

Comparison of the Accuracy of the 7-Item HADS Depression Subscale and 14-Item Total HADS for Screening for Major Depression: A Systematic Review and Individual Participant Data Meta-Analysis

Yin Wu^{1, 2, 3}, Brooke Levis^{1, 3, 4}, Federico M. Daray⁵, John P. A. Ioannidis^{6, 7, 8, 9}, Scott B. Patten¹⁰, Pim Cuijpers¹¹, Roy C. Ziegelstein¹², Simon Gilbody¹³, Felix H. Fischer¹⁴, Suiqiong Fan¹, Ying Sun¹, Chen He¹, Ankur Krishnan¹, Dipika Neupane¹, Parash Mani Bhandari¹, Zelalem Negeri¹, Kira E. Riehm¹, Danielle B. Rice¹, Marleine Azar¹, Xin Wei Yan¹, Mahrukh Imran¹, Matthew J. Chiovitti¹, Jill T. Boruff¹⁵, Dean McMillan¹³, Lorie A. Kloda¹⁶, Sarah Markham¹⁷, Melissa Henry¹, Zahinoor Ismail¹⁸, Carmen G. Loiselle¹⁹, Nicholas D. Mitchell²⁰, Samir Al-Adawi²¹, Kevin R. Beck²², Anna Beraldi²³, Charles N. Bernstein²⁴, Birgitte Boye²⁵, Natalie Büel-Drabe²⁶, Adomas Bunevicius²⁷, Ceyhun Can²⁸, Gregory Carter²⁹, Chih-Ken Chen³⁰, Gary Cheung³¹, Kerrie Clover³², Ronán M. Conroy³³, Gema Costa-Requena³⁴, Daniel Cukor³⁵, Eli Dabscheck³⁶, Jennifer De Souza³⁷, Marina Downing³⁸, Anthony Feinstein³⁹, Panagiotis P. Ferentinos⁴⁰, Alastair J. Flint⁴¹, Pamela Gallagher⁴², Milena Gandy⁴³, Luigi Grassi⁴⁴, Martin Härter⁴⁵, Asuncion Hernando⁴⁶, Melinda L. Jackson⁴⁷, Josef Jenewein⁴⁸, Nathalie Jetté⁴⁹, Miguel Julião⁵⁰, Marie Kjærgaard⁵¹, Sebastian Köhler⁵², Hans-Helmut König⁵³, Lalit K. R. Krishna⁵⁴, Yu Lee⁵⁵, Margrit Löbner⁵⁶, Wim L. Loosman⁵⁷, Anthony W. Love⁵⁸, Bernd Löwe⁵⁹, Ulrik F. Malt⁶⁰, Ruth Ann Marrie⁶¹, Loreto Massardo⁶², Yutaka Matsuoka⁶³, Anja Mehnert⁶⁴, Ioannis Michopoulos⁴⁰, Laurent Misery⁶⁵, Christian J. Nelson⁶⁶, Chong Guan Ng⁶⁷, Meaghan L. O'Donnell⁶⁸, Suzanne J. O'Rourke⁶⁹, Ahmet Öztürk⁷⁰, Alexander Pabst⁷¹, Julie A. Pasco⁷², Jurate Peceliuniene⁷³, Luis Pintor⁷⁴, Jennie L. Ponsford³⁸, Federico Pulido⁷⁵, Terence J. Quinn⁷⁶, Silje E. Reme⁷⁷, Katrin Reuter⁷⁸, Steffi G. Riedel-Heller⁷¹, Alasdair G. Rooney⁷⁹, Roberto Sánchez-González⁸⁰, Rebecca M. Saracino⁶⁶, Melanie P. J. Schellekens⁸¹, Martin Scherer⁸², Marcelo L. Schwarzbald⁸³, Vesile Senturk Cankorur⁸⁴, Louise Sharpe⁸⁵, Michael Sharpe⁸⁶, Sébastien Simard⁸⁷, Susanne Singer⁸⁸, Lesley Stafford⁸⁹, Jon Stone⁹⁰, Natalie A. Strobel⁹¹, Serge Sultan⁹², Antonio L. Teixeira⁹³, Istvan Tiringier⁹⁴, Alyna Turner⁹⁵, Jane Walker⁹⁶, Mark Walterfang⁹⁷, Liang-Jen Wang⁹⁸, Siegfried B. Weyerer⁹⁹, Jennifer White¹⁰⁰, Birgitt Wiese¹⁰¹, Lana J. Williams⁷⁵, Lai-Yi Wong¹⁰², Andrea Benedetti^{3, 103, 104, 105}, and Brett D. Thombs^{1, 2, 3, 105, 106, 107, 108}

¹ Lady Davis Institute for Medical Research, Jewish General Hospital, Montréal, Québec, Canada

² Department of Psychiatry, McGill University

³ Department of Epidemiology, Biostatistics and Occupational Health, McGill University

⁴ Centre for Prognosis Research, School of Medicine, Keele University

⁵ Institute of Pharmacology, School of Medicine, University of Buenos Aires

⁶ Department of Medicine, Stanford University

⁷ Department of Epidemiology and Population Health, Stanford University

⁸ Department of Biomedical Data Science, Stanford University

⁹ Department of Statistics, Stanford University

¹⁰ Department of Community Health Sciences, University of Calgary

¹¹ Department of Clinical, Neuro and Developmental Psychology, Amsterdam Public Health Research Institute, Vrije Universiteit

¹² Department of Medicine, Johns Hopkins University School of Medicine

¹³ Department of Health Sciences, Hull York Medical School, University of York

¹⁴ Department of Psychosomatic Medicine, Charité—Universitätsmedizin Berlin

¹⁵ Schulich Library of Physical Sciences, Life Sciences, and Engineering, McGill University

¹⁶ Library, Concordia University

¹⁷ Department of Biostatistics and Health Informatics, King's College London

¹⁸ Hotchkiss Brain Institute and O'Brien Institute for Public Health, University of Calgary

¹⁹ Ingram School of Nursing, McGill University

²⁰ Department of Psychiatry, University of Alberta

²¹ Department of Behavioural Medicine, College of Medicine & Health Sciences, Sultan Qaboos University

²² Department of Psychiatry, Singapore General Hospital, Singapore

²³ kbo Lech-Mangfall-Klinik für Psychiatrie, Psychotherapie und Psychosomatik, Garmisch-Partenkirchen, Bayern, Germany

