

Package ‘recodeflow’

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Type Package

Title Interface Functions for PMML Creation, and Data Recoding

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Description Contains functions to interface with variables and variable details sheets, including recoding variables and converting them to PMML.

Depends R (>= 3.1.0)

Imports XML (>= 3.98-1.11), sjlabelled, stringr, tidyr, haven, dplyr, magrittr

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URL <https://github.com/Big-Life-Lab/recodeflow>

BugReports <https://github.com/Big-Life-Lab/recodeflow/issues>

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Contents

add_data_field_children_for_start_var	2
attach_apply_nodes	3
attach_cat_value_nodes_for_start_var	4
attach_cont_value_nodes_for_start_var	4

attach_derived_field_child_nodes	5
attach_range_value_nodes	5
build_data_field_for_start_var	6
build_data_field_for_var	6
build_derived_field_node	7
build_derived_field_value_node	7
build_missing_const_node	8
build_numeric_derived_field_apply_node	8
build_ranged_derived_field_apply_node	9
build_trans_dict	9
build_variable_field_ref_node	10
compare_value_based_on_interval	10
create_id_row	11
create_label_list_element	12
example_der_fun	12
format_recoded_value	13
get_data_variable_name	13
get_margins	14
get_margin_closure	14
get_start_var_name	15
get_variable_type_data_type	15
get_var_details_rows	16
get_var_details_row_indices	16
get_var_sheet_row	17
is_equal	17
is_left_open	18
is_numeric	19
is_rec_from_range	19
is_right_open	20
label_data	20
recode_columns	21
recode_to_pmml	21
rec_with_table	23
select_vars_by_role	27
set_data_labels	28

Index **29**

add_data_field_children_for_start_var
Add DataField child nodes for start variable.

Description

Add DataField child nodes for start variable.

Usage

```
add_data_field_children_for_start_var(data_field, var_details_rows)
```

Arguments

`data_field` DataField node to attach child nodes.
`var_details_rows` Variable details rows associated with current variable.

Value

Updated DataField node.

`attach_apply_nodes` *Attach Apply nodes to a parent node.*

Description

Attach Apply nodes to a parent node.

Usage

```
attach_apply_nodes(var_details_rows, parent_node, db_name)
```

Arguments

`var_details_rows` Variable details rows associated with a variable.
`parent_node` An XML node.
`db_name` Database name.

Value

Updated parent node.

attach_cat_value_nodes_for_start_var

Attach categorical value nodes to DataField node for start variable.

Description

Attach categorical value nodes to DataField node for start variable.

Usage

```
attach_cat_value_nodes_for_start_var(var_details_row, data_field)
```

Arguments

var_details_row	Variable details sheet row.
data_field	DataField node to attach Value nodes.

Value

Updated DataField node.

attach_cont_value_nodes_for_start_var

Attach continuous Value nodes for start variable.

Description

Attach continuous Value nodes for start variable.

Usage

```
attach_cont_value_nodes_for_start_var(var_details_row, data_field)
```

Arguments

var_details_row	Variable details sheet row.
data_field	DataField node to attach Value nodes.

Value

Updated DataField node.

attach_derived_field_child_nodes
Attach child nodes to DerivedField node.

Description

Attach child nodes to DerivedField node.

Usage

```
attach_derived_field_child_nodes(  
    derived_field_node,  
    var_details_sheet,  
    var_name,  
    db_name  
)
```

Arguments

derived_field_node	DerivedField node to attach child nodes.
var_details_sheet	Variable details sheet data frame.
var_name	Variable name.
db_name	Database name.

Value

Updated DerivedField node.

attach_range_value_nodes
Attach Value nodes to DataField node. Used when 'recFrom' has a value range.

Description

Attach Value nodes to DataField node. Used when 'recFrom' has a value range.

Usage

```
attach_range_value_nodes(var_details_row, data_field)
```

Arguments

var_details_row Variable details sheet row.
 data_field DataField node to attach Value nodes.

Value

Updated DataField node.

build_data_field_for_start_var
Build DataField node for start variable.

Description

Build DataField node for start variable.

Usage

build_data_field_for_start_var(var_name, var_details_rows)

Arguments

var_name Variable name.
 var_details_rows All variable details rows for the 'var_name' variable.

