl DATA
                                               rrdfqbcrnd0:ADSL_EFFFL
## 12
                      _{\mathtt{ALL}}_{\mathtt{}}
                               factor DATA
                                                                   <NA>
## 13
                  _NONMISS_
                               factor DATA
                                                                   <NA>
## 14
                 proportion
                               factor DATA
                   quantity
## 15
                               factor DATA
## 16
                          Y
                                 ittfl DATA
                                               rrdfqbcrnd0:ADSL_ITTFL
## 17
                                 ittfl DATA
                                               rrdfqbcrnd0:ADSL_ITTFL
                      _ALL_
## 18
                  NONMISS
                                 ittfl DATA
                                               rrdfqbcrnd0:ADSL_ITTFL
## 19
                      count procedure DATA
## 20
                   percent procedure DATA
```

```
## 21
                                saffl DATA
                                               rrdfqbcrnd0:ADSL_SAFFL
## 22
                                saffl DATA
                                               rrdfqbcrnd0:ADSL_SAFFL
                      _ALL_
                  {\tt \_NONMISS}_{\tt \_}
## 23
                                saffl DATA
                                               rrdfqbcrnd0:ADSL_SAFFL
                                              rrdfqbcrnd0:ADSL_TRT01P
## 24
                               trt01p DATA
                   Placebo
                               trt01p DATA
                                              rrdfqbcrnd0:ADSL_TRT01P
                               trt01p DATA
                                              rrdfqbcrnd0:ADSL_TRT01P
## 26 Xanomeline High Dose
       Xanomeline Low Dose
                               trt01p DATA
                                              rrdfqbcrnd0:ADSL_TRT01P
## 27
                               trt01p DATA
                                              rrdfqbcrnd0:ADSL_TRT01P
## 28
                      _ALL_
## 29
                  _NONMISS_
                               trt01p DATA
                                              rrdfqbcrnd0:ADSL_TRT01P
##
                      vnval
## 1
                          N
                          Y
## 2
## 3
                       <NA>
## 4
                       <NA>
## 5
                          Y
## 6
                       <NA>
## 7
                       <NA>
## 8
                         N
## 9
                          Y
## 10
                       <NA>
## 11
                       <NA>
## 12
                       <NA>
## 13
                       <NA>
## 14
                proportion
## 15
                  quantity
## 16
                          Y
## 17
                       <NA>
## 18
                       <NA>
## 19
                      count
## 20
                   percent
## 21
## 22
                       <NA>
## 23
                       <NA>
## 24
## 25
                   Placebo
## 26 Xanomeline High Dose
## 27
       Xanomeline Low Dose
## 28
                       <NA>
## 29
                       <NA>
## crnd-dimension:comp24fl crnd-dimension:comp24fl crnd-dimension:comp24fl crnd-dimension:comp24fl crnd
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): N -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
```

```
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Placebo --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## High Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Xanomeline
## Low Dose -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): ALL --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): count --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): quantity --
## CODING ERROR- no decode value
```

```
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): _ALL_ --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): Y -- CODING
## ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): percent --
## CODING ERROR- no decode value
## Warning in ph.recode(obsData[i, qbdim], recode.list[[qbdim]]): proportion
## -- CODING ERROR- no decode value
cat("TAB1X01 cube stored as ", normalizePath(tab1x01.cube.fn), "\n")
## TAB1X01 cube stored as /tmp/RtmpEOWBli/DC-TAB1X01-R-V-0-0-0.ttl
targetFile<- file.path(targetDir, "CDISC-pilot-TAB1X01.ttl")</pre>
if (file.copy( tab1x01.cube.fn, targetFile, overwrite=TRUE)) {
```

 $\verb|## RDF data cube copied to /home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf/CDISC-pides and the copied to /home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf/CDISC-pides and copied to /home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf/cDISC-pides and copied to /home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf/cDISC-pides and copied to /home/ma/projects/rrdfqbcrnd0/rrdfqbcrndex/inst/extdata/sample-rdf/cDISC-pides and copied to /home/ma/projects/rrdfqbcrndex/inst/extdata/sample-rdf/cDISC-pides and copied to /home/ma/projects/rrdfqbcrndex/rrdf/cDISC-pides and copied to /home/ma/projects/rrdf/cDISC-pides and copi$

Now look at the generated cubes by loading the turtle files.

