

# rempsyc: Convenience functions for psychology

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#### Software

- Review 🗗
- Repository 🗗
- Archive ♂

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# Summary

{rempsyc} is an R package of convenience functions that make the analysis-to-publication workflow faster, easier, and less error-prone. It enables the creation of publication-ready APA (American Psychological Association) tables exportable to Word (via {flextable}) and easily customizable APA-compliant plots (via {ggplot2}). It makes it easy to run statistical tests, check assumptions, and automate various tasks common in psychology research and social sciences more broadly.

### Statement of need

There are many reasons to use R (R Core Team, 2022) for analyzing and reporting data from research studies, such as being compatible with the ideals of open science (Quintana, 2020). However, R has a major downside for novices: its steep learning curve due to its programmatic interface, in contrast to perhaps more user-friendly point-and-click software. Of course, this flexibility is also a strength, as the R community can and does come together to produce packages that make using R increasingly easier and more user-friendly (e.g., the *easystats* ecosystem Lüdecke et al., 2019/2023). The {rempsyc} package (Really Easy Methods for Psychology) contributes to this momentum by providing convenience functions that remove as much friction as possible between your script and your manuscript (in particular, if you are using Microsoft Word).

There are mainly three things that go into a manuscript: text, tables, and figures. {rempsyc} does not generate publication-ready text summarizing analyses; for this, see the {report} package (Makowski et al., 2021/2023). Instead, {rempsyc} focuses on the production of publication-ready tables and figures. Below, I go over a few quick examples of those.

# Examples Features

### Publication-Ready Tables

Many researchers using R still copy-paste the values from the R console to their manuscript, or retype them manually. Yet, this approach increases the risks of copy-paste and retyping errors so common in psychology. This problem is not trivial given that according to some estimates, up to 50% of articles in psychology have at least one statistical error (Nuijten et al., 2016). Ideally, one should be able to format the table directly in R, and to export it to Word directly.

Formatting a table properly in R is already a tedious and time-consuming task, but fortunately several packages take care of this step (e.g., the {broom} or {report} packages, Makowski et al., 2021/2023; Robinson et al., 2022, and there are several others). Exporting these formatted tables to Microsoft Word remains a challenge however. Some packages do export to Word (e.g., the {apaTables} package Stanley & Spence, 2018), but their formatting is often rigid especially when using analyzes or table formats that are not supported by default.



- {rempsyc} solves this problem by allowing maximum flexibility: you manually create the data frame exactly the way you want, and then only use the nice\_table() function on the resulting data frame. nice\_table() works on any data frame, even non-statistical ones like mtcars.
- One of its main benefits however is the automatic formatting of statistical symbols and its integration with other packages. We can for example create a {broom} table and then apply nice\_table() on it. It suits particularly well the pipe workflow.
- 45 library(rempsyc)
  46

```
lm(mpg ~ cyl + wt * hp, mtcars) |>
broom::tidy(conf.int = TRUE) |>
nice_table(broom = "lm")
```

Term	b	SE	t	p	95% CI
(Intercept)	49.49	3.66	13.51	<.001	[41.97, 57.01]
cyl	-0.37	0.51	-0.72	.479	[-1.41, 0.68]
wt	-7.63	1.52	-5.01	< .001	[-10.75, -4.51]
hp	-0.11	0.03	-3.64	.001	[-0.17, -0.05]
$\mathbf{wt} \times \mathbf{hp}$	0.03	0.01	3.23	.003	[0.01, 0.04]

51 We can do the same with a {report} table.

```
52 stats.table <- lm(mpg ~ cyl + wt * hp, mtcars) |>
53 report::report() |>
54 as.data.frame()
```

56 nice\_table(stats.table)



57

my\_table

Parameter	Fit	b	95% CI (b)	t	df	p	β	95% CI (β)
(Intercept)		49.49	[41.97, 57.01]	13.51	27	< .001	-0.18	[-0.36, -0.01]
cyl		-0.37	[-1.41, 0.68]	-0.72	27	.479	-0.11	[-0.42, 0.20]
wt		-7.63	[-10.75, -4.51]	-5.01	27	<.001	-0.62	[-0.85, -0.40]
hp		-0.11	[-0.17, -0.05]	-3.64	27	.001	-0.29	[-0.53, -0.04]
$wt \times hp$		0.03	[0.01, 0.04]	3.23	27	.003	0.29	[0.11, 0.47]
AIC	147.01							
AICc	150.37							
BIC	155.80							
R2	0.89							
R2 (adj.)	0.87							
Sigma	2.17							

The {report} package provides quite comprehensive tables, so one may request an abbreviated table with the 'short' argument. For convenience, it is also possible to highlight significant results for better visual discrimination, using the 'highlight' argument.[1] Once satisfied with the table, we can add a title and note.

my\_table <- nice\_table(
stats.table, short = TRUE, highlight = 0.001,
title = c("Table 1", "A Pretty Regression Model"),
note = c("The data was extracted from the 1974 Motor Trend US magazine.",
"Greyed rows represent statistically significant differences, p < .001."))