Value

DataField node with optype and dataType according to 'fromType'.

build_data_field_for_var
Build DataField node for variable.

Description

Build DataField node for variable.

Usage

build_data_field_for_var(var_name, vars_sheet)

Arguments

var_name Variable name.
vars_sheet Variable sheet data frame.

Value

DataField node for variable.

build_derived_field_node
Build DerivedField node.

Description

Build DerivedField node.

Usage

build_derived_field_node(vars_sheet, var_details_sheet, var_name, db_name)

Arguments

vars_sheet Variables sheet data frame.
var_details_sheet Variable details sheet data frame.
var_name Variable name.
db_name Database name.

Value

DerivedField node.

build_derived_field_value_node
Build Value node for DerivedField node.

Description

Build Value node for DerivedField node.

Usage

build_derived_field_value_node(var_details_row)

Arguments

var_details_row
Variable details sheet row.

Value

Value node.

build_missing_const_node
Build Constant node for a missing value for a variable.

Description

Build Constant node for a missing value for a variable.

Usage

build_missing_const_node(var_details_row)

Arguments

var_details_row
Variable details sheet row.

Value

Constant node.

build_numeric_derived_field_apply_node
Build Apply node with singleton numeric for DerivedField node.

Description

Build Apply node with singleton numeric for DerivedField node.

Usage

build_numeric_derived_field_apply_node(var_details_row, db_name)

Arguments

var_details_row
Variable details sheet row.
db_name
Database name.

Value

Apply node for DerivedField node.

`build_ranged_derived_field_apply_node`
Build Apply node with interval nodes for DerivedField node.

Description

Build Apply node with interval nodes for DerivedField node.

Usage

`build_ranged_derived_field_apply_node(var_details_row, db_name)`

Arguments

`var_details_row` Variable details sheet row.
`db_name` Database name.

Value

Apply node with intervals for DerivedField node.

`build_trans_dict` *Build a TransformationDictionary node.*

Description

Build a TransformationDictionary node.

Usage

`build_trans_dict(vars_sheet, var_details_sheet, var_names, db_name)`

Arguments

`vars_sheet` Variable sheet data frame.
`var_details_sheet` Variable details sheet data frame.
`var_names` Vector of variable names.
`db_name` Database name.

Value

TransformationDictionary node.

`build_variable_field_ref_node`*Build FieldRef node for variable.*

Description

Build FieldRef node for variable.

Usage

```
build_variable_field_ref_node(var_details_row, db_name)
```

Arguments

<code>var_details_row</code>	Variable details sheet row.
<code>db_name</code>	Database name.

Value

FieldRef node.

`compare_value_based_on_interval`*Compare Value Based On Interval*

Description

Compare values on the scientific notation interval

Usage

```
compare_value_based_on_interval(  
    left_boundary,  
    right_boundary,  
    data,  
    compare_columns,  
    interval  
)
```

Arguments

left_boundary	the min value
right_boundary	the max value
data	the data that contains values being compared
compare_columns	The columns inside data being checked
interval	The scientific notation interval

Value

a boolean vector containing true for rows where the comparison is true

create_id_row	<i>ID role creation</i>
---------------	-------------------------

Description

Creates ID row for rec_with_table

Usage

```
create_id_row(data, id_role_name, database_name, variables)
```

Arguments

data	the data that the ID role row is created for
id_role_name	name for the role that ID is created from
database_name	the name of the database
variables	variables sheet containing variable information

Value

data with the ID row attached

create_label_list_element
Create label list element

Description

A data labeling utility function for creating individual variable labels

Usage

```
create_label_list_element(variable_rows)
```

Arguments

variable_rows all variable details rows containing 1 variable information

Value

a list containing labels for the passed variable

example_der_fun *example_der_fun caluclates chol*bili*

Description

example_der_fun caluclates chol*bili

Usage

```
example_der_fun(chol, bili)
```

Arguments

chol the row value for chol
bili the row value for bili

format_recoded_value *Recode NA formatting*

Description

Recodes the NA depending on the var type

Usage

```
format_recoded_value(cell_value, var_type)
```

Arguments

cell_value The value inside the recTo column
var_type the toType of a variable

Value

an appropriately coded tagged NA

get_data_variable_name
Get Data Variable Name

Description

Retrieves the name of the column inside data to use for calculations

Usage

```
get_data_variable_name(  
  data_name,  
  data,  
  row_being_checked,  
  variable_being_checked  
)
```

Arguments

data_name name of the database being checked
data database being checked
row_being_checked
 the row from variable details that contains information on this variable
variable_being_checked
 the name of the recoded variable

Value

the data equivalent of `variable_being_checked`

<code>get_margins</code>	<i>Extract margins from character vector.</i>
--------------------------	---

Description

Extract margins from character vector.