```
dataCubeFile<- tab1x01.cube.fn
```

The rest of the code only depends on the value of dataCubeFile.

cat("RDF data cube copied to ", normalizePath(targetFile), "\n")

```
checkCube <- new.rdf() # Initialize
temp<- load.rdf(dataCubeFile, format="TURTLE", appendTo= checkCube)
summarize.rdf(checkCube)</pre>
```

[1] "Number of triples: 1501"

Get the values in the cube

First set values for accessing the cube.

```
dsdName<- GetDsdNameFromCube( checkCube )
domainName<- GetDomainNameFromCube( checkCube )
forsparqlprefix<- GetForSparqlPrefix( domainName )</pre>
```

The next statement shows the first 10 triples in the cube.

```
observations1Rq<- paste( forsparqlprefix,
,
select *
where {?s ?p ?o .}
limit 10
',
"\n"
)
observations1<- sparql.rdf(checkCube, observations1Rq )
knitr::kable(head(observations1))</pre>
```

s	p	0
rdfs:domain	rdfs:domain	rdf:Property
rdfs:domain	rdfs:range	rdfs:Class
rdfs:comment	rdfs:range	rdfs:Literal
rdf:first	rdf:type	rdf:Property
rdf:first	rdfs:domain	rdf:List
rdf:first	rdfs: subPropertyOf	rdf:first

The next statement shows the first 10 triples in the cube, where the subject is a qb:Observation.

```
observations2Rq<- paste( forsparqlprefix,
,
select *
where { ?s a qb:Observation ; ?p ?o .}
limit 10
,
,
"\n"
)
observations2<- sparql.rdf(checkCube, observations2Rq)
knitr::kable(head(observations2, 10))</pre>
```

\mathbf{s}	p	0
ds:obs49	rdfs:label	49
ds:obs49	crnd-dimension:disconfl	code:CODING ERROR- no decode value
ds:obs49	crnd-dimension: $trt01p$	code:CODING ERROR- no decode value
ds:obs49	crnd-dimension:procedure	code:CODING ERROR- no decode value
ds:obs49	qb:dataSet	ds:dataset-TAB1X01
ds:obs49	crnd-dimension:comp24fl	code:CODING ERROR- no decode value
ds:obs49	crnd-dimension:ittfl	code:CODING ERROR- no decode value
ds:obs49	crnd-attribute:denominator	
ds:obs49	crnd-dimension:efffl	code:CODING ERROR- no decode value
ds:obs49	$\operatorname{crnd-attribute:unit}$	NA

Get cube components

The cube components are shown in the next output.

```
componentsRq<- GetComponentSparqlQuery( forsparqlprefix, dsdName )
components<- as.data.frame(sparql.rdf(checkCube, componentsRq), stringsAsFactors=FALSE)
components$vn<- gsub("crnd-dimension:|crnd-attribute:|crnd-measure:","",components$p)
knitr::kable(components[,c("vn", "label")])</pre>
```

vn	label
comp24fl	comp24fl
disconfl	disconfl
efffl	efffl
factor	Type of procedure (quantity, proportion)
ittfl	ittfl
procedure	Statistical Procedure
saffl	saffl
trt01p	Treatment Arm

The codelists are shown in the next output.

```
codelistsRq<- GetCodeListSparqlQuery( forsparqlprefix, dsdName )
codelists<- as.data.frame(sparql.rdf(checkCube, codelistsRq), stringsAsFactors=FALSE)
codelists$vn<- gsub("crnd-dimension:|crnd-attribute:|crnd-measure:","",codelists$dimension)
codelists$clc<- gsub("code:","",codelists$cl)
knitr::kable(codelists[,c("vn", "clc", "clprefLabel")])</pre>
```

vn	clc	clprefLabel
comp24fl	comp24fl-N	N
comp24fl	comp24fl-Y	Y
comp24fl	${ m comp}24{ m fl} ext{-}ALL$	ALL
comp24fl	${ m comp}$ 24fl- $NONMISS$	NONMISS
disconfl	disconfl-Y	Y
disconfl	disconfl- ALL	ALL
disconfl	disconfl-NONMISS	NONMISS
efffl	efffl-N	N
efffl	efffl-Y	Y
efffl	$\operatorname{efffl-}ALL$	ALL
efffl	efffl-NONMISS	NONMISS
factor	factor- ALL	ALL
factor	factor-NONMISS	NONMISS
factor	factor-proportion	proportion
factor	factor-quantity	quantity
ittfl	ittfl-Y	Y
ittfl	ittfl- ALL	ALL
ittfl	${\it ittfl-NONMISS}$	NONMISS
procedure	procedure-count	count
procedure	procedure-percent	percent
saffl	saffl-Y	Y
saffl	saffl- ALL	ALL

vn	clc	clprefLabel
saffl	saffl-NONMISS	NONMISS
trt01p	trt01p-	
trt01p	trt01p-Placebo	Placebo
trt01p	trt01p-Xanomeline_High_Dose	Xanomeline High Dose
trt01p	trt01p-Xanomeline_Low_Dose	Xanomeline Low Dose
trt01p	${ m trt}01{ m p-}ALL$	ALL
trt01p	${ m trt01p-}NONMISS$	NONMISS

The dimensions are shown in the next output.