Table 1

A Pretty Regression Model

Parameter	b	t	df	p	β	95% CI (β)
(Intercept)	49.49	13.51	27	<.001	-0.18	[-0.36, -0.01]
cyl	-0.37	-0.72	27	.479	-0.11	[-0.42, 0.20]
wt	-7.63	-5.01	27	<.001	-0.62	[-0.85, -0.40]
hp	-0.11	-3.64	27	.001	-0.29	[-0.53, -0.04]
$wt \times hp$	0.03	3.23	27	.003	0.29	[0.11, 0.47]

Note. The data was extracted from the 1974 Motor Trend US magazine.

Greyed rows represent statistically significant differences, p < .001.

- One can then easily save the resulting table to Word with flextable::save\_as\_docx(), specifying the object name and desired path.
- 71 flextable::save\_as\_docx(my\_table, path = "my\_table.docx")
- Additionally, tables created with nice\_table() are {flextable} objects (Gohel & Skintzos,
- <sup>73</sup> 2022), and can be modified as such.[2]

# Formattting Results of Analyses

<sup>75</sup> {rempsyc} also provides its own set of functions to prepare statistical tables before they can be fed to nice\_table() and saved to Word.

# t tests

Dependent Variable	t	df	p	d	95% CI
mpg	-3.77	18.33	.001	-1.48	[-2.27, -0.67]
disp	4.20	29.26	<.001	1.45	[0.64, 2.23]
drat	-5.65	27.20	<.001	-2.00	[-2.86, -1.12]



### Contrasts

```
nice_contrasts(data = mtcars,
response = c("mpg", "disp"),
response =
```

Dependent Variable	Comparison	df	t	p	d	95% CI
	4 - 8	28	3.66	.001	3.59	[2.75, 4.46]
mpg	6 - 8	28	1.29	.207	1.44	[0.78, 2.00]
	4 - 6	28	3.64	.001	2.15	[1.35, 3.06]
	4 - 8	28	-6.04	<.001	-4.80	[-5.75, -3.83]
disp	6 - 8	28	-4.86	<.001	-3.29	[-4.24, -2.25]
	4 - 6	28	-2.70	.012	-1.51	[-2.23, -0.88]

# Regressions

```
data <- lapply(mtcars, scale)
model1 <- lm(mpg ~ disp + wt * hp, data)
model2 <- lm(qsec ~ drat + wt * hp, data)
my.models <- list(model1, model2)

nice_lm(my.models) |>
nice_table(highlight = TRUE)
```

Dependent Variable	Predictor	df	β	t	p	$sr^2$	95% CI
	disp	27	-0.06	-0.35	.728	.00	[0.00, 0.01]
	wt	27	-0.63	-4.39	<.001	.08	[0.00, 0.16]
mpg	hp	27	-0.32	-3.00	.006	.04	[0.00, 0.09]
	wt × hp	27	0.31	3.71	.001	.06	[0.00, 0.12]
	drat	27	-0.03	-0.18	.862	.00	[0.00, 0.01]
	wt	27	0.49	2.37	.025	.07	[0.00, 0.19]
qsec	hp	27	-1.05	-6.92	<.001	.62	[0.40, 0.83]
	$wt \times hp$	27	0.02	0.10	.922	.00	[0.00, 0.00]



### Simple Slopes

nice\_lm\_slopes(my.models, predictor = "wt", moderator = "hp") |>
nice\_table()

Dependent Variable	Predictor (+/-1 SD)	df	β	t	p	sr <sup>2</sup>	95% CI
	wt (LOW-hp)	27	-0.94	-5.60	< .001	.13	[0.02, 0.25]
mpg	wt (MEAN-hp)	27	-0.63	-4.39	< .001	.08	[0.00, 0.16]
	wt (HIGH-hp)	27	-0.32	-1.94	.063	.02	[0.00, 0.05]
	wt (LOW-hp)	27	0.47	1.56	.131	.03	[0.00, 0.10]
qsec	wt (MEAN-hp)	27	0.49	2.37	.025	.07	[0.00, 0.19]
	wt (HIGH-hp)	27	0.51	2.41	.023	.07	[0.00, 0.19]

#### Correlation Matrices

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It is also possible to export a colour-coded correlation matrix to Microsoft Excel. The cormatrix\_excel() function has several benefits over conventional approaches. The base R cor() function for example does not use rounded values and the console is impractical for large matrices. One may manually round values and export it to a .csv file, which is an improvement but still unsatisfying.