Usage

```
get_margins(chars)
```

Arguments

`chars` Character vector.

Value

Margins as character vector.

<code>get_margin_closure</code>	<i>Get closure type for a margin.</i>
---------------------------------	---------------------------------------

Description

Get closure type for a margin.

Usage

```
get_margin_closure(chars)
```

Arguments

`chars` Character vector.

Value

Closure type.

get_start_var_name *Get variable name from variableStart using database name.*

Description

Get variable name from variableStart using database name.

Usage

```
get_start_var_name(var_details_row, db_name)
```

Arguments

var_details_row A variable details row.
db_name Name of database to extract from.

Value

character The name of the start variable.

get_variable_type_data_type
 Get data type for variable type.

Description

Get data type for variable type.

Usage

```
get_variable_type_data_type(var_details_rows, var_type, is_start_var)
```

Arguments

var_details_rows All variable details rows for the variable.
var_type Variable type
is_start_var boolean if the passed variable is variable start

Value

'var_type' data type.

get_var_details_rows *Get all variable details rows for a variable and database combination.*

Description

Get all variable details rows for a variable and database combination.

Usage

```
get_var_details_rows(var_details_sheet, var_name, db_name)
```

Arguments

var_details_sheet	A data frame representing a variable details sheet.
var_name	Variable name.
db_name	Database name.

Value

All variable details rows for the variable and database combination.

get_var_details_row_indices
Get all variable details row indices for a variable.

Description

Get all variable details row indices for a variable.

Usage

```
get_var_details_row_indices(var_details_sheet, var_name)
```

Arguments

var_details_sheet	A data frame representing a variable details sheet.
var_name	Variable name.

Value

All variable details row indices for a variable.

get_var_sheet_row	<i>Get variable row from variable sheet.</i>
-------------------	--

Description

Get variable row from variable sheet.

Usage

```
get_var_sheet_row(var_name, vars_sheet)
```

Arguments

var_name	Variable name.
vars_sheet	Variable sheet data frame.

Value

Variable row.

is_equal	<i>Checks whether two values are equal including NA</i>
----------	---

Description

Compared to the base "==" operator in R, this function returns true if the two values are NA whereas the base "==" operator returns NA

Usage

```
is_equal(v1, v2)
```

Arguments

v1	variable 1
v2	variable 2

Value

boolean value of whether or not v1 and v2 are equal

Examples

```
is_equal(1,2)
# FALSE

is_equal(1,1)
# TRUE

1==NA
# NA

is_equal(1,NA)
# FALSE

NA==NA
# NA

is_equal(NA,NA)
# TRUE
```

is_left_open

Extract margins from character vector.

Description

Extract margins from character vector.

Usage

```
is_left_open(chars)
```

Arguments

chars Character vector.

Value

Whether the left endpoint of an interval is open.

is_numeric	<i>Check if a character object can be converted to a number.</i>
------------	--

Description

Check if a character object can be converted to a number.

Usage

```
is_numeric(chars)
```

Arguments

chars Character object.

Value

Whether 'chars' can be converted to a numeric value.

is_rec_from_range	<i>Check if recFrom is a range for a variable details row.</i>
-------------------	--

Description

Check if recFrom is a range for a variable details row.

Usage

```
is_rec_from_range(var_details_row)
```

Arguments

var_details_row Variable details sheet row.

Value

Whether recFrom is a range.

is_right_open	<i>Extract margins from character vector.</i>
---------------	---

Description

Extract margins from character vector.

Usage

```
is_right_open(chars)
```

Arguments

chars Character vector.

