Simple Tables Demo Doc

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1 Data

Mostly working with this data; but some others come in later to illustrate certain features.

```
library(pmtables)
library(yspec)
spec <- ys_help$spec()</pre>
pmdata <- readRDS("datasets/pmdata.RDS")</pre>
head(pmdata)
## # A tibble: 6 x 9
##
    STUDYf
                DOSEf FORMf WT
                                      CRCL AGE
                                                  N
                                                         ALB
                                                               SCR
     <chr>
                 <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr>
##
## 1 12-DEMO-001 100 mg tablet 71.4 104
                                            33.7 80
                                                        4.20 1.06
## 2 12-DEMO-001 150 mg capsule 89.4 122
                                            24.4 16
                                                        4.63 1.12
## 3 12-DEMO-001 150 mg tablet 81.7 104
                                            34.4 48
                                                        3.83 0.910
## 4 12-DEMO-001 150 mg troche 94.0 93.2 27.4 16
                                                        4.94 1.25
## 5 12-DEMO-001 200 mg tablet 67.9 100
                                            27.5 64
                                                        4.25 1.10
## 6 12-DEMO-001 200 mg troche 76.6 99.2 22.8 16
                                                        4.54 1.15
data <- select(pmdata, STUDY = STUDYf, DOSE = DOSEf, WT, AGE, CRCL, ALB, SCR)
```

2 wrap

There is a wrap function in this document. Don't worry about that; it just puts the output into a table environment and sends the output to get rendered in markdown.

3 Simple table

• Columns are in bold by default (can be turned off)

data %>% stable() %>% pt_wrap(stdout())

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

4 Annotate

- Arguments to identify the name of the generating R script and the output file name
- The output file name is retained as an attribute to be used later when saving the table data
- Arbitrary notes are also allowed (encouraged)

```
stable(
  data,
  r_file = "foo.R",
  output_file = "foo.tex",
  notes = c("Data were analyzed in quadruplicate.", "The results are very clear."),
) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

Data were analyzed in quadruplicate.

The results are very clear.

source code: foo.R source file: foo.tex

5 Align

- Center everything except for
 STUDY (left)
 DOSE and SCR (right)

```
stable(align = cols_center(STUDY = '1', .r = "DOSE,SCR")) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

6 Units

• Automatically put units under the column name

```
units <- ys_get_unit(spec, parens = TRUE)

stable(
  data,
  align = cols_center(STUDY = 'l', .r = "DOSE,SCR"),
  units = units,
) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT (kg)	AGE (years)	CRCL (ml/min)	ALB (g/dL)	SCR (mg/dL)
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

7 Math

- Columns with at least two \$ are "math" and will not be sanitized
- Otherwise the are functions to "prime" the data frame
 - The default is to convert every column to character
 - Then walk the columns, look for non-math columns and sanitize them

```
ptab <- readRDS("datasets/ptab.RDS")</pre>
ptab
## # A tibble: 4 x 6
##
                                            Symbol
                                                                                 SE
     .type
                        Parameter
                                                     Math
                                                                     Estimate
##
     <chr>>
                        <chr>
                                            <chr>
                                                     <chr>
                                                                         <dbl> <dbl>
## 1 Fixed-effects
                                            CL (L/h~ "\\exp(\\thet~
                        Clearance
                                                                         1.22 0.4
## 2 Fixed-effects
                        Volume of distribu~ V2 (L)
                                                     "$\leq(\t \cdot )
                                                                         5.87 0.89
## 3 Fixed-effects
                        Absorption rate co~ KA (1/h~ "$\\theta_3$"
                                                                         1.23 0.1
## 4 Unexplained varia~ Additive_error
                                            RUV
                                                     "$\\sigma_1$"
                                                                         0.02 0.01
stable(
  ptab,
  align = cols_center(Parameter = col_ragged(3), .1 = "Symbol"),
  panel = ".type"
```

8 hline from column

• Use the column to determine where the hline should go

```
stable(
  data,
  hline_from = "STUDY",
) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

9 hline anywhere

• Give row numbers for hline

```
stable(
  data,
  hline_at = c(3,nrow(data))-1,
) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

10 Remove dups

• Discard repeating values in a column

```
stable(
  data,
  hline_from = "STUDY",
  rm_dups = "STUDY",
) %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
	150 mg	89.4	24.4	122	4.63	1.12
	150 mg	81.7	34.4	104	3.83	0.910
	150 mg	94.0	27.4	93.2	4.94	1.25
	200 mg	67.9	27.5	100	4.25	1.10
	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
	100 mg	77.6	29.9	106	4.31	0.981
	50 mg	74.1	37.1	112	4.44	0.900
	50 mg	71.2	34.1	106	4.63	0.868
	75 mg	72.4	38.2	105	3.89	0.900
	75 mg	71.6	34.2	98.9	4.49	0.991
	75 mg	73.6	49.2	103	4.52	0.930

