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1	Depression Prevalence using the HADS-D Compared to SCID Major Depression
2	Classification: an Individual Participant Data Meta-Analysis
3	
4	Running head: Estimating Depression Prevalence using the HADS-D and SCID
5	
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161 ABSTRACT

162 **Objectives:** Validated diagnostic interviews are required to classify depression status and

163 estimate prevalence of disorder, but screening tools are often used instead. We used individual

164 participant data meta-analysis to compare prevalence based on standard Hospital Anxiety and

- 165 Depression Scale depression subscale (HADS-D) cutoffs of ≥ 8 and ≥ 11 versus Structured
- 166 Clinical Interview for DSM (SCID) major depression and determined if an alternative HADS-D
- 167 cutoff could more accurately estimate prevalence.

168 Methods: We searched Medline, Medline In-Process & Other Non-Indexed Citations via Ovid,

169 PsycINFO, and Web of Science (inception-July 11, 2016) for studies comparing HADS-D scores

170 to SCID major depression status. Pooled prevalence and pooled differences in prevalence for

171 HADS-D cutoffs versus SCID major depression were estimated.

172 **Results:** 6,005 participants (689 SCID major depression cases) from 41 primary studies were

included. Pooled prevalence was 24.5% (95% Confidence Interval (CI): 20.5%, 29.0%) for

174 HADS-D \geq 8, 10.7% (95% CI: 8.3%, 13.8%) for HADS-D \geq 11, and 11.6% (95% CI: 9.2%,

- 175 14.6%) for SCID major depression. HADS-D \geq 11 was closest to SCID major depression
- 176 prevalence, but the 95% prediction interval for the difference that could be expected for HADS-
- 177 $D \ge 11$ versus SCID in a new study was -21.1% to 19.5%.

178 **Conclusions:** HADS-D \geq 8 substantially overestimates depression prevalence. Of all possible

179 cutoff thresholds, HADS-D \geq 11 was closest to the SCID, but there was substantial heterogeneity

- in the difference between HADS-D \geq 11 and SCID-based estimates. HADS-D should not be
- 181 used as a substitute for a validated diagnostic interview.
- 182 Key Words: depression, Hospital Anxiety and Depression Scale, individual participant data,
- 183 meta-analysis, screening tools

184 INTRODUCTION

185 Accurately measuring depression prevalence in different populations is important to 186 understand disease burden, interpret research on etiology, and utilize healthcare resources as 187 efficiently as possible (Rogan & Gladen, 1978). In mental health research, diagnostic interviews 188 are required for diagnosis of major depression (First, Spitzer, Gibbon, & Williams, 1995; 189 Wittchen, 1994). These interviews, however, are costly to administer, especially in large groups, 190 due to the time and trained personnel required to conduct them properly. Therefore, self-report 191 screening questionnaires are sometimes used as an inexpensive alternative to evaluate depression 192 prevalence, with the percentage of patients scoring above a cutoff threshold being described as 193 the prevalence of depression (Levis et al., 2019; Thombs, Kwakkenbos, Levis, & Benedetti, 194 2018). Screening tool cutoffs, however, are typically set to cast a wide net and identify many 195 more individuals for further assessment than will meet diagnostic criteria. Thus, commonly used 196 screening tools tend to overestimate depression prevalence, sometimes substantially (Thombs et 197 al., 2018).

198 A previous study used an individual participant data meta-analysis (IPDMA) approach to 199 compare prevalence based on a depression screening tool with prevalence based on a validated 200 diagnostic interview. That meta-analysis examined prevalence based on the Patient Health 201 Questionnaire-9 (PHQ-9) using the standard cutoff of \geq 10 compared to prevalence based on the 202 Structured Clinical Interview for the DSM (SCID) among 9,242 participants from 44 primary 203 studies (Levis et al., 2020). Compared to the SCID, PHQ-9 \geq 10 overestimated prevalence by 204 11.9%; across included studies, the mean and median ratio of PHQ-9 prevalence to SCID-based 205 prevalence were 2.5 and 1.9. In that study, the authors attempted to identify a PHQ-9 cutoff that

would match SCID-based prevalence, but heterogeneity was too high to generate consistentlyaccurate estimates in individual studies for any PHQ-9 cutoff.

208 The Hospital Anxiety and Depression Scale (HADS) is a self-report screening 209 questionnaire designed to be administered to non-psychiatric medical patients. It includes 14 210 items, with 7 assessing symptoms of depression (HADS-D) and 7 assessing symptoms of anxiety 211 (HADS-A) over the past week. To avoid overlap with physical illness, the HADS-D does not 212 include symptoms common to both physical and mental disorders, such as insomnia, loss of 213 appetite, or fatigue. Cutoff thresholds of ≥ 8 and ≥ 11 on the HADS-D are traditionally used as 214 standard cutoffs for identifying people who may have depression (Zigmond & Snaith, 1983). 215 Although not designed for this purpose, the HADS-D is also frequently used to report depression 216 prevalence in primary research studies. A review of recent studies listed in PubMed (2018-2019) 217 identified 32 studies that reported "prevalence" of depression based on a HADS-D cutoff, with \geq 218 8 and \geq 11 used in 66% and 16% of the studies, respectively (see supplementary material 219 eMethods 1 and eTable 1). 220 Although other screening tools and commonly used cutoffs have been shown to 221 overestimate depression prevalence, it is not clear whether this would be the case with the 222 HADS-D. A previous study that investigated prevalence of major depression among survivors of 223 acute myocardial infarction found a prevalence of 20% (10,785 participants, 8 studies) using 224 structured interviews, compared to 16% using a HADS-D cutoff of ≥ 8 (863 participants, 4 225 studies), and 7% using \geq 11 (830 participants, 4 studies) (Thombs et al., 2006). This was a

between-study comparison, however, and no included studies administered both the HADS-D

and a validated diagnostic interview.

