# Analysis for the Nature of our Literature

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## Summary

Within this document I have included a summary of all the analyses included within our manuscript.

```
library(readr)
library(tidyverse)
library(tidyselect)
library(brms)
library(ggdist)
library(distributional)
library(broom)
library(ggpubr)
library(gtsummary)
library(labelled)
library(afex)
library(emmeans)
# Testing out data analysis
# import data -----
df_all = read_csv("df_all.csv") %>%
  mutate(support = factor(support,
                          levels = c("Unclear or not stated",
                                     "Not supported",
                                     "Partial support",
                                     "Full support"))) %>%
  # Coding error in 20 cases
  mutate(sig_test = ifelse(is.na(sig_test), "No", sig_test)) %>%
  mutate(sig_test = structure(factor(sig_test, levels = c("Yes", "No")),
                              label = "Significance Testing"),
         hypo_tested = structure(factor(hypo_tested, levels = c("Yes", "No")),
                                 label = "Hypothesis Tested"),
         effect_size = structure(factor(effect_size, levels = c("Yes","No")),
                                 label = "Effect Size"))
# Get hypothesis tested set -----
hyp tested = df all %>%
  filter(hypo_tested == "Yes")
```

#### Introduction

In this study we collected data on 300 sport and exercise science research articles (100 from 3 journals). Based on the work of Büttner et al. (2020), we anticipated at least 150 (50%) of the articles would include a hypothesis that was tested. Based on the work of Fanelli (2010), Scheel, Schijen, and Lakens (2021), and Büttner et al. (2020) we hypothesized that the percentage of articles that find support for their hypothesis was greater than 80%.

# Hypothesis

For this study, we hypothesized that the rate of positive results (i.e., studies that find at least partial support for their hypothesis) was greater than 80%. Therefore, the null hypothesis ( $H_0$ ) was that the proportion of positive results was less than .8 and our alternative was greater than .8. There was no other effect being estimated in this study therefore the intercept of the model is what will be tested.

 $H_0: Intercept \leq 0.8$ 

 $H_1: Intercept > 0.8$ 

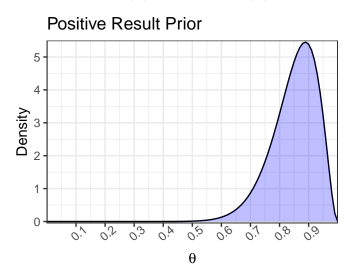
We also hypothesized that more than 60% of studies would test a hypothesis.

 $H_0: Intercept \leq 0.6$ 

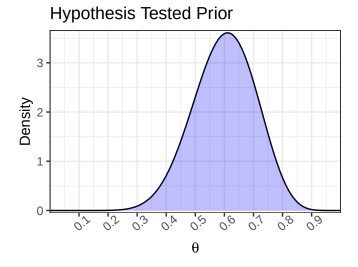
 $H_1: Intercept > 0.6$ 

## **Prior Choice**

The prior we selected for this analysis was informed by the previous studies assuming the true positive rate is approximately 85% (Fanelli 2010). However, we would like to avoid "spiking" the prior in favor of our hypothesis and therefore want a skeptical prior. Based on the work of Scheel, Schijen, and Lakens (2021) and Büttner et al. (2020) the estimated positive rates in original research investigations ranged from 82%-92%, and even some fields included in the survey by Fanelli (2010) observed rates as low as ~70%. Therefore, we selected a prior of  $\beta(17,3)$ , and is visualized it below. This prior is centered around .85, but includes the possibility of higher (.9) and much lower (.7) proportions as compatible parameter estimates.



Similarly, we used prior centered at 60% for the secondary hypothesis test.



## **Bayesian Models**

We constructed our Bayesian models with the following code.

```
# ANALYSIS 1: Positive Result Rate -----
#Set prior
prior_1 = set_prior("beta(17, 3)", class = "b", lb = 0, ub = 1)
#transform data
hyp_pos = hyp_tested %>%
 summarize(
   pos = sum(di_sup == "Y"),
   N = length(di_sup),
   rate = pos/N
  )
#Build model
m_final <- brm(</pre>
 pos | trials(N) ~ 0 + Intercept,
 family = binomial(link = "identity"),
 prior = prior_1,
  data = hyp_pos, sample_prior = "yes", refresh = 0
write_rds(m_final, "m_final.rds")
# ANALYSIS 2: Hypothesis Test Rate -----
#Set prior
prior_2 = set_prior("beta(12, 8)", class = "b", lb = 0, ub = 1)
#Generate test data
hyp_test = df_all %>%
  summarize(
   pos = sum(hypo_tested == "Yes"),
   N = length(hypo_tested),
   rate = pos/N
#Build model
m_final2 <- brm(</pre>
 pos | trials(N) ~ 0 + Intercept,
 family = binomial(link = "identity"),
 prior = prior_2,
  data = hyp_test, sample_prior = "yes", refresh = 0
```

Now, we can import the Bayesian models to get the output for the manuscript.

```
# Import brms analysis 1 (positive) ------
m_final = read_rds("m_final.rds")
# Import brms analysis 2 (prop. of hyp. tested) ------
m_final2 = read_rds("m_final2.rds")

h_test <- hypothesis(m_final, "Intercept > 0.8")
knitr::kable(h_test$hypothesis, caption = "Hypothesis Test #1")
```

Table 1: Hypothesis Test #1

Hypothesis	Estimate	Est.Error	CI.Lower	CI.Upper	Evid.Ratio	Post.Prob	Star
$\overline{\text{(Intercept)-(0.8)} > 0}$	0.0142668	0.0269525	-0.0319772	0.0568162	2.427592	0.70825	

Table 2: Hyp Test #1: 95% Posterior C.I.

