



Universiteit
Leiden
The Netherlands

Common measures or common metrics? A plea to harmonize measurement results

Beurs, E. de; Boehnke, J.R.; Fried, E.I.

Citation

Beurs, E. de, Boehnke, J. R., & Fried, E. I. (2022). Common measures or common metrics?: A plea to harmonize measurement results. *Clinical Psychology And Psychotherapy*, 29(5), 1755-1767. doi:10.1002/cpp.2742

Version: Publisher's Version

License: [Creative Commons CC BY-NC-ND 4.0 license](https://creativecommons.org/licenses/by-nc-nd/4.0/)

Downloaded from: <https://hdl.handle.net/1887/3515169>

Note: To cite this publication please use the final published version (if applicable).

RESEARCH ARTICLE

WILEY

Common measures or common metrics? A plea to harmonize measurement results

Edwin de Beurs¹  | Jan R. Boehnke²  | Eiko I. Fried³ 

¹Department of Clinical Psychology, Leiden University & Arkin GGZ, Amsterdam, The Netherlands

²School of Health Sciences, University of Dundee, Dundee, UK

³Department of Clinical Psychology, Leiden University, Leiden, Zuid-Holland, The Netherlands

Correspondence

Edwin de Beurs, PhD, Department of Clinical Psychology, Leiden University & Arkin GGZ, Amsterdam, The Netherlands.

Email: edwin.de.beurs@arkin.nl

Abstract

Objective: There is a great variety of measurement instruments to assess similar constructs in clinical research and practice. This complicates the interpretation of test results and hampers the implementation of measurement-based care.

Method: For reporting and discussing test results with patients, we suggest converting test results into universally applicable common metrics. Two well-established metrics are reviewed: *T* scores and percentile ranks. Their calculation is explained, their merits and drawbacks are discussed, and recommendations for the most convenient reference group are provided.

Results: We propose to express test results as *T* scores with the general population as reference group. To elucidate test results to patients, *T* scores may be supplemented with percentile ranks, based on data from a clinical sample. The practical benefits are demonstrated using the published data of four frequently used instruments for measuring depression: the CES-D, PHQ-9, BDI-II and the PROMIS depression measure.

Discussion: Recent initiatives have proposed to mandate a limited set of outcome measures to harmonize clinical measurement. However, the selected instruments are not without flaws and, potentially, this directive may hamper future instrument development. We recommend using common metrics as an alternative approach to harmonize test results in clinical practice, as this will facilitate the integration of measures in day-to-day practice.

KEYWORDS

common metrics, depression, percentile rank, self-report measures, *T* score, test result

1 | INTRODUCTION OF THE PROBLEM

Measurement is the most basic building block of scientific research. Since the days of Wundt and Thurstone, measurement has played a prominent role in psychology and a separate branch is dedicated to its research: psychometrics. The past decades have seen a proliferation

of measurement instruments for a broad range of psychological constructs. On the one hand, the vast number of available measurement instruments for psychological concepts reflect the importance of assessment and psychometrics in clinical psychology. Novel measures often incorporate new theoretical perspectives on the phenomenology or treatment of psychopathology (e.g., by stressing cognitive

This is an open access article under the terms of the [Creative Commons Attribution-NonCommercial-NoDerivs](https://creativecommons.org/licenses/by-nc-nd/4.0/) License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

© 2022 The Authors. *Clinical Psychology & Psychotherapy* published by John Wiley & Sons Ltd.

aspects), or signify a change in key features of a disorder (Böhnke & Croudace, 2015). Measures are often validated for different populations (general population vs. clinical sample) or different purposes (diagnosis vs. tracking treatment progress) (McPherson & Armstrong, 2021; Patalay & Fried, 2020), or simply reflect innovations in measurement technology itself. On the other hand, the number of available measurement instruments also poses a challenge to psychological research. There are at least 19 instruments to measure anger (Weidman et al., 2017), and over 280 instruments have been developed to measure depression, many of which are still in use (Fried, 2017; Santor et al., 2006). This is also reflected in little consistency in use of measures in therapy outcome research as noted by Ogles (2013): across 163 studies, authors used 435 unique outcome measures, of which 371 were used only once in a study. This plethora of instruments, in combination with questionable measurement practices, such as outcome switching (Weston et al., 2016) and lack of transparency on how scores are derived from these measures (Weidman et al., 2017), hampers the establishment of a consistent body of knowledge. It complicates the comparison of results from various studies, and unnecessarily slows down scientific progress (Flake & Fried, 2020).

In clinical practice, measurement instruments are increasingly used to aid the initial evaluation of patients and to assess outcomes. Measurement-based care has been called for repeatedly in numerous editorials and reviews (Boehnke & Rutherford, 2021; Fortney et al., 2017; Harding et al., 2011; Lambert, 2007; Lewis et al., 2019; Snyder et al., 2012). Paradoxically, the abundance of measurement instruments poses barriers to their use in clinical practice (Fried, 2017; Santor et al., 2006). Firstly, the use of a wide variety of instruments, each having its own metric, complicates communication among professionals. For instance, severity of depression symptomatology expressed in a score on the Beck Depression Inventory (BDI-II; Beck et al., 1996), with scores ranging from 0 to 63, is not compatible with severity expressed in a score on the Patient Health Questionnaire (PHQ-9; Kroenke & Spitzer, 2002), with scores ranging from 0 to 27. This complicates matters when a clinician using the BDI-II refers a patient to a colleague more familiar with the PHQ-9. Secondly, communication about test results between therapists and patients is complicated by the use of various metrics (Snyder et al., 2019). Moreover, the knowledge gap between clinicians and patients widens, when only clinicians are able to properly interpret test results. More patient engagement, involvement, and shared decision making about the course of treatment is called for (Patel et al., 2008), in part because this leads to better outcomes (Lambert & Harmon, 2018). The knowledge gap may hinder patients in their attempts to obtain a more equal role in the therapeutic process on their journey to recovery. Clear information on the meaning of a test result and how much progress is made towards treatment goals, will help to better involve patients, will grant them a more active role, and will strengthen engagement with the information provided (Goetz, 2010). Finally, the knowledge of measurement instruments and the specific psychometric properties required for the interpretation of test scores, may hinder implementation in daily clinical practice, and allow

Key Practitioner Message

- Converting raw scores from neuro-psychological assessments to common metrics will ease the interpretation of test results among professionals and will facilitate communication with patients receiving mental healthcare.
- Common metrics will clarify the severity of patients' conditions and increase their engagement with outcome data.
- Treatment progress (or lack thereof) can be more easily monitored, understood and communicated to by using common metrics.

practitioners to maintain beliefs that standardized measures are not as accurate as their clinical judgement (de Beurs et al., 2011).

A strategy to facilitate the interpretation of test results in clinical practice has been around for many decades and is well-known: *use of common metrics*. With a metric we mean a system or standard of measurement. Metrics describe the units in which measurements are provided and interpreted. Examples would be the imperial and the metric system which are used across a range of measurement domains as diverse as distance, volume, and weight to describe amounts. In the medical field, common metrics are usually based on physical entities, such as the metre, kilogramme, mol and Kelvin. In psychology we mostly measure non-physical entities, abstract constructs or latent traits, such as depression, which do not permit direct observation (Flake & Fried, 2020; Kellen et al., 2021). Consequently, we cannot fall back on these physical entities and have to express scores on metrics that have no physical counterpart, for example, when expressing severity as the distance from the mean score of a reference group. In the case of psychological assessments, use of a common metric would involve the conversion of raw test scores into universally applicable common metrics, such as standardized scores and percentile ranks, based on normative samples (Kendall et al., 1999). Converting raw scores to common metrics is discussed in many textbooks on psychological testing (Anastasi, 1968; Cronbach, 1984), but it is underused in clinical research and practice. An example of how use of a common metric has worked out well is available in another area of psychological assessment: the measurement of intellectual capacity. The IQ metric was developed more than a century ago by Binet and Simon (1907) and has found universal application; professional psychologists as well as the lay public understand the meaning of IQ-scores, no matter the specific measurement instrument that was used. Interpretation of test results in clinical practice would be much easier if their meaning was as self-evident as IQ test scores.

The remainder of the paper is structured as follows. First, we describe two candidate metrics for universal application: *T* scores and percentile ranks. Both are commonly used in neuropsychological and educational research and are gaining popularity in clinical assessment. Second, we describe the *T* score metric and percentile ranks metric in more detail, and present their main merits and drawbacks. We also

describe how they can be derived by various approaches, illustrate their use with measures for depression, and discuss the choice of a proper reference group for these metrics. Finally, we propose further steps required for widespread implementation and suggest various subjects for future research.

2 | CANDIDATE METRICS

Two options have been proposed: (1) Standard scores (z scores and their more convenient alternatives, such as *T* scores, stanines and stens) and (2) percentile rank scores (Crawford & Garthwaite, 2009; Ley, 1972).

Table 1 presents an overview of these metrics, their calculation, and some of their properties. Figure 1, adapted from Seashore (1955), shows the various metrics under the normal distribution and provides labels for the interpretation of scores when used as a severity indicator or screener. For example, in order to label levels of severity for depression, the PROMIS group has followed the convention of dividing the general population up in four segments (from bottom to

top: 69.1%, 15.0%, 13.6% and 2.3%, respectively). Labels and corresponding *T* scores are: 'within normal limits' ($T < 55.0$), 'mild' ($T = 55.0-59.9$), 'moderate' ($T = 60.0-69.9$) and 'severe' ($T \geq 70.0$) (healthmeasures, n.d.) (see <https://www.healthmeasures.net/score-and-interpret/interpret-scores/promis/promis-score-cut-points>). As an illustration, Figure 1 also includes raw scores on the PHQ-9 (Kroenke & Spitzer, 2002), CES-D (Radloff, 1977) and BDI-II from the US population (Choi et al., 2014) to demonstrate how they would translate into these metrics. For all three instruments, the raw score units are stretched at the lower end, illustrating the negative skewness of the frequency distribution of raw scores in the general population: Low scores are much more common than high scores.