The objectives of the present study were to use an IPDMA approach to (1) compare

229 pooled prevalence based on HADS-D cutoffs of ≥ 8 and ≥ 11 with major depression prevalence

based on the SCID; and (2) use a prevalence-matching approach to determine if any cutoff

threshold on the HADS-D matches prevalence based on the SCID with sufficiently low

heterogeneity that it could be used to accurately measure depression prevalence in future studies.

233 METHODS

This study used a subset of data collected for an IPDMA of the diagnostic accuracy of the HADS-D for screening to detect major depression. Detailed methods of the IPDMA were registered in PROSPERO (CRD42015016761), and a protocol was published (Thombs et al., 2016). The present analysis was not included in the original IPDMA protocol, which focused only on diagnostic accuracy. A protocol for the present study was published on the Open Science Framework prior to initiating the study (<u>https://osf.io/n5a3e/</u>).

240 Study Selection

241 In the main IPDMA, datasets from studies in any language were eligible for inclusion if 242 (1) they included HADS-D scores; (2) they included diagnostic classifications for current Major Depressive Episode (MDE) or Major Depressive Disorder (MDD) based on the Diagnostic and 243 244 Statistical Manual (DSM) or International Classification of Diseases criteria, using a validated 245 semi-structured or fully structured interview; (3) the HADS-D and diagnostic interview were 246 administered within two weeks of each other, since diagnostic criteria for major depression are 247 for symptoms experienced in the last two weeks; (4) participants were ≥ 18 years and not 248 recruited from youth or school-based settings, since the main IPDMA was designed for adult 249 screening, and although there are some adults in schools, the pathways for identification and 250 management are likely very different from other adult settings; and (5) participants were not

251 recruited from psychiatric settings or because they were identified as having symptoms of depression, since screening is done to identify unrecognized cases. Datasets where not all 252 253 participants were eligible were included if primary data allowed selection of eligible participants. 254 For the present study, we included only primary studies that based diagnoses on the SCID 255 (First et al., 1995). The SCID is a semi-structured diagnostic interview designed to be conducted 256 by an experienced clinician; it requires professional judgment and allows rephrasing questions 257 and probes to follow up responses. The reason for including only studies that used the SCID is 258 that in recent analyses using three large IPDMA databases (Levis et al., 2018, Levis et al., 2019, 259 Wu et al., 2020) we found that, compared to semi-structured interviews, fully structured 260 interviews, which are designed for administration by lay interviewers, may identify more patients 261 with low-level symptoms as depressed but fewer patients with high-level symptoms. These 262 results are consistent with the idea that semi-structured interviews most closely replicate clinical interviews done by trained professionals, whereas fully structured interviews are less rigorous 263 264 reference standards; they are less resource-intensive options that can be administered by research 265 staff without diagnostic skills but may misclassify major depression in substantial numbers of 266 patients. An important feature of the SCID is that it allows the interviewer to probe to determine 267 whether a symptom is merely a manifestation of a physical illness. In the HADS IPDMA 268 database, the SCID was the most commonly used semi-structured interview; out of 83 studies, 45 269 used semi-structured interviews, and 41 of the 45 used the SCID. In sensitivity analyses, we also 270 included the 4 studies from the IPDMA database that used semi-structured interviews other than 271 the SCID.

272 Data Sources and Searches

A medical librarian searched Medline, Medline In-Process & Other Non-Indexed
Citations via Ovid, PsycINFO, and Web of Science from inception to July 11, 2016, using a
peer-reviewed search strategy (McGowan et al., 2016) (see supplementary material eMethods 2).
We also reviewed reference lists of relevant reviews and queried contributing authors about nonpublished studies. Search results were uploaded into RefWorks (RefWorks-COS, Bethesda, MD,
USA). After de-duplication, unique citations were uploaded into DistillerSR (Evidence Partners,
Ottawa, Canada) for tracking search results.

280 Two investigators independently reviewed studies by title and abstract for eligibility. If

either deemed a study potentially eligible, a full-text review was done by both investigators

independently. Any disagreements were resolved by consensus and consulting a third

investigator when necessary. For languages other than those in which team members were fluent,translators were consulted.

285 Data Contribution and Synthesis

Authors of eligible datasets were invited to contribute de-identified primary data,

287 including HADS-D scores and major depression classification status. We emailed corresponding

authors of eligible primary studies at least three times, as necessary, with at least two weeks

between each email. If we did not receive a response, we emailed co-authors and attempted to

290 contact corresponding authors by phone.

291Before integrating individual datasets into our synthesized dataset, we compared292published participant characteristics and diagnostic accuracy results with results from raw

293 datasets and resolved any discrepancies in consultation with the original investigators.

294 Data Analysis

295 *Comparison of HADS-D* \geq 8 and \geq 11 *Prevalence with SCID Major Depression Prevalence*

296	For each primary study, we estimated 7 values: (1) the percentage of participants who
297	scored \geq 8 on the HADS-D, (2) the percentage of participants who scored \geq 11 on the HADS-D,
298	(3) the percentage of participants classified as having major depression based on the SCID, (4)
299	the difference between HADS-D \geq 8 percentage and SCID percentage, (5) the ratio for HADS-D
300	\geq 8 percentage versus SCID percentage, and the corresponding (6) difference and (7) ratio for
301	HADS-D \geq 11 versus the SCID. Then, across all studies, we pooled prevalence for HADS-D \geq 8,
302	HADS-D \geq 11, and SCID, and we pooled the HADS-D versus SCID differences in prevalence
303	from each study.
304	Prevalence Matching
305	To identify which HADS-D cutoff best matches SCID-based prevalence, we estimated
306	the pooled difference in prevalence for each possible HADS-D cutoff compared to the SCID.
307	The HADS-D cutoff with the smallest pooled difference was chosen to be the "prevalence-
308	matched cutoff." Then, for each included study, we estimated the difference and ratio in
309	prevalence based on the prevalence-matched cutoff versus SCID major depression. We
310	determined the mean and median absolute difference and the range of differences across all
311	studies. To illustrate the range of difference values that would be expected if a new study were to
312	compare prevalence based on the prevalence-matched cutoff to prevalence based on the SCID,
313	we estimated a 95% prediction interval for the difference.
314	All meta-analyses were conducted in R (R version R 3.4.1 and R Studio version 1.0.143)
315	using the lme4 package. To estimate pooled prevalence values, generalized linear mixed-effects

models with a logit link function were fit using the glmer function. To estimate pooled difference
values, linear mixed-effects models were fit using the lmer function. To account for correlation
between subjects within the same primary study, random intercepts were fit for each primary

study. To quantify heterogeneity, for each analysis, we calculated τ^2 , which is the estimate of between-study variance, and I², which quantifies the proportion of total variability due to the between-study heterogeneity.

