Running	head:	${\bf EXPRESSIVE}$	WRITING

A Meta-Analysis of Expressive Writing on Positive Psychology Variables and Traumatic

2 Stress

research are discussed.

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Abstract

Emotional expression has been shown to be beneficial for promoting both positive psychological and physical health outcomes. Unfortunately, inhibiting emotions can lead to impairments in physical and psychological health. James Pennebaker showed that expressive writing is an effective form of emotional expression, and he and others have used expressive writing as an experimental manipulation to gauge its efficacy in treating a wide variety of health-related and psychological outcomes. While many studies have been conducted that examine the efficacy of expressive writing across such outcomes, a considerable amount of 10 these studies tend to neglect necessary considerations such as different levels of 11 symptomology, power, and meaningfulness of respective effect sizes. Six previous 12 meta-analyses have been conducted that examine expressive writing's effect on psychological 13 outcomes. However, these studies focus on the experimental versus control group effect size. 14 Thus, our meta-analysis sought to examine the efficacy of an expressive writing task on only 15 the experimental conditions in studies measuring posttraumatic growth, posttraumatic stress, and quality of life using random effects models. Results indicated a small overall effect size for posttraumatic stress and negligible to small effect sizes for posttraumatic growth and 18 quality of life. However, those studies requiring a diagnosis of PTSD exhibited a medium to 19 large effect size. Implications for future research design and interpretation of published

Keywords: meta-analysis, posttraumatic stress, expressive writing

A Meta-Analysis of Expressive Writing on Positive Psychology Variables and Traumatic

Stress

26 Emotional Expression

Emotional expression relating to negative emotions or trauma has been shown to 27 enhance both mental and physical health outcomes (Esterling, Antoni, Kumar, & 28 Schneiderman, 1990; Fawzy et al., 1993; Lieberman & Goldstein, 2006; Rachman, 1980; Scheff, 1979). For example, the disclosure of traumatic or stressful events has been shown to reduce stress and lead to positive health outcomes in those with diabetes (Bodor, 2002) and breast cancer (Stanton et al., 2002), among others. Inhibiting thoughts or emotions, rather, may be detrimental to both physical and psychological health (H. S. Goldstein, Edelberg, Meier, & Davis, 1988; Gross & Levenson, 1997; Larson & Chastain, 1990). While some studies suggest that emotional expression in the form of "truth telling" may cause 35 psychological harm to individuals (Brounéus, 2010), the literature presents evidence 36 confirming the negative effects of a lack of emotional expression, such as social concerns, 37 overall psychological dysfunction, and lack of value-congruent behaviors (Frankl, 1959; 38 Pennebaker, 1989; Pennebaker & Beall, 1986; Schulenberg, Hutzell, Nassif, & Rogina, 2008; Wilson & DuFrene, 2009). These resulting negative outcomes may lead to detrimental effects on health (Pennebaker & Beall, 1986). Individuals having experienced a traumatic or 41 stressful life event are significantly more likely to repress thoughts and feelings about their experience compared to individuals who have not experienced such events, thereby subjecting them to potential negative outcomes caused by a lack of emotional expression (Bodor, 2002).

Expressive Writing as Effective Emotional Expression and Meta-Analytic

46 Techniques

Pennebaker and Beall (1986) first experimentally manipulated expressive writing. In their seminal study, Pennebaker and Beall randomly assigned participants to writing groups, including writing about a "stressful or traumatic" life event or a neutral event. Individuals assigned to the trauma/stressor condition exhibited a reduction in health visits after writing
about the emotions surrounding the trauma or stress. This protocol has been termed written
emotional disclosure (WED). Since then, the WED protocol have been employed across a
wide variety of contexts. Although many studies have produced positive outcomes associated
with expressive writing, some of these studies tend to neglect important questions, the most
important of which is whether or not the effect sizes are meaningful (Smyth, 1998). For a
review of the effiacy of WED, we turn to previously conducted meta-analyses.

Meta-analyses allow researchers the opportunity to collectively examine the efficacy of
different psychological interventions/tasks on outcome variables by calculating an overall,
weighted, population effect (Borenstein, Hedges, & Rothstein, 2007; Glass, 1976; Hedges,
1982). Several meta-analyses of WED have been conducted that warrant explanation: Smyth
(1998); Frisina, Borod, and Lepore (2004); Frattaroli (2006); Mogk, Otte, Reinhold-Hurley,
and Kröner-Herwig (2006); van Emmerik, Reijntjes, and Kamphuis (2013); and Reinhold,
Bürkner, and Holling (2018). These meta-analyses have laid a foundation for exploring the
effects of WED on psychological outcomes.

Smyth (1998) conducted the seminal meta-analysis examining the effiacy of expressive writing on psychological well-being, general health, and physical functioning. They included studies employing an expressive writing group and control group (neutral topic). In sum, thirteen studies/effect sizes were included, and the authors found an overall medium effect size, d = 0.47, for the experimental group compared to the control group. A later meta-analysis conducted by Frisina et al. (2004) expanded these analyses and included studies with clinical samples. This meta-analysis included nine studies and found an effect size of d = 0.19 for health-related outcomes and d = 0.07 for psychological outcomes. Mogk et al. (2006) conducted the next expressive writing meta-analysis to update the state of the literature regarding expressive writing. Studies employing Pennebaker's paradigm on experimental and control groups were included. Further, inclusion criteria were methodological techniques that included a four-week follow up and at least 10 participants.

Thirty studies met inclusion criteria. Effects relating to somatic and psychological health outcomes were nonsigificant, corroborating findings from Frisina et al. (2004).

Frattaroli (2006) conducted perhaps the most notable meta-analysis to date examining 79 the efficacy of emotional disclosure on the following constructs using only randomized and 80 control conditions: psychological health, physiological functioning, reported health, health behaviors, and general functioning/life outcomes. Additionally, this meta-analysis was the first to employ random effects models, which estimate the mean of a proposed distribution of population effect sizes. Prior meta-analyses employed fixed effects models, which assume that all studies assess the same "true" population effect size. This assumption may be 85 untenable across different populations (Borenstein et al., 2007). They included a wide range of studies N = 146. Individual studies were again collapsed into one publication effect size, 87 although these effects were also examined separately by health outcome. Overall, Frattaroli 88 (2006) found a weighted r effect size of .08 for all outcomes combined, which would be 89 considered small. Additionally, they examined potential moderators and found larger effect 90 sizes for the following samples: those with physical health problems, those with a history of 91 having experienced traumatic or stressful events, samples not including college students, samples where expressive writing tasks were conducted at home and in private settings, paid participants, more male participants, and fewer participants (see Frattaroli, 2006 for a complete list of moderators). A recent analysis conducted by van Emmerik et al. (2013) 95 employing Pennebaker's paradigm included six eligible studies that compared treatment to control groups. In regards to inclusion criteria, they included studies where participants had a diagnosis of Acute Stress Disorder (ASD) or PTSD. They found that those who participated in the expressive writing group experienced short-term reductions in PTS and comorbid depressive symptoms, combined Hedges' g = 0.81. The most recently published 100 meta-analysis was conducted by Reinhold et al. (2018) and examined the effects of 101 expressive writing on depression by randomizing participants to conditions (expressive 102 writing vs. control). They included thirty-nine randomized controlled trials and excluded 103

individuals with diagnoses of PTSD. This study did not support utilizing expressive writing for depression outcome measures for the specified sample, $g_{post} = -0.09$. Further, they found that expressive writing did not yield any type of long-term effect on depression outcomes.