	2.5%	97.5%
b_Intercept	0.7577946 $0.6781015$	0.8629953 0.9662099
prior_b lp	-5.5422325	-3.1629022

```
h_test2 <- hypothesis(m_final2, "Intercept > 0.6")
knitr::kable(h_test$hypothesis, caption = "Hypothesis Test #2")
```

Table 3: Hypothesis Test #2

Hypothesis	Estimate	Est.Error	CI.Lower	CI.Upper	Evid.Ratio	Post.Prob	Star
$\overline{\text{(Intercept)-(0.8)} > 0}$	0.0142668	0.0269525	-0.0319772	0.0568162	2.427592	0.70825	

Table 4: Hyp Test #2: 95% Posterior C.I.

	2.5%	97.5%
b_Intercept prior b	0.5811895 $0.3766667$	0.6897217 $0.8015812$
lp	-5.9064835	-3.2418816

```
# Main Figures
dat mfinal = posterior samples(m final, "b") %>%
  mutate(Test = "Positive Result Rate")
dat_mfinal2 = posterior_samples(m_final2, "b") %>%
  mutate(Test = "Rate of Hypothesis Tests")
df_mfinal = rbind(dat_mfinal, dat_mfinal2)
# figure 1
p_f1a = df_mfinal %>%
  ggplot(aes(x=b_Intercept,
             fill = Test)) +
  stat_halfeye(alpha = .75) +
  labs(fill = "Interval",
       x = "Probability",
       y = "") +
  theme_bw() +
  facet_wrap(~Test) +
  scale_fill_manual(values =c("lightgreen","skyblue2")) +
  theme(legend.position = "none",
        axis.text.y=element_blank(),
       axis.ticks.y=element_blank(),
        text = element_text(size = 14,
                            face = "bold"))
fig_1b = df_all %>%
  select(support) %>%
  drop_na() %>%
  ggplot(aes(support,
            fill = support)) +
  geom_bar(aes(y = (..count..) / sum(..count..)),
           color = "black") +
  scale_y_continuous(labels = scales::percent,
                     limits = c(0,.5),
                     breaks = c(0,.1,.2,.3,.4,.5),
                     expand = c(0,0)) +
  labs(x = "Level of Hypothesis Support",
       y = "Relative Frequency",
       fill = "Hypothesis Support") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "none")
fig_1c = df_all %>%
  select(hypo_tested) %>%
  drop_na() %>%
  ggplot(aes(hypo_tested,
             fill = hypo_tested)) +
  geom_bar(aes(y = (..count..) / sum(..count..)),
           color = "black") +
  scale_y_continuous(labels = scales::percent,
                     limits = c(0, .75),
```

# Levels of Support

```
tab_sup = df_all %>%
  select(hypo_tested, support) %>%
  drop_na() %>%
  group_by(support) %>%
  summarize(n = n())
tots_sup = sum(tab_sup$n)
tab_sup$percent = tab_sup$n/tots_sup*100
knitr::kable(tab_sup)
```

support	n	percent
Unclear or not stated	13	6.806283
Not supported	34	17.801047
Partial support	55	28.795811
Full support	89	46.596859

# **Exploratory Analyses**

### Statistics by Hypothesis Tested

```
ctab1 = df_all %>%
  select(effect_size, sig_test,
        hypo_tested, pval_type,
        n_just) %>%
  mutate(pval_type = structure(pval_type, label = "P-value reported"),
        n_just = structure(n_just,
                            label = "Sample Size Justification")) %>%
  tbl_summary(by = hypo_tested,
              type = list(sig_test ~ "categorical",
                          effect_size ~ "categorical",
                          n_just ~ "categorical")) %>%
  # add_p(pvalue_fun = ~style_pvalue(.x, digits = 2)) %>%
  add_overall() %>% add_n() %>% modify_header(label ~ "**Variable**") %>%
  modify_spanning_header(c("stat_1", "stat_2") ~ "**Hypothesis Tested**") %>%
  bold_labels()
ctab1
```

Variable	$\mathbf{N}$	Overall, $N = 300$	$\mathbf{Yes},  N = 191$	$\mathbf{No}, N = 109$
Effect Size	300			
Yes		238 (79%)	159 (83%)	79 (72%)
No		62 (21%)	32 (17%)	30 (28%)
Significance Testing	300			
Yes		270 (90%)	181 (95%)	89 (82%)
No		30 (10%)	10(5.2%)	20 (18%)
P-value reported	270	, ,	, ,	, ,
A mix: some exact and relative p-value(s)		151 (56%)	113 (62%)	38 (43%)
Exact (e.g., $p = .049$ )		91 (34%)	52 (29%)	39 (44%)
Relative (e.g., $p < .05$ )		28 (10%)	16 (8.8%)	12 (13%)
Unknown		30	10	20
Sample Size Justification	300			
No		232 (77%)	140 (73%)	92 (84%)
Yes		68 (23%)	51 (27%)	17 (16%)