3 | T SCORES

T scores, so named by McCall (1922) to honour the psychometric pioneers Thorndike, Terman and Thurstone, are based on standardized or *Z* scores. *Z* scores are raw scores converted to a standard scale with $M = 0$ and $SD = 1$ and are calculated based on the mean and standard deviation of a reference group. Standardization to *Z* scores yields inconvenient scores (with a range of -3 to 3 implying negative scores with several decimals to denote sufficient precision) and alternatives have been put forth with a more convenient format, such as stans, stanines and *T* scores. Stans and stanines yield a rather crude categorization of score levels and we left them out from further consideration. *T* scores are *Z* scores multiplied by 10 with 50 points added. They have a mean of 50, a standard deviation of $SD = 10$, and range, in practice, from 20 to 80. Figure 1 shows that a *T* score of 80 is three standard deviations above the mean, a score obtained by only 0.13% of the population, according to the cumulative normal distribution. Thus, 99.7% of the population will score in the 20–80 range. *T* scores have become the metric of choice for commonly used measures in clinical assessment. To cite Cronbach (1984): 'Confusion

TABLE 1 Metrics that can be used for test score standardization and harmonization

	Calculation:	<i>M</i>	<i>SD</i>	Usual range
<i>Z</i> score	$Z = (x - m)/s$	0	1	−3 to 3
Stanine	$S = Z * 2 + 5$	5	2	1 to 9
Sten	$S = Z * 2 + 5$	5.5	2	1 to 10
<i>T</i> score	$T = Z * 10 + 50$	50	10	20 to 80
Percentile rank	$pR = \frac{\text{CumF} - (.5 * F)}{N}$	50	NA	0 to 100

Note: N.B.: x = test raw score; m = average of the test raw scores; s = standard deviation of the raw scores; CumF = Cumulative frequency; F = Frequency; N = total sample; NA = not applicable.

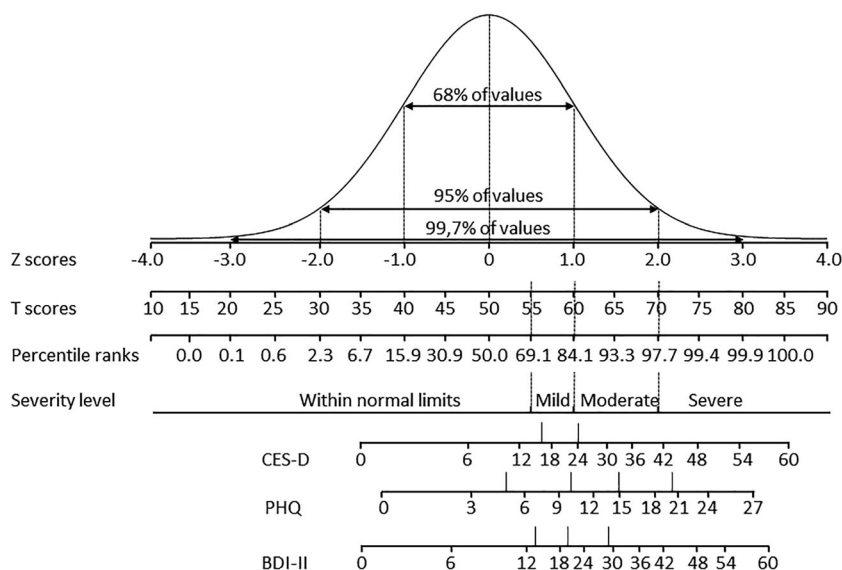


FIGURE 1 The normal distribution with standard scores, *T* scores, percentile rank scores, labels (Ba = below average, M = average, aa = above average) and scores in the general US population on the CES-D, PHQ and BDI-II

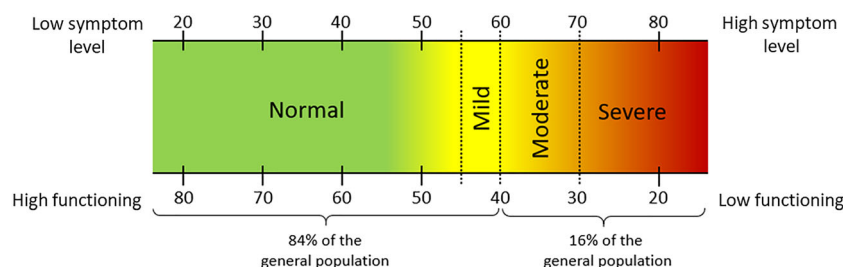


FIGURE 2 A heat map representation of the meaning of *T* scores for measures of symptom severity level and measures of functioning (adapted from www.scientia.global/wp-content/uploads/2017/07/HealthMeasures.pdf) [Colour figure can be viewed at wileyonlinelibrary.com]

results from the plethora of scales. In my opinion, test developers should use the system with mean 50 and s.d. 10 unless there are strong reasons for adopting a less familiar scale. (p. 100)'. Practical guidelines regarding the interpretation of *T* score levels have been established. At the onset of treatment, most patients will have a *T* score in the 65–75 range, and with treatment one may aim for a score below 55, a reasonable cut-off on the *T* score metric for recovery on many instruments that measure the severity of psychopathology (Aschenbrand et al., 2005; Cella et al., 2014; Recklitis & Rodriguez, 2007) and research suggests that many patients prefer colour coding of score levels according to a heat map of normed scores (Brundage et al., 2015). Figure 2 illustrates how the meaning of *T* scores can be conveyed to patients or colleagues.

To obtain *T* scores, various methods can be used. The most straightforward approach is a conversion of raw scores to *T* scores with a simple linear formula [$T = 10 * (x - m/sd) + 50$, where *x* is the raw score and *m* and *sd* the mean and standard deviation of the reference population]. However, this is only feasible when the raw scores have a normal distribution. If this condition is not met—which is quite common when clinical measures are administered to the general population, yielding skewed and leptokurtic distributions with an overrepresentation of low scores—the arithmetic mean and standard deviation are inappropriate descriptors of the frequency distribution of scores and the interpretation of the test result in standard scores is no longer straightforward. However, deviations from normality in raw scores can be fixed in most cases by transformations (Box & Cox, 1964; Liu et al., 2009). Another, more thorough approach is to first establish normalized standard scores with regression-based norming (Zachary & Gorsuch, 1985), and to convert these to *T* scores (Lenhard & Lenhard, 2021). Statistical software is available in R (GAMLSS; Stasinopoulos et al., 2018) and Timmerman et al. (2020) offers a detailed tutorial.

Recently, Item Response Theory (IRT)-based approaches to convert item responses to legacy instruments into *T* scores have gained popularity, using EAP-factor scores or alternatives (Fischer & Rose, 2019). The result of the IRT-based approach applied by Choi et al. (2014) to convert scores to the PROMIS metric (a *T* score with the US population as reference group) is illustrated for the CES-D, PHQ-9 and BDI-II in Figure 1. These approaches again require the assumption that the trait under study is normally distributed. However, the distributions of observed test and latent trait scores do not necessarily coincide (Kellen et al., 2021; Liddell & Kruschke, 2018). Their correspondence depends on the assumptions of the underlying measurement model and therefore impacts all test score interpretations

equally, for example, approaches described as ‘classical test theory’ deal only with test scores as an observed property (DeMars, 2018) and latent variable models separate observed test and latent trait scores (Molenaar & Dolan, 2018). For a non-normal distribution of raw scores, commonly found when clinical traits such as depression or anxiety are assessed in general population samples, alternatives have been developed (Reise et al., 2018). As IRT factor scores are often scaled as a standard score ($M = 0.00$, $SD = 1.00$), they can be easily converted into *T* scores with the linear formula in Table 1.

T scores are well established in clinical psychometrics. They were chosen as a metric by the PROMIS initiative, aimed at developing a new set of measures in health research (Cella et al., 2010). Considerable work has been done in this area, for instance by the Prosetta Stone initiative (Choi et al., 2014; Schalet et al., 2015). Here, raw scores have been converted to the PROMIS *T* score metric for many legacy measures, assessing a wide variety of constructs, including depression (Choi et al., 2014; Wahl et al., 2014), anxiety (Schalet et al., 2014) and pain (Cook et al., 2015). Also other researchers have chosen *T* scores as a common metric, for example, for depression (Wahl et al., 2014), anxiety (Rose & Devine, 2014), physical functioning (Oude Voshaar et al., 2019), fatigue (Friedrich et al., 2019) and personality psychopathology (Zimmermann et al., 2020).

When a unidimensional latent variable model holds and distributions are appropriately smoothed (Lord & Wingersky, 1984), conversion based on percentile ranks (equipercetile linking) and IRT based approaches tend to yield very similar results, as demonstrated by Choi et al. (2014), and more recently by Schalet et al. (2021). The *T* scores metric has been linked to expert judgement regarding the severity of pain interference, fatigue, anxiety and depression for oncology patients (Cella et al., 2014). Severity levels matched almost perfectly with the *T* score metric: 60 differentiated mildly from moderately symptomatic and 70 differentiated moderately from severely symptomatic. Furthermore, *T* scores on the Brief Symptom Inventory (Derogatis, 1975) have been compared with the Clinical Global Impression (CGI; Guy, 1976), and an improvement of 5 *T* score points corresponded well with improvement according to the CGI improvement rating (de Beurs et al., 2019).