- ²⁴ Department of Medicine, University of Manitoba
- ²⁵ Department of Behavioural Medicine, University of Oslo
- ²⁶ Department of Psychiatry and Psychotherapy, University Hospital Zürich
- ²⁷ Neuroscience Institute, Lithuanian University of Health Sciences
- ²⁸ Adana City Training and Research Hospital, Adana, Adana, Turkey
- ²⁹ School of Medicine and Public Health, University of Newcastle
- ³⁰ Community Medicine Research Center, Keelung Chang Gung Memorial Hospital, Chang Gung University College of Medicine
- ³¹ Department of Psychological Medicine, University of Auckland
- ³² Centre for Brain and Mental Health Research, University of Newcastle
- ³³ Division of Population Health Sciences, Royal College of Surgeons in Ireland
- ³⁴ Department of Psychiatry, Clinical Psychology, Hospital Universitari Vall d'Hebron, Universitat Autònoma de Barcelona, Centro de Investigación Biomédica en Red de Salud Mental
- ³⁵ Rogosin Institute, New York, New York, United States
- ³⁶ The Alfred Hospital, Prahran, Victoria, Australia
- ³⁷ Birmingham and Solihull Mental Health Foundation Trust, Birmingham, United Kingdom
- ³⁸ School of Psychological Sciences, Monash University
- ³⁹ Department of Psychiatry, University of Toronto
- ⁴⁰ 2nd Department of Psychiatry, Attikon General Hospital, National and Kapodistrian University of Athens
- ⁴¹ University Health Network, Toronto, Ontario, Canada
- ⁴² School of Psychology, Dublin City University
- ⁴³ The School of Psychological Sciences, Macquarie University, Sydney, New South Wales, Australia
- ⁴⁴ Department of Neuroscience and Rehabilitation, Institute of Psychiatry, University of Ferrara
- ⁴⁵ Department of Medical Psychology, University Medical Center Hamburg, University of Hamburg
- ⁴⁶ HIV Unit, Instituto de Investigacion Hospital 12 de Octubre, Madrid, Spain
- ⁴⁷ Turner Institute for Brain and Mental Health, Monash University
- ⁴⁸ Department of Medical Psychology and Psychotherapy, Medical University of Graz
- ⁴⁹ Department of Neurology, Icahn School of Medicine at Mount Sinai
- ⁵⁰ Equipa Comunitária de Suporte em Cuidados Paliativos de Sintra, Sintra, Portugal
- ⁵¹ Endocrinology Research Group, Medical Clinic, University Hospital of North Norway, Tromsø, Norway
- ⁵² Department of Psychiatry and Neuropsychology, School for Mental Health and Neuroscience, Maastricht University
- ⁵³ Department of Health Economics and Health Services Research, University Medical Center Hamburg-Eppendorf, Hamburg, Germany
- ⁵⁴ Department of Palliative Medicine, National Cancer Centre, Singapore
- ⁵⁵ Department of Psychiatry, Kaohsiung Chang Gung Memorial Hospital, Chang Gung University College of Medicine
- ⁵⁶ Institute of Social Medicine, Occupational Health and Public Health, University of Leipzig
- ⁵⁷ Onze Lieve vrouw Gasthuis, Amsterdam, The Netherlands
- ⁵⁸ Department of Psychology, Victoria University
- ⁵⁹ Department of Psychosomatic Medicine and Psychotherapy, University Medical Center Hamburg-Eppendorf, Hamburg, Germany
- ⁶⁰ Department of Research and Education Division of Surgery and Clinical Neuroscience, University of Oslo
- ⁶¹ Department of Medicine, Max Rady College of Medicine, Rady Faculty of Health Sciences, University of Manitoba
- ⁶² Centro de Biología Celular y Biomedicina, Facultad de Medicina y Ciencia, Universidad San Sebastián
- ⁶³ Division of Health Care Research, Center for Public Health Sciences, National Cancer Center, Tokyo, Japan
- ⁶⁴ Department of Medical Psychology and Medical Sociology, University of Leipzig
- ⁶⁵ Department of Dermatology, University Hospital of Brest, Brest, Finistère, France
- ⁶⁶ Department of Psychiatry and Behavioral Sciences, Memorial Sloan Kettering Cancer Center, New York, New York, United States
- ⁶⁷ Department of Psychological Medicine, Faculty of Medicine, University of Malaya
- ⁶⁸ Phoenix Australia, Carlton, Victoria, Australia
- ⁶⁹ School of Health in Social Sciences, University of Edinburgh
- ⁷⁰ Istanbul Sabahattin Zaim University
- ⁷¹ Institute of Social Medicine, Occupational Health and Public Health, Medical Faculty, University of Leipzig
- ⁷² The Institute for Mental and Physical Health and Clinical Translation, School of Medicine, Deakin University
- ⁷³ Faculty of Medicine, Clinic of Internal Diseases, Family Medicine and Oncology, Vilnius University
- ⁷⁴ Instituto de Investigaciones Biomédicas Augusto Pi i Sunyer, Barcelona, Spain
- ⁷⁵ HIV Unit, Hospital 12 de Octubre, imas12, UCM, Madrid, Spain
- ⁷⁶ Institute of Cardiovascular and Medical Sciences, University of Glasgow
- ⁷⁷ Department of Psychology, Faculty of Social Sciences, University of Oslo
- ⁷⁸ Private Practice for Psychotherapy and Psycho-oncology, Freiburg, Baden-Württemberg, Germany
- ⁷⁹ Division of Psychiatry, University of Edinburgh
- ⁸⁰ Department of Psychiatry, Institut de Neuropsiquiatria i Addiccions, Centre Emili Mira, Parc de Salut Mar, Barcelona, Spain
- ⁸¹ Scientific Research Department, Helen Dowling Institute, Bilthoven, The Netherlands
- ⁸² Institute of Primary Medical Care, University Medical Center Hamburg-Eppendorf, Hamburg, Germany
- ⁸³ Department of Internal Medicine, Federal University of Santa Catarina, Florianópolis, Santa Catarina, Brazil

- ⁸⁴ Department of Psychiatry, Faculty of Medicine, Ankara University
⁸⁵ School of Psychology, The University of Sydney
⁸⁶ Department of Psychological Medicine, University of Oxford
⁸⁷ Département des sciences de la santé, Université du Québec à Chicoutimi
⁸⁸ Institute of Medical Biostatistics, Epidemiology and Informatics, University Medical Centre Mainz, Mainz, Germany
⁸⁹ Melbourne School of Psychological Sciences, University of Melbourne, Victoria, Australia
⁹⁰ Centre for Clinical Brain Sciences, University of Edinburgh
⁹¹ Kurongkurl Katitjin, Edith Cowan University
⁹² Département de Psychologie, Faculté des arts et des sciences, Université de Montréal
⁹³ University of Texas Health Science Center at Houston, Houston, Texas, United States
⁹⁴ Medical School, Institute of Behavioral Sciences, Pécs University
⁹⁵ Faculty of Health and Medicine, School of Medicine and Public Health, University of Newcastle
⁹⁶ Department of Psychiatry, University of Oxford
⁹⁷ Neuropsychiatry Unit, Royal Melbourne Hospital, Melbourne, Australia
⁹⁸ Department of Child and Adolescent Psychiatry, Kaohsiung Chang Gung Memorial Hospital, Chang Gung University College of Medicine
⁹⁹ Central Institute of Mental Health, Medical Faculty Mannheim, Heidelberg University, Mannheim, Baden-Württemberg, Germany
¹⁰⁰ Department of Physiotherapy, School of Primary and Allied Health Care, Monash University
¹⁰¹ Institute of General Practice, Hannover Medical School
¹⁰² Kwai Chung Hospital, Hong Kong SAR, China
¹⁰³ Respiratory Epidemiology and Clinical Research Unit, McGill University Health Centre, Montréal, Québec, Canada
¹⁰⁴ Centre for Outcomes Research & Evaluation, Research Institute of the McGill University Health Centre, Montréal, Québec, Canada
¹⁰⁵ Department of Medicine, McGill University
¹⁰⁶ Department of Psychology, McGill University
¹⁰⁷ Department of Educational and Counselling Psychology, McGill University
¹⁰⁸ Biomedical Ethics Unit, McGill University

The seven-item Hospital Anxiety and Depression Scale Depression subscale (HADS-D) and the total score of the 14-item HADS (HADS-T) are both used for major depression screening. Compared to the HADS-D, the HADS-T includes anxiety items and requires more time to complete. We compared the screening accuracy of the HADS-D and HADS-T for major depression detection. We conducted an individual participant data meta-analysis and fit bivariate random effects models to assess diagnostic accuracy among participants with both HADS-D and HADS-T scores. We identified optimal cutoffs, estimated sensitivity and specificity with 95% confidence intervals, and compared screening accuracy across paired cutoffs via two-stage and individual-level models. We used a 0.05 equivalence margin to assess equivalency in sensitivity and specificity. 20,700 participants (2,285 major depression cases) from 98 studies were included. Cutoffs of ≥ 7 for the HADS-D (sensitivity 0.79 [0.75, 0.83], specificity 0.78 [0.75, 0.80]) and ≥ 15 for the HADS-T (sensitivity 0.79 [0.76, 0.82], specificity 0.81 [0.78, 0.83]) minimized the distance to the top-left corner of the receiver operating characteristic curve. Across all sets of paired cutoffs evaluated, differences of sensitivity between HADS-T and HADS-D ranged from -0.05 to 0.01 (0.00 at paired optimal cutoffs), and differences of specificity were within 0.03 for all cutoffs (0.02 – 0.03). The pattern was similar among outpatients, although the HADS-T was slightly (not nonequivalently) more specific among inpatients. The accuracy of HADS-T was equivalent to the HADS-D for detecting major depression. In most settings, the shorter HADS-D would be preferred.

Public Significance Statement

The present study suggests that the accuracy of 14-item Hospital Anxiety and Depression Scale (HADS-D) and the seven-item HADS Depression subscale (HADS-D) are equivalent for detecting major depression. Using the seven-item HADS-D for depression screening instead of the full 14-item HADS-T has minimal influence on performance of the measure but would reduce patient and participant burden in most clinical and research settings.

Keywords: HADS-D, HADS-T, individual participant data meta-analysis, depression screening, diagnostic accuracy

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Brett D. Thombs  <https://orcid.org/0000-0002-5644-8432>

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The 14-item Hospital Anxiety and Depression Scale (HADS; Zigmond & Snaith, 1983) was developed to facilitate the identification of anxiety disorders and major depression in people with a physical illness. The HADS includes two subscales. The seven-item Depression subscale (HADS-D) was designed to assess continuous depressive symptoms and for depression screening, whereas the seven-item Anxiety subscale (HADS-A) was designed to assess and screen for anxiety (Zigmond & Snaith, 1983). Both HADS-D and full HADS total scores (HADS-T) have been used to screen for major depression (Mitchell et al., 2010; Vodermaier & Millman, 2011). The HADS-T takes more time to complete and includes anxiety items not specific to depression. Some have suggested, though, that anxiety symptoms should be considered when assessing depression (Schatzberg, 2019). Furthermore, previous reviews have provided some preliminary evidence that HADS-T may perform better than the HADS-D (Mitchell et al., 2010; Vodermaier & Millman, 2011).