322 We conducted two sets of post hoc analyses. First, some studies had high depression 323 prevalence. Thus, to test whether differences in prevalence between the HADS-D and SCID 324 might be influenced by heterogeneity in depression levels, we repeated the main analysis of 325 prevalence excluding studies with SCID-based prevalence $\geq 20.0\%$. Second, we assessed 326 whether differences in prevalence for the prevalence-matched cutoff and SCID were associated 327 with study or patient characteristics. To do this, we fit an additional linear mixed-effects model 328 for pooled prevalence difference, including age, sex, country human development index category ("very high" [reference group] or "high", based on the United Nation's Human Development 329 330 Index for the year of publication), recruitment setting category (nonmedical care, inpatient care 331 [reference group], outpatient care, or mixed inpatient and outpatient care), and sample size as 332 fixed-effect covariates. For this analysis, we excluded 520 participants (8.7%) who were missing 333 age or sex data. We repeated all analyses including 4 studies that used semi-structured interviews 334 other than the SCID.

335 **RESULTS**

The initial search for the main IPDMA found 10,015 unique titles and abstracts for potential eligibility. Of these, we excluded 9,584 studies after reviewing titles and abstracts and 238 studies after full-text review. There were 193 eligible studies using data from 133 unique samples from which 75 (56.4%) contributed individual participant data. Authors also contributed data from 8 unpublished studies, resulting in a total of 83 datasets. For our main analyses, we excluded 42 studies that used diagnostic interviews other than the SCID. In total, the main

analyses included 41 primary studies involving 6,005 participants (689 SCID major depression
cases; 11.5%; Figure 1). Of 58 eligible primary studies with unique samples that did not
contribute individual participant data, 26 used the SCID (3,096 participants). Thus, the main
analyses in the present study included 61.2% of eligible studies that used the SCID (41 of 67)
and 66.0% of eligible participants (6,005 of 9,101). See Table 1 for characteristics of each
included study.

There were 4 additional studies that used semi-structured diagnostic interviews other than the SCID (635 participants; 65 major depression cases; 10.2%), which we included in sensitivity analyses. Two of these studies used the Monash Interview for Liaison Psychiatry, one used the Schedule for Affective Disorders and Schizophrenia, and one used the Schedules for Clinical Assessment in Neuropsychiatry. Thus, these analyses included 45 primary studies (6,640 participants; 754 major depression cases; 11.4%; Table 1).

Objective 1: Comparison of HADS-D \geq 8, HADS-D \geq 11 and SCID Major Depression

- 355 Prevalence
- 356 *Pooled Prevalence*

The results for individual studies are presented in Table 1. For the 41 studies included in our main analyses, the percentage of participants who scored ≥ 8 on the HADS-D ranged from 4.2% to 82.7%, with a pooled prevalence of 24.5% (95% CI: 20.5% to 29.0%, τ^2 :0.49, I²: 97.2%). The percentage of participants who scored ≥ 11 on the HADS-D ranged from 0.3% to 74.7%, with a pooled prevalence of 10.7% (95% CI: 8.3% to 13.8%, τ^2 : 0.71, I²: 97.1%). The

- 362 percentage of participants classified as having SCID major depression ranged from 0% to 50.0%,
- 363 with a pooled prevalence of 11.6% (95% CI: 9.2% to 14.6%, τ^2 : 0.6, I²:97.1%).

Excluding 8 studies (552 participants; 185 major depression cases; 33.5%) with SCIDbased prevalence of 20.0% or over, prevalence based on the HADS-D \ge 8 was 21.8% (95% CI: 18.4% to 25.6%, τ^2 : 0.31, I²= 96.4). Prevalence based on the HADS-D \ge 11 was 9.2% (95% CI: 7.3% to 11.6%, τ^2 : 0.41, I²= 96.0). Prevalence based on the SCID was 8.9% (95% CI: 7.6% to 10.4%, τ^2 : 0.14, I²= 94.7).

- 369 Results were similar when the 4 studies using interviews other than the SCID were370 included.
- **371** *Pooled Difference and Ratio*
- 372 The difference between HADS-D \geq 8 and SCID-based prevalence in the main analyses
- 373 ranged from -9.5% to 41.3%, and the pooled difference was 12.4% (95% CI: 8.8% to 16%, τ^2 :

374 0.01, I²: 97.2%). The difference between HADS-D \geq 11 and SCID-based prevalence ranged from

- -31.0% to 33.3%, and the pooled difference was -0.8% (95% CI: -4.1% to 2.5%, τ^2 : 0.01, I²:
- 376 97.2%).
- 377 Results were similar in the sensitivity analyses. Pooled difference for HADS-D ≥ 8 was
- 378 11.9% (95% CI: 8.6% to 15.2%, τ^2 : 0.01, I²: 97.4%), and pooled difference for HADS-D ≥ 11
- 379 was -1.0% (95% CI: -4.0% to 2.0%, τ^2 : 0.01, I²: 97.5%)). The ratio of HADS-D \geq 8 prevalence
- to SCID major depression prevalence ranged from 0.4 to 7.7 times (mean: 2.6 times; median: 2
- times). The ratio of HADS-D \geq 11 prevalence to SCID major depression prevalence ranged from
- 382 0 to 3.8 times (mean: 1.2 times; median: 0.8 times).
- 383 *Mean Ratio and Difference in Individual Studies*
- In the main analyses, the mean ratio of HADS-D to SCID-based prevalence was 0.73
- times for the 3 studies with HADS-D \geq 8-based prevalence < 10.0% (mean difference: -2.7%),
- 1.8 times for the 7 studies with HADS-D \geq 8-based prevalence between 11.0% and 19.0% (mean