To help clinicians who want to express a test score of a single patient as a common metric, crosswalk tables have been published for many measures to convert raw scores into *T* scores (e.g., Batterham et al., 2018; Choi et al., 2014; Zimmermann et al., 2020). Several user manuals for measures provide crosswalk tables, including the Minnesota Multiphasic Personality Inventory (MMPI-2; Butcher et al., 1989), the Brief Symptom Inventory (BSI; Derogatis, 1975), and

the Child Behavior Checklist (CBCL; Achenbach, 1991). The Prosetta Stone Initiative (<https://www.prosetta.org>) provides crosswalk tables for many measures to the PROMIS *T* score, which is based on the general US population. *T* scores can also be derived using a webapp (Fischer & Rose, 2016). Further, a crosswalk formula allows for the arithmetic conversion of raw scores to *T* scores or percentile ranks. These formulas can be established from cross-walk tables with statistical software, such as curve fitting in the regression module of SPSS or the *nls* and *nlstools* packages for Non-linear Least Squares modelling in R (Baty et al., 2015). An example of conversion by formula is provided by Roelofs et al. (2013) and de Beurs et al. (Submitted).

4 | PERCENTILE RANK SCORES

Contrasting standard scores, the percentile rank of a score ranges from 0 to 100, indicating the percentage of cases that are less than that score. In other words, percentile ranks denote with a score of 0 to 100 the relative position of the tested person among their peers from a reference group in 101 intervals (Kurtz & Mayo, 1979). In clinical contexts, this may translate into a message to the patient, such as, 'At least 75% of our patients have a lower score at the onset of treatment'. As such, percentile ranks are an easily understood representation of a test result, which helps to convey its meaning to colleagues and patients. This may explain why percentile ranks are widely used in educational assessment. Percentile ranks are also depicted in Figure 1.

Percentile ranks are calculated with the formula in the last column of Table 1. When normative data are available, the frequency of each raw score can be established with $PR = (\text{CumF} - [0.5 * F]) / N$ (Crocker & Algina, 1986). The cumulative frequency (CumF) is the count of all scores less than or equal to the score of interest, *F* is the frequency of the score of interest and *N* is the number of observations. The metric is calculated as the percentage of scores that fall below the score of interest plus half of those obtaining exactly that score. Under the assumption of a normal distribution, percentile ranks can be derived from *z* scores (and *T* scores) according to the formula for the cumulative normal probability, available as a statistical function in many software packages (R, SPSS, STATA and MS Excel). The formula can also be approximated by the logistic function, which describes a sigmoid curve: $PR = 100 / (1 + e^{(-1.75 * Z)})$, where *Z* is the standard score or $PR = 100 / (1 + e^{(-0.175 * T + 8.75)})$, and where *T* is the *T* score.

The literature on using percentile ranks in clinical practice is limited. Crawford and Garthwaite (2009) have propagated their use for clinical (neuro)psychological application. Crawford and colleagues have published percentile rank scores and confidence intervals for several depression and anxiety measures based on Australian samples (Crawford et al., 2011) and UK samples (Crawford et al., 2009) and made a computer programme available to calculate these scores. They also published crosswalk tables to convert raw scores into percentile ranks (Crawford et al., 2011; Crawford & Henry, 2003). Recently, raw scores and percentile ranks were published for the Inventory of Depression and Anxiety Symptoms (IDAS-II; Nelson et al., 2018).

5 | T SCORES AND PERCENTILE RANKS COMPARED

A minimum requirement for a universally applicable metric is that it should be easy to interpret and express in a straightforward manner how common or exceptional a test result is (Snyder et al., 2012). The COSMIN checklist defines interpretability as 'the degree to which one can assign a qualitative meaning' to a score (Prinsen et al., 2018). This would solve many of the issues of idiosyncratic scoring methods mentioned in the introduction. The *T* score denotes the commonness of a test result through its distance from the mean of a reference group in standard units, where 50 represent the mean and 10 points represent a standard deviation. The interpretation of the *T* score requires some knowledge from the test user regarding the normal distribution, such as the '68-95-99.7' rule, the shorthand to remember that 68% of the scores fall within 1 SD from the mean, 95% fall within 2 SDs, and 99.7% fall within 3 SDs (see Figure 1). A score beyond *T* = 80 or below *T* = 20 is quite exceptional and only obtained by the highest 0.13% of the reference group. In contrast, percentile ranks denote the commonness of a test result in a more intuitive way by expressing the score as the percentage of respondents with a lower score.

T scores are interval-scaled, assuming this is a reasonable assumption or given for the original scores. In contrast, percentile rank scores are not equidistant and should be considered as ordinal scores. For instance, a high percentile rank score (*PR* = 75) corresponds to a *T* score of 57, which expresses a modest distance from the mean of 50. In the high range the *T* scores 65, 70, 75 and 80 correspond to 93.3, 97.7, 99.4 and 99.9 (see Figure 1). Thus, percentile rank scores have markedly unequal units, especially at the extremes, which results in underestimation of differences between scores near the mean and overestimation of differences at the extremes.

Figure 3 shows, for a selection of raw BDI-II scores, their relation with *T* scores and clinical percentile ranks (only the even raw scores are shown). Dutch data were used to obtain *T* scores and percentile ranks from a clinical sample (de Beurs et al., 2011). The relation between normal distributed *T* scores and percentile ranks follows a sigmoid curve, which reflects the stretching of the percentile rank metric relative to the *T* score metric at the extremes (due to the bell shape of the normal distribution percentile rank intervals are closer together at the middle of the scale than they are at the extremes, see also Figure 1).

When percentile rank scores are erroneously regarded as equidistant, conclusions about the test result are often wrong. The biased perception of percentile ranks was demonstrated in a study by Bowman (2002) with third-year undergraduate students, who appeared inclined to overestimate percentile ranks above 80 or below 20 as quite extreme test result (whereas under the normal curve *PR* = 80 corresponds to *T* = 58.4, that is, less than 1 SD removed from the mean, see Figure 1). Furthermore, given the prevailing view around admissible transformations and their link with arithmetical and statistical operations with percentile ranks, simple operations are not permissible, such as calculating an average over a set of scores, or calculating a difference score between repeated assessments. For

some discussion of the topic, see Meijer and Oosterloo (2008). Interestingly, this drawback interferes with the advantage of percentile scores that they are generally more intuitive to interpret.

We conclude that for the interpretation of test results and further processing of scores, *T* scores are the best choice, but for communication with colleagues and patients, we recommend supplementing these with percentile ranks. This is because *T* scores are more versatile given their psychometric properties, but percentile ranks are more intuitive to interpret. We further note that, as extreme percentile scores are susceptible to misinterpretation, caution in communication is required.

6 | CHOOSING APPROPRIATE REFERENCE GROUPS

The two common metrics we propose, *T* scores and percentile ranks, have the goal to denote how common or exceptional a test result is. This requires an appropriate reference group. What is appropriate depends on the question at hand: Do we want to know a person's severity level as compared to the general population, or as compared to patients receiving mental healthcare? We provide different recommendations for *T* scores and percentile ranks. For *T* scores used in broad, epidemiological mental healthcare contexts that aim for a universally applicable metric, the general population is the best choice, because if we assume underlying traits, general population reference samples allow comparing any subgroup within, as well as any comparison between subgroups with the same reference standard. For the same reason, in clinical use, a universally applied metric should be normed in reference to the general population. After all, it would be odd to norm IQ scores on persons with impaired intellectual functioning or, at the other extreme, the intellectually gifted.

For percentile ranks, as for *T* scores, the general population is the appropriate reference for experimental, epidemiological or clinical research. However, when it comes to clinical practice, we recommend

using a clinical reference group. This is because patients, especially at the start of treatment, will often score among the highest 5% to 10% of the general population sample (Löwe et al., 2008; Schulte-van Maaren et al., 2013). Consequently, percentile rank scores based on the general population will be quite limited in range when applied to treatment seeking patients, making it difficult to distinguish patients from each other. Furthermore, at the extreme ends of the percentile metric, a substantial change in raw score (or *T* score) translates into a small change in percentile rank. Thus, changes reflecting improvement during treatment will be diminished when we use the percentile metric where most patients score above 90. Expressing the score as a *clinical* percentile (i.e., relative to a comparable clinical group), will yield more useful information. When the same BDI-II raw scores are expressed in clinical percentiles, they range from PR = 51.2 to PR = 100.0.

When clinical percentiles are used, the clinical population they are derived from should be specified (e.g., inpatients, outpatients and outpatients seen in private practice). An illustrative example is the normative data published on the Outcome Questionnaire (OQ-45; Lambert et al., 2004) by Timman et al. (2017), who provide normative data for various patient groups. Also, Tingey et al. (1996) discuss the issue of clinical subgroups varying by severity and propose a solution within the framework of the Jacobson et al. (1984) approach to clinically significant change.

For more detailed communication with subclinical samples, where both reference groups are justifiable (e.g., at the end of successful treatment), one could present both a general population and clinical percentile. A report could state that a patient has a *T* score of 60, which is high compared to the general population (where 84% have a lower score), but below average compared to other patients attending the clinic (only 33% have a lower score). The information offered will depend on the context, but we caution against overloading the lay user with too much information, as this may be confusing and runs counter to the principle of simplicity and minimalism of 'less is more' (Peters et al., 2007).