#### Effect Size

```
ct_effsize = table(df_all$effect_size)
ct_effsize
##
## Yes No
## 238 62
binom_eff = binom.test(ct_effsize[1], sum(ct_effsize))
binom_eff
##
## Exact binomial test
##
## data: ct_effsize[1] and sum(ct_effsize)
## number of successes = 238, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.7430468 0.8377215
## sample estimates:
## probability of success
               0.7933333
eff_pr = paste0(round(binom_eff$estimate*100,2),"\\% [",
                   round(binom_eff$conf.int[1]*100,2),", ",
                   round(binom_eff$conf.int[2]*100,2)
                   ,"]")
eff_pr
```

## Significance Testing

```
ct_sig = table(df_all$sig_test)
ct_sig
##
## Yes No
## 270 30
binom_sig = binom.test(ct_sig[1], sum(ct_sig))
binom_sig
##
##
  Exact binomial test
##
## data: ct_sig[1] and sum(ct_sig)
## number of successes = 270, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.8603267 0.9315083
## sample estimates:
## probability of success
##
sig_pr = paste0(round(binom_sig$estimate*100,2),"\\% [",
                round(binom_sig$conf.int[1]*100,2),", ",
                round(binom_sig$conf.int[2]*100,2)
                ,"]")
sig_pr
```

## Significance Testing with NO Hypothesis Tested

```
ct_sig2 = table(subset(df_all, hypo_tested == "No")$sig_test)
ct_sig2
##
## Yes No
## 89 20
binom_sig2 = binom.test(ct_sig2[1], sum(ct_sig2))
binom_sig2
##
  Exact binomial test
##
##
## data: ct_sig2[1] and sum(ct_sig2)
## number of successes = 89, number of trials = 109, p-value = 1.418e-11
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.7309446 0.8841692
## sample estimates:
## probability of success
                0.8165138
##
sig_pr2 = paste0(round(binom_sig2$estimate*100,2),"\\% [",
                round(binom_sig2$conf.int[1]*100,2),", ",
                round(binom_sig2$conf.int[2]*100,2)
                ,"]")
sig_pr2
```

## Significance Testing with Hypothesis Tested

```
ct_sig3 = table(subset(df_all, hypo_tested == "Yes")$sig_test)
ct_sig3
##
## Yes No
## 181 10
binom_sig3 = binom.test(ct_sig3[2], sum(ct_sig3))
binom_sig3
##
  Exact binomial test
##
##
## data: ct_sig3[2] and sum(ct_sig3)
## number of successes = 10, number of trials = 191, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.02538906 0.09417495
## sample estimates:
## probability of success
               0.05235602
sig_pr3 = paste0(round(binom_sig3$estimate*100,2),"\\% [",
                 round(binom_sig3$conf.int[1]*100,2),", ",
                 round(binom_sig3$conf.int[2]*100,2)
                 ,"]")
sig_pr3
```

## Manuscripts Reporting "Exact" p-values

```
ct_ptype = table(df_all$pval_type)
ct_ptype
##
## A mix: some exact and relative p-value(s)
                      Exact (e.g., p = .049)
##
##
                    Relative (e.g., p < .05)
##
##
binom_ptype = binom.test(ct_ptype[[2]],sum(ct_ptype))
binom_ptype
##
## Exact binomial test
##
## data: ct_ptype[[2]] and sum(ct_ptype)
## number of successes = 91, number of trials = 270, p-value = 9.265e-08
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.2808696 0.3968281
## sample estimates:
## probability of success
                 0.337037
ptype_pr = paste0(round(binom_ptype$estimate*100,2),"\\% [",
                  round(binom_ptype$conf.int[1]*100,2),", ",
                  round(binom_ptype$conf.int[2]*100,2)
                  ,"] of manuscripts reported exact p-values for all results (p = .045) versus only rel
ptype_pr
## [1] "33.7\\% [28.09, 39.68] of manuscripts reported exact p-values for all results (p = .045) versus
```

## Manuscripts Reporting "Exact" or "Mixed" p-value types

```
ct_ptype2 = table(df_all$pval_type)
ct_ptype2
##
## A mix: some exact and relative p-value(s)
##
                      Exact (e.g., p = .049)
##
##
                    Relative (e.g., p < .05)
##
##
binom_ptype2 = binom.test(ct_ptype2[[2]]+ct_ptype2[[1]],sum(ct_ptype2))
binom_ptype2
##
## Exact binomial test
##
## data: ct_ptype2[[2]] + ct_ptype2[[1]] and sum(ct_ptype2)
## number of successes = 242, number of trials = 270, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.8536141 0.9299761
## sample estimates:
## probability of success
                0.8962963
ptype_pr2 = paste0(round(binom_ptype2$estimate*100,2),"\\% [",
                   round(binom_ptype2$conf.int[1]*100,2),", ",
                   round(binom_ptype2$conf.int[2]*100,2)
                   ,"] of manuscripts reported at least *some* exact p-values (e.g., p = .045) versus r
ptype_pr2
## [1] "89.63\\% [85.36, 93] of manuscripts reported at least *some* exact p-values (e.g., p = .045) ve
```

# Preregistration by Study Type

```
ctab2 = df_all %>%
 select(clin_trial, rct, animal, prereg) %>%
 mutate(clin_trial = structure(clin_trial,
                              label = "Clinical Trial"),
        rct = structure(rct,
                      label = "RCT"),
        animal = structure(animal,
                         label = "Animal Study")) %>%
 tbl_summary(by = prereg,
              type = list(clin_trial ~ "categorical",
                         rct ~ "categorical",
                          animal ~ "categorical")) %>%
  # add_p(pvalue_fun = ~style_pvalue(.x, digits = 2)) %>%
 add_overall() %>% add_n() %>% modify_header(label ~ "**Variable**") %>%
 modify_spanning_header(c("stat_1", "stat_2") ~ "**Preregistration**") %>%
 bold_labels()
ctab2
```