Posttraumatic Stress Disorder (PTSD) is a disorder involving re-experiencing thoughts

107 Posttraumatic Stress

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or events after a trauma. This generates a context where individuals are prone to 109 affect-related deficiencies and maladaptive behaviors (American Psychiatric Association, 110 2013). DSM-5 criteria are based on twenty symptoms structured into four different subsets 111 in those having experienced a traumatic event. These subsets are as follows: re-experiencing, 112 avoidance, negative alterations in cognition and mood, and increased arousal (Crespo & 113 Gomez, 2016). While the renewed DSM-5 criteria are now increasingly employed, the current 114 meta-analysis considers studies using DSM-IV criteria. DSM-IV criteria are similar and 115 include the following: exposure to a traumatic event, re-experiencing (intrusion), avoidance, 116 and increased arousal (American Psychiatric Association, 2013). Further, the studies employed in the current meta-analysis are divided according to these subsets (arousal, intrusion, and avoidance). PTSD affects a wide variety of groups, a few of which are sexual 119 assault survivors (Klump, 2008), Iraq and Afghanistan war veterans (Gentes et al., 2014), 120 and those exposed to natural disasters (Wang et al., 2000). 121 Research conducted on the efficacy of expressive writing on PTSD symptoms presents 122 intriguing results. Sloan, Marx, Epstein, and Lexington (2007) examined individuals with at 123 least moderate PTSD symptom severity and found that individuals assigned to an emotional expression writing condition reported fewer PTSD and depression symptoms during follow up. Sloan, Marx, and Greenberg (2011) found that PTSD symptoms decreased after a 126 written emotional disclosure task, although this decrease was not significantly different than 127 a control group change. Di Blasio et al. (2015) recruited women who had just given birth 128 and assessed them a few days after experiencing childbirth along with a three-month 129

follow-up. Results showed that women who had participated in the expressive writing task had lower depression and posttraumatic stress symptoms than the group assigned to a 131 neutral writing condition. Additionally, regression models showed that expressive writing 132 was significantly linked to a reduction of PTSD symptoms across different dimensional levels 133 of symptom severity. Only 20 of the 113 women recruited for this study qualified for a 134 diagnosis of PTSD, but those who reported mild symptomology responded better to the task 135 than those meeting criteria for PTSD. This limitation suggests that those with moderate 136 distress could perhaps benefit more from an expressive writing task than those diagnosed 137 with or meeting the qualifications for PTSD. It may also explain the differences in results in 138 comparing to Sloan et al. (2011), as they found that those with a clinical diagnosis of PTSD 130 did not respond to an emotional disclosure writing task. Perhaps it may be more 140 advantageous to examine effect sizes separately for diagnoses of PTSD and subclinical symptoms. Further Sloan, Marx, Bovin, Feinstein, and Gallagher (2012) adapted a writing protocol to focus primarily on the emotions, meaning, and "hot spots" associated with the 143 trauma. They referred to this procedure as the written exposure therapy (WET) protocol, distinguishable from the paradigm adapted by Pennebaker and Beall (1986). In their seminal 145 study examining the efficacy of WET for motor-vehicle accident related PTSD, they found that those in the WET condition experienced significant reductions in PTSD symptoms 147 throughout the course of the study. Since then, a small number of other studies employing 148 the WET procedure have been employed in those with PTSD. While these will be included 149 in the current review, the newness of this protocol does not allow exclusive examination 150 using meta-analytic techniques. 151

52 Posttraumatic Growth

While the literature mostly discusses potentially harmful outcomes to traumatic events such as emotional distress, traumatic events also provide opportunities for personal growth (Aslam & Kamal, 2013). Traumatic events, either natural or human-inflicted, can lead to

positive outcomes by allowing the individual to take a different perspective (Cobb, Tedeschi, 156 Calhoun, & Cann, 2006; Taku, Calhoun, Cann, & Tedeschi, 2008). The relationship between 157 positive growth after a traumatic event and symptom reduction is unclear, as it is a complex 158 process. Thus, it is necessary to examine how expressive writing might influence each 159 variable separately, which is one of the key goals of this meta-analysis (Slavin-Spenny, Cohen, 160 Oberleitner, & Lumley, 2011). Models receiving empirical support within the last decade 161 suggest that traumatic events offer opportunities for both negative and positive experiences 162 (Tedeschi & Calhoun, 1995; Weiss, 2002). Posttraumatic Growth (PTG) is a positive 163 experience after a traumatic event (Aslam & Kamal, 2013; Yilmaz & Zara, 2016). 164 Specifically, PTG is classified as broad cognitive benefits that are seen after a traumatic 165 experience. These benefits can be categorized into building closer relationships, examining 166 new possibilities, appreciating life, recognizing personal strengths, and undergoing spiritual changes (Dursun, Steger, Bentele, & Schulenberg, 2016; Tedeschi & Calhoun, 2004).

PTG is associated with a variety of desired outcomes (Dursun et al., 2016). PTG has 169 been studied in those experiencing natural disasters, war, and other harms such as sexual 170 assault. Finally, PTG has been studied in those experiencing medical diagnoses such as 171 different types of cancer and diseases. Although the relationship between PTG and symptom 172 reduction is not yet fully understood, perhaps expressive writing allows the individual to 173 fully comprehend the event. Pennebaker and Graybeal (2001) speculated that expressive 174 writing allows an individual to feel more connected with his or her surroundings. Although this speculation does not directly explain positive outcomes after an expressive writing task, perhaps individuals gain a better appreciation for life after gaining a better sense of 177 connectedness with that individual's surroundings. One might expect effect sizes to be larger 178 for those studies requiring a diagnosis of PTSD, as such growth may not be possible in those 179 with subclinical symptomology. 180

Quality of Life

Quality of Life (QOL), according to Theofilou (2013) is an evaluation of the "goodness" 182 that an individual experiences, separated into domains of reactions to life events, disposition, 183 life fulfillment, and satisfaction with life experiences. More generally, QOL refers to an 184 individual's attitude towards the target life situation (Costanza et al., 2007), delineated into 185 objective and subjective components. Objectively, QOL refers to components outside of an 186 individual and measurable by others, while subjective QOL is an individual's assessment of 187 his or her own experiences (Costanza et al., 2007). The current meta-analysis will focus 188 solely on the subjective components of QOL, as it is obtainable through questionnaires. 189 Pennebaker and Graybeal (2001) suggested that expressive writing allows one to feel more 190 connected with their surroundings. Further, they explain that expressive writing allows 191 people to see things in a different way and better understand themselves. By understanding 192 a traumatic or stressful event, one is said to see things differently and perhaps look at the situation with a more positive mindset. The changes that occur after expressive writing may 194 also allow one to find meaning in the traumatic event, thereby increasing the QOL of that 195 individual (Frankl, 1959). Higher QOL may be considered a type of PTG, which is why the 196 current meta-analysis sought to examine the efficacy of studies utilizing expressive writing to 197 improve QOL and PTG in the same study. 198