```
dimensionsRq <- GetDimensionsSparqlQuery( forsparqlprefix )
dimensions<- sparql.rdf(checkCube, dimensionsRq)
knitr::kable(dimensions)</pre>
```

\mathbf{p}

```
crnd-dimension:disconfl
crnd-dimension:saffl
crnd-dimension:trt01p
crnd-dimension:factor
crnd-dimension:procedure crnd-dimension:comp24fl
crnd-dimension:ittfl
crnd-dimension:efffl
```

Then the attributes as shown in the next output.

```
attributesRq<- GetAttributesSparqlQuery( forsparqlprefix )
attributes<- sparql.rdf(checkCube, attributesRq)
knitr::kable(attributes)</pre>
```

\mathbf{p}

crnd-attribute:denominator crnd-attribute:unit

Get observations

And finally the SPARQL query for observations, showing only the first 10 observations.

observationsRq<- GetObservationsSparqlQuery(forsparqlprefix, domainName, dimensions, attributes) cat(observationsRq)

```
## prefix rdf: <http://www.w3.org/1999/02/22-rdf-syntax-ns#>
## prefix skos: <http://www.w3.org/2004/02/skos/core#>
## prefix prov: <http://www.w3.org/ns/prov#>
## prefix rdfs: <http://www.w3.org/2000/01/rdf-schema#>
## prefix dcat: <http://www.w3.org/ns/dcat#>
## prefix owl: <http://www.w3.org/2002/07/owl#>
## prefix xsd: <http://www.w3.org/2001/XMLSchema#>
```