388 or greater (mean difference: 15.2%). The mean ratio was 0.7 times for the 19 studies with 389 HADS-D \geq 11-based prevalence < 10.0% (mean difference: -4.4%), 1.5 times for the 15 studies 390 with HADS-D \geq 11-based prevalence between 11.0% and 19.0% (mean difference: -1.3), and 2 391 times for the 7 studies with HADS-D \geq 11-based prevalence of 20.0% or greater (mean 392 difference: 9.8%). Results were similar when the 4 additional studies were included. **Objective 2: Prevalence Matching** 393 394 Of all possible HADS-D cutoffs, ≥ 11 produced the pooled prevalence estimate that most 395 closely matched SCID major depression prevalence (HADS-D \geq 11: 10.7%, SCID: 11.6%) 396 (Figure 2). This cutoff underestimated depression prevalence compared to the SCID, but only 397 slightly (pooled difference: -0.8%). HADS-D \geq 10 produced a pooled prevalence of 14.7% 398 (pooled difference: 3.1%), and HADS-D \geq 12 a pooled prevalence of 7.9% (pooled difference: -399 3.7%). The mean absolute difference between HADS-D \geq 11 and SCID was 8.2%, and the 400 median absolute difference was 6.7%. The 95% prediction interval for the difference between HADS-D > 11 and SCID-based prevalence was -21.1% to 19.5%. Results were similar in 401 402 sensitivity analyses. In the post-hoc analysis, no participant or study characteristics were

difference: 6.1%), and 2.9 times for the 31 studies with HADS-D \geq 8-based prevalence of 20.0%

403 significantly associated with differences in prevalence for the HADS-D prevalence-match cutoff

404 compared to the SCID.

405 **DISCUSSION**

387

406 Previous research has demonstrated that there may be substantial differences between
407 screening tools and diagnostic tools in estimating depression prevalence (Levis et al., 2020,

408 Thombs et al., 2018, Levis et al., 2019). In the present study, we found that the most commonly

409 used HADS-D cutoff threshold for reporting depression prevalence of ≥ 8 overestimated

410 depression prevalence (24.5%) substantially compared to SCID major depression prevalence 411 (11.6%). A HADS-D cutoff of \geq 11 underestimated prevalence only slightly in aggregate 412 compared to the SCID (10.7%), but heterogeneity in the difference between HADS-D \geq 11 and 413 SCID-based estimates in individual studies was high. The 95% prediction interval for difference 414 between HADS-D \geq 11 and SCID-based prevalence ranged from approximately -20% to 20%, 415 which suggests that any single new study using HADS-D \geq 11 may over or underestimate 416 depression prevalence by up to 20%.

Results from the present study are partially consistent with what might be expected 417 418 theoretically when comparing screening tools and diagnostic tools (Thombs et al., 2018). Since 419 screening tools are designed to cast a wide net and identify individuals who might be depressed, 420 they generally tend to overestimate depression prevalence when compared to diagnostic 421 interviews, which are designed to determine who meets diagnostic criteria. This was indeed the 422 case in our study for results from the HADS-D \geq 8, which were in line with those from a 423 previous study that found that the PHQ-9 similarly overestimated prevalence (Levis et al., 2020). 424 A finding that was unique to the present study was that estimates based on another commonly 425 used cutoff threshold, HADS \geq 11, were in aggregate consistent with major depression 426 prevalence based on the SCID. The findings from the present study differed from those in a 427 previous synthesis of evidence from post-acute myocardial infarction patients in which 428 depression prevalence estimates based on HADS-D ≥ 8 and ≥ 11 were both lower than estimates 429 based on structured interviews (Thombs et al., 2006). This discrepancy may be due to the 430 specific clinical population eligible for the review or because none of the studies included in that review administered both the HADS-D and a structured interview to the same group of 431 432 individuals.

Identifying a HADS-D cutoff that consistently matches the SCID would allow researchers to use screening questionnaires rather than diagnostic interviews for prevalence estimation, thus conserving time and resources. However, when we used a prevalence-matching approach and identified the closest HADS-D cutoff (≥ 11) to the SCID, although the aggregate estimates were similar, heterogeneity between studies was too high to suggest that HADS-D ≥ 11 would accurately estimate prevalence in any particular future study. In fact, it may substantially under or overestimate prevalence in individual studies.

Researchers often describe the proportion of individuals scoring at or above a cutoff 440 441 threshold as prevalence of "depressive symptoms" or "clinically significant depressive 442 symptoms" rather than prevalence of "depression". However, this does not resolve the problem. 443 There is no evidence that impairment becomes meaningful at or above these thresholds, which 444 have been set for the purpose of screening, and not for impairment delineation. While individuals 445 scoring above these thresholds have greater impairment on average than those scoring below the 446 threshold, this would be the case for any threshold that is set. Reporting the proportion of 447 individuals scoring above a threshold may be useful for comparisons between samples. However, 448 it should not be characterized as "prevalence" or as the percentage of individuals who have 449 "symptoms of depression" versus no symptoms.

Ideally, semi-structured interviews should be used for prevalence estimation, since they
provide patient-specific details that help interviewers determine whether the diagnostic criteria
for depression are met. They also most closely replicate full assessments done by trained
professionals (Wu et al., 2020). However, these interviews are not always feasible as they are
time-intensive compared to screening questionnaires. Diagnostic interviews also require trained
research staff or mental health professionals to conduct them properly. Hiring clinicians or

training research staff to do this can be costly and time-consuming, especially when assessing
large numbers of study participants. When determining which diagnostic interview to use,
researchers should consider the advantages and disadvantages of each, including performance,
cost, and required training (Wu et. al., 2020). When publishing studies, researchers should
discuss their reasons for selecting a particular interview, as well as the implications of their
selection.