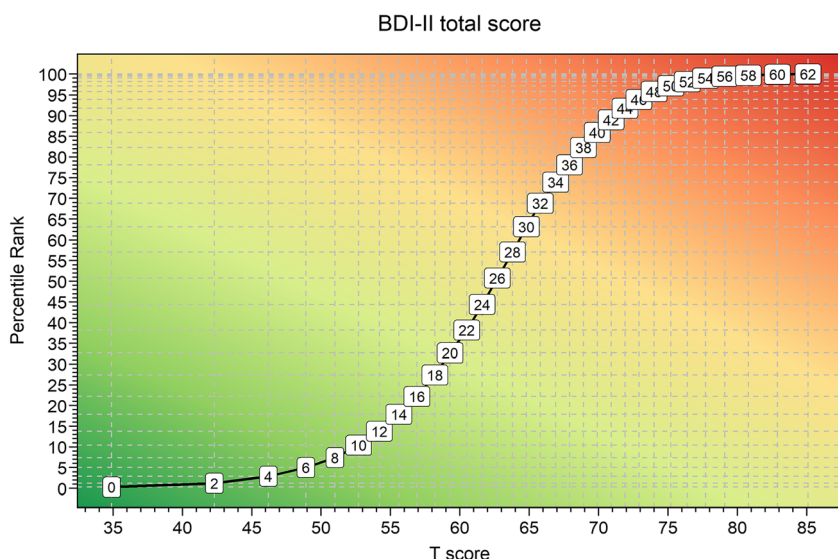


FIGURE 3 Selected raw scores on the BDI-II with *T* scores (x axis) and percentile rank scores relative to the clinical population (y axis); horizontal and vertical grid lines are based on raw scores [Colour figure can be viewed at wileyonlinelibrary.com]

7 | ILLUSTRATION

To demonstrate the utility of common metrics, we provide as an example, the crosswalk tables in Table 2. Data for Table 2 stem from tab. A1–A3 of Choi et al. (2014). They provide crosswalk tables to convert raw scores on three commonly used depression measures, the Centre for Epidemiologic Studies Depression scale CES-D, the PHQ-9 and the BDI-II to the PROMIS depression scale *T* score. For a

selection of raw scores correspondence with the *T* score metric is shown, based on the US general population sample of PROMIS. Table 2 shows the conversion of raw test scores to *T* scores and percentile ranks, and the difference between percentile rank scores based on the general population and the clinical population. Clinical percentile ranks were based on Dutch data from de Beurs et al. (2011). According to interpretation guidelines for the CES-D (Radloff, 1977), a summed item raw score of 0–15 indicates absent to mild depression,

TABLE 2 Raw scores, PROMIS *T* scores, on the CESD, PHQ-9 and BDI-II (from Choi et al., 2014) and BDI-II percentile scores based on Dutch population and clinical samples

CES-D			PHQ-9			BDI-II			PR-pop	PR-clin
RS		<i>T</i>	RS		<i>T</i>	RS		<i>T</i>		
0	no to mild	34.5	0	minimal	37.4	0	minimal	34.9	10.1	0.5
2	no to mild	41.1	1	minimal	42.7	2	minimal	42.3	26.7	1.7
4	no to mild	44.7	2	minimal	45.9	4	minimal	46.2	38.5	3.4
6	no to mild	47.5	3	minimal	48.3	6	minimal	48.9	48.6	5.5
8	no to mild	49.8	4	minimal	50.5	8	minimal	51.0	57.3	7.9
10	no to mild	51.7	5	mild	52.5	10	minimal	52.7	64.1	10.8
12	no to mild	53.4	6	mild	54.2	12	minimal	54.2	69.7	14.5
14	no to mild	54.8	7	mild	55.8	14	mild	55.6	74.4	18.4
16	moderate	56.2	8	mild	57.2	16	mild	56.9	78.2	22.6
18	moderate	57.4	9	mild	58.6	18	mild	58.2	81.3	27.7
20	moderate	58.6	10	moderate	59.9	20	moderate	59.3	84.0	33.3
22	moderate	59.7	11	moderate	61.1	22	moderate	60.5	86.3	38.7
24	severe	60.8	12	moderate	62.3	24	moderate	61.6	88.5	44.8
26	severe	61.8	13	moderate	63.5	26	moderate	62.7	90.3	51.2
28	severe	62.9	14	moderate	64.7	28	moderate	63.8	91.6	57.5
30	severe	63.9	15	mod. Severe	65.8	30	severe	64.8	92.7	63.6
32	severe	64.9	16	mod. Severe	66.9	32	severe	65.8	94.0	69.2
34	severe	66.0	17	mod. Severe	68.0	34	severe	65.8	95.2	74.0
36	severe	67.0	18	mod. Severe	69.2	36	severe	67.6	96.0	78.3
38	severe	68.1	19	mod. Severe	70.3	38	severe	68.9	96.8	82.5
40	severe	69.2	20	severe	71.5	40	severe	69.9	97.4	86.3
42	severe	70.4	21	severe	72.7	42	severe	70.9	98.0	89.5
44	severe	71.7	22	severe	74.0	44	severe	71.9	98.5	92.3
46	severe	73.0	23	severe	75.3	46	severe	72.9	98.9	94.6
48	severe	74.4	24	severe	76.7	48	severe	74.0	99.2	96.3
50	severe	76.0	25	severe	78.3	50	severe	75.2	99.5	97.7
52	severe	77.7	26	severe	80.0	52	severe	76.4	99.8	98.6
54	severe	79.7	27	severe	82.3	54	severe	77.7	99.8	99.2
56	severe	82.0				56	severe	79.1	99.9	99.6
58	severe	84.3				58	severe	80.8	99.9	99.8
60	severe	86.4				60	severe	82.9	100.0	99.9
						62	severe	85.1	100.0	100.0
						63	severe	86.3	100.0	100.0

Note: N.B.: RS = raw score; *T* = *T* score is PROMIS Depression scores (based on the US general population). *T* scores can be approximated by applying a rational function to the raw score (RS): ($T = 35.7 + (3.83 * RS - 0.0023 * RS^2)/(1 + 0.13 * RS - 0.0012 * RS^2)$); Percentile ranks can be approximated from *T* scores with: PR-pop = $-2.9 + 103.7/(1 + \exp(-0.162[T - 49.6]))$; PR-cl = $1.0 + 100.7/(1 + \exp(-0.232 * [T - 62.7]))$.

16–23 moderate and 24–60 or more indicates severe depression. The cut-off for caseness is ≥ 16 ; this corresponds to a T score of 56.2. For the PHQ-9, scores from 0 to 4 indicate minimal depression, 5–9 mild, 10–14 moderate, 15–19 moderately severe and 20–27 severe depression (Kroenke & Spitzer, 2002). The proposed cut-off for caseness is ≥ 10 , corresponding with 59.9 on the T score metric. Interpretation guidelines from the manual of the BDI-II denote a total score of 0–13 as within the minimal range, 14–19 as mild, 20–28 as moderate and 29–63 as severe. A recent meta-analysis suggested ≥ 18.2 as optimal cut-off for clinical caseness (Von Glischinski et al., 2019). For these three measures, caseness starts at $T = 56$ to 60, and the cut-off values between mild and moderate depression generally coincide with $T = 60$, which was also proposed for the T score metric by the PROMIS group (www.promis.org). When linking multiple depression measures to one common construct and metric, Wahl et al. (2014) suggested $T = 54.0$, $T = 66.2$ and $T = 72.9$ as thresholds for mild, moderate and severe depression in the BDI-II. These cut-off values are similar, but not equal which may be due to differences in operationalization of the construct or differences between the normative samples.

For the BDI-II, we also added percentile ranks to Table 2 from a sample ($N = 7500$) of the Dutch general population (Roelofs et al., 2013) and a sample of patients ($N = 9844$) seeking psychiatric care who participated in Routine Outcome Monitoring (de Beurs et al., 2011). This clearly illustrates the difference between percentile ranks based on the general population and on a clinical sample. A person with a raw score of 20 on the BDI-II has a population-based percentile rank of PR-pop = 84.0 (among the highest 16% of the general population), but a PR-clin of 33.3 (the lowest 33% of the clinical population). Someone else with a score of 26 has a population percentile rank of PR-pop = 90.3 (the highest 10% of the population), but the clinical percentile rank is PR-clin = 51.2, which indicates that a BDI-II score of 26 is close to the clinical median.

8 | DISCUSSION

The current practice of using a wide diversity of instruments for similar constructs in psychological research contributes to weakly defined measurement, as opposed to the strongly defined measurement of the physical sciences (Finkelstein, 2003). This threatens the quality of our research and hampers progress. Among clinical professionals, it may lead to a Babylonian confusion of tongues in communications about the meaning of clinical test results. Finally, and the focus of our paper, it hampers crucial communication between clinicians and their patients about the meaning of test scores.

Common metrics are ready for implementation as the long history of references for their justification above shows. In fact, they have already been implemented in Dutch clinical practice (de Beurs, 2010), where 'Delta T ' has become a commonly used metric to denote patients' progress from pretest to posttest. Yet, we see two crucial areas of further development, each with several suggestions. The first area is practical: Common metrics should be used universally, and

several steps ought to be taken to make common metrics a reality. These include a better understanding of the need for common metrics, and a demonstration of the utility of such metrics. We hope our manuscript contributes in this regard. Another crucial step for practical implementation is that crosswalk tables and figures become more easily available to demonstrate the conversion of raw scores to T scores, for instance, as a standard element of the documentation of a measure. Furthermore, we need to establish crosswalk formulas that can be built into software which administers and scores questionnaires.