199 Current Meta-Analysis

The purpose of the current meta-analysis is to examine studies employing expressive writing procedures using Pennebaker's paradigm (WED) and the more recent WET protocol on variables relevant to the field of positive psychology (PTG and QOL) and PTS, with effect sizes separated by the paper's indication of PTSD diagnosis when sample sizes are large enough. Based on recently published literature regarding efficacy of expressive writing for different levels of PTSD symptoms, this marker is an important facet to consider (Di Blasio et al., 2015; Reinhold et al., 2018; Sloan et al., 2011). No review has examined the effects of

expressive writing on PTS separated by diagnosis. Additionally, no meta-analysis has been 207 conducted that examines the efficacy of expressive writing on positive outcome variables such 208 as PTG and QOL, in line with the field of positive psychology. The meta-analyses described 209 sequentially above also focused on experimental versus control group effect sizes or p-values, 210 rather than emphasizing change for the expressive writing group. This focus is likely because 211 of the analyses provided in these publications, especially when using randomized controlled 212 trial research designs. While this design is the gold standard for medicine, the current 213 meta-analysis sought to examine the magnitude of change for participants who experienced 214 an expressive writing task. For example, a comparison group may increase their quality of 215 life scores by two points in a controlled study, while the experimental group increases their 216 quality of life scores by four points; thus, creating a significant difference in change between 217 the two groups. This information is valuable, but it does not tell the reader the magnitude of 218 the change for the writing group, wherein four points might only be a small effect when 219 examined within the group who received the writing task.

This analysis will also focus on changes across time for groups who received the 221 expressive writing task to determine what size of effects one might expect given a specific 222 measurement schedule (i.e., one to three months, three months to six months, etc.). This 223 analysis should present researchers with a renewed examination of the efficacy of expressive 224 writing on the aforementioned variables using newer meta-analytic techniques. Newer 225 methods of meta-analysis, including p-curve (Simonsohn, Nelson, & Simmons, 2014; 226 Simonsohn, Simmons, & Nelson, 2015), p-uniform (van Aert, Wicherts, & van Assen, 2016), 227 PET-PEESE (Stanley & Doucouliagos, 2014), selection models (Vevea & Hedges, 1995), and trim and fill methods (Carter & McCullough, 2014) allow for better estimation of meta-analytic effect sizes. These analyses would be best performed by examining each potential effect separately, rather than averaging effects of each publication into one study 231 effect size (a common trend in the previously mentioned meta-analysis). In addition to an 232 estimate of overall effect sizes using updated techniques, the current meta-analysis estimates 233

power for effects on writing groups, as research has shown a consistent under powering of psychological studies, combined with a misunderstanding of the sample size needed for adequately powering one's work (Bakker, Hartgerink, Wicherts, & van der Maas, 2016).

Studies were collected through online databases, such as PsycINFO and Google

237 Method

238 Data Collection

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Scholar, using the following search terms and their combinations: Posttraumatic Growth, 240 PTG, Quality of Life, QOL, Posttraumatic Stress, PTS, Expressive Writing, Emotional Disclosure, Written Emotional Disclosure (WED), Written Exposure Therapy (WET). Within these articles, the change in outcome variables (PTS, PTG, QOL) from pre- to 243 post-test was the dependent variable of interest. Generally, groups were separated into an experimental and control group and then examined at different time points. For purposes of 245 this meta-analysis, only participants assigned to the experimental condition were examined 246 due to having received the expressive writing task. If a study included multiple assessment 247 time points, then these measurements were examined sequentially (i.e., time 1 to time 2, 248 time 2 to time 3) to determine change across time for the dependent variable. 249 264 citations focusing on PTS, PTG, and QOL were identified through the literature 250 search and previous meta-analyses. After screening these studies, 53 articles were retained for 251 containing the appropriate information for this meta-analysis. This manuscript was written 252 with papaja in R (Aust & Barth, 2017) with the analyses inline with the text. The complete 253 set of data, excluded article list with reasoning, and other relevant information can be found at: https://osf.io/4mjqt. Generally, studies were included if they utilized WED or WET, 255 included relevant numbers to compute an effect size, and included the relevant outcome variables. After having two reviewers independently code articles, 223 effect sizes were 257 calculated. On average, each study represented M = 4.21, SD = 3.31 effects, ranging from 1 258 to 16 effects. 163 effects were calculated for PTS, 21 for PTG, and 37 for QOL. Studies were 259

coded for PTSD diagnosis as no (not mentioned or not included), mixed (mentioned number of participants but all included), and yes (included as criteria). After examining the number of effects in each of these categories for each variable, only the PTS results will be split by PTSD diagnosis with 16 no mention, 16 in the mixed category, and 86 yeses.

²⁶⁴ Calculations for Effect Size, Variance, and Confidence Intervals

For our purposes, we used Cohen's (1988) standards for nomenclature for small (0.20), 265 medium (0.50), and large (0.80) d values, although it is important to note that Cohen 266 himself suggested that these values should be based on the area of study. Generally, however, 267 these effect size criteria are used within the social sciences. Each study implemented a 268 pre-test to post-test style repeated measures design, usually with paired t-tests, ANOVA, or 269 regression analyses. The means, standard deviations, and N values were collected from each 270 study. In general, Cohen's d values were calculated using the following formula for paired t 271 using means and standard deviations: 272

$$d_{av} = \frac{M_1 - M_2}{\frac{SD_1 + SD_2}{2}}$$

This equation is described in detail in Cumming (2012) as an alternative to the traditional calculation of d for paired samples t, wherein the denominator is the standard deviation of the difference scores:

$$d_z = \frac{M_1 - M_2}{SD_{diff}}$$

This equation for d_{av} not only allows for calculations from published articles that do not include SD_{diff} (i.e., most articles included), but also has been shown to be less upwardly biased than d_z . Alternative formulas include controlling for r between paired levels, as described in Lakens (2013); however, these values were not available in the selected articles, and Lakens also recommends d_{av} as an effect size for paired designs. When only mean differences and standard deviation of the difference scores were available, the second equation for d_z was used.