Variable	$\mathbf{N}$	Overall, $N = 300$	No, N = 273	$\mathbf{Yes},  N = 27$
Clinical Trial	300			
No		260 (87%)	256 (94%)	4(15%)
Yes		40 (13%)	17 (6.2%)	23 (85%)
RCT	300	, ,	, ,	, ,
No		236 (79%)	227 (83%)	9 (33%)
Yes		64 (21%)	46 (17%)	18 (67%)
Animal Study	300	,	, ,	, ,
No		291 (97%)	264 (97%)	27 (100%)
Yes		9 (3.0%)	9 (3.3%)	0 (0%)

#### Preregistration Rate

```
ct_prereg = table(df_all$prereg)
ct_prereg
##
## No Yes
## 273 27
binom_prereg = binom.test(ct_prereg[[2]],sum(ct_prereg))
binom_prereg
##
## Exact binomial test
##
## data: ct_prereg[[2]] and sum(ct_prereg)
## number of successes = 27, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.06014953 0.12824251
## sample estimates:
## probability of success
prereg_pr = paste0(round(binom_prereg$estimate*100,2),"\\% [",
                   round(binom_prereg$conf.int[1]*100,2),", ",
                   round(binom_prereg$conf.int[2]*100,2)
                   ,"] of manuscripts reporting preregistration or clinical trial registration informat
prereg_pr
```

## [1] "9\\% [6.01, 12.82] of manuscripts reporting preregistration or clinical trial registration info

# Sample Size by Journal

Variable	N	Overall, N = 300	<b>EJSS</b> , N = 100	<b>JSAMS</b> , N = 100	<b>MSSE</b> , N = 100
Sample Size Justification	300				
No		232~(77%)	81 (81%)	86 (86%)	65~(65%)
Yes		68~(23%)	19 (19%)	14 (14%)	35 (35%)
Sample Size Information	300				
No (sample size(s) unclear/unreported)		5 (1.7%)	0 (0%)	4 (4.0%)	1 (1.0%)
Partial (some information missing)		2(0.7%)	0 (0%)	1 (1.0%)	1(1.0%)
Yes (e.g., total and group sample sizes provided)		293 (98%)	100 (100%)	95 (95%)	98 (98%)

## Sample Size Justification

```
ct_njust = table(df_all$n_just)
ct_njust
##
## No Yes
## 232 68
binom_njust = binom.test(ct_njust[[2]], sum(ct_njust))
binom_njust
##
## Exact binomial test
##
## data: ct_njust[[2]] and sum(ct_njust)
## number of successes = 68, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.1805308 0.2782908
## sample estimates:
## probability of success
##
                0.2266667
njust_pr = paste0(
 round(binom_njust$estimate * 100, 2),
  "% [",
  round(binom_njust$conf.int[1] * 100, 2),
  round(binom_njust$conf.int[2] * 100, 2)
  "]"
)
njust_pr
```

```
## [1] "22.67% [18.05, 27.83]"
```

### Sample Size Information

```
ct_samp = table(df_all$sample_info)
ct samp
##
              No (sample size(s) unclear/unreported)
##
##
##
                  Partial (some information missing)
##
## Yes (e.g., total and group sample sizes provided)
binom_samp = binom.test(ct_samp[[3]], sum(ct_samp))
binom_samp
##
##
   Exact binomial test
##
## data: ct_samp[[3]] and sum(ct_samp)
## number of successes = 293, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.9525174 0.9905685
## sample estimates:
## probability of success
                0.9766667
samp_pr = paste0(
  round(binom_samp$estimate * 100, 2),
  "% [",
 round(binom_samp$conf.int[1] * 100, 2),
 round(binom_samp$conf.int[2] * 100, 2)
  "] of manuscripts reported all the required sample size information (total and group sample sizes)."
samp_pr
```

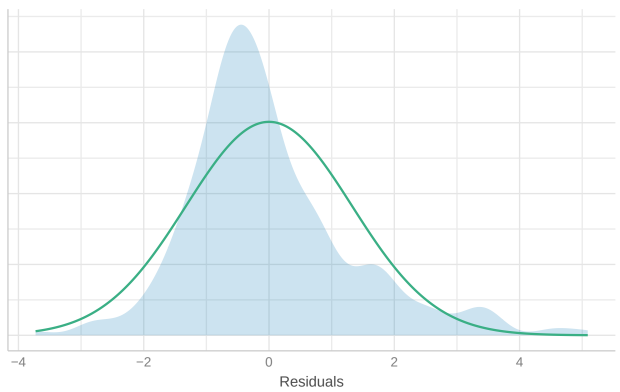
## Sample Size by Discipline

```
aov_1way = df_all %>%
  select(n, sci_cat, doi) %>% mutate(n = as.numeric(n)) %>% drop_na() %>%
  afex::aov_4(formula = log(n) ~ sci_cat + (1|doi))
# Not perfect fit but close
# fit is best with log transform compared to others
# See MASS::boxcox(aov_1way); very close to best fit
plot(performance::check_normality(aov_1way))
```

## Warning: Non-normality of residuals detected (p < .001).

#### Normality of Residuals

Distribution should be close to the normal curve



knitr::kable(nice(aov\_1way))