Value

Whether the right endpoint of an interval is open.

label_data	<i>label_data</i>
------------	-------------------

Description

Attaches labels to the data_to_label to preserve metadata

Usage

```
label_data(label_list, data_to_label)
```

Arguments

label_list the label list object that contains extracted labels from variable details
data_to_label The data that is to be labeled

Value

Returns labeled data

recode_columns *recode_columns*

Description

Recodes columns from passed row and returns just table with those columns and same rows as the data

Usage

```

recode_columns(
  data,
  variables_details_rows_to_process,
  data_name,
  log,
  print_note,
  else_default
)

```

Arguments

data The source database
variables_details_rows_to_process rows from variable details that are applicable to this DB
data_name Name of the database being passed
log The option of printing log
print_note the option of printing the note columns
else_default default else value to use if no else is present

Value

Returns recoded and labeled data

recode_to_pmml *Creates a PMML document from variable and variable details sheets for specified database.*

Description

Creates a PMML document from variable and variable details sheets for specified database.

Usage

```

recode_to_pmml(var_details_sheet, vars_sheet, db_name, vars_to_convert = NULL)

```

Arguments

<code>var_details_sheet</code>	A data frame representing a variable details sheet.
<code>vars_sheet</code>	A data frame representing a variables sheet.
<code>db_name</code>	A string containing the name of the database that holds the start variables. Should match up with one of the databases in the <code>databaseStart</code> column.
<code>vars_to_convert</code>	A vector of strings containing the names of variables from the variable column in the variable details sheet that should be converted to PMML. Passing in an empty vector will convert all the variables.

Value

A PMML document.

Examples

```
var_details_sheet <-
data.frame(
  "variable" = rep(c("A", "B", "C"), each = 3),
  "dummyVariable" = c("AY", "AN", "ANA", "BY", "BN", "BNA", "CY", "CN", "CNA"),
  "toType" = rep("cat", times = 9),
  "databaseStart" = rep("tester", times = 9),
  "variableStart" = rep(
    c("tester::startA", "tester::startB", "tester::startC"),
    each = 3
  ),
  "fromType" = rep("cat", times = 9),
  "recTo" = rep(c("1", "2", "NA::a"), times = 3),
  "numValidCat" = rep("2", times = 9),
  "catLabel" = rep(c("Yes", "No", "Not answered"), times = 3),
  "catLabelLong" = rep(c("Yes", "No", "Not answered"), times =
    3),
  "recFrom" = rep(c("1", "2", "9"), times = 3),
  "catStartLabel" = rep(c("Yes", "No", "Not answered"), times =
    3),
  "variableStartShortLabel" = rep(c("Group A", "Group B", "Group C"), each =
    3),
  "variableStartLabel" = rep(c("Group A", "Group B", "Group C"), each =
    3),
  "units" = rep("NA", times = 9),
  "notes" = rep("This is not real data", times = 9)
)
vars_sheet <-
data.frame(
  "variable" = c("A", "B", "C"),
  "label" = c("Group A", "Group B", "Group C"),
  "labelLong" = c("Group A", "Group B", "Group C"),
  "section" = rep("tester", times=3),
  "subject" = rep("tester", times = 3),
  "variableType" = rep("Categorical", times=3),
```

```

      "databaseStart" = rep("tester", times = 3),
      "units" = rep("NA", times = 3),
      "variableStart" = c("tester::startA", "tester::startB", "tester::startC")
    )
db_name <- "tester"
vars <- c("A", "B", "C")

actual_pmml <- recode_to_pmml(
  var_details_sheet,
  vars_sheet,
  db_name,
  vars
)

```

rec_with_table	<i>Recode with Table</i>
----------------	--------------------------

Description

Creates new variables by recoding variables in a dataset using the rules specified in a variables details sheet

Usage

```

rec_with_table(
  data,
  variables = NULL,
  database_name = NULL,
  variable_details = NULL,
  else_value = NA,
  append_to_data = FALSE,
  log = FALSE,
  notes = TRUE,
  var_labels = NULL,
  custom_function_path = NULL,
  attach_data_name = FALSE,
  id_role_name = NULL,
  name_of_environment_to_load = NULL,
  append_non_db_columns = FALSE
)