11 Panel

- Divide the table using column contents
- there's a bug somewhere there

```
stable(
  data,
  align = cols_center(DOSE = "l", SCR = "r"),
  panel = vars("Study number:" = STUDY)
) %>% pt_wrap(stdout())
```

DOSE	WT	AGE	CRCL	ALB	SCR				
12-DEMO-001									
100 mg	71.4	33.7	104	4.20	1.06				
150 mg	89.4	24.4	122	4.63	1.12				
150 mg	81.7	34.4	104	3.83	0.910				
150 mg	94.0	27.4	93.2	4.94	1.25				
200 mg	67.9	27.5	100	4.25	1.10				
200 mg	76.6	22.8	99.2	4.54	1.15				
12-DEM	O-002								
100 mg	61.3	38.3	113	4.04	1.28				
100 mg	77.6	29.9	106	4.31	0.981				
50 mg	74.1	37.1	112	4.44	0.900				
50 mg	71.2	34.1	106	4.63	0.868				
75 mg	72.4	38.2	105	3.89	0.900				
75 mg	71.6	34.2	98.9	4.49	0.991				
75 mg	73.6	49.2	103	4.52	0.930				

12 Colspan

• Group columns

```
stable(
  data,
  align = cols_left(.c = "WT,AGE,CRCL"),
  span = colgroup("In final model", WT:CRCL),
) %>% pt_wrap(stdout())
```

		In	final m	odel		
STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

13 Colspan - multiple

- Multiple groupings
- Multiple levels

```
stable(
  data,
  panel = list(Study = "STUDY"),
  align = cols_left(.c = "WT,AGE,CRCL"),
  units = ys_get_unit(spec, parens = TRUE),
  r_file = "foo.R", output_file = "foo.tex",
  span = list(
    colgroup("Meh", DOSE:WT),
    colgroup("Hrm", AGE:CRCL),
    colgroup("Huh", ALB:SCR),
    colgroup("Expert opinion", CRCL:SCR, level = 2)
  )
) %>% pt_wrap(stdout())
```

		ert opini	ion			
Mel	1	ŀ	Irm	Huh		
DOSE	WT (kg)	AGE (years)	CRCL (ml/min)	ALB (g/dL)	SCR (mg/dL)	
12-DEM	O-001					
100 mg	71.4	33.7	104	4.20	1.06	
150 mg	89.4	24.4	122	4.63	1.12	
150 mg	81.7	34.4	104	3.83	0.910	
150 mg	94.0	27.4	93.2	4.94	1.25	
200 mg	67.9	27.5	100	4.25	1.10	
200 mg	76.6	22.8	99.2	4.54	1.15	
12-DEM	O-002					
100 mg	61.3	38.3	113	4.04	1.28	
100 mg	77.6	29.9	106	4.31	0.981	
50 mg	74.1	37.1	112	4.44	0.900	
50 mg	71.2	34.1	106	4.63	0.868	
75 mg	72.4	38.2	105	3.89	0.900	
75 mg	71.6	34.2	98.9	4.49	0.991	
75 mg	73.6	49.2	103	4.52	0.930	

source code: foo.R source file: foo.tex

14 Colspan - from cols

• we have some columns of the form tag.name

```
dotdata <- readRDS("datasets/with-dots.RDS")</pre>
head(dotdata, n=2)
## # A tibble: 2 x 7
   STUDY
               Normal.WT Normal.CRCL Normal.ALB ESRD.WT ESRD.CRCL ESRD.ALB
##
     <chr>
                <chr>
                           <chr>
                                       <chr>
                                                  <chr>
                                                          <chr>
                                                                    <chr>>
## 1 12-DEMO-001 71.4
                                       4.20
                           104
                                                  78.5
                                                          26.0
                                                                    2.10
## 2 12-DEMO-001 81.7
                           104
                                       3.83
                                                  89.9
                                                          26.1
                                                                    1.92
stable(
  dotdata,
  panel = c(Study = "STUDY"),
 span_split = colsplit(sep = '.'),
 align = cols_left(.c = "WT,AGE,CRCL"),
 r_file = "foo.R", output_file = "foo.tex"
) %>% pt_wrap(stdout())
```