Commonly used HADS-D cutoff thresholds of ≥ 8 for “possible” depression and ≥ 11 for “probable” depression were established in the original validation study, which included only 100 participants (11 depression cases; Zigmond & Snaith, 1983). A recent individual participant data meta-analysis (IPDMA) on HADS-D accuracy to screen for major depression (101 studies; 22,574 participants; 2,549 major depression cases) found that a cutoff of ≥ 7 maximized combined sensitivity and specificity across reference standards; standard cutoffs of ≥ 8 and ≥ 11 were less sensitive but more specific (Wu, Levis, Sun, et al., 2021). There is not a standard cutoff for screening to detect major depression with the HADS-T.

Two previous meta-analyses, both done with studies of cancer patients, have indirectly compared the HADS-D and HADS-T for detecting major depression (Mitchell et al., 2010; Vodermaier & Millman, 2011). Both searched through October 2009 for eligible studies. One evaluated nine studies that used the HADS-D with a

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cutoff of 8 or greater and six studies that used the HADS-T with a cutoff of 15 (number of participants not reported; Mitchell et al., 2010), whereas the other included 2–5 studies each in analyses of HADS-D cutoffs of 7, 9, and 11 and HADS-T cutoffs of 15, 17, 19, and 20 (470–872 participants per analysis; Vodermaier & Millman, 2011). Both meta-analyses suggested that the HADS-T may perform better than the HADS-D, but there was a high level of uncertainty due to indirect comparisons between participants from different studies that reported HADS-D and HADS-T results, the small number of total participants, and possible selective outcome reporting bias (Levis et al., 2017; Neupane et al., 2021; Rice & Thombs, 2016; Thombs et al., 2011; Thombs & Rice, 2016) since not all primary studies reported results from the same cutoffs.

Using the full 14-item HADS-T for depression screening would be warranted if it is sufficiently more accurate than the shorter seven-item HADS-D to justify the additional time and patient burden involved. We previously assessed the accuracy of the HADS-D using IPDMA (Wu, Levis, Sun, et al., 2021). IPDMA involves a standard systematic review, followed by synthesis of original research data from primary studies, rather than extracting summary data (Riley et al., 2010). In that IPDMA, we found that diagnostic accuracy of HADS-D was not significantly different for any cutoffs across reference standards based on participant characteristics, including age, sex, cancer diagnosis, country human development index (HDI) levels, participant recruitment settings, or the study's risk of bias ratings (Wu, Levis, Sun, et al., 2021). In the present

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study, we included studies from the HADS-D IPDMA where HADS-T scores were provided or could be calculated from individual item scores. Our objectives were to (a) directly compare screening accuracy of the HADS-T and HADS-D for major depression detection using the same participant data across all studies regardless of reference standard, and (b) replicate the comparison among studies that used a semistructured diagnostic interview [e.g., *Diagnostic and Statistical Manual of Mental Disorders* Structured Clinical Interview for the (DSM SCID; First, 1995)] as a reference standard, since semistructured interviews more closely reflect the actual diagnostic process than fully structured interviews.

Method

The present study used a subset of studies and participants from our previously conducted HADS-D IPDMA (Wu, Levis, Sun, et al., 2021) for which HADS-T scores were also available. Analyses of

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Data contribution agreements with primary study authors do not include permission to make their data publicly available, although the data set used in this study will be archived through a McGill University repository (Borealis, <https://borealisdata.ca/dataverse/depressdproject/>). The R codes used for the analysis will be made publicly available through the same repository. Requests to access the data set to verify study results but not for other purposes can be sent to the corresponding authors via the "Access Data Set" function on the repository website.

Yin Wu played lead role in formal analysis, investigation, project administration, software, visualization and writing of original draft, supporting role in funding acquisition and equal role in conceptualization, data curation, methodology, resources and writing of review and editing. Brooke Levis played supporting role in conceptualization, data curation and formal analysis and equal role in methodology and writing of review and editing. Federico M. Daray played supporting role in writing of review and editing and equal role in conceptualization. John P. A. Ioannidis played supporting role in conceptualization and methodology and equal role in writing of review and editing. Scott B. Patten played supporting role in conceptualization and methodology. Pim Cuijpers played supporting role in conceptualization and methodology. Roy C. Ziegelstein played supporting role in conceptualization, methodology and writing of review and editing. Simon Gilbody played supporting role in conceptualization and methodology. Felix H. Fischer played supporting role in formal analysis, methodology and writing of review and editing. Suiqiong Fan played supporting role in data curation and formal analysis. Ying Sun played equal role in data curation and project administration. Chen He played supporting role in writing of review and editing and data contribution. Ankur Krishnan played supporting role in writing of review and editing and data contribution. Dipika Neupane played supporting role in writing of review and editing and data contribution. Parash Mani Bhandari played supporting role in writing of review and editing and data contribution. Zelalem Negeri played supporting role in writing of review and editing and data contribution. Kira E. Riehm played supporting role in writing of review and editing and data contribution. Danielle B. Rice played supporting role in writing of review and editing and data contribution. Marleine Azar played supporting role in writing of review and editing and data contribution. Xin Wei Yan played supporting role in writing of review and editing and data contribution. Mahrukh Imran played supporting role in writing of review and editing and data contribution. Matthew J. Chiovitti

HADS-D and HADS-T diagnostic accuracy were conducted according to the HADS-D IPDMA methods (Wu, Levis, Sun, et al., 2021) with the addition of analyses to directly compare HADS-D and HADS-T accuracy.

Data Set Eligibility

For the main HADS-D meta-analysis, data sets from articles in any language were eligible for inclusion if (a) they included diagnostic classification for current Major Depressive Disorder (MDD) or Major Depressive Episode (MDE) using *DSM* (American Psychiatric Association, 1987, 1994, 2000, 2013) or *International Classification of Diseases* (ICD; World Health Organization, 1992) criteria based on a validated semistructured or fully structured interview; (b) they included total scores for the HADS-D; (c) the diagnostic interview and HADS-D were administered within 2 weeks of each other, because *DSM* and ICD major

played supporting role in writing of review and editing and data contribution. Jill T. Boruff played supporting role in writing of review and editing and data contribution. Dean McMillan played supporting role in writing of review and editing and data contribution. Lorie A. Kloda played supporting role in writing of review and editing and data contribution. Sarah Markham played supporting role in writing of review and editing and data contribution. Melissa Henry played supporting role in writing of review and editing and data contribution. Zahinoor Ismail played supporting role in writing of review and editing and data contribution. Carmen G. Loiselle played supporting role in writing of review and editing and data contribution. Nicholas D. Mitchell played supporting role in writing of review and editing and data contribution. Samir Al-Adawi played supporting role in writing of review and editing and data contribution. Kevin R. Beck played supporting role in writing of review and editing and data contribution. Anna Beraldi played supporting role in writing of review and editing and data contribution. Charles N. Bernstein played supporting role in writing of review and editing and data contribution. Birgitte Boye played supporting role in writing of review and editing and data contribution. Natalie Büel-Drabe played supporting role in writing of review and editing and data contribution. Adomas Bunevicius played supporting role in writing of review and editing and data contribution. Ceyhun Can played supporting role in writing of review and editing and data contribution. Gregory Carter played supporting role in writing of review and editing and data contribution. Chih-Ken Chen played supporting role in writing of review and editing and data contribution. Gary Cheung played supporting role in writing of review and editing and data contribution. Kerrie Clover played supporting role in writing of review and editing and data contribution. Ronán M. Conroy played supporting role in writing of review and editing and data contribution. Gema Costa-Requena played supporting role in writing of review and editing and data contribution. Daniel Cukor played supporting role in writing of review and editing and data contribution. Eli Dabscheck played supporting role in writing of review and editing and data contribution. Jennifer De Souza played supporting role in writing of review and editing and data contribution. Marina Downing played supporting role in writing of review and editing and data contribution. Anthony Feinstein played supporting role in writing of review and editing and data contribution. Panagiotis P. Ferentinos played supporting role in writing of review and editing and data contribution. Alastair J. Flint played supporting role in writing of review and editing and data contribution. Pamela Gallagher played supporting role in writing of review and editing and data contribution. Milena Gandy played supporting role in writing of review and editing and data contribution. Luigi Grassi played supporting role in writing of review and editing and data contribution. Martin Härter played supporting role in writing of review and editing and data contribution. Asuncion Hernando played supporting role in writing of review and editing

depression diagnostic criteria specify that symptoms must have been present in the last 2 weeks; (d) participants were ≥ 18 years of age and not recruited from youth or psychiatric settings; and (e) participants were not recruited because they were identified as having symptoms of depression, since screening is done to identify previously unrecognized cases. We focused on MDD and MDE because major guidelines on depression screening have focused on screening for major depression but have not considered screening for less severe conditions, such as dysthymia or persistent depressive disorder, for which treatment options and effectiveness are much less well delineated (Joffres et al., 2013; National Collaborating Centre for Mental Health [UK], 2010; Siu et al., 2016). Consistent with this, few primary studies collect or report

diagnostic status for dysthymia or persistent depressive disorder. Data sets where not all participants were eligible were included if primary data allowed selection of eligible participants. For the present study, we only included primary data sets from the HADS-D IPDMA that also provided HADS-T scores or item scores to calculate HADS-T scores.