Another area of improvement is psychometric research. While a detailed review of remaining obstacles is beyond the scope of this article, we list four challenges here. First, conversion of raw scores to T scores based on the general population requires normative data from community samples. Currently, such data are not available for all measures. However, when using IRT methods, there are work-arounds, such as the common-item approach or planned missing data designs, applicable when the scales or datasets to be linked share sets of items. Furthermore, various approaches to establish normalized T scores (equipercentile linking, regression-based norming, IRT) should be evaluated and compared. Additional research is needed to investigate whether even more complex alternatives, such as calibrated projection (Schalet et al., 2021; Thissen et al., 2015) are justified by a greater accuracy of the resulting T scores. Second, existing normative data are limited to specific countries. The PROMIS T score metric, for instance, is based on normative data of the US population. To test whether we can apply this metric internationally requires collecting data in other countries and comparing scores (Terwee et al., 2021). This will ease the international comparison of treatment outcome research and make such data more relevant for practice contexts as well. Third, the possible influence of translation of measures and their adaptations to other languages and cultures needs to be investigated (Sousa & Rojanasrirat, 2011; Van Widenfelt et al., 2005). We also need to study whether measures are invariant for gender, age, education level, and/or socio-economic status, as influence of these factors would imply that different norms (and T scores and percentile ranks) may be used for subgroups of respondents (Teresi et al., 2009). To accommodate this, additional conversion formulas should be provided that include coefficients for gender and age, which may result from regression-based norming (Timmerman et al., 2020). Furthermore, other external variables may need to be taken into account, such as the cultural background of clients or aspects of their psychopathology (Böhne & Croudace, 2015). Finally, conversion to T scores should correct for non-normality of raw test scores, in order to end up with T scores with genuine equal intervals. To do so, conversion tables and formulas could be based on IRT approaches, the frequency distribution of summed item scores (rank order-based normalization), or on other normalization approaches. The approach that yields the best T score metric warrants further study (Kolen & Brennan, 2014).

The area of harmonizing mental health assessment is a key methodological issue and finding solutions has recently been put on the policy agenda far beyond practical solutions for clinical reporting and discussion, as is presented here. For instance, mandating a limited

set of instruments has been suggested as a solution (Wolpert, 2020). Plans were launched by the National Institute of Mental Health and the Wellcome Trust (Farber et al., 2020) to prescribe a limited set of measurement instruments for research and outcomes assessment: the Patient Health Questionnaire (PHQ-9) for depression, the General Anxiety Disorder self-report scale (GAD-7; Spitzer et al., 2006) for anxiety, the Revised Child Anxiety and Depression Scale (RCADS-22; Chorpita et al., 2000) for depression and anxiety in children and adolescents and the World Health Organization Disability Assessment Schedule (WHODAS; Üstün et al., 2010) for impact of disabilities on adult functioning. However, while increasing comparability and interpretability of the results of research, this may have unintended negative consequences (Patalay & Fried, 2020). The selected instruments are not without flaws or drawbacks. For instance, the PHQ-9 and GAD-7 were developed to screen for depression and anxiety in the general population, and not to track progress during treatment. Both are quite brief, which limits their scope and diminishes their reliability and accuracy, making them less suitable to monitor (statistical) reliable change (Jacobson & Truax, 1991) in clinical practice, potentially leading to a premature conclusion that no clinical change has been achieved. Also, mandated use of only a few measures may hamper progress in measurement development, as it diminishes the incentive to develop alternative assessment techniques, such as the PROMIS initiative with Computer Adapted Testing (Cella et al., 2010), or improve upon prior measures. Still, some harmonization of outcome measures in clinical practice is in order and has stimulated important initiatives, such as the International Consortium for Health Outcome Measurement (ICHOM), which proposes standard outcome sets for medical conditions, including depression and anxiety (Obbarius et al., 2017). This will help to limit the proliferation of measurement-based care and may enhance the quality of mental healthcare (Fortney et al., 2017; Kilbourne et al., 2018).

Another part of the literature builds on the key to a proper understanding of what we aim for with common metrics: the difference between *equating* or merely *expressing* scores on a common metric, which should not be confused with each other. Equating test scores assumes that the same construct is being measured (Kolen & Brennan, 2014). In contrast, expressing scores on a common metric does not imply this assumption and merely aims to align test results with a common metric for the measurement of constructs that remain distinct, even if they are expressed on the same metric. For example, a *T* score of 60 for depression does not denote the same thing as a *T* score of 60 for anxiety, as the two constructs differ. In fact, given the considerable differences in content of various depression instruments, such as the CES-D and the BDI, similar *T* scores on these measures may not denote the same thing (Fried, 2017). Consequently, *T* scores for different constructs should not be directly compared, beyond the fact that they express how *extraordinary* the score is in comparison to a reference group such as the general population. While such scores carry crucial information, we caution against comparing or equating *T* scores (or percentile scores, for that matter) stemming from different constructs. For a person who scores *T* = 65 on depression and *T* = 55 on anxiety, the proper interpretation and

message would be that the depression score deviates more from the general population mean (1.5 SD) than the anxiety score (0.5 SD) or the depression score is more exceptional than the anxiety score. This differs slightly but meaningfully from the erroneous inference that the patient is more depressed than anxious.

Confusion between common *metrics* and common *constructs* may arise from the research literature on the PROMIS measures and PROMIS *T* score metric. Various legacy measures have been linked to PROMIS measures and can be converted with crosswalk tables into the preferred metric of PROMIS (*T* scores). This research is usually done per construct, such as, depression (Choi et al., 2014), anxiety (Schalet et al., 2014), pain (Cook et al., 2015) and physical functioning (Schalet et al., 2015), all titled 'Establishing a common metric for ...'. The aim of these studies was equating test results from various measures and mapping them per construct on the *T* score metric. For PROMIS measures this implies that all are scaled on a common metric: US population-based *T* scores. Moreover, for convenience, scores on PROMIS measures all share the same direction (a higher score represents more of the measured construct, such as depression, pain, mobility, functioning, or quality of life), and are similarly colour coded in graphical output (green is normal, red is severe, see Figure 2). However, using the same metric does not imply that the same construct is measured. The metric merely expresses conveniently for each construct the exceptionality of the score (e.g., depression), but it does not imply that these scores are comparable across constructs. For more studies using this approach, see, for example, for depression (Wahl et al., 2014), physical functioning (Oude Voshaar et al., 2019), personality pathology (Zimmermann et al., 2020), psychological distress (Batterham et al., 2018) and fatigue (Lai et al., 2014).

Finally, it is important to note that converting scale scores to *T* scores does not imply that the original scores are lost. The sum scores metric can still be calculated and reported in research, for instance, to compare scores with previously published research findings that used the traditional way of scoring.

To conclude, use of common metrics, particularly population-based *T* scores and clinical percentile ranks, may aid to harmonize measurement practices in clinical psychology research, will be helpful for a proper interpretation of test results, enhance the communication about tests results among professionals and ease explanation of their meaning to patients in mental healthcare. To bring this about, a first step is to provide easy ways to express test results as *T* scores, for instance, in user manuals for assessment instruments with cross walk tables and formulas. Also, nowadays many self-report questionnaires are administered via computer or smart phone, and expressing results in *T* scores can be accomplished easily within the software. Next steps are educating professionals, helping them gain experience with *T* scores and asking them to help educate their clients on the meaning of *T* scores. The common use of IQ scores and understanding by the general public on what IQ scores convey shows the feasibility of this approach. We hope that our paper stimulates the use of *T* scores as the main way to give meaning to (neuro-)psychological test results.

CONFLICT OF INTEREST

The authors have no conflict of interest to declare.

AUTHOR CONTRIBUTIONS

EdeB wrote the first draft of the manuscript and analysed the data that were added to Table 2. EIF and JRB commented and improved the manuscript.

DATA AVAILABILITY STATEMENT

Predominantly previously published data were used for this manuscript. A Pre-test Routine Outcome Monitoring data from a clinical sample of a mental healthcare institute (Rivierduinen in Leiden) was used to add a column to Table 2. The Medical Ethical Boards of the Leiden University Medical Center approved use of anonymized data that were collected routinely in clinical practice. Patients gave informed consent for anonymized use of their data.