The {apaTables} package (Stanley & Spence, 2018) allows exporting the correlation matrix to Word in an APA format, and in many cases this already meets the formal requirements of APA style. However, the Word format is not suitable for large matrices, as it will often spread beyond the document's margin limits.

Another approach is to export the matrix to an image, like the {correlation} package does (Makowski et al., 2020).[3] For very small matrices, this works extremely well, and the colour is an immense help to quickly identify which correlations are strong or weak, positive or negative, and significant or non-significant. Again, however, this does not work so well for large matrices because labels might overlap or navigating the large figure becomes difficult.

When the goal is more exploratory in nature, and one has large matrices, it can be beneficial to export them to Excel. {rempsyc} combines the idea of using a coloured correlation matrix from the {correlation} package with the idea of exporting to Excel using {openxlsx2} (Barbone & Garbuszus, 2023).

{rempsyc} also provides some usability improvements, like freezing the first row and column so as to be able to easily see which variables correlate with which other variables, regardless of how far or deep those variables are located within the matrix.

The colour represents the strength of the correlation, whereas the stars represent different significance thresholds for the p value is.[4] The exact p values are provided in a second tab for reference purposes, so all information is readily available in just one function call.

```
cormatrix_excel(data = infert,
filename = "cormatrix1",
select = c("age", "parity", "induced", "case", "spontaneous",
"stratum", "pooled.stratum"))
```



	1	А	В	С	D	Е	F	G	н	1
	1	Paramete	age	parity	induced	case	spontaneo	stratum	pooled.str	atum
	2	age	1.0	.08	10	.0	08	21 ***	17 *	
	3	parity	.08	1.0	.45 ***	.01	.31 ***	31 ***	.12	
	4	induced	10	.45 ***	1.0	.02	27 ***	10	.16 *	
	5	case	.0	.01	.02	1.0	.36 ***	.0	.0	
	6	spontaneo	08	.31 ***	27 ***	.36 ***	1.0	.06	.21 ***	
	7	stratum	21 ***	31 ***	10	.0	.06	1.0	.75 ***	
	8	pooled.str	17*	.12	.16 *	.0	.21 ***	.75 ***	1.0	
	9		_							
122			r_value	p_valu	ues	$\oplus$				
133										
133	4	А	В	С	D	E	F	G	н	1
133	1	A Paramete	В				F spontaneo		H pooled.str	I ratum
133	1 2		В	С	D	Е				l ratum
133		Paramete	B age	C parity	D induced	E case	spontaneo	stratum	pooled.st	I ratum
133	2	Paramete age	B age .0	c parity .194	D induced .113	E case .956	spontaneo	stratum .001	pooled.str	ratum
133	2	Paramete age parity	B age .0 .194	C parity .194 .0	D induced .113	case .956 .889	spontaneo .186 .0	stratum .001 .0	,006 ,059	ratum
133	2 3 4 5	Paramete age parity induced	B age .0 .194 .113	c parity .194 .0	D induced .113 .0 .0	E case .956 .889 .789	spontaneo .186 .0 .0	stratum .001 .0 .113	,006 ,059 ,010	ratum
133	2 3 4 5 6	Paramete age parity induced case	B age .0 .194 .113	c parity .194 .0 .0	D induced .113 .0 .0	E case .956 .889 .789	.186 .0 .0	.001 .0 .113 .952	,006 ,059 ,010 ,939	ratum
133	2 3 4 5 6	Paramete age parity induced case spontaneo	B age .0 .194 .113 .956 .186 .001	C parity .194 .0 .0 .889	D induced .113 .0 .0 .789	E case .956 .889 .789 .0	.186 .0 .0 .0 .0	.001 .0 .113 .952	,006 ,059 ,010 ,939	ratum
133	2 3 4 5 6 7	Paramete age parity induced case spontaneo	B age .0 .194 .113 .956 .186 .001	C parity .194 .0 .0 .0 .889 .0 .0	D induced .113 .0 .0 .789 .0 .113	Case .956 .889 .789 .0 .0 .0 .952	spontaneo .186 .0 .0 .0 .0 .0	stratum .001 .0 .113 .952 .341	006 .059 .010 .939 .001 .0	ratum

# **Publication-Ready Figures**

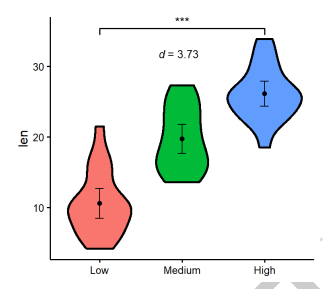
Preparing figures according to APA style, having them look good, and being able to save them in high-resolution with the proper ratios is often challenging. Working with {ggplot2} (Wickham, 2016) provides tremendous flexibility, but an unintended consequence is that doing even trivial operations can at times be daunting.