```
## prefix pav: <http://purl.org/pav>
## prefix dc: <http://purl.org/dc/elements/1.1/>
## prefix dct: <http://purl.org/dc/terms/>
## prefix mms: <http://rdf.cdisc.org/mms#>
## prefix cts: <a href="http://rdf.cdisc.org/ct/schema">
## prefix cdiscs: <http://rdf.cdisc.org/std/schema#>
## prefix cdash-1-1: <http://rdf.cdisc.org/std/cdash-1-1#>
## prefix cdashct: <http://rdf.cdisc.org/cdash-terminology#>
## prefix sdtmct: <a href="http://rdf.cdisc.org/sdtm-terminology">http://rdf.cdisc.org/sdtm-terminology</a>
## prefix sdtm-1-2: <http://rdf.cdisc.org/std/sdtm-1-2#>
## prefix sdtm-1-3: <http://rdf.cdisc.org/std/sdtm-1-3#>
## prefix sdtms-1-3: <a href="http://rdf.cdisc.org/sdtm-1-3/schema">
## prefix sdtmig-3-1-2: <http://rdf.cdisc.org/std/sdtmig-3-1-2#>
## prefix sdtmig-3-1-3: <http://rdf.cdisc.org/std/sdtmig-3-1-3#>
## prefix sendct: <http://rdf.cdisc.org/send-terminology#>
## prefix sendig-3-0: <a href="http://rdf.cdisc.org/std/sendig-3-0">http://rdf.cdisc.org/std/sendig-3-0">
## prefix adamct: <a href="http://rdf.cdisc.org/adam-terminology">http://rdf.cdisc.org/adam-terminology</a>
## prefix adam-2-1: <a href="http://rdf.cdisc.org/std/adam-2-1#">http://rdf.cdisc.org/std/adam-2-1#</a>
## prefix adamig-1-0: <a href="http://rdf.cdisc.org/std/adamig-1-0#">http://rdf.cdisc.org/std/adamig-1-0#</a>
## prefix adamvr-1-2: <a href="http://rdf.cdisc.org/std/adamvr-1-2">http://rdf.cdisc.org/std/adamvr-1-2">
## prefix qb: <http://purl.org/linked-data/cube#>
## prefix rrdfqbcrnd0: <http://www.example.org/rrdfqbcrnd0/>
## prefix code: <http://www.example.org/dc/code/>
## prefix dccs: <a href="http://www.example.org/dc/tab1x01/dccs/">
## prefix ds: <http://www.example.org/dc/tab1x01/ds/>
## prefix crnd-dimension: <a href="http://www.example.org/dc/dimension">http://www.example.org/dc/dimension">
## prefix crnd-attribute: <http://www.example.org/dc/attribute#>
## prefix crnd-measure: <http://www.example.org/dc/measure#>
##
## select * where {
## ?s a qb:Observation ;
##
        qb:dataSet ds:dataset-TAB1X01 ;
##
        crnd-dimension:disconfl ?disconfl;
##
        crnd-dimension:saffl ?saffl;
##
        crnd-dimension:trt01p ?trt01p;
##
        crnd-dimension:factor ?factor;
##
        crnd-dimension:procedure ?procedure;
##
       crnd-dimension:comp24fl ?comp24fl;
##
        crnd-dimension:ittfl ?ittfl;
##
       crnd-dimension:efffl ?efffl;
##
        crnd-attribute:denominator ?denominator;
##
        crnd-attribute:unit ?unit;
        crnd-measure:measure ?measure .
## optional{ ?disconfl skos:prefLabel ?disconflvalue . }
## optional{ ?saffl skos:prefLabel ?safflvalue . }
## optional{ ?trt01p skos:prefLabel ?trt01pvalue . }
## optional{ ?factor skos:prefLabel ?factorvalue . }
## optional{ ?procedure skos:prefLabel ?procedurevalue . }
## optional{ ?comp24fl skos:prefLabel ?comp24flvalue . }
## optional{ ?ittfl skos:prefLabel ?ittflvalue . }
## optional{ ?efffl skos:prefLabel ?effflvalue . }
## }
## order by ?s
```

disconflyalue	safflvalue	trt01pvalue	factorvalue	procedurevalue	comp24flvalue	ittflvalue	effflvalue	denominato
NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	ittfl
NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	ittfl
NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	ittfl
NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	ittfl
NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	saffl

Get observations with labels

The SPARQL query for observations with labels for variables, showing only the first 10 observations.

observationsDescriptionRq<- GetObservationsWithDescriptionSparqlQuery(forsparqlprefix, domainName, dimecat(observationsDescriptionRq)

```
## prefix rdf: <http://www.w3.org/1999/02/22-rdf-syntax-ns#>
## prefix skos: <http://www.w3.org/2004/02/skos/core#>
## prefix prov: <http://www.w3.org/ns/prov#>
## prefix rdfs: <http://www.w3.org/2000/01/rdf-schema#>
## prefix dcat: <http://www.w3.org/ns/dcat#>
## prefix owl: <http://www.w3.org/2002/07/owl#>
## prefix xsd: <http://www.w3.org/2001/XMLSchema#>
## prefix pav: <http://purl.org/pav>
## prefix dc: <http://purl.org/dc/elements/1.1/>
## prefix dct: <http://purl.org/dc/terms/>
## prefix mms: <http://rdf.cdisc.org/mms#>
## prefix cts: <http://rdf.cdisc.org/ct/schema#>
## prefix cdiscs: <a href="http://rdf.cdisc.org/std/schema">
## prefix cdash-1-1: <http://rdf.cdisc.org/std/cdash-1-1#>
## prefix cdashct: <a href="http://rdf.cdisc.org/cdash-terminology">http://rdf.cdisc.org/cdash-terminology</a>
## prefix sdtmct: <http://rdf.cdisc.org/sdtm-terminology#>
## prefix sdtm-1-2: <http://rdf.cdisc.org/std/sdtm-1-2#>
## prefix sdtm-1-3: <http://rdf.cdisc.org/std/sdtm-1-3#>
## prefix sdtms-1-3: <a href="http://rdf.cdisc.org/sdtm-1-3/schema">
## prefix sdtmig-3-1-2: <http://rdf.cdisc.org/std/sdtmig-3-1-2#>
## prefix sdtmig-3-1-3: <http://rdf.cdisc.org/std/sdtmig-3-1-3#>
## prefix sendct: <http://rdf.cdisc.org/send-terminology#>
## prefix sendig-3-0: <a href="http://rdf.cdisc.org/std/sendig-3-0#">http://rdf.cdisc.org/std/sendig-3-0#</a>>
## prefix adamct: <http://rdf.cdisc.org/adam-terminology#>
## prefix adam-2-1: <http://rdf.cdisc.org/std/adam-2-1#>
## prefix adamig-1-0: <http://rdf.cdisc.org/std/adamig-1-0#>
## prefix adamvr-1-2: <a href="http://rdf.cdisc.org/std/adamvr-1-2#">http://rdf.cdisc.org/std/adamvr-1-2#</a>
```