```

Arguments

data	A dataframe containing the variables to be recoded. Can also be a named list of dataframes.
variables	Character vector containing the names of the new variables to recode to or a dataframe containing a variables sheet.

database_name	A String containing the name of the database containing the original variables which should match up with a database from the databaseStart column in the variables details sheet. Should be a character vector if data is a named list where each vector item matches a name in the data list and also matches with a value in the databaseStart column of a variable details sheet.
variable_details	A dataframe containing the specifications for recoding.
else_value	Value (string, number, integer, logical or NA) that is used to replace any values that are outside the specified ranges (no rules for recoding).
append_to_data	Logical, if TRUE (default), the newly created variables will be appended to the original dataset.
log	Logical, if FALSE (default), a log containing information about the recoding will not be printed.
notes	Logical, if FALSE (default), will not print the content inside the ‘Note‘ column of the variable being recoded.
var_labels	labels vector to attach to variables in variables
custom_function_path	string containing the path to the file containing functions to run for derived variables. This file will be sourced and its functions loaded into the R environment.
attach_data_name	logical to attach name of database to end table
id_role_name	name for the role to be used to generate id column
name_of_environment_to_load	Name of package to load variables and variable_details from
append_non_db_columns	boolean determining if data not present in this cycle should be appended as NA

Details

The **variable_details** dataframe needs the following columns:

variable Name of the new variable created. The name of the new variable can be the same as the original variable if it does not change the original variable definition

toType type the new variable *cat = categorical, cont = continuous*

databaseStart Names of the databases that the original variable can come from. Each database name should be separated by a comma. For eg., "cchs2001_p, cchs2003_p, cchs2005_p, cchs2007_p"

variableStart Names of the original variables within each database specified in the databaseStart column. For eg. , "cchs2001_p::RACA_6A, cchs2003_p::RACC_6A, ADL_01". The final variable specified is the name of the variable for all other databases specified in databaseStart but not in this column. For eg., ADL_01 would be the original variable name in the cchs2005_p and cchs2007_p databases.

fromType variable type of start variable. *cat = categorical or factor variable cont = continuous variable (real number or integer)*

recTo Value to recode to

recFrom Value/range being recoded from

Each row in the *variables details* sheet encodes the rule for recoding value(s) of the original variable to a category in the new variable. The categories of the new variable are encoded in the *recTo* column and the value(s) of the original variable that recode to this new value are encoded in the *recFrom* column. These recode columns follow a syntax similar to the *sjmisc::rec()* function. Whereas in the *sjmisc::rec()* function the recoding rules are in one string, in the variables details sheet they are encoded over multiple rows and columns (*recFrom* and *recTo*). For eg., a recoding rule in the *sjmisc* function would like like "1=2;2=3" whereas in the variables details sheet this would be encoded over two rows with *recFrom* and *recTo* values of the first row being 1 and 2 and similarly for the second row it would be 2 and 3. The rules for describing recoding pairs are shown below:

recode pairs Each recode pair is a row

multiple values Multiple values from the old variable that should be recoded into a new category of the new variable should be separated with a comma. e.g., *recFrom* = "1,2"; *recTo* = 1 will recode values of 1 and 2 in the original variable to 1 in the new variable

value range A value range is indicated by a colon, e.g. *recFrom*= "1:4"; *recTo* = 1 will recode all values from 1 to 4 into 1

min and max minimum and maximum values are indicated by *min* (or *lo*) and *max* (or *hi*), e.g. *recFrom* = "min:4"; *recTo* = 1 will recode all values from the minimum value of the original variable to 4 into 1

"else" All other values, which have not been specified yet, are indicated by *else*, e.g. *recFrom* = "else"; *recTo* = NA will recode all other values (not specified in other rows) of the original variable to "NA")

"copy" the *else* token can be combined with *copy*, indicating that all remaining, not yet recoded values should stay the same (are copied from the original value), e.g. *recFrom* = "else"; *recTo* = "copy"

NA's NA values are allowed both for the original and the new variable, e.g. *recFrom* "NA"; *recTo* = 1. or *recFrom* = "3:5"; *recTo* = "NA" (recodes all NA into 1, and all values from 3 to 5 into NA in the new variable)

Value

a dataframe that is recoded according to rules in *variable_details*.