462 To our knowledge, this is the first study to synthesize evidence and directly compare 463 depression prevalence based on HADS-D scores versus the SCID. Strengths of this study are that 464 we examined data from 41 primary research studies including 6,005 participants, and that we 465 directly compared status based on HADS-D scores to status based on a validated diagnostic 466 interview. A limitation is that we did not incorporate data from 39% of eligible studies that used 467 the SCID (26 of 61) and 34% of eligible participants (3,096 of 9,101), since they did not provide 468 individual participant data. Furthermore, since not all studies described the qualifications of the 469 individuals administering the SCID, it is possible that interviewer skill-level contributed to 470 heterogeneity. Since the objective of our study was to determine how accurate the HADS-D is 471 for estimating depression prevalence, we did not evaluate whether the correct individuals were 472 identified; that is beyond the scope of this study. Since diagnostic criteria for major depression 473 are for symptoms experienced in the last two weeks, we ensured that all studies administered the 474 HADS-D and SCID within two-weeks of each other. However, studies may not have 475 administered the HADS-D and SCID on the same day. This may have contributed to variability in responses to the SCID and the HADS-D, but it would not be expected to contribute to bias. 476 477 We included studies where diagnoses were based on DSM or ICD criteria, but only one study 478 used ICD (De Souza et. al., 2009). This study did not use the SCID and was included only in

479 sensitivity analyses. Finally, this study considered only the HADS-D, which is one screening tool 480 out of many that are commonly used in clinical practice. As shown in this study, the degree to 481 which the use of screening tools may accurately estimate prevalence depends on the specific 482 screening tool and cutoff threshold used. 483 In conclusion, we found that the standard HADS-D cutoff of ≥ 8 , which is most 484 commonly used by researchers to estimate depression prevalence, resulted in overestimation when compared to the SCID. The other standard screening cutoff of ≥ 11 most closely matched 485 486 SCID prevalence, but heterogeneity in the difference between HADS-D and SCID-based 487 estimates in individual studies was high and not associated with study or participant 488 characteristics. Findings are consistent with evidence demonstrating that depression screening 489 tools should not be used for diagnostic purposes. Studies should only report prevalence of 490 depression if they used a validated diagnostic interview designed for case classification. 491 Clinicians and researchers should be aware that the prevalence of depression reported in studies 492 using depression screening tools may not be accurate.

Contributors:

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495		CGL, NDM, MT (DEPRESSD Knowledge Users), ABenedetti, and BDT (DEPRESSD
496		Directors) were responsible for the conception, design and oversight of the main IPDMA
497		project of which the present study is a part.
498	•	EB, DN, BLevis, JPAI, ABenedetti, and BDT were responsible for the conception and
499		design of the present study
500	•	JTB and LAK designed and conducted database searches to identify eligible studies.
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506		and BDT contributed to data extraction and coding for the meta-analysis.
507	•	EB, DN, BLevis, ABenedetti, and BDT contributed to the data analysis and
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509	•	EB, DN, BLevis, YW, and BDT contributed to drafting the manuscript.
510	•	All authors provided a critical review and approved the final manuscript. ABenedetti and
511		BDT are the guarantors; they had full access to all the data in the study and take
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513		

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515 All authors have completed the Unified Competing Interest form at

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Table 1. Characteristics of included studies.

Author, year	Country	Population	N total	N (%) Major Depression	Mean Age	% Female	N (%) HADS-D ≥8	% Difference: HADS-D≥8 - Major Depression	Ratio: HADS-D ≥ 8 / Major Depressi on	N (%) HADS-D ≥ 11	% Difference: HADS-D≥11 - Major Depression	Ratio: HADS-D ≥11 / Major Depressi on	
Studies from IPDMA that used the SCID and were included in main analyses													
Akechi, 2006	Japan	Outpatients with cancer in palliative care	223	17 (8.0%)	61.1	65.0%	97 (43.0%)	35.9%	5.7	43 (19.0%)	11.7%	2.5	
Amoozegar, 2017	Canada	Patients with migraines	102	51 (50.0%)	42.5	81.4%	53 (52.0%)	2.0%	1.0	32 (31.0%)	-18.6%	0.6	
Beraldi, 2014	Germany	Patients of haemato- oncology	120	10 (8.0%)	52.1	32.5%	32 (27.0%)	18.3%	3.2	16 (13.0%)	5.0%	1.6	
Braeken, 2010	Netherla nds	Dutch cancer patients in radiotherapy	13	1 (8.0%)	69.4	NR	4 (31.0%)	23.1%	4.0	2 (15.0%)	7.7%	2.0	
Cukor, 2008	USA	Patients with end-stage renal disease	70	14 (20.0%)	53.3	52.9%	18 (26.0%)	5.7%	1.3	7 (10.0%)	-10.0%	0.5	
da Rocha e Sil va, 2013	Brazil	Patients with stroke	47	14 (30.0%)	59.8	51.1%	16 (34.0%)	4.3%	1.1	7 (15.0%)	-14.9%	0.5	
Ferentinos, 2011	Greece	Patients with amyotrophic	36	8 (22.0%)	62.0	41.7%	11 (31.0%)	8.3%	1.4	6 (17.0%)	-5.6%	0.7	

		lateral sclerosis										
Fiest, 2014	Canada	Patients with epilepsy	180	30 (17.0%)	41.1	51.4%	31 (17.0%)	0.6%	1.0	18 (10.0%)	-6.7%	0.6
Fischer, 2014	Germany	Patients with heart failure	194	11 (6.0%)	65.9	20.6%	49 (25.0%)	19.6%	4.5	25 (13.0%)	7.2%	2.3
Gagnon, 2005	Canada	Patients admitted to hospital due to fall	108	14 (13.0%)	78.1	87.0%	22 (20.0%)	7.4%	1.6	7 (6.0%)	-6.5%	0.5
Goebel, 2011	Germany	Patients with brain tumors	26	0 (0.0%)	58.3	50.0%	5 (19.0%)	19.2%	-	1 (4.0%)	3.8%	_
Golden, 2006	Ireland	Outpatients with Hepatitis C	86	7 (8.0%)	37.7	25.6%	24 (28.0%)	19.8%	3.4	11 (13.0%)	4.7%	1.6
Gould, 2011	Australia	Patients with traumatic brain injury	189	15 (8.0%)	35.7	21.7%	35 (19.0%)	10.6%	2.3	12 (6.0%)	-1.6%	0.8
Honarmand, 2009	Canada	Patients with multiple sclerosis	140	9 (6.0%)	43.9	74.3%	26 (19.0%)	12.1%	2.9	10 (7.0%)	0.7%	1.1
Juliao, 2013	Portugal	Patients with advanced disease	75	31 (41.0%)	NR	NR	62 (83.0%)	41.3%	2.0	56 (75.0%)	33.3%	1.8
Keller, 2004	Germany	Inpatients with cancer at the department of surgery	76	4 (5.0%)	56.7	38.2%	22 (29.0%)	23.7%	5.5	15 (20.0%)	14.5%	3.8