```
## prefix rrdfqbcrnd0: <a href="http://www.example.org/rrdfqbcrnd0/">
## prefix code: <http://www.example.org/dc/code/>
## prefix dccs: <http://www.example.org/dc/tab1x01/dccs/>
## prefix ds: <http://www.example.org/dc/tab1x01/ds/>
## prefix crnd-dimension: <a href="http://www.example.org/dc/dimension">
## prefix crnd-attribute: <http://www.example.org/dc/attribute#>
## prefix crnd-measure: <http://www.example.org/dc/measure#>
## select * where {
## ?s a qb:Observation ;
## qb:dataSet ds:dataset-TAB1X01 ;
## crnd-dimension:disconfl ?disconfl;
## crnd-dimension:saffl ?saffl;
## crnd-dimension:trt01p ?trt01p;
## crnd-dimension:factor ?factor;
## crnd-dimension:procedure ?procedure;
## crnd-dimension:comp24fl ?comp24fl;
## crnd-dimension:ittfl ?ittfl;
## crnd-dimension:efffl ?efffl;
## crnd-attribute:denominator ?denominator;
## crnd-attribute:unit ?unit;
## crnd-measure:measure
                              ?measure .
## optional{ ?disconfl skos:prefLabel ?disconflvalue . }
## optional{ ?saffl skos:prefLabel ?safflvalue . }
## optional{ ?trt01p skos:prefLabel ?trt01pvalue . }
## optional{ ?factor skos:prefLabel ?factorvalue . }
## optional{ ?procedure skos:prefLabel ?procedurevalue . }
## optional{ ?comp24fl skos:prefLabel ?comp24flvalue . }
## optional{ ?ittfl skos:prefLabel ?ittflvalue . }
## optional{ ?efffl skos:prefLabel ?effflvalue . }
## optional{ crnd-dimension:disconfl rdfs:label ?disconfllabel . }
## optional{ crnd-dimension:saffl rdfs:label ?saffllabel . }
## optional{ crnd-dimension:trt01p rdfs:label ?trt01plabel . }
## optional{ crnd-dimension:factor rdfs:label ?factorlabel . }
## optional{ crnd-dimension:procedure rdfs:label ?procedurelabel . }
## optional{ crnd-dimension:comp24fl rdfs:label ?comp24fllabel . }
## optional{ crnd-dimension:ittfl rdfs:label ?ittfllabel . }
## optional{ crnd-dimension:efffl rdfs:label ?efffllabel . }
## BIND( IRI(crnd-dimension:disconfl) as ?disconflIRI)
## BIND( IRI(crnd-dimension:saffl) as ?safflIRI)
## BIND( IRI(crnd-dimension:trt01p) as ?trt01pIRI)
## BIND( IRI(crnd-dimension:factor) as ?factorIRI)
## BIND( IRI(crnd-dimension:procedure) as ?procedureIRI)
## BIND( IRI(crnd-dimension:comp24f1) as ?comp24f1IRI)
## BIND( IRI(crnd-dimension:ittfl) as ?ittflIRI)
## BIND( IRI(crnd-dimension:efffl) as ?effflIRI)
## BIND( IRI( ?s ) AS ?measureIRI)
## }
observationsDesc<- as.data.frame(sparql.rdf(checkCube, observationsDescriptionRq), stringsAsFactors=FA
knitr::kable(observationsDesc[ 1:10 ,
     c(paste0(rep(sub("crnd-dimension:|crnd-attribute:|crnd-measure:", "", dimensions),each=3),
       sub("crnd-dimension:|crnd-attribute:|crnd-measure:", "", attributes), "measure", "measureIRI"
```

prefix qb: <http://purl.org/linked-data/cube#>

)]
)	

disconfllabel	disconflyalue	disconfIRI	saffllabel	safflvalue	saffIRI	trt01plabel	trt(
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA
disconfl	NA	crnd-dimension:disconfl	saffl	NA	crnd-dimension:saffl	Treatment Arm	NA

Reproduce the metadata for the workbook from cube

Here is an example of roundtripping: make the metadata used for the workbook from RDF data cube.

First get the dimensions, measure and attribute