We planned to use traditional and newer methods of meta-analysis, following guidelines from Cooper, Hedges, and Valentine (2009) and Borenstein et al. (2007), as well as van Aert et al. (2016). Sampling variance of the effect sizes were estimated using the escalc() function from the metafor package in R (Viechtbauer, 2010). The variance formula was originally published in S. B. Morris and DeShon (2002) and is shown below:

$$v = \frac{1}{n} \left(\frac{n-1}{n-3}\right) (1 + n * d^2) - \frac{d^2}{[c(n-1)]^2}$$

In this formula, n is the number of paired observations, d is the calculated effect size, and c is a correction factor, wherein df are n-1 (Hedges, 1982):

$$c = 1 - \frac{3}{4 * df - 1}$$

We used the metagen() function in the metafor package to calculate both fixed and 290 random effects models, which uses standard error of the effect to calculate overall estimates 291 of an effect and their confidence intervals. Thus, we took the square root of the variance 292 estimate for standard error. Given these calculations, the goal of this analysis was to 293 calculate a combined effect size, along with a confidence interval for study planning and an assessment of the literature. A fixed effects model requires the assumption that there is a 295 true population effect size across all studies. By including multiple measures of psychological outcomes, this assumption may be tenuous, and therefore, a random effects model was also calculated. In random effects models, the true effect is assumed to vary across studies 298 (Borenstein et al., 2007). For a fixed effects model, the effect sizes are weighted by their 299 inverse variance (v; Sánchez-Meca & Marín-Martínez, 2008), which is calculated 300 automatically in *metafor* by: 301

$$w_i^{FE} = \frac{1}{v}$$

The advantage to this procedure is that analyses are weighted by their precision, that is, that studies with more information (often, larger samples), are given larger weights in the overall estimated effect size (Borenstein et al., 2007). Random effects models are also weighted by inverse variance, with an additional correction for variance between studies, τ_{DL}^2 , as described by DerSimonian and Laird (1986):

$$w_i^{RE} = \frac{1}{v + \tau_{DL}^2}$$

Confidence intervals were calculated in two ways for this study. Cumming (2012), 307 Kelley (2007), and Smithson (2001) have shown that the distribution of d values are 308 non-normal, and thus, CIs should be estimated using the non-centrality parameter and a 309 non-normal distribution. These values were calculated using the functions in the MOTE 310 library which iteratively estimates the appropriate non-centrality parameter and converts 311 back to d values (i.e., non-centrality parameter divided by the square root of n; Buchanan, 312 Valentine, & Scofield, 2017; Smithson, 2001, 2003). However, the metafor package in R uses 313 central distributions to estimate CIs for each study and overall effect sizes. Therefore, we 314 present both sets of values for the interested reader, as meta-analytic procedures have not 315 implemented non-central distributions of effect sizes. 316

317 Additional Meta-Analytic Techniques

p-Curve and p-Uniform. We used *p*-curve.com to conduct a *p*-curve analysis

(Simonsohn et al., 2014). The purpose of this type of analysis is to detect true effects.

Specifically, *p*-curve is used to reveal possible *p*-hacking in published literature in order to

decipher whether or not a true effect exists. Broadly, *p*-hacking occurs when researchers use

questionable research practices to create significant results by manipulating dependent

variables or covariates. Additionally, authors may add participants if the initial findings are

not significant (Bruns & Ioannidis, 2016). Researchers may also decide to exclude

participants for final analyses if that exclusion leads to a significant difference (L. K. John,

Loewenstein, & Prelec, 2012). Thus, it is necessary to distinguish between true and false effects in order to effectively interpret effect sizes corresponding to those p-values. p-curve 327 accomplishes this task by examining the distributions of the published p-values. If an effect 328 exists, or rather the results should be interpreted as presented, the distribution of p-values 329 will be positively skewed (Simonsohn et al., 2014). If, however, no effect exists, then the 330 distribution of p-values will be flat. p-curve analyses ultimately provide evidence of 331 p-hacking in groups of studies and has become an important tool for interpreting 332 meta-analyses. In order to accurately estimate effect sizes because of scrutiny associated 333 with effect size estimation of p-curve, we also conducted p-uniform. p-uniform analyses, too, 334 are interpreted by examining the distribution of p-values in a set of studies (van Aert et al., 335 2016). However, it is assumed that the population effect size equals the effect size from the 336 dataset. Because of this assumption, the population effect size is referred to as uniform. This analysis also examines for publication bias and presents the researcher with a corrected effect 338 size. Publication bias occurs when only select studies are published, usually only significant 339 studies, although many factors can bias a study's publication (McShane, Böckenholt, & Hansen, 2016). p-uniform was calculated from code provided by van Aert (2017) on GitHub. 341

Originally, meta-analyses relied on the calculation of Egger's PET-PEESE. 342 regression test which examined the relationship of the standard error (predictor) to the effect 343 size estimates (criterion). In this regression, the intercept values were used to determine if 344 effect size measures were different than zero, by providing a meta-analytic estimate (Egger, 345 Davey Smith, Schneider, & Minder, 1997; Stanley, 2005). PET-PEESE analyses examine for 346 publication bias by adapting parts from Egger's traditional regression tests: PET (Precision 347 Effect Test) and PEESE (Precision Effect Estimate with Standard Error, Carter & 348 McCullough, 2014). PET is a more reliable test of publication bias with effect size estimates 340 of zero, $b_0 = 0$, while PEESE is more accurate with non-zero effect size estimates, $b_0 \neq 0$ 350 (Stanley & Doucouliagos, 2014). PET-PEESE was calculated using Hilgard's (2016) code 351 provided on GitHub. 352

Selection Models. Selection model analyses provide the researcher with a test of publication bias and effect size estimates using maximum likelihood estimation (Vevea & Hedges, 1995; Vevea & Woods, 2005). Using selection models, researchers are able to discover effect size estimates as well as evidence of publication bias (McShane et al., 2016) by using a mixed general linear model to estimate these values. Selection models were calculated with the weightr package in R (Coburn & Vevea, 2017).

Trim and Fill. Trim and Fill analyses, in contrast to PET-PEESE, regress standard 359 error (criterion) and effect size estimates (predictor). Specifically, the purpose of Trim and 360 Fill techniques is to examine whether or not publication bias may influence the regression 361 equation (Carter & McCullough, 2014). Effect sizes and standard error terms are graphically 362 displayed on x and y-axes, respectively, in a funnel plot. If this graphical representation 363 indicates asymmetry, considered a gap of missing data points in the lower center area of the 364 plot, the study set can be assumed to have studies that are both non-significant and small in 365 sample size (van Assen, van Aert, & Wicherts, 2015). This funnel is then trimmed until 366 symmetry is achieved. Missing studies from the symmetrical graph are imputed (filled) while 367 maintaining the given symmetry (Duval & Tweedie, 2000). The meta-analytic effect size is 368 then estimated from the trimmed and filled funnel plot. Trim and fill analyses, as well as 369 funnel plots included below, were calculated with the *metafor* package. 370

Results Results

PTS

Overall Effect Size. As described above, both fixed effects and random effects
models with centralized confidence intervals are presented in Table 1. Studies were examined
for potential outliers using the *metafor* package in *R*. This package calculates traditional
regression influence values, such as Cook's and hat values (J. Cohen, 1988). These values
indicate change in overall meta-analytic model with and without the effect; thus,
determining their impact on the pooled effect size (Viechtbauer, 2010). Because published

studies likely represent the range of the sampling distribution of effect sizes, we included the analyses with and without outliers to present evidence for both paths a researcher might take when examining an overall effect.