Kjaergaard, 2014	Norway	Healthy population	357	20 (6.0%)	52.5	100.0%	15 (4.0%)	-1.4%	0.8	1 (0.3%)	-5.3%	0
Kugaya, 2000	Japan	Inpatients with Cancer	81	3 (4.0%)	61.2	25.9%	23 (28.0%)	24.7%	7.7	9 (11.0%)	7.4%	3.0
Lambert, 2015	Australia	Patients with cancer	164	25 (15.0%)	58.5	65.9%	33 (20.0%)	4.9%	1.3	16 (10.0%)	-5.5%	0.6
Löwe, 2002	Germany	Medical outpatients	497	64 (13.0%)	41.8	66.4%	193 (39.0%)	26.0%	3.0	100 (20.0%)	7.2%	1.6
Meyer, 2008	Germany	Patients undergoing laryngectom y	102	4 (4.0%)	60.4	93.1%	25 (25.0%)	20.6%	6.2	13 (13.0%)	8.8%	3.2
Michopoulos, 2010	Greece	Elderly inpatients	194	27 (14.0%)	74.0	47.9%	83 (43.0%)	28.9%	3.1	47 (24.0%)	10.3%	1.7
Navines, 2012	Spain	Patients with chronic hepatitis C	500	32 (6.0%)	43.4	30.6%	74 (15.0%)	8.4%	2.3	31 (6.0%)	-0.2%	1.0
Öztürk, 2013	Turkey	Patients with acne	45	7 (16.0%)	20.9	80.0%	14 (31.0%)	15.6%	2.0	5 (11.0%)	-4.4%	0.7
Patten, 2015	Canada	Patients with multiple sclerosis	42	20 (48.0%)	NR	28.6%	16 (38.0%)	-9.5%	0.8	7 (17.0%)	-31%	0.4
Pintor, 2006	Spain	Patients on waiting list for heart transplantati on	73	13 (18.0%)	55.2	16.4%	15 (21.0%)	2.7%	1.2	8 (11.0%)	-6.8%	0.6
Rooney, 2013	UK	Adults with cerebral glioma	133	15 (11.0%)	53.7	42.9%	20 (15.0%)	3.8%	1.3	9 (7.0%)	-4.5%	0.6

Ryan, 2012	Ireland	Patients with advanced cancer	203	8 (4.0%)	61.6	49.3%	46 (23.0%)	18.7%	5.8	16 (8.0%)	3.9%	2.0
Sanchez- Gistau, 2012	Spain	Patients with epilepsy	296	35 (12.0%)	36.1	55.7%	74 (25.0%)	13.2%	2.1	40 (14.0%)	1.7%	1.1
Sanchez, 2012	Spain	Patients undergoing heart transplantati on	22	3 (14.0%)	54.2	9.1%	6 (27.0%)	13.6%	2.0	2 (9.0%)	-4.5%	0.7
Sanchez, 2014	Spain	Candidates for heart transplantati on	120	8 (7.0%)	55.6	22.5%	26 (22.0%)	15%	3.2	7 (6.0%)	-0.8%	0.9
Schwarzbold, 2014	Brazil	Patients with severe traumatic brain injury	44	14 (32.0%)	32.8	18.2%	12 (27.0%)	-4.5%	0.9	8 (18.0%)	-13.6%	0.6
Simard 2015	Canada	Survivors of cancer	60	7 (12.0%)	60.3	43.3%	3 (5.0%)	-6.7%	0.4	1 (2.0%)	-10%	0.1
Singer, 2008	Germany	Patients with laryngeal cancer	141	8 (6.0%)	63.7	8.5%	38 (27.0%)	21.3%	4.8	16 (11.0%)	5.7%	2.0
Singer, 2009	UK	Patients with cancer in acute care	580	55 (9.0%)	59.4	38.4%	200 (34.0%)	25%	3.6	101 (17.0%)	7.9%	1.8
Stone, 2004	UK	Outpatients after stroke	35	4 (11.0%)	71.2	31.4%	5 (14.0%)	2.9%	1.2	3 (9.0%)	-2.9%	0.8
Tung, 2015	China	Patients with diabetes	136	33 (24.0%)	39.8	56.6%	32 (24.0%)	-0.7%	1.0	12 (9.0%)	-15.4%	0.4