This is why {rempsyc} setups a few default plot types, ready to be saved to your preferred format (.pdf, .tiff, or .png).

### **Violin Plots**

```
nice_violin(data = ToothGrowth,
143
                 group = "dose",
144
                 response = "len"
145
                 xlabels = c("Low", "Medium", "High"),
146
                 comp1 = 1,
147
                 comp2 = 3,
148
                 has.d = TRUE,
149
                 d.y = 30)
150
```





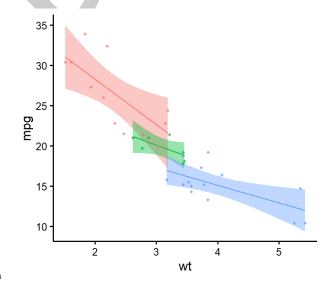
For an example of such use in publication, see Thériault et al. (2021).

One can easily save the resulting figure with ggplot2::ggsave(), specifying the desired file name, extension, and resolution.

Recommended dimensions for saving {rempsyc} figures is 7 inches wide and 7 inches high at 300 dpi, which makes sure that the resolution is high enough even if saving to non-vector graphics formats like .png. That said, scalable vector graphics formats like .pdf or .eps are still recommended for high-resolution submissions to scientific journals.

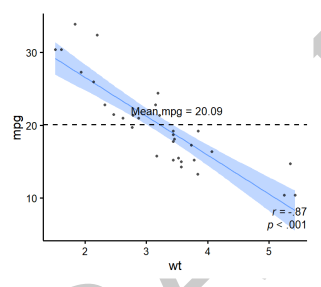
# Scatter Plots

Figures are {ggplot2} objects (Wickham, 2016), and can be modified as such.





```
nice_scatter(data = mtcars,
169
                  predictor = "wt",
170
                  response = "mpg",
171
                  has.confband = TRUE,
172
                  has.r = TRUE,
173
                  has.p = TRUE) +
174
      ggplot2::geom_hline(yintercept = mean(mtcars$mpg), colour = "black",
175
                           linewidth = 1.4, linetype = "dashed") +
176
      ggplot2::annotate("text", x = 3.5, y = 22, size = 7,
177
                         label = paste("Mean mpg =", round(mean(mtcars$mpg), 2)))
178
```



For an example of such use in publication, see Krol et al. (2020).

### Overlapping Circles

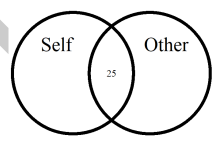
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For psychologists using the Inclusion of Other in the the Self Scale (Aron et al., 1992), it can be useful to interpolate the original discrete scores (1 to 7) into a group average representation of the conceptual self-other overlap. For example, assuming the group mean is 3.5 on the 1 to 7 scale, overlap\_circle() will draw a 25% overlap from interpolation:

86 overlap\_circle(3.5)



For an example of such use in publication, see Thériault et al. (2021).



### **Testing assumptions**

When comes time to test assumptions of a linear model, the best option is the check\_model()
function from easystats' {performance} package, which allows direct visual evaluation of
assumptions (Lüdecke, Ben-Shachar, et al., 2021). Indeed, visual assessment of diagnostic
plots is recommended over statistical tests since they are overpowered in large samples and
underpowered in small samples (Kozak & Piepho, 2018).

That said, if for whatever reason one wants to check objective asumption tests for a linear model, rempsyc makes this easy with the nice\_assumptions() function, which provide p values for normality (Shapiro-Wilk), homoscedasticity (Breusch-Pagan) and autocorrelation of residuals (Durbin-Watson) in one call.