Examples

```
var_details <-
data.frame(
  "variable" = c("time", rep("status", times = 3), rep("trt", times = 2),
  "age", rep("sex", times = 2), rep("ascites", times = 2),
  rep("hepato", times = 2), rep("spiders", times = 2),
  rep("edema", times = 3),
  "bili", "chol", "albumin", "copper", "alk.phos", "ast",
  "trig", "platelet", "protime", rep("stage", times = 4)),
  "dummyVariable" = c("NA", "status0", "status1", "status2", "trt1", "trt2",
  "NA", "sexM", "sexF", "ascites0", "ascites1", "hepato0", "hepato1",
  "spiders0", "spiders1", "edema0.0", "edema0.5", "edema1.0",
  rep("NA", times = 9), "stage1", "stage2", "stage3", "stage4"),
  "toType" = c("cont", rep("cat", times = 3), rep("cat", times = 2),
  "cont", rep("cat", times = 2), rep("cat", times = 2),
```

```

rep("cat", times = 2), rep("cat", times = 2), rep("cat", times = 3),
rep("cont", times = 9), rep("cat", times = 4)),
"databaseStart" = rep("tester1, tester2", times = 31),
"variableStart" = c("[time]", rep("[status]", times = 3), rep("[trt]",
times = 2), "[age]", rep("[sex]", times = 2), rep("[ascites]",
times = 2), rep("[hepato]", times = 2), rep("[spiders]", times = 2),
rep("[edema]", times = 3), "[bili]", "[chol]", "[albumin]", "[copper]",
"[alk.phos]", "[ast]", "[trig]", "[platelet]", "[protime]",
rep("[stage]", times = 4)), "fromType" = c("cont", rep("cat", times = 3),
rep("cat", times = 2), "cont", rep("cat", times = 2),
rep("cat", times = 2), rep("cat", times = 2), rep("cat", times = 2),
rep("cat", times = 3), rep("cont", times = 9), rep("cat", times = 4)),
"recTo" = c("copy", "0", "1", "2", "1", "2", "copy", "m", "f", "0", "1", "0",
"1", "0", "1", "0.0", "0.5", "1.0", rep("copy", times = 9), "1", "2", "3", "4"),
"catLabel" = c("", "status 0", "status 1", "status 2", "trt 1", "trt 2", "",
"sex m", "sex f", "ascites 0", "ascites 1", "hepato 0", "hepato 1",
"spiders 0", "spiders 1", "edema 0.0", "edema 0.5", "edema 1.0",
rep("", times = 9), "stage 1", "stage 2", "stage 3", "stage 4"),
"catLabelLong" = c("", "status 0", "status 1", "status 2", "trt 1",
"trt 2", "", "sex m", "sex f", "ascites 0", "ascites 1", "hepato 0",
hepato 1", "spiders 0", "spiders 1", "edema 0.0", "edema 0.5", "edema 1.0",
rep("", times = 9), "stage 1", "stage 2", "stage 3", "stage 4"),
"recFrom" = c("else", "0", "1", "2", "1", "2", "else", "m", "f", "0", "1", "0",
"1", "0", "1", "0.0", "0.5", "1.0", rep("else", times = 9), "1", "2", "3", "4"),
"catStartLabel" = c("", "status 0", "status 1", "status 2", "trt 1",
"trt 2", "", "sex m", "sex f", "ascites 0", "ascites 1", "hepato 0",
"hepato 1", "spiders 0", "spiders 1", "edema 0.0", "edema 0.5", "edema 1.0",
rep("", times = 9), "stage 1", "stage 2", "stage 3", "stage 4"),
"variableStartShortLabel" = c("time", rep("status", times = 3),
rep("trt", times = 2), "age", rep("sex", times = 2),
rep("ascites", times = 2), rep("hepato", times = 2),
rep("spiders", times = 2), rep("edema", times = 3), "bili", "chol",
"albumin", "copper", "alk.phos", "ast", "trig", "platelet", "protime",
rep("stage", times = 4)),
"variableStartLabel" = c("time", rep("status", times = 3),
rep("trt", times = 2), "age", rep("sex", times = 2),
rep("ascites", times = 2), rep("hepato", times = 2),
rep("spiders", times = 2), rep("edema", times = 3), "bili", "chol",
"albumin", "copper", "alk.phos", "ast", "trig", "platelet", "protime",
rep("stage", times = 4)),
"units" = rep("NA", times = 31),
"notes" = rep("This is sample survival pbc data", times = 31)
)
var_sheet <-
data.frame(
"variable" = c("time", "status", "trt", "age", "sex", "ascites", "hepato",
"spiders", "edema", "bili", "chol", "albumin", "copper", "alk.phos",
"ast", "trig", "platelet", "protime", "stage"),
"label" = c("time", "status", "trt", "age", "sex", "ascites", "hepato",
"spiders", "edema", "bili", "chol", "albumin", "copper", "alk.phos",
"ast", "trig", "platelet", "protime", "stage"),
"labelLong" = c("time", "status", "trt", "age", "sex", "ascites", "hepato",
"spiders", "edema", "bili", "chol", "albumin", "copper", "alk.phos",