Turner, 2012	Australia	Patients after stroke	72	13 (18.0%)	66.7	47.2%	18 (25.0%)	6.9%	1.4	5 (7.0%)	-11.1%	0.4
Turner, Unpublished	Australia	Patients undergoing cardiac rehabilitatio n	52	4 (8.0%)	60.3	86.5%	4 (8.0%)	0%	1.0	3 (6.0%)	-1.9%	0.8
Walker, 2007	UK	Patients with cancer	361	30 (8.0%)	NR	23.5%	45 (12.0%)	4.2%	1.5	14 (4.0%)	-4.4%	0.5
Walterfang, 2007	Australia	Sample of Australian Patients with Adrenomyel oneuropathy	10	1 (10.0%)	43.8	10.0%	3 (30.0%)	20%	3.0	2 (20.0%)	10%	2.0
			Studi	es that used otl	her semi-	structured	interviews an	d were included in	sensitivity	analyses		
Love, 2002 ¹	Australia	Outpatients with breast cancer	Studi	es that used oth 28 (9.0%)	her semi- 46.3	structured 100.0%	interviews an 35 (12.0%)	d were included in 2.3%	1.2	analyses 8 (3.0%)	-6.60%	0.3
Love, 2002 ¹ Love, 2004 ²	Australia Australia	Outpatients with breast cancer Outpatients with breast cancer	Studi 302 227	es that used oth 28 (9.0%) 16 (7.0%)	her semi- 46.3 51.7	structured 100.0% 100.0%	interviews an 35 (12.0%) 43 (19.0%)	d were included in 2.3% 11.9%	1.2 2.7	analyses 8 (3.0%) 16 (7.0%)	-6.60% 0%	0.3
Love, 2002 ¹ Love, 2004 ² O'Rourke, 1998 ³	Australia Australia UK	Outpatients with breast cancer Outpatients with breast cancer Patients with stroke	Studi 302 227 56	es that used oth 28 (9.0%) 16 (7.0%) 9 (16.0%)	her semi- 46.3 51.7 67.1	structured 100.0% 100.0% 33.9%	interviews an 35 (12.0%) 43 (19.0%) 13 (23.0%)	d were included in 2.3% 11.9% 7.1%	1.2 2.7 1.4	analyses 8 (3.0%) 16 (7.0%) 7 (13.0%)	-6.60% 0% -3.60%	0.3 1.0 0.8

Author, year	Country	Population	N total	N (%) Major Depression	Mean Age	Percent Female	N (%) HADS-D ≥8	% Difference: HADS-D≥8 - Major Depression	Ratio: HADS-D ≥ 8 / Major Depressi on	N (%) HADS-D≥ 11	% Difference: HADS-D≥11 - Major Depression	Ratio: HADS-D ≥11 / Major Depressi on
			Stud anal	lies from IPDM yses	A that us	sed the SCI	D and were in	cluded in main				
Akechi, 2006	Japan	Outpatients with cancer in palliative care	223	17 (8.0%)	61.1	65.0	97 (43.0%)	35.9%	5.7	43 (19.0%)	11.7%	2.5
Amoozegar, 2017	Canada	Patients with migraines	102	51 (50.0%)	42.5	81.4	53 (52.0%)	2.0%	1.0	32 (31.0%)	-18.6%	0.6
Beraldi, 2014	Germany	Patients of haemato- oncology	120	10 (8.0%)	52.1	32.5	32 (27.0%)	18.3%	3.2	16 (13.0%)	5.0%	1.6
Braeken, 2010	Netherla nds	Dutch cancer patients in radiotherapy	13	1 (8.0%)	69.4	NA	4 (31.0%)	23.1%	4.0	2 (15.0%)	7.7%	2.0
Cukor, 2008	USA	Patients with end-stage renal disease	70	14 (20.0%)	53.3	52.9	18 (26.0%)	5.7%	1.3	7 (10.0%)	-10.0%	0.5
da Rocha e Sil va, 2013	Brazil	Patients with stroke	47	14 (30.0%)	59.8	51.1	16 (34.0%)	4.3%	1.1	7 (15.0%)	-14.9%	0.5
Ferentinos, 2011	Greece	Patients with amyotrophic lateral sclerosis	36	8 (22.0%)	62.0	41.7	11 (31.0%)	8.3%	1.4	6 (17.0%)	-5.6%	0.7

Fiest, 2014	Canada	Patients with epilepsy	180	30 (17.0%)	41.1	51.4	31 (17.0%)	0.6%	1.0	18 (10.0%)	-6.7%	0.6
Fischer, 2014	Germany	Patients with heart failure	194	11 (6.0%)	65.9	20.6	49 (25.0%)	19.6%	4.5	25 (13.0%)	7.2%	2.3
Gagnon, 2005	Canada	Patients admitted to hospital due to fall	108	14 (13.0%)	78.1	87.0	22 (20.0%)	7.4%	1.6	7 (6.0%)	-6.5%	0.5
Goebel, 2011	Germany	Patients with brain tumors	26	0 (0.0%)	58.3	50.0	5 (19.0%)	19.2%	_	1 (4.0%)	3.8%	_
Golden, 2006	Ireland	Outpatients with Hepatitis C	86	7 (8.0%)	37.7	25.6	24 (28.0%)	19.8%	3.4	11 (13.0%)	4.7%	1.6
Gould, 2011	Australia	Patients with traumatic brain injury	189	15 (8.0%)	35.7	21.7	35 (19.0%)	10.6%	2.3	12 (6.0%)	-1.6%	0.8
Honarmand, 2009	Canada	Patients with multiple sclerosis	140	9 (6.0%)	43.9	74.3	26 (19.0%)	12.1%	2.9	10 (7.0%)	0.7%	1.1
Juliao, 2013	Portugal	Patients with advanced disease	75	31 (41.0%)	NA	NA	62 (83.0%)	41.3%	2.0	56 (75.0%)	33.3%	1.8
Keller, 2004	Germany	Inpatients with cancer at the department of surgery	76	4 (5.0%)	56.7	38.2	22 (29.0%)	23.7%	5.5	15 (20.0%)	14.5%	3.8
Kjaergaard, 2014	Norway	Healthy population	357	20 (6.0%)	52.5	100.0	15 (4.0%)	-1.4%	0.8	1 (0.3%)	-5.3%	0