Effect	df	MSE	F	ges	p.value
sci_cat	9, 285	1.81	21.81 ***	.408	<.001

```
emm_samps = emmeans::emmeans(aov_1way, ~ sci_cat, type = "response")
knitr::kable(emm_samps)
```

sci_cat	response	SE	df	lower.CL	upper.CL
Applied exercise physiology (human)	30.06059	4.285230	285	22.705895	39.79756
Basic physiology (animal and cell physiology)	35.03300	15.704629	285	14.496810	84.66075

sci_cat	response	SE	df	lower.CL	upper.CL
Biomechanics	25.68742	5.093465	285	17.386780	37.95089
Clinical research	41.54535	10.375167	285	25.412325	67.92044
Environmental physiology (heat, cold, &	15.86997	6.749130	285	6.871256	36.65356
altitude)					
Epidemiology	1161.57881	306.361126	285	691.179304	1952.12057
Motor learning/control/behavior	68.17904	21.035155	285	37.145928	125.13838
Other	165.97110	42.955841	285	99.721555	276.23320
Sport performance	50.48799	12.831597	285	30.614795	83.26161
Sport/exercise psychology	66.11571	25.667626	285	30.792118	141.96125

# Non-Parametric Analysis

```
kruskal.test(as.numeric(n) ~ sci_cat, data = df_all)
```

```
##
## Kruskal-Wallis rank sum test
##
## data: as.numeric(n) by sci_cat
## Kruskal-Wallis chi-squared = 93.078, df = 9, p-value = 3.919e-16
```

## Other Open Science Practices

#### **Data Availability Statements**

```
ct_datstat = table(df_all$data_state)
ct_datstat
##
## No Yes
## 293
binom_datstat = binom.test(ct_datstat[[2]],300)
binom_datstat
##
## Exact binomial test
##
## data: ct_datstat[[2]] and 300
## number of successes = 7, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.009431502 0.047482565
## sample estimates:
## probability of success
               0.02333333
datstat_pr = paste0(round(binom_datstat$estimate*100,2),"% [",
                 round(binom_datstat$conf.int[1]*100,2),", ",
                 round(binom_datstat$conf.int[2]*100,2)
                 ,"] of manuscripts had a data accessibility statement")
datstat_pr
```

## [1] "2.33% [0.94, 4.75] of manuscripts had a data accessibility statement"

## Open Data Availability

```
ct_odat = table(df_all$open_data)
ct_odat
##
## No Yes
##
    5
binom_odat = binom.test(ct_odat[[2]],300)
binom_odat
##
## Exact binomial test
##
## data: ct_odat[[2]] and 300
## number of successes = 2, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.0008083864 0.0238734976
## sample estimates:
## probability of success
              0.006666667
odat_pr = paste0(round(binom_odat$estimate*100,2),"% [",
                 round(binom_odat$conf.int[1]*100,2),", ",
                 round(binom_odat$conf.int[2]*100,2)
                 ,"] of manuscripts reported some form of data sharing or open data")
odat_pr
```

## [1] "0.67% [0.08, 2.39] of manuscripts reported some form of data sharing or open data"

# **Replication Studies**

```
ct_replic = table(df_all$replic)
ct_replic
##
## No
## 300
binom_replic = binom.test(0,300)
binom_replic
##
## Exact binomial test
##
## data: 0 and 300
## number of successes = 0, number of trials = 300, p-value < 2.2e-16
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.0000000 0.01222097
## sample estimates:
## probability of success
##
replic_pr = paste0(
  round(binom_replic$estimate * 100, 2),
  round(binom_replic$conf.int[1] * 100, 2),
 round(binom_replic$conf.int[2] * 100, 2)
  "] of manuscripts explicitly stated they were replicating a previous study."
replic_pr
```

## [1] "0% [0, 1.22] of manuscripts explicitly stated they were replicating a previous study."

# Analysis by Journal

## **Hypothesis Support**

```
## Hypothesis Support ----
tab_jhyp = table(df_all$journal,df_all$support)
tab_jhyp
##
           Unclear or not stated Not supported Partial support Full support
##
##
    EJSS
                                                            19
##
     JSAMS
                               2
                                            8
                                                            17
                                                                         19
    MSSE
                               6
                                            13
                                                            19
                                                                         36
##
chisq_support = chisq.test(tab_jhyp)
chisq_support
##
## Pearson's Chi-squared test
##
## data: tab_jhyp
## X-squared = 2.4004, df = 6, p-value = 0.8794
```

# Hypothesis Tested

```
## Hypothesis Tested -----
tab_jtest = table(df_all$journal,df_all$hypo_tested)
tab_jtest
##
##
          Yes No
## EJSS 71 29
##
    JSAMS 46 54
    MSSE 74 26
##
chisq_jtest = chisq.test(tab_jtest)
chisq_jtest
## Pearson's Chi-squared test
##
## data: tab_jtest
## X-squared = 20.433, df = 2, p-value = 3.656e-05
# BayesFactor::contingencyTableBF(tab_jtest, sampleType = "poisson")
```

# Significance Testing

```
## Significance Testing -----
tab_jsig = table(df_all$journal,df_all$sig_test)
tab_jsig
##
##
         Yes No
## EJSS 92 8
##
    JSAMS 84 16
##
    MSSE 94 6
chisq_jsig = chisq.test(tab_jsig)
chisq_jsig
## Pearson's Chi-squared test
##
## data: tab_jsig
## X-squared = 6.2222, df = 2, p-value = 0.04455
```