ORCID

Edwin de Beurs  <https://orcid.org/0000-0003-3832-8477>

Jan R. Boehnke  <https://orcid.org/0000-0003-0249-1870>

Eiko I. Fried  <https://orcid.org/0000-0001-7469-594X>

REFERENCES

- Achenbach, T. M. (1991). *Manual for the child behavior checklist 4-18 and 1991 profiles*. Department of Psychiatry, University of Vermont.
- Anastasi, A. (1968). *Psychological testing*. Macmillan.
- Aschenbrand, S. G., Angelosante, A. G., & Kendall, P. C. (2005). Discriminant validity and clinical utility of the CBCL with anxiety-disordered youth. *Journal of Clinical Child & Adolescent Psychology*, 34(4), 735-746. https://doi.org/10.1207/s15374424jccp3404_15
- Batterham, P. J., Sunderland, M., Slade, T., Cleave, A. L., & Carragher, N. (2018). Assessing distress in the community: Psychometric properties and crosswalk comparison of eight measures of psychological distress. *Psychological Medicine*, 48(8), 1316-1324. <https://doi.org/10.1017/S0033291717002835>
- Baty, F., Ritz, C., & Baty, M. F. (2015). Package 'nlstools'. In *Tools for nonlinear regression analysis* (p. 66). <https://doi.org/10.18637/jss.v066.i05>
- Beck, A. T., Steer, R. A., & Brown, G. K. (1996). *BDI-II Beck depression inventory manual*. The Psychological Corporation.
- Binet, A., & Simon, T. (1907). Le développement de l'intelligence chez les enfants. *L'Année Psychologique*, 14(1), 1-94. https://www.persee.fr/doc/psy_0003-5033_1907_num_14_1_3737
- Boehnke, J. R., & Rutherford, C. (2021). Using feedback tools to enhance the quality and experience of care. *Quality of Life Research*, 30(11), 3007-3013. <https://doi.org/10.1007/s11136-021-03008-8>
- Böhnke, J., & Croudace, T. J. (2015). Factors of psychological distress: Clinical value, measurement substance, and methodological artefacts. *Social Psychiatry and Psychiatric Epidemiology*, 50(4), 515-524. <https://doi.org/10.1007/s00127-015-1022-5>
- Bowman, M. L. (2002). The perfidy of percentiles. *Archives of Clinical Neuropsychology*, 17(3), 295-303. [https://doi.org/10.1016/S0887-6177\(01\)00116-0](https://doi.org/10.1016/S0887-6177(01)00116-0)
- Box, G. E., & Cox, D. R. (1964). An analysis of transformations. *Journal of the Royal Statistical Society: Series B (Methodological)*, 26(2), 211-243. <https://www.jstor.org/stable/2984418>
- Brundage, M. D., Smith, K. C., Little, E. A., Bantug, E. T., Snyder, C. F., & The P. R. O. D. P. S. A. B. (2015). Communicating patient-reported outcome scores using graphic formats: Results from a mixed-methods evaluation. *Quality of Life Research*, 24(10), 2457-2472. <https://doi.org/10.1007/s11136-015-0974-y>
- Butcher, J. N., Dahlstrom, W. G., Graham, J. R., Tellegen, A., & Kaemmer, B. (1989). *MMPI-2 (Minnesota multiphasic personality inventory-2): Manual for administration and scoring*. University of Minnesota Press.
- Cella, D., Riley, W., Stone, A., Rothrock, N., Reeve, B., Yount, S., ... Hays, R. (2010). The Patient-Reported Outcomes Measurement Information System (PROMIS) developed and tested its first wave of adult self-reported health outcome item banks: 2005-2008. *Journal of Clinical Epidemiology*, 63(11), 1179-1194. <https://doi.org/10.1016/j.jclinepi.2010.04.011>
- Cella, D., Choi, S., Garcia, S., Cook, K. F., Rosenbloom, S., Lai, J.-S., Tatum, D. S., & Gershon, R. (2014). Setting standards for severity of common symptoms in oncology using the PROMIS item banks and expert judgment. *Quality of Life Research*, 23(10), 2651-2661. <https://doi.org/10.1007/s11136-014-0732-6>
- Choi, S. W., Schalet, B., Cook, K. F., & Cella, D. (2014). Establishing a common metric for depressive symptoms: Linking the BDI-II, CES-D, and PHQ-9 to PROMIS depression. *Psychological Assessment*, 26(2), 513-527. <https://doi.org/10.1037/a0035768>
- Chorpita, B. F., Yim, L., Moffitt, C., Umemoto, L. A., & Francis, S. E. (2000). Assessment of symptoms of DSM-IV anxiety and depression in children: A revised child anxiety and depression scale. *Behaviour Research and Therapy*, 38(8), 835-855. [https://doi.org/10.1016/S0005-7967\(99\)00130-8](https://doi.org/10.1016/S0005-7967(99)00130-8)
- Cook, K. F., Schalet, B. D., Kallen, M. A., Rutsohn, J. P., & Cella, D. (2015). Establishing a common metric for self-reported pain: Linking BPI pain interference and SF-36 bodily pain subscale scores to the PROMIS pain interference metric. *Quality of Life Research*, 24(10), 2305-2318. <https://doi.org/10.1007/s11136-014-0790-9>
- Crawford, J., Cayley, C., Lovibond, P. F., Wilson, P. H., & Hartley, C. (2011). Percentile norms and accompanying Interval estimates from an Australian general adult population sample for self-report mood scales (BAI, BDI, CRSD, CES-D, DASS, DASS-21, STAI-X, STAI-Y, SRDS, and SRAS). *Australian Psychologist*, 46(1), 3-14. <https://doi.org/10.1111/j.1742-9544.2010.00003.x>
- Crawford, J. R., & Garthwaite, P. H. (2009). Percentiles please: The case for expressing neuropsychological test scores and accompanying confidence limits as percentile ranks. *The Clinical Neuropsychologist*, 23(2), 193-204. <https://doi.org/10.1080/13854040801968450>
- Crawford, J. R., & Henry, J. D. (2003). The Depression Anxiety Stress Scales (DASS): Normative data and latent structure in a large non-clinical sample. *British Journal of Clinical Psychology*, 42(2), 111-131. <https://doi.org/10.1348/014466503321903544>
- Crawford, J. R., Garthwaite, P. H., Lawrie, C. J., Henry, J. D., MacDonald, M. A., Sutherland, J., & Sinha, P. (2009). A convenient method of obtaining percentile norms and accompanying interval estimates for self-report mood scales (DASS, DASS-21, HADS, PANAS, and sAD). *British Journal of Clinical Psychology*, 48(2), 163-180. <https://doi.org/10.1348/014466508X377757>
- Crocker, L. M., & Algina, J. (1986). *Introduction to classical and modern test theory*. Holt, Rinehart, and Winston. <https://books.google.nl/books?id=tfkgAQAAIAAJ>
- Cronbach, L. J. (1984). *Essentials of psychological testing* (4th ed.). Harper & Row.
- de Beurs, E. (2010). De genormaliseerde T-score, een 'euro' voor testuitslagen [The normalised T-score: A euro for test results]. *Maandblad Geestelijke Volksgezondheid*, 65, 684-695. www.sbggz.nl
- de Beurs, E., Den Hollander-Gijsman, M. E., Van Rood, Y. R., Van der Wee, N. J., Giltay, E. J., Van Noorden, M. S., Van der Lem, R., Van Fenema, E., & Zitman, F. G. (2011). Routine outcome monitoring in the Netherlands: Practical experiences with a web-based strategy for the assessment of treatment outcome in clinical practice. *Clinical Psychology & Psychotherapy*, 18(1), 1-12. <https://doi.org/10.1002/cpp.696>