2 outliers were detected with this procedure, all showing very large effect sizes, average 382 d=2.81. The fixed and random effects estimates without these points are also included in 383 Table 1. Figures 1, 2, 3, and 4 portray the effect sizes for PTS studies, separated by intrusions, avoidance, hyperarousal, and total scores for easier viewing (i.e., over 100+ effect sizes did not fit easily on one combined graph). Although these categories are not reflective of updated DSM-5 criteria, researchers have not yet conducted enough studies using 387 expressive writing on PTS with updated PTSD criteria to warrant a meta-analysis. Name 388 acronym coding can be found in the data online. This forest plot includes the non-centralized 389 confidence interval calculated from the MOTE library (Buchanan et al., 2017). Shape size 390 indicates study weight, and these values were taken from the overall random effects 391 meta-analysis and normalized by dividing by the mean weight. The dashed lines indicate the 392 average non-weighted lower and upper confidence interval limit for the non-centralized 393 estimates. Overall, PTS studies include a small effect size that appears to be significantly 394 greater than zero across all estimate types (fixed, random, with or without outliers). 395

We further calculated the overall effect sizes by PTSD diagnosis category using a 396 random effects model. Studies only including individuals with a diagnosis of PTSD exhibited 397 a medium effect size (before and after outlier exclusion): with outliers d = 0.64 [0.48, 0.79]; 398 without outliers d = 0.55 [0.41, 0.69], while studies not requiring (or listing) a PTSD 399 diagnosis showed a small to medium effect size: d = 0.32 [0.24, 0.40]. Similarly, the mixed 400 category showed a small to medium effect size : d = 0.35 [0.16, 0.54]. Complete estimates of 401 all the following analyses split by diagnosis are included online at https://osf.io/4mjqt/, and 402 their pattern of results is similar to the overall pattern here. 403

Homogeneity. A prerequisite for newer meta-analytic techniques includes the
assessment of homogeneity of the effects (van Aert et al., 2016). Using the *metafor* package

in R, we calculated the Q-statistic and the I^2 index (Cochran, 1954; Huedo-Medina, 406 Sánchez-Meca, Marín-Martínez, & Botella, 2006). Significant values imply inconsistencies 407 across the variable or variables of interest and are represented by Q. In contrast, I^2 indicates 408 the percentage of heterogeneity along with a 95% CI. Both can, however, be biased with a 409 small number of experiments included for analyses (Higgins, Thompson, Deeks, & Altman, 410 2003; Huedo-Medina et al., 2006). Thus, we sought to calculate an overall level of 411 heterogeneity after examining each variable separately before and after excluding outliers. 412 For PTS studies including outliers, we found significant heterogeneity, Q(162) = 776.74, p < 100413 .001 and $I^2 = 79.1$, 95% CI[75.9 - 81.9]. These values were reduced slightly with the 414 exclusion of outliers, Q(160) = 677.98, p < .001 and $I^2 = 76.4$, 95% CI[72.6 - 79.7]. 415

Power was calculated in two different ways using the pwr package in R 416 (Champely, 2016). Post hoc power was first calculated using sample size and effect size 417 statistics from each individual study. Additionally, we calculated power using the study 418 sample size and estimated overall effect size from the random effects model with and without 419 outliers, as explained by Francis (2012) and Francis (2014). The first estimate indicates the 420 likelihood of finding an effect from our sample statistics, while the second indicates the 421 likelihood of finding the true population effect size. If each study had been conducted on only the change in the experimental group, 46.6% of studies would have been considered significant at $\alpha < .05$. The average power of these studies based on their original study 424 characteristics was .48 (SD = .36). Power for the random-effects meta-analytic effect size 425 with outliers was .52 (SD = .25) and without outliers was .49 (SD = .25). Therefore, power 426 consistently was around 40-50% for studies examining PTS, regardless of outlier effects. In 427 these studies, only 28.8% achieved recommended 80% power for their found effect size, a 428 smaller 24.5% for the random-effect outlier effect size, and even smaller 20.2% for power 429 calculations on the random-effect size without the outliers. 430

Other Meta-Analytic Estimates. As noted in van Aert et al. (2016), p-curve and p-uniform analyses are upwardly biased when heterogeneity is high. Therefore, we use

caution when interpreting these analyses on PTS outcomes. As seen in Table 1, the 433 estimates for p-uniform were higher than other techniques, likely because of the focus on 434 significant p-values and the great degree of heterogeneity described earlier. P-curve pictures 435 can be found at https://osf.io/4mjqt/online, and this analysis indicated evidentiary value at 436 p < .001. Additionally, the p-uniform analysis indicated that there was likely no publication 437 bias present, Z = -5.71, p = 1.000. When examining the PET analysis, we found that the 438 intercept was significant, which indicated that PEESE was likely a better estimator of the 439 meta-analytic effect size. PEESE estimates were lower than the original meta-analytic estimate, but confidence intervals indicated that the effect is small to medium, and still 441 larger than zero. Selection models indicated a larger effect size, especially with the 442 random-effects models, and these effects were influenced by the outliers found in the 443 published studies. Trim and fill models are shown in Table 1, and figures are included online. Nineteen missing studies were imputed for both models with and without outliers. Across all these effect size estimates, we found that expressive writing was likely to decrease PTS symptoms in a small to moderate way. The correlation of effect size with time between measurement times was r = -.01, 95% CI [-.17, .14], t(161) = -0.15, p = .879, and448 r = -.08, 95% CI [-.23, .08], t(159) = -1.00, p = .320 without outliers. This result indicated that the effect of expressive writing slightly decreased across time. 450

451 **PTG**

Overall Effect Size. Both fixed and random effects models with centralized
confidence intervals for PTG are presented in Table 2. When examining expressive writing
on PTG, no outliers were detected. Fixed and random effects estimates are included in Table
kyper to the property of the study. Dashed lines indicate non-weighted lower and upper
confidence intervals for non-centralized estimates. Overall, PTG studies indicated a
negligible to small effect size across both random and fixed effects models, and the

non-centralized confidence intervals indicated an effect that crossed zero.

Homogeneity. Using the metafor package in R, we calculated both a Q statistic and I^2 index. Since PTG studied did not contain any outliers, we did not calculate two separate analyses to examine heterogeneity both with and without outliers. We did not find significant heterogeneity across PTG studies, Q(20) = 14.18, p = .821 and $I^2 = 0.0$, 95% CI[0.0 - 25.3].

Power. First, we calculated *post hoc* power using both sample and effect size statistics from individual studies. Individual studies examining change in experimental groups showed that 9.5% of studies would have been considered significant at $\alpha < .05$.

Average power of PTG studies was .15 (SD = .16). 0.0% achieved recommended 80% power for their found effect size. Additionally, we calculated power using study sample size and estimated effect size from our random effects model. Power for the true effect size was .08 (SD = .02). Again, 0.0% achieved recommended 80% power.

Other Meta-Analytic Estimates. Due to no heterogeneity across PTG studies, 471 we can use both p-curve and p-uniform analyses with more confidence. A pictorial 472 representation of p-curve can be found at https://osf.io/4mjqt/. This analysis did not 473 indicate evidentiary value, p = .75, as only two of the results would be considered significant 474 at $\alpha < .05$. p-uniform estimates are presented in Table 2. Specifically, these analyses 475 indicated that there was no publication bias present, Z = 0.70, p = .243. The p-uniform 476 estimates of the effect size for PTG were negative, in contrast to the fixed and random 477 effects overall model. The confidence interval for this analysis indicates a wide range of 478 possible effects. In examining PET-PEESE analyses, we did not find a significant intercept, 479 indicating that PET is most likely a better effect size estimator. PET analyses indicated that the effect size is negligible to small, with our confidence interval crossing zero. These results corroborated our original effect size calculations. Selection models indicated negligible to small effect sizes, again wherein the confidence interval includes zero effect. Trim and fill 483 models are shown in Table 2, and figures are included online. Zero studies were imputed for 484 our model, and thus, the effect size estimate is the same as the overall model. Across 485

techniques, we found that expressive writing has little to no effect on PTG. The correlation of effect size across measurement times in PTG studies at subsequent time points was r = .09, 95% CI [-.36, .50], t(19) = 0.38, p = .707, and no change over time was found.