Search Strategy and Study Selection

A medical librarian searched Medline, Medline In-Process and other Non-Indexed Citations and PsycINFO via OvidSP, and Web of Science via the Institute for Scientific Information Web of Knowledge from inception to October 25, 2018 using a peer-reviewed

and data contribution. Melinda L. Jackson played supporting role in writing of review and editing and data contribution. Josef Jenewein played supporting role in writing of review and editing and data contribution. Nathalie Jetté played supporting role in writing of review and editing and data contribution. Miguel Julião played supporting role in writing of review and editing and data contribution. Marie Kjærgaard played supporting role in writing of review and editing and data contribution. Sebastian Köhler played supporting role in writing of review and editing and data contribution. Hans-Helmut König played supporting role in writing of review and editing and data contribution. Lalit K.R. Krishna played supporting role in writing of review and editing and data contribution. Yu Lee played supporting role in writing of review and editing and data contribution. Margrit Löbner played supporting role in writing of review and editing and data contribution. Wim L. Loosman played supporting role in writing of review and editing and data contribution. Anthony W. Love played supporting role in writing of review and editing and data contribution. Bernd Löwe played supporting role in writing of review and editing and data contribution. Ulrik F. Malt played supporting role in writing of review and editing and data contribution. Ruth Ann Marie played supporting role in writing of review and editing and data contribution. Loreto Massardo played supporting role in writing of review and editing and data contribution. Yutaka Matsuoka played supporting role in writing of review and editing and data contribution. Anja Mehnert played supporting role in writing of review and editing and data contribution. Ioannis Michopoulos played supporting role in writing of review and editing and data contribution. Laurent Misery played supporting role in writing of review and editing and data contribution. Christian J. Nelson played supporting role in writing of review and editing and data contribution. Chong Guan Ng played supporting role in writing of review and editing and data contribution. Meaghan L. O'Donnell played supporting role in writing of review and editing and data contribution. Suzanne J. O'Rourke played supporting role in writing of review and editing and data contribution. Ahmet Öztürk played supporting role in writing of review and editing and data contribution. Alexander Pabst played supporting role in writing of review and editing and data contribution. Julie A. Pasco played supporting role in writing of review and editing and data contribution. Jurate Peceliuniene played supporting role in writing of review and editing and data contribution. Luis Pintor played supporting role in writing of review and editing and data contribution. Jennie L. Ponsford played supporting role in writing of review and editing and data contribution. Federico Pulido played supporting role in writing of review and editing and data contribution. Terence J. Quinn played supporting role in writing of review and editing and data contribution. Silje E. Reme played supporting role in writing of review and editing and data contribution. Katrin Reuter played supporting role in writing of review and editing and data contribution. Steffi G. Riedel-Heller played supporting role in writing of review and editing and data contribution. Alasdair G. Rooney played supporting role in writing of review and editing and data contribution. Roberto Sánchez-González played supporting role in writing of review and editing and data contribution. Rebecca M. Saracino played supporting role in

writing of review and editing and data contribution. Melanie P.J. Schellekens played supporting role in writing of review and editing and data contribution. Martin Scherer played supporting role in writing of review and editing and data contribution. Marcelo L. Schwarzbald played supporting role in writing of review and editing and data contribution. Vesile Senturk Cankorur played supporting role in writing of review and editing and data contribution. Louise Sharpe played supporting role in writing of review and editing and data contribution. Michael Sharpe played supporting role in writing of review and editing and data contribution. Sébastien Simard played supporting role in writing of review and editing and data contribution. Susanne Singer played supporting role in writing of review and editing and data contribution. Lesley Stafford played supporting role in writing of review and editing and data contribution. Jon Stone played supporting role in writing of review and editing and data contribution. Natalie A. Strobel played supporting role in writing of review and editing and data contribution. Serge Sultan played supporting role in writing of review and editing and data contribution. Antonio L. Teixeira played supporting role in writing of review and editing and data contribution. Istvan Tiringier played supporting role in writing of review and editing and data contribution. Alyna Turner played supporting role in writing of review and editing and data contribution. Jane Walker played supporting role in writing of review and editing and data contribution. Mark Walterfang played supporting role in writing of review and editing and data contribution. Liang-Jen Wang played supporting role in writing of review and editing and data contribution. Siegfried B. Weyerer played supporting role in writing of review and editing and data contribution. Jennifer White played supporting role in writing of review and editing and data contribution. Birgitt Wiese played supporting role in writing of review and editing and data contribution. Lana J. Williams played supporting role in writing of review and editing and data contribution. Lai-Yi Wong played supporting role in writing of review and editing and data contribution. Andrea Benedetti played lead role in funding acquisition and writing of review and editing and equal role in conceptualization, formal analysis, methodology and supervision. Brett D. Thombs played lead role in funding acquisition and supervision and equal role in conceptualization, methodology, project administration and writing of review and editing.

The main HADS-D individual participant data meta-analysis (IPDMA) was registered in the International Prospective Register of Systematic Reviews (CRD42015016761), and a protocol was published (Thombs et al., 2016). The present study was not included in the protocol for the main HADS-D IPDMA, but a separate protocol was developed and posted online prior to initiating the study (<https://osf.io/438ak/>).

Correspondence concerning this article should be addressed to Brett D. Thombs, Lady Davis Institute for Medical Research, Jewish General Hospital, 3755 Cote Ste Catherine Road, Montréal, QC, H3T 1E2, Canada or Andrea Benedetti, Centre for Outcomes Research & Evaluation, Research Institute of the McGill University Health Centre, 5252 Boulevard de Maisonneuve, Montréal, QC, H4A 3S5, Canada. Email: brett.thombs@mcgill.ca or andrea.benedetti@mcgill.ca

(McGowan et al., 2016) search strategy (Supplemental Methods A). 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 deduplication, unique citations were uploaded into DistillerSR (Evidence Partners, Ottawa, Canada) for tracking search results.

Pairs of investigators independently reviewed titles and abstracts for eligibility. If either deemed a study potentially eligible, full-text review was done by two investigators, independently, with disagreements resolved by consensus, consulting a third investigator when necessary. Translators were consulted for languages other than those for which team members were fluent.

Data Contribution, Extraction, and Synthesis

Authors of eligible data sets were invited to contribute deidentified primary data. We emailed corresponding authors of eligible primary studies at least three times, as necessary. If we did not receive a response, we emailed coauthors and attempted to contact corresponding authors by phone.

Diagnostic interview and country were extracted from published reports by pairs of investigators independently, with disagreements resolved by consensus. Countries were categorized as “very high,” “high” or “low-medium” development based on the United Nation’s HDI for the country for the year of the study publication. The HDI is a statistical composite index that includes indicators of life expectancy, education, and income (United Nations Development Programme, 2020). Participant-level data included age, sex, participant recruiting setting, HADS-D scores, HADS-T scores, and major depression status (case or noncase). For defining major depression, we considered MDD or MDE based on the *DSM* or *ICD*. If more than one was reported, we prioritized MDE over MDD (because screening would attempt to detect depressive episodes and further interview would determine if the episode is related to MDD, bipolar disorder or persistent depressive disorder). We also prioritized *DSM* over *ICD* because most studies use *DSM* criteria.

Individual participant data were converted to a standard format and synthesized into a single data set with study-level data. We compared published participant characteristics and diagnostic accuracy estimates with results from raw data sets and resolved any discrepancies in consultation with primary study investigators.

Risk of Bias Assessment

Risk of bias of included studies was assessed by two investigators independently using the Quality Assessment of Diagnostic Accuracy Studies-2 tool (QUADAS-2; Supplemental Methods B; Whiting et al., 2011). Any discrepancies were resolved via consensus with a third investigator involved as necessary. Risk of bias was coded at both study and participant levels since some classifications (e.g., the time between index test and reference standard) may have differed among participants from the same study. The QUADAS-2 results were used to describe the risk of bias of each included study.

Statistical Analyses

To compare the screening accuracy of the HADS-D and HADS-T across relevant cutoffs to detect major depression, we first estimated overall sensitivity and specificity for HADS-D and HADS-T by

combining all studies regardless of reference standard. Reference standards used in primary studies included semistructured interviews (e.g., SCID; First, 1995), fully structured interviews (the Mini International Neuropsychiatric Interview [MINI] excluded; e.g., Composite International Diagnostic Interview; Robins et al., 1988), and the MINI (Lecrubier et al., 1997; Sheehan et al., 1997). Different types of reference standards have different design and performance characteristics (Levis et al., 2018; Levis, McMillan, et al., 2019; Wu, Levis, Ioannidis, et al., 2021; Wu, Levis, Sun, et al., 2020), and estimates of sensitivity and specificity differ by type (Levis, Benedetti, et al., 2019; Levis et al., 2020; Negeri et al., 2021; Wu, Levis, Sun, et al., 2021). It is reasonable to assume, though, that differences in sensitivity and specificity between HADS-D and HADS-T accuracy among the same participants are not associated with reference standard type, since in each primary study the HADS-D and HADS-T were compared to the same reference standard. Thus, our main analysis included all studies regardless of reference standard.