- de Beurs, E., Carlier, I. V., & Van Hemert, A. M. (2019). Approaches to denote treatment outcome: Clinical significance and clinical global impression compared. *International Journal of Methods in Psychiatric Research*, 28, e1797. <https://doi.org/10.1002/mpr.1797>
- de Beurs, E., Oudejans, S., & Terluin, B. (Submitted). A common measurement scale for scores from self-report instruments in mental health care: T scores with a normal distribution. *European Journal of Psychological Assessment*.
- DeMars, C. E. (2018). Classical Test Theory and Item Response Theory. In *The Wiley handbook of psychometric testing* (pp. 49–73). <https://doi.org/10.1002/9781118489772.ch2>
- Derogatis, L. R. (1975). *The brief symptom inventory*. Clinical Psychometric Research.
- Farber, G., Wolpert, M., & Kemmer, D. (2020). Common measures for mental health science: Laying the foundations. Wellcome Trust. Retrieved 11-1-2021 from: <https://wellcome.ac.uk/sites/default/files/CMB-and-CMA-July-2020-pdf.pdf>
- Finkelstein, L. (2003). Widely, strongly and weakly defined measurement. *Measurement*, 34(1), 39–48. [https://doi.org/10.1016/S0263-2241\(03\)00018-6](https://doi.org/10.1016/S0263-2241(03)00018-6)
- Fischer, H. F., & Rose, M. (2016). www.common-metrics.org: A web application to estimate scores from different patient-reported outcome measures on a common scale. *BMC Medical Research Methodology*, 16(1), 142. <https://doi.org/10.1186/s12874-016-0241-0>
- Fischer, H. F., & Rose, M. (2019). Scoring depression on a common metric: A comparison of EAP estimation, plausible value imputation, and full Bayesian IRT modeling. *Multivariate Behavioral Research*, 54(1), 85–99. <https://doi.org/10.1080/00273171.2018.1491381>
- Flake, J. K., & Fried, E. I. (2020). Measurement Schmeasurement: Questionable measurement practices and how to avoid them. *Advances in Methods and Practices in Psychological Science*, 3(4), 456–465. <https://doi.org/10.1177/2515245920952393>
- Fortney, J. C., Unützer, J., Wrenn, G., Pyne, J. M., Smith, G. R., Schoenbaum, M., & Harbin, H. T. (2017). A tipping point for measurement-based care. *Psychiatric Services*, 68(2), 179–188. <https://doi.org/10.1176/appi.ps.201500439>
- Fried, E. I. (2017). The 52 symptoms of major depression: Lack of content overlap among seven common depression scales. *Journal of Affective Disorders*, 208, 191–197. <https://doi.org/10.1016/j.jad.2016.10.019>
- Friedrich, M., Hinz, A., Kuhnt, S., Schulte, T., Rose, M., & Fischer, F. (2019). Measuring fatigue in cancer patients: A common metric for six fatigue instruments. *Quality of Life Research*, 28(6), 1615–1626. <https://doi.org/10.1007/s11136-019-02147-3>
- Goetz, T. (2010). *It's time to redesign medical data [Presentation]*. TEDMED.
- Guy, W. (1976). "Clinical Global Impressions" ECDEU Assessment Manual for Psychopharmacology, Revised. DHEW publication ADM 76-338. Government Printing Office.
- Harding, K. J., Rush, A. J., Arbuckle, M., Trivedi, M. H., & Pincus, H. A. (2011). Measurement-based care in psychiatric practice: A policy framework for implementation. *Journal of Clinical Psychiatry*, 72(8), 1136–1143. <https://doi.org/10.4088/JCP.10r06282whi>
- healthmeasures. (n.d.). <https://www.healthmeasures.net/score-and-interpret/interpret-scores/promis/promis-score-cut-points>
- Jacobson, N. S., & Truax, P. (1991). Clinical significance: A statistical approach to defining meaningful change in psychotherapy research. *Journal of Consulting and Clinical Psychology*, 59(1), 12–19. <https://doi.org/10.1037//0022-006X.59.1.12>
- Jacobson, N. S., Follette, W. C., & Revenstorf, D. (1984). Psychotherapy outcome research: Methods for reporting variability and evaluating clinical significance. *Behavior Therapy*, 15, 336–352. (In File. [https://doi.org/10.1016/S0005-7894\(84\)80002-7](https://doi.org/10.1016/S0005-7894(84)80002-7)
- Kellen, D., Davis-Stober, C. P., Dunn, J. C., & Kalish, M. L. (2021). The problem of coordination and the pursuit of structural constraints in psychology. *Perspectives on Psychological Science*, 16(4), 767–778. <https://doi.org/10.1177/1745691620974771>
- Kendall, P. C., Marrs-Garcia, A., Nath, S. R., & Sheldrick, R. C. (1999). Normative comparisons for the evaluation of clinical significance. *Journal of Consulting and Clinical Psychology*, 67(3), 285–299. <https://doi.org/10.1037/0022-006X.67.3.285>
- Kilbourne, A. M., Beck, K., Spaeth-Rublee, B., Ramanuj, P., O'Brien, R. W., Tomoyasu, N., & Pincus, H. A. (2018). Measuring and improving the quality of mental health care: A global perspective. *World Psychiatry*, 17(1), 30–38. <https://doi.org/10.1002/wps.20482>
- Kolen, M. J., & Brennan, R. L. (2014). *Test equating, scaling, and linking: Methods and practices* (3rd ed.). Springer Science & Business Media. <https://doi.org/10.1007/978-1-4939-0317-7>
- Kroenke, K., & Spitzer, R. L. (2002). The PHQ-9: A new depression diagnostic and severity measure. *Psychiatric Annals*, 32(9), 509–515. <https://doi.org/10.3928/0048-5713-20020901-06>
- Kurtz, A. K., & Mayo, S. T. (1979). Percentiles and Percentile Ranks. In *Statistical methods in education and psychology* (pp. 145–163). Springer. https://doi.org/10.1007/978-1-4612-6129-2_6
- Lai, J.-S., Cella, D., Yanez, B., & Stone, A. (2014). Linking fatigue measures on a common reporting metric. *Journal of Pain and Symptom Management*, 48(4), 639–648. <https://doi.org/10.1016/j.jpainsymman.2013.12.236>
- Lambert, M. J. (2007). Presidential address: What we have learned from a decade of research aimed at improving psychotherapy outcome in routine care. *Psychotherapy Research*, 17(1), 1–14. <https://doi.org/10.1080/10503300601032506>
- Lambert, M. J., & Harmon, K. L. (2018). The merits of implementing routine outcome monitoring in clinical practice. *Clinical Psychology: Science and Practice*, 25(4), e12268. <https://doi.org/10.1111/cpsp.12268>
- Lambert, M. J., Gregersen, A. T., & Burlingame, G. M. (2004). The Outcome Questionnaire-45. In M. E. Maruish (Ed.), *The use of psychological testing for treatment planning and outcomes assessment: Instruments for adults* (3rd ed.) (pp. 191–234). Lawrence Erlbaum Associates Publishers. <http://search.ebscohost.com/login.aspx?direct=true&db=psyh&AN=2004-14941-006&site=ehost-live>
- Lenhard, W., & Lenhard, A. (2021). Improvement of norm score quality via regression-based continuous norming. *Educational and Psychological Measurement*, 81(2), 229–261. <https://doi.org/10.1177/0013164420928457>
- Lewis, C. C., Boyd, M., Puspitasari, A., Navarro, E., Howard, J., Kassab, H., Hoffman, M., Scott, K., Lyon, A., & Douglas, S. (2019). Implementing measurement-based care in behavioral health: A review. *JAMA Psychiatry*, 76(3), 324–335. <https://doi.org/10.1001/jamapsychiatry.2018.3329>
- Ley, P. (1972). *Quantitative aspects of psychological assessment*. Duckworth.
- Liddell, T. M., & Kruschke, J. K. (2018). Analyzing ordinal data with metric models: What could possibly go wrong? *Journal of Experimental Social Psychology*, 79, 328–348. <https://doi.org/10.1016/j.jesp.2018.08.009>
- Liu, H., Lafferty, J., & Wasserman, L. (2009). The nonparanormal: Semiparametric estimation of high dimensional undirected graphs. *Journal of Machine Learning Research*, 10(10). <https://doi.org/10.1145/1577069.1755863>
- Lord, F. M., & Wingersky, M. S. (1984). Comparison of IRT true-score and equipercentile observed-score "equatings". *Applied Psychological Measurement*, 8(4), 453–461. <https://doi.org/10.1177/014662168400800409>
- Löwe, B., Decker, O., Müller, S., Brähler, E., Schellberg, D., Herzog, W., & Herzberg, P. Y. (2008). Validation and standardization of the Generalized Anxiety Disorder Screener (GAD-7) in the general population. *Medical Care*, 45(3), 266–274. <https://doi.org/10.1097/MLR.0b013e318160d093>
- McCall, W. A. (1922). *How to measure in education*. MacMillan.
- McPherson, S., & Armstrong, D. (2021). Psychometric origins of depression. *History of the Human Sciences*, 09526951211009085. <https://doi.org/10.1177/09526951211009085>