Kugaya, 2000	Japan	Inpatients with Cancer	81	3 (4.0%)	61.2	25.9	23 (28.0%)	24.7%	7.7	9 (11.0%)	7.4%	3.0
Lambert, 2015	Australia	Patients with cancer	164	25 (15.0%)	58.5	65.9	33 (20.0%)	4.9%	1.3	16 (10.0%)	-5.5%	0.6
Löwe, 2002	Germany	Medical outpatients	497	64 (13.0%)	41.8	66.4	193 (39.0%)	26.0%	3.0	100 (20.0%)	7.2%	1.6
Meyer, 2008	Germany	Patients undergoing laryngectom y	102	4 (4.0%)	60.4	93.1	25 (25.0%)	20.6%	6.2	13 (13.0%)	8.8%	3.2
Michopoulos, 2010	Greece	Elderly inpatients	194	27 (14.0%)	74.0	47.9	83 (43.0%)	28.9%	3.1	47 (24.0%)	10.3%	1.7
Navines, 2012	Spain	Patients with chronic hepatitis C	500	32 (6.0%)	43.4	30.6	74 (15.0%)	8.4%	2.3	31 (6.0%)	-0.2%	1.0
Öztürk, 2013	Turkey	Patients with acne	45	7 (16.0%)	20.9	80.0	14 (31.0%)	15.6%	2.0	5 (11.0%)	-4.4%	0.7
Patten, 2015	Canada	Patients with multiple sclerosis	42	20 (48.0%)	NA	28.6	16 (38.0%)	-9.5%	0.8	7 (17.0%)	-31%	0.4
Pintor, 2006	Spain	Patients on waiting list for heart transplantati on	73	13 (18.0%)	55.2	16.4	15 (21.0%)	2.7%	1.2	8 (11.0%)	-6.8%	0.6
Rooney, 2013	UK	Adults with cerebral glioma	133	15 (11.0%)	53.7	42.9	20 (15.0%)	3.8%	1.3	9 (7.0%)	-4.5%	0.6

Ryan, 2012	Ireland	Patients with advanced cancer	203	8 (4.0%)	61.6	49.3	46 (23.0%)	18.7%	5.8	16 (8.0%)	3.9%	2.0
Sanchez- Gistau, 2012	Spain	Patients with epilepsy	296	35 (12.0%)	36.1	55.7	74 (25.0%)	13.2%	2.1	40 (14.0%)	1.7%	1.1
Sanchez, 2012	Spain	Patients undergoing heart transplantati on	22	3 (14.0%)	54.2	9.1	6 (27.0%)	13.6%	2.0	2 (9.0%)	-4.5%	0.7
Sanchez, 2014	Spain	Candidates for heart transplantati on	120	8 (7.0%)	55.6	22.5	26 (22.0%)	15%	3.2	7 (6.0%)	-0.8%	0.9
Schwarzbold, 2014	Brazil	Patients with severe traumatic brain injury	44	14 (32.0%)	32.8	18.2	12 (27.0%)	-4.5%	0.9	8 (18.0%)	-13.6%	0.6
Simard 2015	Canada	Survivors of cancer	60	7 (12.0%)	60.3	43.3	3 (5.0%)	-6.7%	0.4	1 (2.0%)	-10%	0.1
Singer, 2008	Germany	Patients with laryngeal cancer	141	8 (6.0%)	63.7	8.5	38 (27.0%)	21.3%	4.8	16 (11.0%)	5.7%	2.0
Singer, 2009	UK	Patients with cancer in acute care	580	55 (9.0%)	59.4	38.4	200 (34.0%)	25%	3.6	101 (17.0%)	7.9%	1.8
Stone, 2004	UK	Outpatients after stroke	35	4 (11.0%)	71.2	31.4	5 (14.0%)	2.9%	1.2	3 (9.0%)	-2.9%	0.8
Tung, 2015	China	Patients with diabetes	136	33 (24.0%)	39.8	56.6	32 (24.0%)	-0.7%	1.0	12 (9.0%)	-15.4%	0.4

Turner, 2012	Australia	Patients after stroke	72	13 (18.0%)	66.7	47.2	18 (25.0%)	6.9%	1.4	5 (7.0%)	-11.1%	0.4
Turner, Unpublished	Australia	Patients undergoing cardiac rehabilitatio n	52	4 (8.0%)	60.3	86.5	4 (8.0%)	0%	1.0	3 (6.0%)	-1.9%	0.8
Walker, 2007	UK	Patients with cancer	361	30 (8.0%)	NA	23.5	45 (12.0%)	4.2%	1.5	14 (4.0%)	-4.4%	0.5
Walterfang, 2007	Australia	Sample of Australian Patients with Adrenomyel oneuropathy	10	1 (10.0%)	43.8	10.0	3 (30.0%)	20%	3.0	2 (20.0%)	10%	2.0
			Stud 39.8	71.2ies that us	ed other	semi-struct	ured interview	vs and were inclu	led in sensit	ivity analyses		
Love, 2002 ¹	Australia	Outpatients with breast cancer	Stud 39.8 302	71.2ies that us 28 (9.0%)	ed other 46.3	semi-struct	ured interview 35 (12.0%)	2.3%	ded in sensit	ivity analyses 8 (3.0%)	-6.60%	0.3
Love, 2002 ¹ Love, 2004 ²	Australia Australia	Outpatients with breast cancer Outpatients with breast cancer	Stud 39.8 302 227	71.2ies that us 28 (9.0%) 16 (7.0%)	ed other 46.3 51.7	semi-struct 100.0 100.0	ured interview 35 (12.0%) 43 (19.0%)	2.3% 11.9%	ded in sensit	ivity analyses 8 (3.0%) 16 (7.0%)	-6.60% 0%	0.3 1.0
Love, 2002 ¹ Love, 2004 ² O'Rourke, 1998 ³	Australia Australia UK	Outpatients with breast cancer Outpatients with breast cancer Patients with stroke	Stud 39.8 302 227 56	71.2ies that us 28 (9.0%) 16 (7.0%) 9 (16.0%)	ed other 46.3 51.7 67.1	semi-struct 100.0 100.0 33.9	ured interview 35 (12.0%) 43 (19.0%) 13 (23.0%)	2.3% 11.9% 7.1%	ded in sensit 1.2 2.7 1.4	ivity analyses 8 (3.0%) 16 (7.0%) 7 (13.0%)	-6.60% 0% -3.60%	0.3 1.0 0.8

^{1, 2} Diagnostic interview = Monash Interview for Liaison Psychiatry

³ Diagnostic interview = Schedule for Affective Disorders and Schizophrenia

⁴ Diagnostic interview = Schedules for Clinical Assessment in Neuropsychiatry

NR= Not reported