Separately, as a sensitivity analysis, to ensure that results would not differ by clinical interview, we repeated all analyses for only studies that used a semistructured interview as the reference standard. Semistructured interviews (e.g., SCID; First, 1995, Schedules for Clinical Assessment in Neuropsychiatry; World Health Organization, 1994, Schedule for Affective Disorders and Schizophrenia; Endicott & Spitzer, 1987, and Monash Interview for Liaison Psychiatry; Clarke et al., 1998) are intended to be administered by experienced diagnosticians and are considered to more closely reflect clinical diagnostic procedures than fully structured interviews or the MINI (Brugha et al., 1999, 2001; Nosen & Woody, 2008). We did not conduct additional sensitivity analyses with fully structured interviews or the MINI.

Overall and separately, for studies that used a semistructured reference standard, for all possible cutoffs 0–21 of the HADS-D and 0–42 of the HADS-T, we fitted bivariate random effects models via Gauss-Hermite quadrature (Riley et al., 2008). This is a two-stage meta-analytic approach that models sensitivity and specificity simultaneously and accounts for the correlation between them and the precision of estimates within studies. We also constructed empirical receiver operating characteristic (ROC) plots based on pooled sensitivity and specificity estimates and calculated area under the curves (AUC) for the two tests.

To investigate heterogeneity across studies, overall and for studies with a semistructured reference standard, we generated forest plots for the differences in sensitivity and specificity estimates between the HADS-D and HADS-T for the optimal cutoffs based on pooled results. We also quantified heterogeneity at the optimal cutoffs for the HADS-D and HADS-T by reporting the estimated variances of the random effects for the differences in the HADS-D and HADS-T sensitivity and specificity (τ^2 ; Fagerland et al., 2014; Higgins & Thompson, 2002).

To compare the diagnostic accuracy of the HADS-D and HADS-T, using the analyses that pooled across reference standards and within semistructured reference standard category, we first calculated the differences of the AUCs with 95% confidence intervals (CIs). Second, we compared the ROC plots visually to determine if one measure consistently perform better than the other across cutoffs. Third, we compared differences in sensitivity and specificity for optimal cutoffs and other cutoffs close to the optimal cutoff to determine if there were differences and the

magnitude of any differences. To do this, we identified the optimal cutoff that minimized the values of the distance to the top-left corner of the ROC curves (NCSS Statistical Software, 2017) for both HADS-D and HADS-T and a set of other cutoffs that were close to the optimal cutoff. The distance to the top-left corner of the ROC curve for each cutoff value is calculated by $d = \sqrt{(1 - \text{Sensitivity})^2 + (1 - \text{Specificity})^2}$ (NCSS Statistical Software, 2017). Since there is no a priori method to align cutoffs on the HADS-D and HADS-T that perform most similarly in terms of sensitivity and specificity, we did this based on examination of results and consensus among investigators. Then, we compared the sensitivity and specificity between the HADS-D and HADS-T for pairs of optimal cutoffs and four other pairs of cutoffs close to the optimal; the interval between cutoffs for HADS-T was 2 instead of 1 because HADS-T doubled the length and the total score of HADS-D. For all cutoffs on the HADS-D and HADS-T, 95% CIs for the differences between HADS-D and HADS-T sensitivity and specificity were constructed via a cluster bootstrap approach (van der Leeden et al., 1997, 2008) with resampling at the study and subject level. For each comparison, we ran 1,000 iterations of the bootstrap. For each bootstrap iteration, the bivariate random effects model was fitted to the HADS-D and HADS-T data, and the pooled sensitivities and specificities were computed separately, as described above, for all cutoffs of HADS-D and HADS-T.

In addition to comparing the HADS-D and HADS-T with pooling of study-level results, as a sensitivity analysis, we compared sensitivity and specificity of the HADS-D and HADS-T across cutoffs via an individual-level analysis. For the individual-level analysis, for each pair of matched HADS-D and HADS-T cutoffs, we fitted a linear mixed model with the difference between the HADS-D and HADS-T screening results as the outcome. The screening result is dichotomous, either positive = 1 or negative = 0. If the HADS-T screening result was positive (which was 1), but HADS-D was negative (which was 0), the outcome, that is, the difference between HADS-T and HADS-D results, was $1 - 0 = 1$; if both screening results were positive or negative, the outcome was $0 (1 - 1 \text{ or } 0 - 0)$; and if the HADS-T screening result was negative, but HADS-D was positive, the outcome was $-1 (0 - 1 = -1)$. This model modeled the differences in sensitivity and specificity simultaneously and included random effects both at the study level. From this model, for each set of HADS-D and HADS-T paired cutoffs, we estimated the difference in sensitivity and specificity between the two tests and associated CIs. These CIs from the bootstrap approach and individual-level analysis allowed us to test whether the sensitivity and specificity of the HADS-T is equivalent to that of the HADS-D based on a prespecified equivalence margin of $\delta = 0.05$ (Walker & Nowacki, 2011), as we have done in previous studies (Harel et al., 2021; Ishihara et al., 2019; Wu, Levis, Riehm, et al., 2020).

As a sensitivity analysis, we compared accuracy of HADS-D and HADS-T results stratified by subgroups based on inpatient and outpatient care settings (we planned to conduct sensitivity analysis in each participant recruit setting, separately, but we were able to do this only for inpatient and outpatient medical settings because there were too few participants from nonmedical and mixed inpatient/outpatient settings). In addition, we conducted a subgroup analysis only among patients from cancer studies because meta-analyses (Mitchell et al., 2010; Vodermaier & Millman, 2011) of studies from cancer care settings reported that the HADS-T may perform better than the HADS-D in those settings. We did not conduct the

sensitivity analysis to assess whether inclusion of published results from the eligible studies that did not provide raw data influenced results because we did this in the main HADS-D IPDMA and found no differences (Wu, Levis, Sun, et al., 2021).

To examine whether measurement differences across participant characteristics, including country, may have influenced our results, we assessed whether sensitivity and specificity differed for the HADS-D based on these characteristics, and then, we reexamined HADS-D and HADS-T differences for any variables where differences were found. To assess possible influences on sensitivity and specificity, we conducted one-stage metaregressions. In the first step, we repeated the analysis that we did in the main HADS-D IPDMA by interacting all subgrouping variables (age [measured continuously], sex [reference category = female]), country HDI level [reference category = very high], cancer diagnosis [reference category = no], participant recruiting setting [reference category = inpatient specialty care], interactions of QUADAS-2 signaling item responses [reference category = low risk] with logit (sensitivity) and logit (1—specificity) of the HADS-D (Wu, Levis, Sun, et al., 2021). We conducted these analyses separately by reference standards (semistructured interview, fully structured interview, MINI), since these types of interviews have been shown to identify different individuals (Wu, Levis, Sun, et al., 2021). In the second step, we added country/language variables to the model (Germany, Spain, Lithuania, Norway, Korea, Japan [reference category = English speaking countries]). These models were restricted to the subset of the studies from countries with more than 500 participants that had complete data for all relevant variables and used a semistructured interview or the MINI (there were not enough data for the studies that used a fully structured reference standard). Country HDI level was dropped from the model because all countries included in this analysis had very high HDI. For any variables that were found to be associated with the sensitivity or specificity across all cutoffs, we compared accuracy of HADS-D and HADS-T results stratified by subgroups based on these variables.

All analyses were run in R; R Version R 3.5.0 (R Core Team, 2020) and R Studio Version 1.1.423 (R Studio Team, 2020) using the lme4 package (Bates et al., 2015).

Registration and Protocol

The main HADS-D IPDMA was registered in the International Prospective Register of Systematic Reviews (CRD42015016761), and a protocol was published (Thombs et al., 2016). The present study was not included in the protocol for the main HADS-D IPDMA, but a separate protocol was developed and posted online prior to initiating the study (<https://osf.io/438ak/>).

Data Availability

Data contribution agreements with primary study authors do not include permission to make their data publicly available, although the data set used in this study will be archived through a McGill University repository (Borealis, <https://borealisdata.ca/dataverse/depressproject/>). The R codes used for the analysis will be made publicly available through the same repository. Requests to access the data set to verify study results but not for other purposes can be sent to the corresponding authors via the “Access Data Set” function on the repository website.

Results

Search Results and Inclusion of Primary Data

For the main HADS-D IPDMA, of 14,465 unique titles and abstracts identified from the database search, 13,895 were excluded after title and abstract review and 330 after full text (Supplemental Table A), leaving 240 eligible articles with data from 165 unique participant samples (Supplemental Figure A). Of the 165 unique samples, 93 (56%) contributed data (66% of eligible participants). In addition, authors of included studies contributed data from 10 studies that were unpublished or did not come up in the search, for a total of 103 HADS-D data sets contributed to our IPDMA. Five studies without HADS individual item scores or separate total scores for the HADS-D and HADS-T were excluded from the present study (see Supplemental Table B2). Thus, 20,700 participants (2,285 major depression cases) from 98 studies were analyzed (91% of 22,755 participants from the 103 HADS-D data sets). Included study characteristics are shown in Supplemental Table B1. Characteristics of eligible studies that did not provide data, including the five studies excluded because they only provided HADS-D or HADS-T total scores, are shown in Supplemental Table B2.