- Meijer, R. R., & Oosterloo, S. J. (2008). A note on measurement scales and statistical testing. *Measurement: Interdisciplinary Research and Perspectives*, 6(3), 198–204. <https://doi.org/10.1080/15366360802324446>
- Molenaar, D., & Dolan, C. V. (2018). Nonnormality in Latent Trait Modeling. In *The Wiley handbook of psychometric testing* (pp. 347–373). <https://doi.org/10.1002/9781118489772.ch13>
- Nelson, G. H., O'Hara, M. W., & Watson, D. (2018). National norms for the expanded version of the inventory of depression and anxiety symptoms (IDAS-II). *Journal of Clinical Psychology*, 74(6), 953–968. <https://doi.org/10.1002/jclp.22560>
- Obbarius, A., van Maasakkers, L., Baer, L., Clark, D. M., Crocker, A. G., de Beurs, E., Emmelkamp, P. M. G., Furukawa, T. A., Hedman-Lagerlöf, E., Kangas, M., Langford, L., Lesage, A., Mwesigire, D. M., Nolte, S., Patel, V., Pilkonis, P. A., Pincus, H. A., Reis, R. A., Rojas, G., ... Rose, M. (2017). Standardization of health outcomes assessment for depression and anxiety: Recommendations from the ICHOM Depression and Anxiety Working Group. *Quality of Life Research*, 26, 1–15. <https://doi.org/10.1007/s11136-017-1659-5>
- Ogles, B. M. (2013). Measuring Change in Psychotherapy Research. In M. J. Lambert (Ed.), *Bergin and Garfield's handbook of psychotherapy and behavior change* (pp. 134–166).
- Oude Voshaar, M., Vonkeman, H., Courvoisier, D., Finckh, A., Gossec, L., Leung, Y., Michaud, K., Pinheiro, G., Soriano, E., & Wulfrat, N. (2019). Towards standardized patient reported physical function outcome reporting: Linking ten commonly used questionnaires to a common metric. *Quality of Life Research*, 28(1), 187–197. <https://doi.org/10.1007/s11136-018-2007-0>
- Patalay, P., & Fried, E. I. (2020). Editorial perspective: Prescribing measures: Unintended negative consequences of mandating standardized mental health measurement. *Journal of Child Psychology and Psychiatry*, 62, 1032–1036. <https://doi.org/10.1111/jcpp.13333>
- Patel, S. R., Bakken, S., & Ruland, C. (2008). Recent advances in shared decision making for mental health. *Current Opinion in Psychiatry*, 21(6), 606–6012. <https://doi.org/10.1097/YCO.0b013e32830eb6b4>
- Peters, E., Dieckmann, N., Dixon, A., Hibbard, J. H., & Mertz, C. K. (2007). Less is more in presenting quality information to consumers. *Medical Care Research and Review*, 64(2), 169–190. <https://doi.org/10.1177/10775587070640020301>
- Prinsen, C. A. C., Mokkink, L. B., Bouter, L. M., Alonso, J., Patrick, D. L., De Vet, H. C. W., & Terwee, C. B. (2018). COSMIN guideline for systematic reviews of patient-reported outcome measures. *Quality of Life Research*, 27(5), 1147–1157. <https://doi.org/10.1007/s11136-018-1798-3>
- Radloff, L. S. (1977). The CES-D scale: A self-report depression scale for research in the general population. *Applied Psychological Measurement*, 1, 385–401. <https://doi.org/10.1177/014662167700100306>
- Recklitis, C. J., & Rodriguez, P. (2007). Screening childhood cancer survivors with the brief symptom inventory-18: Classification agreement with the symptom checklist-90-revised. *Psycho-Oncology*, 16(5), 429–436. <https://doi.org/10.1002/pon.1069>
- Reise, S. P., Rodriguez, A., Spritzer, K. L., & Hays, R. D. (2018). Alternative approaches to addressing non-Normal distributions in the application of IRT models to personality measures. *Journal of Personality Assessment*, 100(4), 363–374. <https://doi.org/10.1080/00223891.2017.1381969>
- Roelofs, J., Van Breukelen, G., De Graaf, L. E., Beck, A. T., Arntz, A., & Huibers, M. J. H. (2013). Norms for the Beck depression inventory (BDI-II) in a large Dutch community sample. *Journal of Psychopathology and Behavioral Assessment*, 35(1), 93–98. <https://doi.org/10.1007/s10862-012-9309-2>
- Rose, M., & Devine, J. (2014). Assessment of patient-reported symptoms of anxiety. *Dialogues in Clinical Neuroscience*, 16(2), 197–211. <https://doi.org/10.31887/DCNS.2014.16.2/mrose>
- Santor, D. A., Gregus, M., & Welch, A. (2006). FOCUS ARTICLE: Eight decades of measurement in depression. *Measurement: Interdisciplinary Research and Perspectives*, 4(3), 135–155. https://doi.org/10.1207/s15366359mea0403_1
- Schalet, B. D., Cook, K. F., Choi, S. W., & Cella, D. (2014). Establishing a common metric for self-reported anxiety: Linking the MASQ, PANAS, and GAD-7 to PROMIS anxiety. *Journal of Anxiety Disorders*, 28(1), 88–96. <https://doi.org/10.1016/j.janxdis.2013.11.006>
- Schalet, B. D., Revicki, D. A., Cook, K. F., Krishnan, E., Fries, J. F., & Cella, D. (2015). Establishing a common metric for physical function: Linking the HAQ-DI and SF-36 PF subscale to PROMIS® physical function. *Journal of General Internal Medicine*, 30(10), 1517–1523. <https://doi.org/10.1007/s11606-015-3360-0>
- Schalet, B. D., Lim, S., Cella, D., & Choi, S. W. (2021). Linking scores with patient-reported health outcome instruments: A validation study and comparison of three linking methods. *Psychometrika*, 86(3), 717–746. <https://doi.org/10.1007/s11336-021-09776-z>
- Schulte-van Maaren, Y. W., Carlier, I. V., Zitman, F. G., Van Hemert, A. M., De Waal, M. W., Van der Does, A. W., Van Noorden, M. S., & Giltay, E. J. (2013). Reference values for major depression questionnaires: The Leiden routine outcome monitoring study. *Journal of Affective Disorders*, 149(1–3), 342–349. <https://doi.org/10.1016/j.jad.2013.02.009>
- Seashore, H. G. (1955). Methods of expressing test scores. *Test Service Bulletin*, 48, 7–10. <https://eric.ed.gov/?id=ED079347>
- Snyder, C., Smith, K., Holzner, B., Rivera, Y. M., Bantug, E., Brundage, M., Weber, D., Basch, E., Aaronson, N., Reeve, B., Velikova, G., Heckert, A., Stotsky-Himelfarb, E., Chauhan, C., Hoffman, V., Ganz, P., Barbera, L., Frank, E., Smith, M. L., & PRO Data Presentation Delphi Panel. (2019). Making a picture worth a thousand numbers: Recommendations for graphically displaying patient-reported outcomes data. *Quality of Life Research*, 28(2), 345–356. <https://doi.org/10.1007/s11136-018-2020-3>
- Snyder, C. F., Aaronson, N. K., Choucair, A. K., Elliott, T. E., Greenhalgh, J., Halyard, M. Y., Hess, R., Miller, D. M., Reeve, B. B., & Santana, M. (2012). Implementing patient-reported outcomes assessment in clinical practice: A review of the options and considerations. *Quality of Life Research*, 21(8), 1305–1314. <https://doi.org/10.1007/s11136-011-0054-x>
- Sousa, V. D., & Rojjanasrirat, W. (2011). Translation, adaptation and validation of instruments or scales for use in cross-cultural health care research: A clear and user-friendly guideline. *Journal of Evaluation in Clinical Practice*, 17(2), 268–274. <https://doi.org/10.1111/j.1365-2753.2010.01434.x>
- Spitzer, R. L., Kroenke, K., Williams, J. B., & Löwe, B. (2006). A brief measure for assessing generalized anxiety disorder: The GAD-7. *Archives of Internal Medicine*, 166(10), 1092–1097. <https://doi.org/10.1001/archinte.166.10.1092>
- Stasinopoulos, M. D., Rigby, R. A., & Bastiani, F. D. (2018). GAMLSS: a distributional regression approach. *Statistical Modelling*, 18(3–4), 248–273. <https://doi.org/10.1177/1471082X18759144>
- Teresi, J. A., Ocepek-Welikson, K., Kleinman, M., Eimicke, J. P., Crane, P. K., Jones, R. N., Lai, J. S., Choi, S. W., Hays, R. D., Reeve, B. B., Reise, S. P., Pilkonis, P. A., & Cella, D. (2009). Analysis of differential item functioning in the depression item bank from the patient reported outcome measurement information system (PROMIS): An item response theory approach. *Psychological Science Quarterly*, 51(2), 148–180. <http://www.ncbi.nlm.nih.gov/pubmed/20336180>
- Terwee, C. B., Crins, M. H. P., Roorda, L. D., Cook, K. F., Cella, D., Smits, N., & Schalet, B. D. (2021). International application of PROMIS computerized adaptive tests: US versus country-specific item parameters can be consequential for individual patient scores. *Journal of Clinical Epidemiology*, 134, 1–13. <https://doi.org/10.1016/j.jclinepi.2021.01.011>
- Thissen, D., Liu, Y., Magnus, B., & Quinn, H. (2015). Extending the use of multidimensional IRT calibration as projection: Many-to-one linking

- and linear computation of projected scores. In L. A. Van der Ark, D. M. Bolt, W.-C. Wang, J. A. Douglas, & S.-M. Chow (Eds.), *Quantitative psychology research* (pp. 1–16). Springer. https://doi.org/10.1007/978-3-319-19977-1_1
- Timman, R., De Jong, K., & De Neve-Enthoven, N. (2017). Cut-off scores and clinical change indices for the Dutch outcome questionnaire (OQ-45) in a large sample of normal and several psychotherapeutic populations. *Clinical Psychology & Psychotherapy*, 24(1), 72–81. <https://doi.org/10.1002/cpp.1979>
- Timmerman, M. E., Voncken, L., & Albers, C. J. (2020). A tutorial on regression-based norming of psychological tests with GAMLSS. *Psychological Methods*, 26, 357–373. <https://doi.org/10.1037/met0000348>
- Tingey, R., Lambert, M., Burlingame, G., & Hansen, N. (1996). Assessing clinical significance: Proposed extensions to method. *Psychotherapy Research*, 6(2), 109–123. <https://doi.org/10.1080/10503309612331331638>
- Üstün, T. B., Kosstanjek, N., Chatterji, S., & Rehm, J. (2010). Measuring health and disability: Manual for WHO disability assessment schedule WHODAS 2.0.
- Van Widenfelt, B. M., Treffers, P. D., de Beurs, E., Siebelink, B. M., & Koudijs, E. (2005). Translation and cross-cultural adaptation of assessment instruments used in psychological research with children and families. *Clinical Child and Family Psychology Review*, 8(2), 135–147. <https://doi.org/10.1007/s10567-005-4752-1>
- Von Glischinski, M., Von Brachel, R., & Hirschfeld, G. (2019). How depressed is “depressed”? A systematic review and diagnostic meta-analysis of optimal cut points for the Beck Depression Inventory revised (BDI-II). *Quality of Life Research*, 28(5), 1111–1118. <https://doi.org/10.1007/s11136-018-2050-x>
- Wahl, I., Löwe, B., Bjorner, J. B., Fischer, F., Langs, G., Voderholzer, U., Aita, S. A., Bergemann, N., Brähler, E., & Rose, M. (2014). Standardization of depression measurement: A common metric was developed for 11 self-report depression measures. *Journal of Clinical Epidemiology*, 67(1), 73–86. <https://doi.org/10.1016/j.jclinepi.2013.04.019>
- Weidman, A. C., Steckler, C. M., & Tracy, J. L. (2017). The jingle and jangle of emotion assessment: Imprecise measurement, casual scale usage, and conceptual fuzziness in emotion research. *Emotion*, 17(2), 267–295. <https://doi.org/10.1037/emo0000226>
- Weston, J., Dwan, K., Altman, D., Clarke, M., Gamble, C., Schroter, S., Williamson, P., & Kirkham, J. (2016). Feasibility study to examine discrepancy rates in prespecified and reported outcomes in articles submitted to *The BMJ*. *BMJ Open*, 6(4), e010075. <https://doi.org/10.1136/bmjopen-2015-010075>
- Wolpert, M. (2020). Funders agree first common metrics for mental health science. <https://www.linkedin.com/pulse/funders-agree-first-common-metrics-mentalhealth-science-wolpert>
- Zachary, R. A., & Gorsuch, R. L. (1985). Continuous norming: Implications for the WAIS-R. *Journal of Clinical Psychology*, 41(1), 86–94. [https://doi.org/10.1002/1097-4679\(198501\)41:1<86::AID-JCLP2270410115>3.0.CO;2-W](https://doi.org/10.1002/1097-4679(198501)41:1<86::AID-JCLP2270410115>3.0.CO;2-W)
- Zimmermann, J., Müller, S., Bach, B., Hutsebaut, J., Hummelen, B., & Fischer, F. (2020). A common metric for self-reported severity of personality disorder. *Psychopathology*, 53(3), 168–178. <https://doi.org/10.1159/000507377>

How to cite this article: de Beurs, E., Boehnke, J. R., & Fried, E. I. (2022). Common measures or common metrics? A plea to harmonize measurement results. *Clinical Psychology & Psychotherapy*, 1–13. <https://doi.org/10.1002/cpp.2742>