QOL

Overall Effect Size. Finally, for QOL, both fixed and random effects models with centralized confidence intervals are presented in Table 3. Two outliers were detected with this procedure, average d = -0.07. While the average effect of these outliers indicates a small number, it is important to note that these two outliers were the largest positive and negative effects found from the Possemato, Ouimette, and Geller (2010) study. Fixed and random effects estimates without these points are also included in Table 3, while Figure 6 shows effect sizes for QOL studies. Overall, QOL studies indicated a negligible to small effect that showed a non-significant decrease in quality of life as a result of expressive writing.

Homogeneity. For QOL studies including outliers, we found significant heterogeneity from our random effects model, $Q(36)=200.09,\ p<.001$ and $I^2=82.0,\ 95\%$ CI[75.9 - 86.5]. After excluding outliers, our random effects model still indicated heterogeneity, $Q(34)=93.18,\ p<.001$ and $I^2=63.5,\ 95\%$ CI[47.6 - 74.6].

Power. In conducting post hoc power using sample and effect size statistics from individual studies, we found that 21.6% of studies would have been considered significant at $\alpha < .05$. Average power based on actual study characteristics was .33 (SD = .32). Power for the random effects meta-analytic effect size with outliers was .05 (SD = .00) and without outliers was .05 (SD = .00). Unfortunately, power was around 5% for both random effects models with and without outliers. In these studies, 18.9% achieved adequate power of 80% on their found effect size, while 0.0% achieved 80% power for our random effects model with outliers. Finally, without outliers, 0.0% achieved 80% power.

Other Meta-Analytic Estimates. We exert caution in interpreting p-curve and p-uniform analyses on QOL outcomes with and without outliers due to heterogeneity. As

seen in Table 1, p-uniform estimates were stronger and positive than other techniques 512 because of the high degree of heterogeneity recently described. p-curve pictures can be found 513 at the following OSF Link: https://osf.io/4mjqt. Eight studies were significant at $\alpha < .05$, 514 and the studies indicated evidentiary value, p = .004. p-uniform analyses did not indicate 515 publication bias, Z = -2.75, p = .997. In PET-PEESE analyses, we found that the intercept 516 was not significant, and therefore, PET was a better estimator of the meta-analytic effect. 517 Table 1 indicates that both of these analyses estimate the effect size around zero, with a 518 confidence interval that includes zero. Selection models correspondingly show small effects 519 crossing zero, except for random effects models with outliers, that appear to be heavily 520 influenced by the outliers. Trim and fill models are shown in Table 3, and figures are 521 included online. No studies were imputed for these analyses, and therefore, the effect size 522 estimates match the original meta-analysis. Overall, these results appear to point to no effects, ranging across zero with several negative estimates. Interestingly, the correlation of effect sizes across measurement times with outliers was r = -.37, 95% CI [-.62, -.05],525 t(35) = -2.33, p = .026 and r = -.64, 95% CI [-.80, -.39], t(33) = -4.75, p < .001 without 526 outliers. The effect of expressive writing appears to be positive at short time intervals and 527 decreases into negative effects at longer time intervals.

Discussion

In examining pre- to post-test comparisons across each variable separately, we found
that PTS studies indicated a small effect size across all meta-analytic estimates.

Interestingly, those studies requiring a diagnosis of PTSD for inclusion resulted in a medium
effect size, while those studies not requiring a PTSD diagnosis resulted in a small to medium
effect size. These results suggest that those with clinical symptoms of PTSD may benefit
more from expressive writing interventions. Further, these results are in contrast to
recently-conducted studies, which suggest that those with subclinical symptoms benefit more
from expressive writing tasks (Di Blasio et al., 2015; Sloan et al., 2011). Both QOL and

PTG studies indicated a negligible to small effect size using random effects models.

Although the PTG effect in our overall meta-analysis estimate was significant, other methods indicate this small effect is likely not different from zero. We also examined the relationship of time between measurements of the dependent variables and the corresponding effect size to determine if effects change over time. For both PTS and PTG, there was no relationship between effect size and time. For QOL studies, a medium to large negative correlation was found. A negative relationship between time and effect size implies that writing tasks were more effective in the initial time points, and effects decreased over longer time spans.

The authors note several limitations. Generally, ineffective emotional expression may 546 be a contributing factor. If participants/clients are not deeply engaged with the material, an 547 expressive writing task may not be effective, as Pennebaker and Graybeal (2001) imply that 548 connectedness is an important factor for the task. However, it may be difficult to implement 540 a check for engagement in these types of research designs. Doing so may also set a context 550 that will inhibit emotional processing and general responses. Research on expressive writing 551 has found a wide range of outcomes for different variables (Frattaroli, 2006), and these 552 various results may explain the large heterogeneity found in this study. Encouragingly, we 553 did not find much evidence of publication bias, and therefore, these estimates may represent a true population effect size. Regardless, methodology of expressive writing studies is variable, as it is applied in different forms across different contexts. Ideally, it would be 556 possible to control for these varied instructions and protocols. However, this is simply not 557 feasible, as most studies do not use measures that examine how engaged an individual is 558 with the material. As such, this current meta-analysis sought to provide readers with a 559 global effect of expressive writing on the aforementioned outcome variables. More studies are 560 needed to examine potential moderating effects of participant engagement. 561

The authors also note limitations in regards to the specific outcome variables. The nature of the construct of PTG makes it difficult to analyze rigorously. For example, on the Posttraumatic Growth Inventory (commonly used to study PTG), one could respond 0 to

the item "I have a greater appreciation for the value in my own life" because they already 565 had a high level of appreciation in their life (i.e., ceiling effect). This conceptual issue may 566 account for the non-effect of expressive writing on PTG. Logically, it would be difficult to 567 determine whether or not an individual experiences growth from trauma without having 568 experienced trauma. In conducting the literature search for the present meta-analysis, an 560 insufficient number of studies requiring a diagnosis of PTSD employed PTG as an outcome 570 variable. Thus, it was difficult to determine whether participants in the studies employed 571 had experienced trauma in line with DSM-IV criteria. For PTS, studies not specifying 572 whether or not participants had a diagnosis of PTSD were included. It is possible that 573 studies included in the subclinical symptom category did in fact include participants without 574 PTSD diagnosis (perhaps it was simply not assessed by means of a structured clinical 575 interview). It is also crucial to consider mainstream issues not specific to expressive writing and the outcome variables utilized in the present study. 577