Of 98 included studies, 58 used semistructured interviews to assess major depression (10,311 participants), including 54 that used the SCID (9,676 participants); 31 used the MINI (7,445 participants); and 9 used other. Participant characteristics are shown in Table 1.

Supplemental Table C shows QUADAS-2 ratings for included studies. There were only 11 studies with “low” risk of bias rating across all QUADAS-2 domains.

Table 1
Participant Data by Subgroups

Participant subgroup	<i>N</i> studies ^a	<i>N</i> participants	<i>N</i> (%) major depression
All participants	98	20,700	2,285 (11)
Participants not currently diagnosed with a mental disorder or receiving treatment for a mental health problem	38	6,995	495 (7)
Age <60	92	11,795	1,452 (12)
Age ≥60	92	8,741	779 (9)
Women	96	11,111	1,342 (12)
Men	89	9,494	911 (10)
Very high country human development index	90	20,088	2,130 (11)
High country human development index	8	612	155 (25)
Participants diagnosed with cancer ^b	27	5,767	433 (8)
Inpatient specialty care	38	8,827	1,047 (12)
Outpatient specialty care	54	9,547	1,072 (11)
Nonmedical	7	1,908	116 (6)
Inpatient/outpatient mixed	3	418	50 (12)

^a Some variables were coded at the study level, while others were coded at the participant level. Thus, number of studies does not always add up to the total number. ^b The statistics here were from individual-level variable of cancer diagnosis, slight different from what we used in the subgroup analysis which based on the study-level care setting variable.

Comparison of Screening Accuracy Between the HADS-D and HADS-T

ROC plots comparing sensitivity and specificity estimates for all cutoffs between the HADS-D (0–21) and HADS-T (0–42) among all included studies are shown in Figure 1. A large part of the plots for the HADS-D and HADS-T were overlapping. The HADS-T performed better than HADS-D at some cutoffs, but this pattern was not consistent across cutoffs. The AUCs for the HADS-D and HADS-T were similar among all studies (0.853 vs. 0.872). We also compared the ROCs among studies that used a semistructured reference standard and found a similar pattern (Supplemental Figure B).

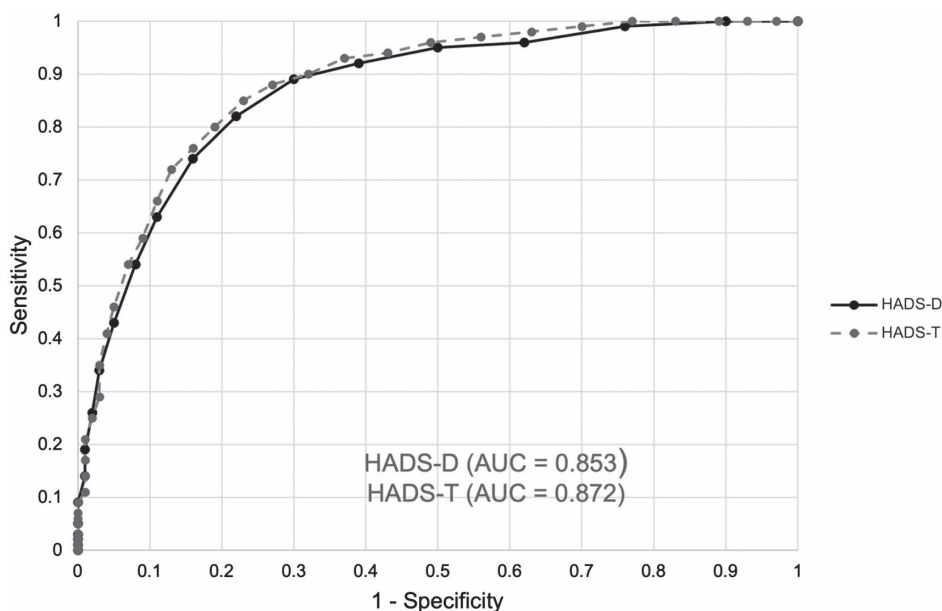
Based on the pooled sensitivity and specificity across all HADS-D and HADS-T cutoffs, among all studies, the cutoff that minimized the values of the distance to the top-left corner of the ROC curves was ≥7 for the HADS-D (sensitivity [95% CI] = 0.79 [0.75, 0.83], specificity [95% CI] = 0.78 [0.75, 0.80]) and ≥15 for the HADS-T (sensitivity [95% CI] = 0.79 [0.76, 0.82], specificity [95% CI] = 0.81 [0.78, 0.83]; Table 2).

The comparison of sensitivity and specificity between the HADS-D and HADS-T for the optimal cutoffs (HADS-D ≥7 vs. HADS-T ≥15) and other cutoffs close to the optimal cutoffs (≥5 vs. ≥11; ≥6 vs. ≥13; ≥8 vs. ≥17; ≥9 vs. ≥19; ≥10 vs. ≥21; and ≥11 vs. ≥23) are presented in Table 2. Overall, for the pairs of optimal cutoffs or other cutoffs close to the optimal, the differences in sensitivity and specificity between HADS-D and HADS-T using the bootstrapping approach across all 98 primary studies were small. Precision of estimates was high, and the width of 95% CIs ranged from 5% to 9% for sensitivity and 2% to 4% for specificity across all cutoffs examined. For sensitivity, the differences of HADS-T – HADS-D for all pairs of cutoffs were not statistically significant (the differences were between –0.05 and 0.01, CIs were within or overlapped with the range of –0.05 and 0.05). Therefore, at five pairs of optimal cutoffs or other cutoffs close to the optimal, the sensitivity of the HADS-T was equivalent to that of the HADS-D; the equivalency was indeterminate on the other two pairs, based on the prespecified equivalence margin of $\delta = 0.05$. For specificity, estimates of HADS-T were equivalent to HADS-D for all seven pairs of cutoffs (the differences of HADS-T – HADS-D were between 0.02 and 0.03; CIs were all within –0.05 and 0.05). Relevant results among studies that used a semistructured reference standard were consistent with overall estimates (Supplemental Table D1).

The comparison of results via individual-level analysis are presented in Table 3. For each pair of matched HADS-D and HADS-T cutoffs, the differences in sensitivity and specificity between the two tests were similar to those from the bivariate random effects models. This was also true among studies that used a semistructured reference standard (Supplemental Table D2).

Among participants in inpatient care settings (Table 4; 8,827 participants from 38 studies), the comparison results of HADS-T – HADS-D in sensitivity were similar to the overall estimates; the differences in specificity were slightly larger than overall estimates, however, the 95% CIs generally overlapped with –0.05 and 0.05 and were classified as indeterminate to equivalency, with one exception (HADS-D ≥6 vs. HADS-T ≥13) for which HADS-T specificity was greater than for the HADS-D. The comparison results among participants in outpatient care settings (Table 5; 9,547 participants from 54 studies) and participants from studies done in cancer care settings (Supplemental Table E; 5,608

Figure 1
ROC Curve for HADS-D and HADS-T Across All Studies



Note. ROC = receiver operating characteristic; HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; AUC = area under the curves.

participants from 23 studies) were similar to overall estimates. Within the semistructured reference standard category, similar patterns were found (Supplemental Tables D3 and D4).

The metaregression results indicated no significant differences in sensitivity and specificity were found for any individual participant characteristics or risk of bias ratings (Supplemental Tables F1–F3). After adding the country/language variables to the model, the sensitivity and specificity of HADS-D was invariant based on all variables across reference standards except that specificity estimates of the HADS-D were associated with Germany and Spain among studies that used a semistructured reference standard; specifically, the HADS-D had lower specificity among participants from Germany and Spain compared to studies done with participants from English speaking countries (Supplemental Tables G1 and G2).

Therefore, we conducted subgroup analysis of our comparisons of HADS-D and HADS-T accuracy for participants from Germany or Spain. For each pair of matched HADS-D and HADS-T cutoffs among participants from Germany (Supplemental Table H1), the comparison results of HADS-T – HADS-D in sensitivity and specificity were similar to the overall estimates; among participants from Spain (Supplemental Table H2), differences in specificity were slightly larger than overall estimates, however, the 95% CIs all overlapped with -0.05 and 0.05 and were classified as indeterminate to equivalent, and differences in sensitivity were similar to the overall estimates.

A forest plot of the differences of sensitivity and specificity estimates for HADS-D ≥ 7 versus HADS-T ≥ 15 across all studies is shown in Figure 2. At the optimal cutoffs, there was low heterogeneity in the differences between HADS-D and HADS-T across the 98 studies with estimated interstudy heterogeneity

(τ^2) < 0.01 for sensitivity and < 0.01 for specificity. The forest plot of the differences of sensitivity and specificity estimates at optimal cutoffs for the HADS-D and HADS-T among studies that used a semistructured reference standard is shown in Supplemental Figure C.