```
workbookDimAttrMeasRq<- GetDimAttrMeasInWorkbookFormatSparqlQuery( forsparqlprefix )
dimensionsattr<- sparql.rdf(checkCube, workbookDimAttrMeasRq )
knitr::kable(dimensionsattr)</pre>
```

$\overline{\text{compType}}$	compName	codeType	nciDomainValue
dimension	crnd-dimension:disconfl	NA	NA
dimension	crnd-dimension:saffl	NA	NA
dimension	crnd-dimension: $trt01p$	NA	NA
dimension	crnd-dimension:factor	NA	NA
dimension	crnd-dimension:procedure	NA	NA
dimension	crnd-dimension:comp24fl	NA	NA
dimension	crnd-dimension:ittfl	NA	NA
dimension	crnd-dimension:efffl	NA	NA
attribute	crnd-attribute:denominator	NA	NA
attribute	crnd-attribute:unit	NA	NA
measure	crnd-measure:measure	NA	NA

Secondly, get the metadata for the workbook. To get the metadata element "cubeVersion" a workaround is needed. The cubeversion is not directly available but from dcat:distribution derived as the result of paste0("DC-", domainName,"-R-V-",cubeVersion, ".ttl").

```
workbookMetadataRq<- GetMetaDataInWorkbookFormatSparqlQuery( forsparqlprefix )
metadata<- sparql.rdf(checkCube, workbookMetadataRq)
cubeVersion<- gsub("-",".", gsub("DC-.*-R-V-([^\\.]+).ttl", "\\1", metadata[ metadata[,2]=="distribution", metadataX<- rbind(metadata, cbind(compType="metadata", compName="cubeVersion", compLabel=cubeVersion))
knitr::kable(metadataX)</pre>
```

	compType	compName	compLabel
	metadata metadata metadata	title distribution comment	Demographics Analysis Results DC-TAB1X01-R-V-0-0-0.ttl
compLabel	metadata metadata metadata	label description obsFileName cubeVersion	Table 14-1.01 Summary of Populations Data from adsl1.sas program tab1x01.csv 0.0.0

Session information

sessionInfo()

```
## R version 3.2.3 (2015-12-10)
## Platform: x86_64-redhat-linux-gnu (64-bit)
## Running under: Fedora 23 (Workstation Edition)
##
## locale:
                                      LC NUMERIC=C
  [1] LC_CTYPE=en_GB.UTF-8
  [3] LC_TIME=en_GB.UTF-8
                                      LC_COLLATE=en_GB.UTF-8
##
   [5] LC MONETARY=en GB.UTF-8
                                      LC MESSAGES=en GB.UTF-8
##
##
  [7] LC PAPER=en GB.UTF-8
                                      LC NAME=en GB.UTF-8
## [9] LC ADDRESS=en GB.UTF-8
                                      LC TELEPHONE=en GB.UTF-8
## [11] LC_MEASUREMENT=en_GB.UTF-8
                                      LC_IDENTIFICATION=en_GB.UTF-8
##
## attached base packages:
## [1] methods
                          graphics grDevices utils
                stats
                                                         datasets base
## other attached packages:
## [1] rrdfqbcrndex_0.2.2 rrdfqbcrnd0_0.2.2
                                                rrdfqb_0.2.2
## [4] xlsx_0.5.7
                            xlsxjars_0.6.1
                                                rrdfcdisc_0.2.2
## [7] devtools_1.11.1
                            RCurl_1.95-4.8
                                                bitops_1.0-6
## [10] rrdfancillary_0.2.2 rrdf_2.1.2
                                                rrdflibs_1.4.0
## [13] rJava_0.9-8
##
## loaded via a namespace (and not attached):
                       knitr_1.12.3
## [1] Rcpp_0.12.4
                                       magrittr_1.5
                                                        roxygen2_5.0.1
  [5] highr_0.5.1
                        stringr_1.0.0
                                        tools_3.2.3
                                                        withr_1.0.1
## [9] htmltools_0.3.5 yaml_2.1.13
                                        digest_0.6.9
                                                        formatR 1.3
## [13] memoise 1.0.0
                        evaluate_0.8.3 rmarkdown_0.9.5 stringi_1.0-1
```