```

```

    "ast", "trig", "platelet", "protime", "stage"),
    "section" = rep("tester", times=19),
    "subject" = rep("tester", times = 19),
    "variableType" = c("cont", "cat", "cat", "cont", "cat", "cat", "cat",
    "cat", "cat", rep("cont", times = 9), "cat"),
    "databaseStart" = rep("tester1, tester2", times = 19),
    "units" = rep("NA", times = 19),
    "variableStart" = c("[time]", "[status]", "[trt]", "[age]", "[sex]",
    "[ascites]", "[hepato]", "[spiders]", "[edema]", "[bili]", "[chol]",
    "[albumin]", "[copper]", "[alk.phos]", "[ast]", "[trig]", "[platelet]",
    "[protime]", "[stage]")
  )
library(survival)
tester1 <- survival::pbc[1:209,]
tester2 <- survival::pbc[210:418,]
db_name1 <- "tester1"
db_name2 <- "tester2"

rec_sample1 <- rec_with_table(data = tester1,
  variables = var_sheet,
  variable_details = var_details,
  database_name = db_name1)

rec_sample2 <- rec_with_table(data = tester2,
  variables = var_sheet,
  variable_details = var_details,
  database_name = db_name2)

```

select_vars_by_role *Vars selected by role*

Description

Selects variables from variables sheet based on passed roles

Usage

```
select_vars_by_role(roles, variables)
```

Arguments

roles	a vector containing a single or multiple roles to match by
variables	the variables sheet containing variable info

Value

a vector containing the variable names that match the passed roles

set_data_labels	<i>Set Data Labels</i>
-----------------	------------------------

Description

sets labels for passed database, Uses the names of final variables in variable_details/variables_sheet as well as the labels contained in the passed dataframes

Usage

```
set_data_labels(data_to_label, variable_details, variables_sheet = NULL)
```

Arguments

data_to_label newly transformed dataset
variable_details
 variable_details.csv
variables_sheet
 variables.csv

Value

labeled data_to_label

Index

add_data_field_children_for_start_var, 2
attach_apply_nodes, 3
attach_cat_value_nodes_for_start_var, 4
attach_cont_value_nodes_for_start_var, 4
attach_derived_field_child_nodes, 5
attach_range_value_nodes, 5

build_data_field_for_start_var, 6
build_data_field_for_var, 6
build_derived_field_node, 7
build_derived_field_value_node, 7
build_missing_const_node, 8
build_numeric_derived_field_apply_node, 8
build_ranged_derived_field_apply_node, 9
build_trans_dict, 9
build_variable_field_ref_node, 10

compare_value_based_on_interval, 10
create_id_row, 11
create_label_list_element, 12

example_der_fun, 12

format_recoded_value, 13

get_data_variable_name, 13
get_margin_closure, 14
get_margins, 14
get_start_var_name, 15
get_var_details_row_indices, 16
get_var_details_rows, 16
get_var_sheet_row, 17
get_variable_type_data_type, 15

is_equal, 17
is_left_open, 18

is_numeric, 19
is_rec_from_range, 19
is_right_open, 20

label_data, 20

rec_with_table, 23
recode_columns, 21
recode_to_pmml, 21

select_vars_by_role, 27
set_data_labels, 28