#### Effect Sizes

```
## Effect Size ----
tab_jes = table(df_all$journal,df_all$effect_size)
tab_jes
##
##
         Yes No
## EJSS 90 10
##
    JSAMS 72 28
    MSSE 76 24
##
chisq_jes = chisq.test(tab_jes)
chisq_jes
## Pearson's Chi-squared test
##
## data: tab_jes
## X-squared = 10.897, df = 2, p-value = 0.004302
```

# Sample Size Justification

```
## Sample Size Justification ----
tab_jjust = table(df_all$journal,df_all$n_just)
tab_jjust
##
##
         No Yes
## EJSS 81 19
##
    JSAMS 86 14
    MSSE 65 35
chisq_jjust = chisq.test(tab_jjust)
chisq_jjust
## Pearson's Chi-squared test
##
## data: tab_jjust
## X-squared = 13.73, df = 2, p-value = 0.001044
```

# Analysis among Clinical Trials

### Hypothesis Support

```
# Clinical Trial breakdown -----
df_clin = subset(df_all, clin_trial == "Yes")
## Hypothesis Support (di)
tab_clindisup = table(df_clin$di_sup)
tab_clindisup
##
## N Y
## 9 16
binom_clindisup = binom.test(tab_clindisup[2], sum(tab_clindisup),
binom_clindisup
##
## Exact binomial test
##
## data: tab_clindisup[2] and sum(tab_clindisup)
## number of successes = 16, number of trials = 25, p-value = 0.07416
## alternative hypothesis: true probability of success is not equal to 0.8
## 95 percent confidence interval:
## 0.4252063 0.8202832
## sample estimates:
## probability of success
                     0.64
tab_clinsup = table(df_clin$support)
knitr::kable(tab_clinsup)
```

Var1	Freq
Unclear or not stated	$\frac{}{2}$
Not supported	9
Partial support	6
Full support	10

### Hypothesis Tested

```
## Hypothesis Tested
tab_clinhypo = table(df_clin$hypo_tested)
tab_clinhypo
##
## Yes No
## 27 13
binom_clinhypo = binom.test(tab_clinhypo[2], sum(tab_clinhypo),
binom_clinhypo
##
## Exact binomial test
##
## data: tab_clinhypo[2] and sum(tab_clinhypo)
## number of successes = 13, number of trials = 40, p-value = 0.0005468
## alternative hypothesis: true probability of success is not equal to 0.6
## 95 percent confidence interval:
## 0.1857290 0.4912949
## sample estimates:
## probability of success
##
                    0.325
Sample Size Justification
## Sample Size Just -----
tab_clinjust = table(df_clin$n_just)
tab_clinjust
##
##
  No Yes
## 15 25
binom_clinjust = binom.test(tab_clinjust[2], sum(tab_clinjust))
binom clinjust
##
## Exact binomial test
## data: tab_clinjust[2] and sum(tab_clinjust)
## number of successes = 25, number of trials = 40, p-value = 0.1539
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.4580148 0.7727373
## sample estimates:
## probability of success
```

0.625

### Preregistration

```
tab_clinreg = table(df_clin$prereg)
binom_clinreg = binom.test(tab_clinreg[2], sum(tab_clinreg))
### by journal ----
tab_clinregj = table(df_clin$prereg, df_clin$journal)
```

## Analysis by RCT

#### Hypothesis Support

```
# RCT breakdown -----
df_rct = subset(df_all, rct == "Yes")
## Hypothesis Support (di)
tab_rctdisup = table(df_rct$di_sup)
tab_rctdisup
##
## N Y
## 11 33
binom_rctdisup = binom.test(tab_rctdisup[2], sum(tab_rctdisup),
                            p = .8
binom_rctdisup
##
## Exact binomial test
##
## data: tab_rctdisup[2] and sum(tab_rctdisup)
## number of successes = 33, number of trials = 44, p-value = 0.4493
## alternative hypothesis: true probability of success is not equal to 0.8
## 95 percent confidence interval:
## 0.5966196 0.8680726
## sample estimates:
## probability of success
##
                     0.75
tab_rctsup = table(df_rct$support)
knitr::kable(tab_rctsup)
```

Var1	Freq
Unclear or not stated	3
Not supported	11
Partial support	13
Full support	20

# Hypothesis Tested

## Sample Size Justification

```
## Sample Size Just -----
tab_rctjust = table(df_rct$n_just)
tab_rctjust
##
## No Yes
## 32 32
binom_rctjust = binom.test(tab_rctjust[2], sum(tab_rctjust))
binom_rctjust
##
## Exact binomial test
## data: tab_rctjust[2] and sum(tab_rctjust)
## number of successes = 32, number of trials = 64, p-value = 1
## alternative hypothesis: true probability of success is not equal to 0.5
## 95 percent confidence interval:
## 0.3723229 0.6276771
## sample estimates:
## probability of success
##
                     0.5
```