	Normal			ESRD						
WT	CRCL	ALB	WT	CRCL	ALB					
12-D	12-DEMO-001									
71.4	104	4.20	78.5	26.0	2.10					
81.7	104	3.83	89.9	26.1	1.92					
89.4	122	4.63	98.4	30.6	2.32					
94.0	93.2	4.94	103	23.3	2.47					
67.9	100	4.25	74.7	25.1	2.13					
76.6	99.2	4.54	84.2	24.8	2.27					
12-D	EMO-00	2								
77.6	106	4.31	85.4	26.4	2.16					
61.3	113	4.04	67.4	28.2	2.02					
71.2	106	4.63	78.3	26.4	2.32					
74.1	112	4.44	81.5	28.0	2.22					
71.6	98.9	4.49	78.8	24.7	2.25					
72.4	105	3.89	79.6	26.3	1.94					
73.6	103	4.52	81.0	25.8	2.26					

source code: foo.R source file: foo.tex

15 Handle wide columns

15.1 descr is taking over the table

```
ptable <- readRDS("datasets/ptable.RDS")
stable(ptable) %>% pt_wrap(stdout())
```

symbol	descr	estimate	standard.erro
CL (L)	Metabolic clearance in adults who graduated high school before 1973 and live in Muncie.	100	200

15.2 Limit descr to 5 cm

```
stable(
  ptable,
  align = cols_center(descr = col_ragged(5))
) %>% pt_wrap(stdout())
```

symbol	descr	estimate	standard.error	rse
CL (L)	Metabolic clearance in adults who graduated high school before 1973 and live in Muncie.	100	200	2

16 Identify a summary row

We can point to one or more rows and style it up as a "summary row"

```
df.total <- readRDS(file = "datasets/with-total.RDS")</pre>
df.total
## # A tibble: 3 x 6
              STUDY WT AGE CRCL ALB SCR <chr> <chr > <chr > <chr > <chr > <ch > 
##
## 1 12-DEMO-001 80.2 28.4 104 4.40 1.10
## 2 12-DEMO-002 71.7 37.3 106 4.33 0.979
## 3 all
                                          75.9 32.8 105 4.37 1.04
stable(
        df.total,
        units = ys_get_unit(spec, parens = TRUE),
        align = cols_center(STUDY = '1'),
        r_file = "foo.R", output_file = "output.tex",
        col_rename = c("Protocol no." = "STUDY"),
        notes = "WT: weight; CRCL: creatinine clearance",
         sumrows = sumrow(
                df.total$STUDY == "all",
               label = "All Studies",
                bold = TRUE
        )
) %>% pt_wrap(stdout())
```

Protocol no.	WT (kg)	AGE (years)	CRCL (ml/min)	ALB (g/dL)	SCR (mg/dL)
12-DEMO-001	80.2	28.4	104	4.40	1.10
12-DEMO-002	71.7	37.3	106	4.33	0.979
All Studies	75.9	32.8	105	4.37	1.04

WT: weight; CRCL: creatinine clearance

source code: foo.R source file: output.tex

17 Save the table to file

• There is a output file attribute on the text that is returned

```
tab <- stable(data, output_file = "foo.tex")
str(tab)

## 'stable' chr [1:27] "{\\def\\arraystretch{1.4}\\tabcolsep=5pt" ...
## - attr(*, "stable_file")= chr "foo.tex"

This can be used to save
stable_save(tab)

file.exists("foo.tex")

## [1] TRUE
Oops...
try(stable_save(as.character(tab)))</pre>
```

Error : bad input - there is no stable_file attribute; maybe this object was corrupted or it wasn't

18 Fontsize

• Where's my glasses?

data %>% stable(fontsize = "tiny") %>% pt_wrap(stdout())

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930

19 Row space

```
data %>% slice(1:3) %>% stable() %>% pt_wrap(stdout())
```

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910

data %>% slice(1:3) %>% stable(row_space = 0.9) %>% pt_wrap(stdout())

STUDY	DOSE				ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910

data %>% slice(1:3) %>% stable(row_space = 2) %>% pt_wrap(stdout())

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910

20 Col space

data %>% stable(col_space = 20) %>% pt_wrap(stdout())

STUDY	DOSE	WT	AGE	CRCL	ALB	SCR
12-DEMO-001	100 mg	71.4	33.7	104	4.20	1.06
12-DEMO-001	150 mg	89.4	24.4	122	4.63	1.12
12-DEMO-001	150 mg	81.7	34.4	104	3.83	0.910
12-DEMO-001	150 mg	94.0	27.4	93.2	4.94	1.25
12-DEMO-001	200 mg	67.9	27.5	100	4.25	1.10
12-DEMO-001	200 mg	76.6	22.8	99.2	4.54	1.15
12-DEMO-002	100 mg	61.3	38.3	113	4.04	1.28
12-DEMO-002	100 mg	77.6	29.9	106	4.31	0.981
12-DEMO-002	50 mg	74.1	37.1	112	4.44	0.900
12-DEMO-002	50 mg	71.2	34.1	106	4.63	0.868
12-DEMO-002	75 mg	72.4	38.2	105	3.89	0.900
12-DEMO-002	75 mg	71.6	34.2	98.9	4.49	0.991
12-DEMO-002	75 mg	73.6	49.2	103	4.52	0.930