### Categorical Predictors

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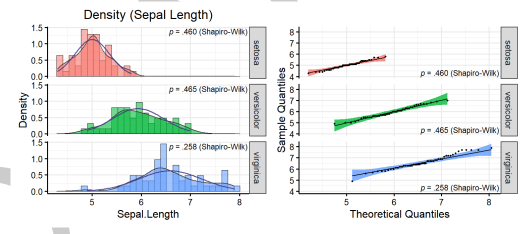
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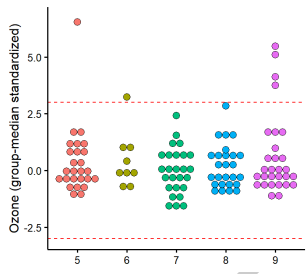
nice\_normality() makes it easy to visually check normality in the case of categorical predictors (i.e., when using groups), through a combination of quantile-quantile plots, density plots, and histograms.

```
nice_normality(data = iris,
variable = "Sepal.Length",
group = "Species",
shapiro = TRUE,
histogram = TRUE,
title = "Density (Sepal Length)")
```



Similarly for univariate outliers using median absolute deviations from the median (MAD, Leys et al., 2013).

```
plot_outliers(airquality,
group = "Month",
response = "Ozone")
```



```
215
    Univariate outliers based on the median/MAD can also be simply requested with find_mad().[5]
216
    find_mad(airquality, names(airquality), criteria = 3)
217
218
   ## 8 outlier(s) based on 3 median absolute deviations for variable(s):
219
   ##
        Ozone, Solar.R, Wind, Temp, Month, Day
220
   ##
221
   ## Outliers per variable:
   ##
223
       $0zone
   ##
224
   ##
         Row Ozone_mad
225
          30
              3.218284
   ## 1
226
          62
               3.989131
227
          99
              3.488081
       3
228
              3.025573
       4 101
229
   ## 5 117
               5.261028
   ## 6 121 3.333911
231
   ##
232
   ## $Wind
         Row Wind_mad
234
           9 3.049871
   ## 1
235
          48 3.225825
236
   Homoscedasticity can also be checked numerically with nice_var() or visually with
    nice_varplot().
238
   nice_var(data = iris,
239
              variable = names(iris[1:4]),
240
              group = "Species") |>
241
```

nice\_table()



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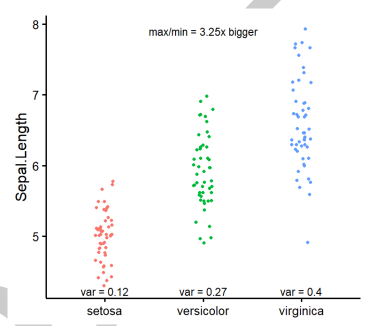
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Species	Setosa	Versicolor	Virginica	Variance.ratio	Criteria	Heteroscedastic
Sepal.Length	0.12	0.27	0.40	3.30	4.00	FALSE
Sepal.Width	0.14	0.10	0.10	1.50	4.00	FALSE
Petal.Length	0.03	0.22	0.30	10.20	4.00	TRUE
Petal.Width	0.01	0.04	0.07	6.80	4.00	TRUE

```
nice_varplot(data = iris,
variable = "Sepal.Length",
group = "Species")
```



# **Utility functions**

Finally, with the idea of making the analysis workflow easier in mind, {rempsyc} also provides a few other utility functions. nice\_na() allows reporting item-level missing values per scale, as well as participant's maximum number of missing items by scale, as per recommendations (Parent, 2013).

extract\_duplicates() creates a data frame of only observations with a duplicated ID or participant number, so they can be investigated more thoroughly. best\_duplicate() allows to follow-up on this investigation and only keep the "best" duplicate, meaning those with the fewer number of missing values, and in case of ties, the first one.

nice\_reverse() permits the automatic reverse-coding of scores so common for psychology questionnaires, provided the minimum and maximum score values are known.

There are other functions that the reader can explore at their leisure on the package official website. However, hopefully, this overview has given the reader a gentle introduction to this package.



# Licensing and Availability

The {rempsyc} package is licensed under the GNU General Public License (GPL v3.0). It is available on CRAN, and can be installed using install.packages("rempsyc"). The full tutorial website can be accessed at: https://rempsyc.remi-theriault.com/. All code is open-source and hosted on GitHub, and bugs can be reported at https://github.com/rempsyc/rempsyc/issues/.

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- [1] This argument can be used logically, as 'TRUE' or 'FALSE', but can also be provided with
- $_{324}$  a numeric value representing the cut-off threshold for the p value
- 225 [2] A great resource for this is the {flextable} e-book: https://ardata-fr.github.io/
- flextable-book/
- [3] Exporting the correlation matrix to an image through the {correlation} package also requires
- the {see} package (Lüdecke, Patil, et al., 2021)
- p [4] For convenience, colours are only used when the corresponding p value is at least smaller
- 330 than .05
- $_{331}$  [5] Once one has identified outliers, it is also possible to winsorize them with the
- winsorize\_mad() function.