# Preregistation

```
tab_rctreg = table(df_rct$prereg)
binom_rctreg = binom.test(tab_rctreg[2], sum(tab_rctreg))
```

# Analysis by Discipline

```
tab_dissupp = table(df_all$sci_cat, df_all$support)
tab_dissupp
```

## ## ## ## ## ## ##	Applied exercise physiology (human) Basic physiology (animal and cell physiology) Biomechanics Clinical research Environmental physiology (heat, cold, & altitude) Epidemiology Motor learning/control/behavior Other Sport performance Sport/exercise psychology	Unclear or not stated 3 3 5 1 0 0 0 0 1
##		
##		Not supported
##	Applied exercise physiology (human)	18
##	Basic physiology (animal and cell physiology)	0
## ##	Biomechanics Clinical research	4 4
##	Environmental physiology (heat, cold, & altitude)	0
##	Epidemiology	1
##	Motor learning/control/behavior	0
##	Other	1
##	Sport performance	4
##	Sport/exercise psychology	2
##	1 7 50	
##		Partial support
##	Applied exercise physiology (human)	16
##	Basic physiology (animal and cell physiology)	1
##	Biomechanics	14
##	Clinical research	4
##	Environmental physiology (heat, cold, & altitude)	4
##	Epidemiology	1
##	Motor learning/control/behavior	3
##	Other	3
##	Sport performance	6
##	Sport/exercise psychology	3
## ##		
## ##		Full support
##	Applied evercise physiology (human)	Full support
##	Applied exercise physiology (human)  Rasic physiology (animal and cell physiology)	27
## ##	Applied exercise physiology (human) Basic physiology (animal and cell physiology) Biomechanics	

```
Environmental physiology (heat, cold, & altitude)
##
##
    Epidemiology
                                                                 5
    Motor learning/control/behavior
                                                                 9
##
##
    Other
                                                                 8
    Sport performance
                                                                 7
##
    Sport/exercise psychology
                                                                 2
##
chisq_dissupp = chisq.test(tab_dissupp)
chisq_dissupp
##
## Pearson's Chi-squared test
##
## data: tab_dissupp
## X-squared = 40.022, df = 27, p-value = 0.051
```

#### Hypothesis Tested

```
tab_dishypop = table(df_all$sci_cat, df_all$hypo_tested)
tab_dishypop
##
##
                                                      Yes No
     Applied exercise physiology (human)
##
                                                       64 25
    Basic physiology (animal and cell physiology)
##
                                                       8 1
##
    Biomechanics
                                                       38 12
##
    Clinical research
                                                       18 11
    Environmental physiology (heat, cold, & altitude)
##
                                                       7 3
##
                                                        7 19
    Epidemiology
    Motor learning/control/behavior
##
                                                       12 7
##
    Other
                                                       12 15
    Sport performance
##
                                                       17 12
    Sport/exercise psychology
##
                                                        8 4
chisq_dishypop = chisq.test(tab_dishypop)
chisq_dishypop
##
## Pearson's Chi-squared test
##
## data: tab_dishypop
```

## X-squared = 28.438, df = 9, p-value = 0.0008056

#### Plots

```
p_dissup = df_all %>%
  group_by(support, sci_cat) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(support)) %>%
  ggplot( aes(fill=support, y=count, x=sci_cat)) +
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "",
       y = "Relative Frequency",
       fill = "") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "bottom") +
  coord_flip()+
  theme(text = element_text(face = "bold"))
p_dishypo = df_all %>%
  group_by(hypo_tested, sci_cat) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(hypo_tested)) %>%
  ggplot( aes(fill=hypo_tested, y=count, x=sci_cat)) +
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "",
       y = "Relative Frequency",
       fill = "Hypothesis Tested") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "bottom") +
  coord_flip() +
  theme(text = element_text(face = "bold"))
emm_plot = plot(emm_samps) +
  scale_x_continuous(trans = "log",
                     breaks = c(10,15,20,30,40,50,65,80,110,150,1095)) +
  labs(x = "Estimated Mean Sample Size (log scale)",
       v = "") +
  theme bw() +
  theme(text = element_text(face = "bold"))
fig3 = ggarrange(p_dishypo,p_dissup,emm_plot,
                 ncol = 1,
                 labels = "AUTO")
```

```
# Main Figures -
p_2a = df_all %>%
  group_by(journal, support) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(support)) %>%
  ggplot( aes(fill=support, y=count, x=journal)) +
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "Journal",
       y = "Relative Frequency",
       fill = "Hypothesis Support") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "top",
        text = element_text(face = "bold"))
p_2b = df_all %>%
  group_by(journal, hypo_tested) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(hypo_tested)) %>%
  ggplot( aes(fill=hypo_tested, y=count, x=journal)) +
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "Journal",
       y = "Relative Frequency",
       fill = "Hypothesis Tested") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "top",
        text = element text(face = "bold"))
p_2c = df_all %>%
  group_by(journal, effect_size) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(effect_size)) %>%
  ggplot( aes(fill=effect_size, y=count, x=journal)) +
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "Journal",
       y = "Relative Frequency",
       fill = "Effect Size Reported") +
  theme_classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "top",
        text = element text(face = "bold"))
```

```
p_2d = df_all %>%
  group_by(journal, n_just) %>%
  summarize(count = n(),
            .groups = 'drop') %>%
  filter(!is.na(n_just)) %>%
 {\tt ggplot(\ aes(fill=n\_just,\ y=count,\ x=journal))\ +}
  geom_bar(position="fill", stat="identity",
           color = "black")+
  scale_y_continuous(labels = scales::percent) +
  labs(x = "Journal",
      y = "Relative Frequency",
       fill = "Sample Size Justification") +
  theme classic() +
  scale_fill_viridis_d(option = "E") +
  theme(legend.position = "top",
        text = element_text(face = "bold"))
fig_2 = ggarrange(p_2b, p_2a, p_2c, p_2d,
                   ncol = 1,
                   labels = "AUTO")
```