Discussion

We assessed the equivalency of screening accuracy of the HADS-D and HADS-T across all cutoffs to detect major depression and compared accuracy across paired optimal cutoffs and other cutoffs close to the optimal cutoffs to test whether the HADS-T is superior to HADS-D for major depression detection. There were two main findings. First, among all 98 included studies the values of the distance to the top-left corner of the ROC curves (Riley et al., 2008) were minimized at a HADS-D cutoff ≥ 7 (sensitivity = 0.79, specificity = 0.78) and at a HADS-T cutoff ≥ 15 (sensitivity = 0.79, specificity = 0.81). Second, at paired optimal cutoffs and six other cutoffs close to the optimal cutoffs, the HADS-D was similarly accurate compared to the HADS-T overall and among studies that used a semistructured reference standard.

Overall, for all 98 primary studies, across all sets of paired cutoffs, the sensitivity and specificity of the HADS-T were classified as equivalent to that of the HADS-D based on the prespecified equivalency margin. Although the HADS-T was slightly more specific (range 0.02–0.03), all the 95% CIs for differences in sensitivity and specificity of HADS-T – HADS-D were within or overlapped with the range of -0.05 and 0.05 . When we analyzed data separately among studies that used a semistructured reference standard, differences in sensitivity and specificity between the HADS-D and HADS-T were similar to the overall estimates.

Table 2
Comparison of Sensitivity and Specificity Estimates Between HADS-D and HADS-T for Pairs of Optimal Cutoffs and Cutoffs Close to the Optimal Cutoffs Across All Studies

Cutoff	HADS-D ^a			HADS-T			HADS-T—HADS-D			
	Sensitivity	95% CI	Specificity	Cutoff	Sensitivity	95% CI	Specificity	95% CI	Specificity	95% CI
5	0.90	[0.87, 0.92]	0.61	11	0.91	[0.89, 0.93]	0.63	[0.60, 0.66]	0.01	[-0.01, 0.04]
6	0.86	[0.82, 0.88]	0.70	13	0.86	[0.83, 0.88]	0.73	[0.70, 0.75]	0.00	[-0.03, 0.03]
7 ^b	0.79	[0.75, 0.83]	0.78	15 ^c	0.79	[0.76, 0.82]	0.81	[0.78, 0.83]	0.00	[-0.05, 0.02]
8	0.70	[0.66, 0.74]	0.84	17	0.70	[0.66, 0.74]	0.87	[0.85, 0.89]	0.00	[-0.05, 0.04]
9	0.60	[0.55, 0.64]	0.89	19	0.58	[0.54, 0.61]	0.91	[0.90, 0.93]	-0.02	[-0.07, 0.02]
10	0.50	[0.45, 0.54]	0.92	21	0.45	[0.41, 0.49]	0.95	[0.94, 0.95]	-0.05	[-0.10, -0.01]
11	0.39	[0.35, 0.43]	0.95	23	0.34	[0.31, 0.37]	0.97	[0.96, 0.97]	-0.05	[-0.10, -0.01]

Note. HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; CI = confidence interval; ROC = receiver operating characteristic.
^a N studies = 98; N participants = 20,700; N major depression = 2,285. ^b The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-D. ^c The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-T.

Table 3

Comparison of Sensitivity and Specificity Estimates Between HADS-D and HADS-T for Pairs of Optimal Cutoffs and Cutoffs Close to the Optimal Cutoffs Across All Studies via Individual-Level Model

HADS-D ^a Cutoff	HADS-T Cutoff	HADS-T—HADS-D	
		Sensitivity	Specificity
5	11	0.02 (-0.00, 0.03)	0.01 (-0.00, 0.03)
6	13	0.01 (-0.01, 0.03)	0.03 (0.01, 0.04)
7 ^b	15 ^c	0.00 (-0.02, 0.03)	0.02 (0.01, 0.04)
8	17	0.00 (-0.03, 0.03)	0.03 (0.02, 0.04)
9	19	-0.02 (-0.05, 0.01)	0.03 (0.02, 0.04)
10	21	-0.05 (-0.08, -0.02)	0.03 (0.02, 0.03)
11	23	-0.05 (-0.09, -0.02)	0.02 (0.02, 0.03)

Note. HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; ROC = receiver operating characteristic.
^a N participants = 20,700; N major depression = 2,285. ^b The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-D. ^c The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-T.

Furthermore, similar to overall estimates, there were no substantive differences in performance between the HADS-D and HADS-T in detecting major depression among medical outpatients. Among inpatients, the HADS-T and HADS-D were also equivalent in sensitivity. The HADS-T performed slightly better than HADS-D in terms of specificity, and equivalency was indeterminate based on the prespecified equivalence margin, except for one pair of cutoffs. This finding is possibly related to the greater presence of anxiety symptoms in inpatients versus outpatients and its relationship to depression (Schatzberg, 2019).

Previous conventional meta-analyses of results from cancer patients (Mitchell et al., 2010; Vodermaier & Millman, 2011) suggested that the HADS-T may perform better than the HADS-D, but that conclusion was highly uncertain given the limitations of the samples and methods. Through our IPDMA, with its large data set and more rigorous comparison methods including both bivariate random effects models and individual-level models, a two-level bootstrap approach (Fagerland et al., 2014; Higgins & Thompson, 2002), and subgroup analysis, we found there was no consistent evidence that the HADS-T is superior to HADS-D for major depression detection, including in cancer care settings. In addition, we did not identify any differences between HADS-D and HADS-T accuracy that were associated with individual participant characteristics or countries. Therefore, in research and clinical general practice, using the full 14-item HADS-T for depression screening would likely result in no to minimal gain in screening accuracy but would add unnecessary burden to patients compared to the seven-item HADS-D.

To our knowledge, this is the first meta-analysis that directly compared the HADS-D and HADS-T for screening for depression using the same large individual participant data set for both screening tools. Strengths of this study included the large overall sample size and high precision of estimates of differences, the ability to compare results for HADS-D and HADS-T across all cutoffs from all studies, and the ability to assess screening accuracy overall and by inpatient and outpatient subgroups. There are also limitations to

Table 4
Comparison of Sensitivity and Specificity Estimates Between HADS-D and HADS-T for Pairs of Optimal Cutoffs and Cutoffs Close to the Optimal Cutoffs Among Participants Recruited From Inpatient Care Settings

Cutoff	HADS-D ^a			HADS-T			HADS-T—HADS-D						
	Sensitivity	95% CI	Specificity	95% CI	Cutoff	Sensitivity	95% CI	Specificity	95% CI	Sensitivity	95% CI	Specificity	95% CI
5	0.90	[0.87, 0.93]	0.55	[0.49, 0.60]	11	0.90	[0.87, 0.92]	0.62	[0.56, 0.68]	0.00	[-0.03, 0.03]	0.07	[0.04, 0.11]
6	0.86	[0.83, 0.89]	0.64	[0.58, 0.69]	13	0.85	[0.81, 0.88]	0.72	[0.67, 0.77]	-0.01	[-0.07, 0.02]	0.08	[0.06, 0.12]
7 ^b	0.80	[0.75, 0.83]	0.73	[0.68, 0.78]	15 ^{c,d}	0.79	[0.74, 0.82]	0.81	[0.76, 0.85]	-0.01	[-0.08, 0.02]	0.08	[0.05, 0.11]
8	0.73	[0.68, 0.78]	0.80	[0.76, 0.84]	17	0.69	[0.64, 0.74]	0.87	[0.83, 0.90]	-0.04	[-0.11, 0.03]	0.07	[0.04, 0.09]
9	0.63	[0.58, 0.69]	0.86	[0.82, 0.89]	19	0.59	[0.54, 0.64]	0.91	[0.88, 0.93]	-0.04	[-0.14, 0.01]	0.05	[0.03, 0.07]
10	0.55	[0.49, 0.61]	0.90	[0.87, 0.93]	21	0.46	[0.41, 0.51]	0.95	[0.92, 0.96]	-0.09	[-0.19, -0.03]	0.05	[0.03, 0.06]
11	0.45	[0.39, 0.51]	0.93	[0.91, 0.95]	23	0.36	[0.32, 0.41]	0.97	[0.95, 0.98]	-0.09	[-0.18, -0.02]	0.04	[0.02, 0.05]