The psychological scientific community has shifted focus to reproducibility and 578 research design in the last several years (Nelson, Simmons, & Simonsohn, 2018), and much of 570 this discussion has focused on adequately powering studies for publication (Bakker et al., 580 2016; S. E. Maxwell, Lau, & Howard, 2015). S. E. Maxwell et al. (2015) and Open Science 581 Collaboration (2015) have shown that the "replication crisis" may be attributed to low power 582 in published studies. The power found in the current meta-analysis was very poor, with very 583 few studies reaching the suggested 80% criterion to adequately power their study. This result 584 was the same when considering individual study characteristics or the estimate true 585 population effect size. Research by Bakker et al. (2016) indicates that researchers' intuitions about power are particularly poor, and many studies could benefit from more informed power analyses. Anderson, Kelley, and Maxwell (2017) recently published a primer on power, 588 with an online application to help with sample size planning for many types of research 589 designs. Additionally, we encourage researchers to report power analyses of studies in order 590 to better understand methodology for replication and reproducibility. 591

Meta-analyses, while useful tools to pool for population effect sizes, contain various 592 limitations to their usefulness (van Elk et al., 2015). As mentioned previously, these analyses 593 can be affected by high heterogeneity, which was found in this study (van Aert et al., 2016). 594 Selection models have been criticized when using a smaller number of studies (van Assen et 595 al., 2015), and trim and fill analyses may not always estimate accurate confidence intervals 596 and funnel plots may be biased with heterogeneity (Terrin, Schmid, Lau, & Olkin, 2003). 597 When focusing on improving the psychological sciences, van Elk et al. (2015) suggest that 598 the reliability and size of effects may be best elucidated by conducting large preregistered 599 studies. This suggestion will also improve the outlook for power in published studies, and 600 projects such as Many Labs can aide in subsidizing large samples (Klein et al., 2014). Even 601 with limitations, meta-analyses allow researchers to examine the state of a research area, and 602 we find potential with expressive writing on reducing PTS symptoms, and an overall need for 603 better sample size and power planning for studies.

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Table 1 Effect Size Estimates for PTS Results

Model	Fixed Effects	Random Effects	Fixed No Outliers	Random No Outliers
Overall Effects	0.36 [0.34, 0.39]	0.42 [0.35, 0.49]	0.36 [0.33, 0.38]	0.40 [0.33, 0.46]
Z Values	24.64, p < .001	12.35, p < .001	23.97, p < .001	12.38, p < .001
<i>p</i> -Uniform	0.63 [0.54, 0.72]	-	$0.61 \ [0.52, \ 0.70]$	-
PET	0.09 [0.01, 0.18]	-	$0.14 \ [0.06, \ 0.22]$	-
PEESE	0.24 [0.20, 0.29]	-	$0.26 \ [0.22, \ 0.31]$	-
Selection Models	0.33 [0.28, 0.37]	$0.45 \ [0.33, \ 0.57]$	$0.29 \ [0.24, \ 0.33]$	$0.39 \ [0.27, \ 0.50]$
Trim and Fill	$0.28 \ [0.25, \ 0.31]$	0.28 [0.21, 0.36]	$0.28 \ [0.25, \ 0.31]$	$0.28 \ [0.21, \ 0.35]$

Note. [] indicates the 95 percent confidence interval for each effect size estimate.

 $\begin{tabular}{ll} Table 2 \\ Effect Size Estimates for PTG Results \end{tabular}$

Model	Fixed Effects	Random Effects
Overall Effects	0.10 [0.02, 0.17]	0.10 [0.02, 0.17]
Z Values	2.45, p = .014	2.45, p = .014
<i>p</i> -Uniform	-0.11 [-1.43, 0.42]	-
PET	0.06 [-0.20, 0.32]	-
PEESE	0.08 [-0.04, 0.20]	-
Selection Models	0.09 [-0.01, 0.18]	0.09 [-0.03, 0.20]
Trim and Fill	$0.10 \ [0.02, \ 0.17]$	$0.10 \ [0.02, \ 0.17]$

Note. [] indicates the 95 percent confidence interval for each effect size estimate.

Table 3 ${\it Effect Size Estimates for QOL Results}$

Model	Fixed Effects	Random Effects	Fixed No Outliers	Random No Outliers
Overall Effects	-0.01 [-0.07, 0.05]	-0.01 [-0.16, 0.13]	-0.01 [-0.07, 0.05]	-0.01 [-0.11, 0.09]
Z Values	-0.33, p = .745	-0.18, p = .860	-0.25, p = .805	-0.20, p = .838
<i>p</i> -Uniform	0.79 [0.33, 1.61]	-	$0.62 \ [0.10, \ 0.96]$	-
PET	0.05 [-0.26, 0.36]	-	0.05 [-0.29, 0.38]	-
PEESE	0.00 [-0.17, 0.17]	-	0.00 [-0.19, 0.19]	-
Selection Models	-0.06 [-0.12, 0.01]	0.51 [-0.09, 1.12]	-0.04 [-0.11, 0.03]	$0.05 \ [-0.15, \ 0.24]$
Trim and Fill	-0.01 [-0.07, 0.05]	-0.01 [-0.16, 0.13]	-0.01 [-0.07, 0.05]	-0.01 [-0.11, 0.09]

 $\it Note.$ [] indicates the 95 percent confidence interval for each effect size estimate.

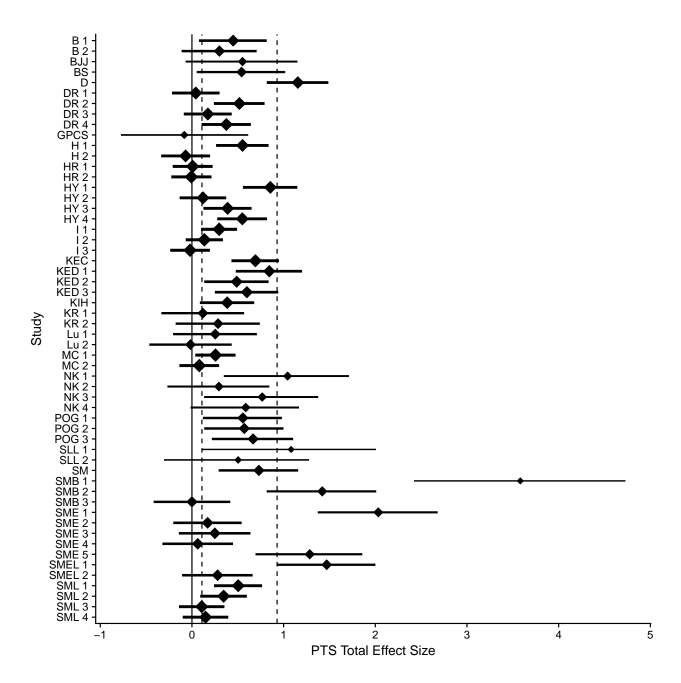


Figure 1. Effect sizes and their non-centralized confidence interval for PTS total scores. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.