# Figure 1

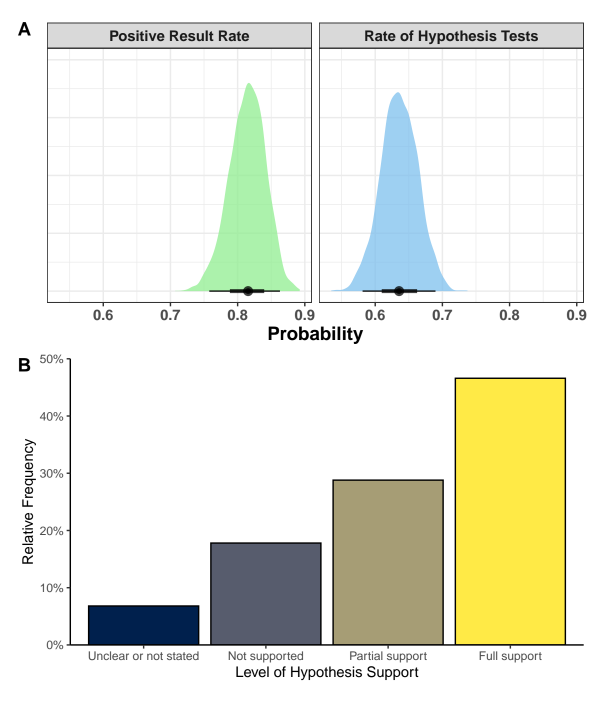


Figure 1. A) Posterior distributions from Bayesian model with the 50% and 95% percent compatibility intervals represented by the error bars at the bottom and B) Relative frequencies of the level of support reported for manuscripts with hypotheses (N = 191) with 17.8% report no support, 28.8% stating partial support, 46.6% stating full support, and 6.81% for which support was unclear or not stated.

# Figure 2

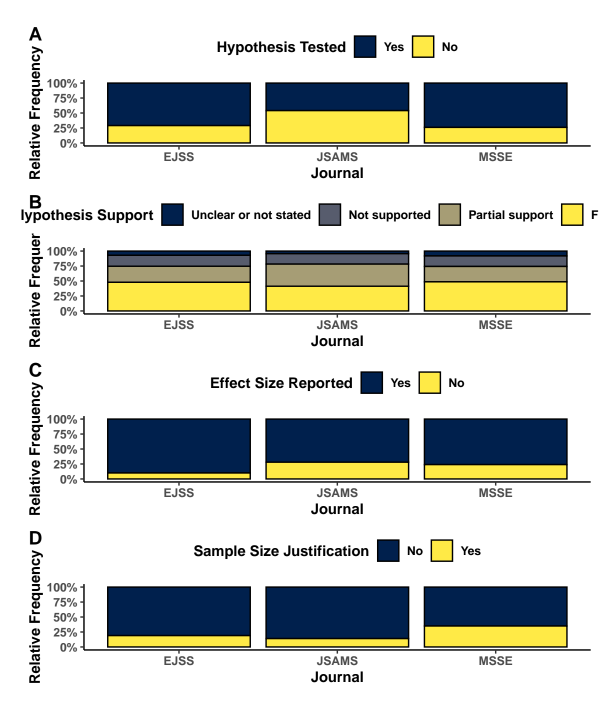


Figure 2. Relative frequencies, by journal, for A) level of reported support for hypotheses, B) indication of whether a hypothesis was tested, C) indication of whether an effect size was reported, or D) indication of if sample size was justified by the authors. Journals included the European Journal of Sport Science (EJSS), the Journal of Science and Medicine in Sport (JSAMS), and Medicine and Science in Sport and Exercise (MSSE),

# Figure 3

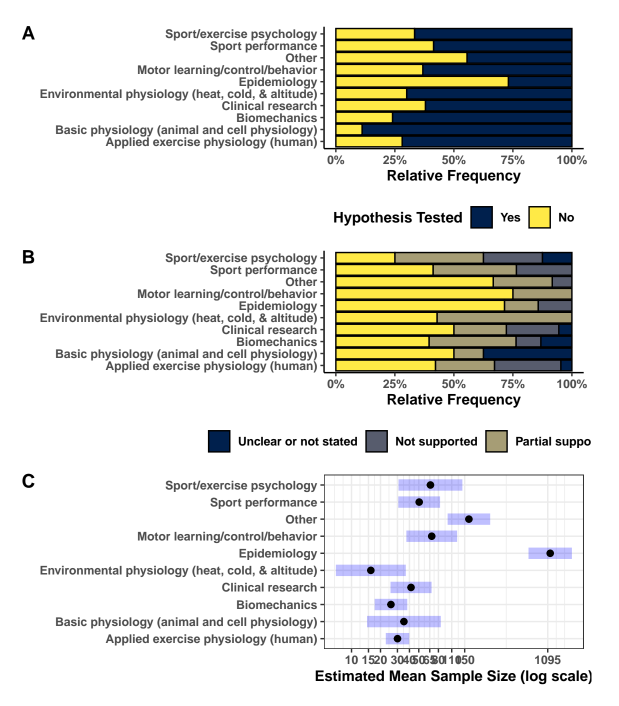


Figure 3. The breakdown, by discipline, for A) indication of whether a hypothesis was tested B) level of reported support for hypotheses, and C) the estimated total sample size (grey bands indicate 95% confidence intervals).

#### References

- Büttner, Fionn, Elaine Toomey, Shane McClean, Mark Roe, and Eamonn Delahunt. 2020. "Are Questionable Research Practices Facilitating New Discoveries in Sport and Exercise Medicine? The Proportion of Supported Hypotheses Is Implausibly High." *British Journal of Sports Medicine*. https://doi.org/10.1136/bjsports-2019-101863.
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