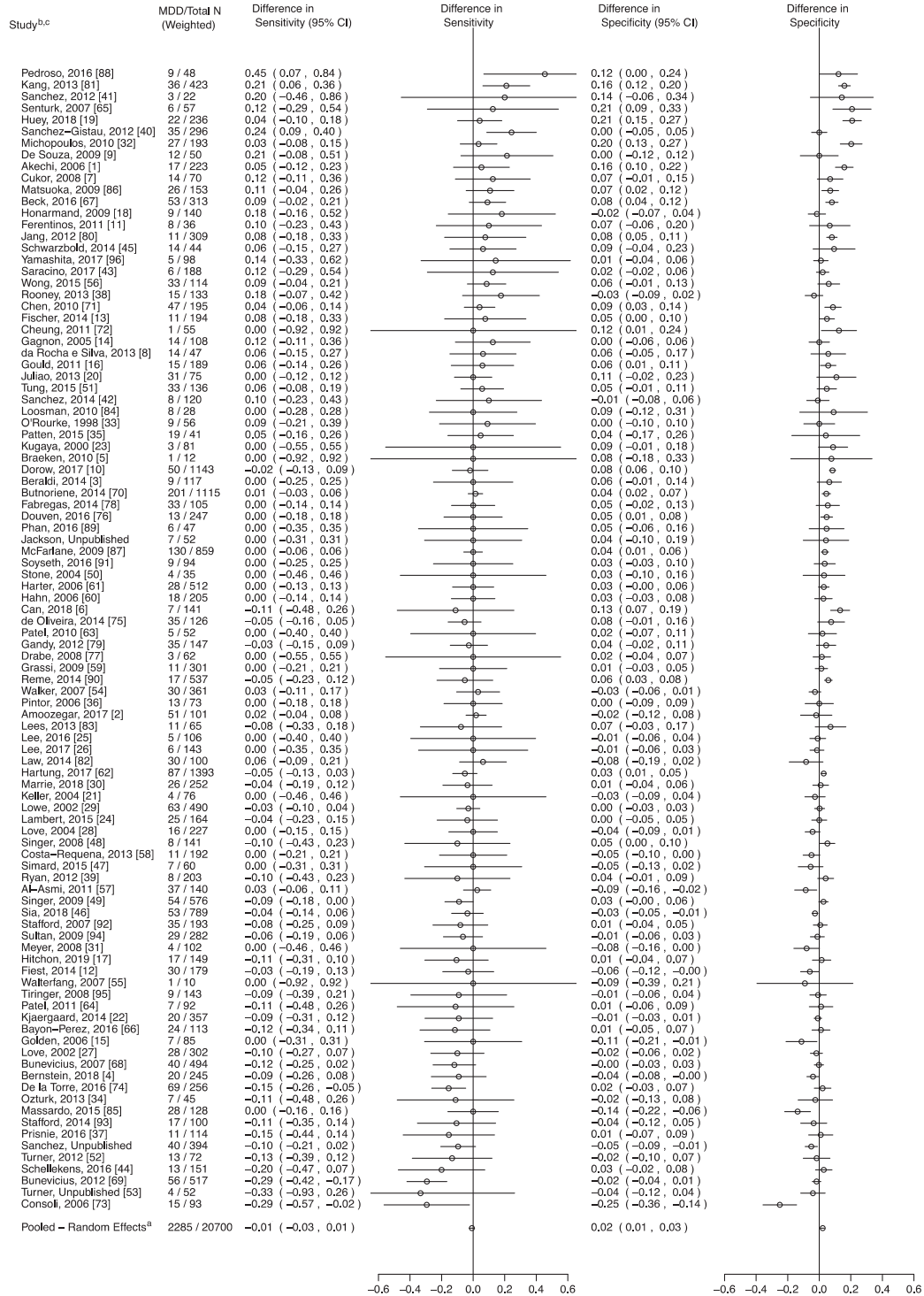
Note. HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; CI = confidence interval; ROC = receiver operating characteristic.
^a *N* studies = 38; *N* participants = 8,827; *N* major depression = 1,047. ^b The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-D. ^c The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-T. ^d On this cutoff of HADS-T, the model convergence code was 0 when using the default optimizer in glmer, but there were meaningful CIs.

Table 5
Comparison of Sensitivity and Specificity Estimates Between HADS-D and HADS-T for Pairs of Optimal Cutoffs and Cutoffs Close to the Optimal Cutoffs Among Participants Recruited From Outpatient Care Settings

Cutoff	HADS-D ^a			HADS-T			HADS-T—HADS-D						
	Sensitivity	95% CI	Specificity	95% CI	Cutoff	Sensitivity	95% CI	Specificity	95% CI	Sensitivity	95% CI	Specificity	95% CI
5	0.91	[0.87, 0.94]	0.63	[0.60, 0.67]	11	0.92	[0.89, 0.95]	0.62	[0.59, 0.66]	0.01	[-0.02, 0.04]	-0.01	[-0.03, 0.01]
6	0.87	[0.82, 0.91]	0.72	[0.69, 0.75]	13	0.88	[0.84, 0.91]	0.72	[0.69, 0.75]	0.01	[-0.02, 0.05]	0.00	[-0.01, 0.02]
7 ^b	0.82	[0.75, 0.86]	0.79	[0.76, 0.81]	15 ^c	0.81	[0.76, 0.84]	0.80	[0.77, 0.82]	-0.01	[-0.07, 0.04]	0.01	[-0.01, 0.03]
8	0.71	[0.65, 0.77]	0.85	[0.83, 0.87]	17	0.73	[0.67, 0.78]	0.86	[0.84, 0.88]	0.02	[-0.04, 0.07]	0.01	[-0.00, 0.03]
9	0.60	[0.54, 0.66]	0.90	[0.88, 0.91]	19	0.59	[0.53, 0.65]	0.91	[0.90, 0.92]	-0.01	[-0.08, 0.04]	0.01	[0.00, 0.03]
10	0.49	[0.43, 0.55]	0.93	[0.91, 0.94]	21	0.45	[0.39, 0.52]	0.94	[0.93, 0.95]	-0.04	[-0.11, 0.02]	0.01	[0.00, 0.03]
11	0.38	[0.32, 0.44]	0.95	[0.94, 0.96]	23	0.34	[0.29, 0.39]	0.96	[0.95, 0.97]	-0.04	[-0.10, 0.01]	0.01	[0.00, 0.02]

Note. HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; CI = confidence interval; ROC = receiver operating characteristic.
^a *N* studies = 54; *N* participants = 9,547; *N* major depression = 1,072. ^b The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-D. ^c The cutoff minimizes the values of the distance to the top-left corner of the ROC curves for HADS-T.

Figure 2
Forest Plots of the Difference in Sensitivity and Specificity Estimates at the Optimal Cutoff (HADS-D: ≥7; HADS-T: ≥15) Between HADS-D and HADS-T Across All Studies



Note. N Studies = 98; N Participants = 20,700; N major depression = 2,285. HADS-D = seven-item Hospital Anxiety and Depression Scale Depression subscale; HADS-T = 14-item Hospital Anxiety and Depression Scale Depression subscale; MDD = Major Depressive Disorder; CI = confidence interval.

a tau^2 for the difference of sensitivity and specificity were both <0.001. b References for all included studies are marked with an asterisk in the reference list. The reference numbers refer to Supplemental Material References. c The studies were sorted by the sum of difference in sensitivity and difference in specificity in descending order.

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consider. First, for the full IPDMA data, primary data from 72 of 165 published eligible data sets (44% of data sets, 34% of participants) were not included, and only those data sets with complete data for all individual HADS item scores (91% of available data) were included in this study. Nonetheless, this sample was much larger than the few primary studies that have previously compared the HADS-D and HADS-T. Second, we did not conduct analyses restricted to studies with “low” risk of bias ratings across QUADAS-2 domains. However, in sensitivity analysis in this study and in our main IPDMA on the HADS-D (Wu, Levis, Sun, et al., 2021), risk of bias ratings were not associated with screening accuracy. Third, the present study used a subset of studies and participants from our previously conducted HADS-D IPDMA (Wu, Levis, Sun, et al., 2021). This IPDMA project was designed to assess the accuracy of the HADS-D for detecting major depression. Diagnoses of other mental disorders, including, anxiety disorders, were not collected in most of the included primary studies. Thus, we were not able to evaluate the sensitivity and specificity of the HADS-D, HADS-Anxiety, or HADS-T for detecting mental disorders generally. Forth, we did not record interrater reliability for risk of bias ratings; however, all ratings were done by trained reviewers and any disagreements were addressed by consensus, including a third investigator as necessary.

Conclusions

In summary, this study found that sensitivity and specificity of the HADS-T were not superior to the HADS-D for detecting major depression in a large individual participant data set. Using the seven-item HADS-D for depression screening instead of the full 14-item HADS-T has minimal influence on performance of the measure but would reduce patient and participant burden in clinical and research settings. Both HADS-D and HADS-T have only modest screening ability and discussion of their exact indications for use and related caveats are beyond the scope of this article. However, there were no substantive differences in performance between the HADS-D and HADS-T in detecting major depression among medical outpatients, although there was a slight advantage in specificity of indeterminate equivalency for the HADS-T among medical inpatients, for whom adding the anxiety items of HADS-A may improve accuracy.

Ethical Approval

As this study involved secondary analysis of anonymized previously collected data, the Research Ethics Committee of the Jewish General Hospital declared that this project did not require research ethics approval. However, for each included data set, we confirmed that the original study received ethics approval and that all patients provided informed consent.

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