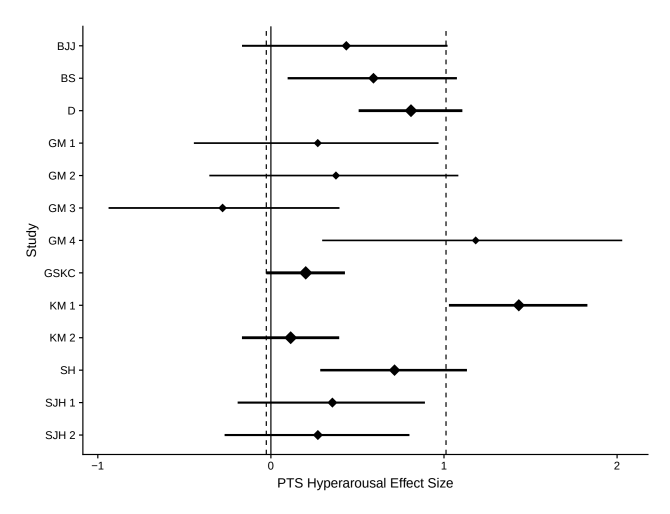


Figure 2. Effect sizes and their non-centralized confidence interval for PTS Hyperarousal. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.

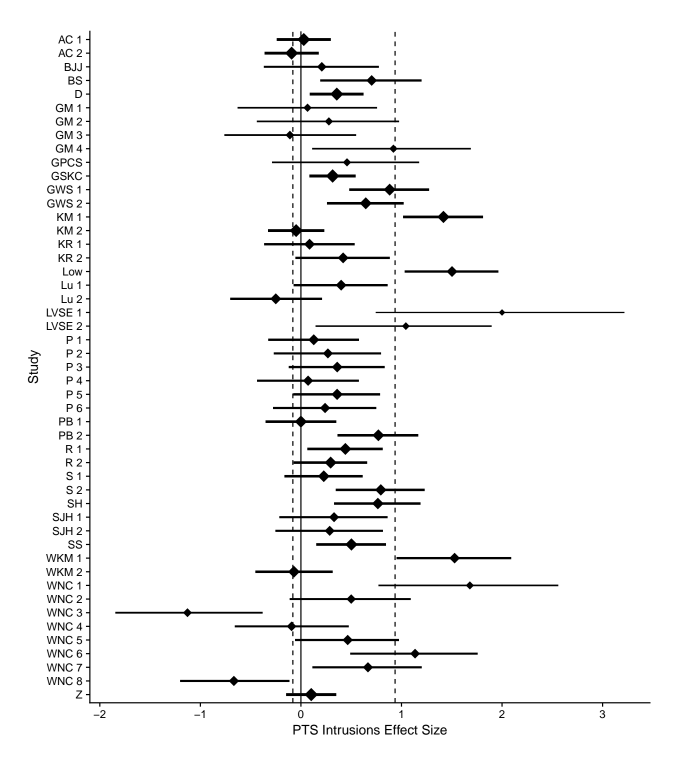


Figure 3. Effect sizes and their non-centralized confidence interval for PTS Intrusion scores. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.

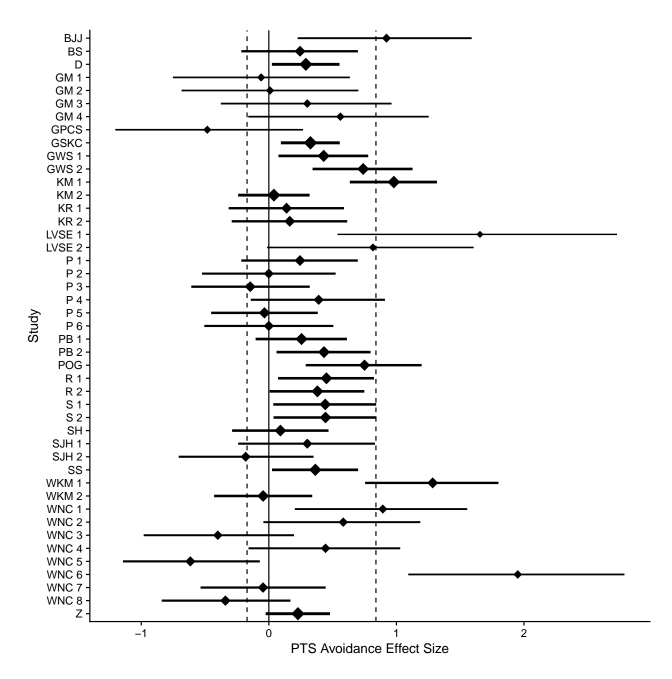


Figure 4. Effect sizes and their non-centralized confidence interval for PTS Avoidance Scores. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.

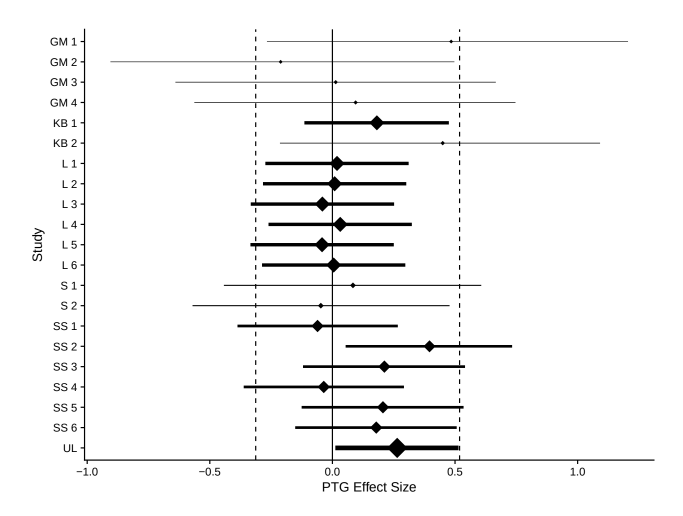


Figure 5. Effect sizes and their non-centralized confidence interval for PTG outcome variables. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.

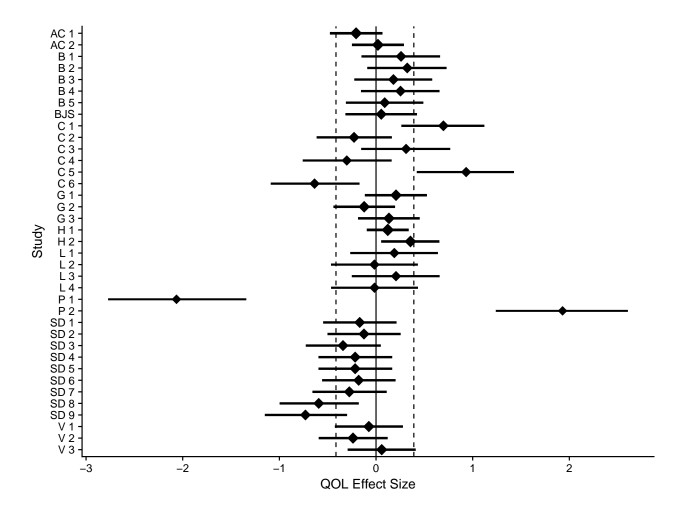


Figure 6. Effect sizes and their non-centralized confidence interval for QOL outcome variables. Dashed lines indicated average non-weighted lower and upper confidence interval limits. Diamond size indicates normalized study weight from a random effects model.