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# Author manuscript

Depress Anxiety. Author manuscript; available in PMC 2020 September 01.

Published in final edited form as:

Depress Anxiety. 2019 September; 36(9): 813-823. doi:10.1002/da.22940.

# Measurement Invariance of the Patient Health Questionnaire-9 (PHQ-9) Depression Screener in U.S. Adults Across Sex, Race/ Ethnicity, and Education Level: NHANES 2005-2016

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#### **Abstract**

**Background:** Despite its popularity, little is known about the measurement invariance of the Patient Health Questionnaire-9 (PHQ-9) across U.S. sociodemographic groups. Use of a screener shown not to possess measurement invariance could result in under-detection/treatment of depression, potentially exacerbating sociodemographic disparities in depression. Therefore, we assessed the factor structure and measurement invariance of the PHQ-9 across major U.S. sociodemographic groups.

**Methods:** U.S. population representative data came from the 2005–2016 National Health and Nutrition Examination Survey (NHANES) cohorts We conducted a measurement invariance analysis of 31,366 respondents across sociodemographic factors of sex, race/ethnicity, and education level.

**Results:** Considering results of single-group confirmatory factor analyses (CFAs), depression theory, and research utility, we justify a two-factor structure for the PHQ-9 consisting of a cognitive/affective factor and a somatic factor (RMSEA=0.034, TLI=0.985, CFI=0.989). Based on multiple-group CFAs testing configural, scalar, and strict factorial invariance, we determined that invariance held for sex, race/ethnicity, and education level groups, as all models demonstrated close model fit (RMSEA=0.025–0.025, TLI=0.985–0.992, CFI=0.986–0.991). Finally, for all steps CFI was < -0.004, and RMSEA was < 0.01.

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**Conclusions:** We demonstrate that the PHQ-9 is acceptable to use in major U.S. sociodemographic groups and allows for meaningful comparisons in total, cognitive/affective, and somatic depressive symptoms across these groups, extending its use to the community. This knowledge is timely as medicine moves towards alternative payment models emphasizing high-quality and cost-efficient care, which will likely incentivize behavioral and population health efforts. We also provide a consistent, evidence-based approach for calculating PHQ-9 subscale scores.

#### Introduction

Depression is a top public health concern due to its high prevalence, chronicity, and grave ramifications. The lifetime prevalence of major depressive disorder (MDD) in the U.S. is 16% (Kessler et al., 2003). Depression also disproportionately affects people with lower socioeconomic status (Lorant et al., 2003). The course of MDD is often chronic, with a 15-year recurrence rate of 35% in the general population (Hardeveld, Spijker, De Graaf, Nolen, & Beekman, 2010). Its grave ramifications include increased disability, mortality, and societal costs. Depression is the second leading cause of disability (Ferrari et al., 2013), and associated with increased mortality risk (Cuijpers et al., 2014). Moreover, the total annual cost of depression has increased from \$83 billion in 2000 to \$210 billion in 2010 (Greenberg, Fournier, Sisitsky, Pike, & Kessler, 2015; Greenberg et al., 2003).

These alarming observations have motivated efforts to improve the detection and management of depression by routinely administering screeners. In 2016, the U.S. Preventive Services Task Force (USPSTF) recommended depression screening in the general adult population (Siu et al., 2016). Accompanying editorials underscored the importance of "population screening" (Thase, 2016) while also critiquing the USPSTF statement for not specifying "the ideal screening interval and the settings with highest potential yield" (Reynolds & Frank, 2016, pg. 189). One widely-used, self-report depression screener that was designed for primary care is the Patient Health Questionnaire-9 (PHQ-9; Hirschtritt & Kroenke, 2017). The PHQ-9 was identified as the first choice screener for adults in the accompanying editorial (Thase, 2016). In addition to a depression screener, the PHQ-9 has been validated as a continuous measure of depressive symptom severity (Kroenke, Spitzer, & Williams, 2001). The push for increased depression screening is occurring in the context of medicine's movement toward alternative payment models that emphasize high-quality and cost-efficient care (Ganguli & Ferris, 2018). It has been posited that behavioral health integration and population health management will be incentivized and thus more widely adopted under these emerging payment models (Burwell, 2015; Joynt Maddox, 2018; "Smarter Spending. Healthier People: Paying Providers for Value, Not Volume," 2015). These potentially profound changes to the healthcare system will amplify the need for brief depression screeners known to operate in an unbiased manner in major sociodemographic groups in the U.S., especially considering depression is associated with high healthcare utilization and costs (Bock et al., 2014; Kroenke & Unutzer, 2017; Luber et al., 2000; Prina et al., 2015). Moreover, measurement-based care is a key pillar in evidence-based behavioral health integration programs that have not only improved depression outcomes but also have proven cost-effective (Kroenke & Unutzer, 2017).

Despite its widespread use in research and clinical practice, surprisingly little is known about the psychometric performance of the PHQ-9 across major U.S. sociodemographic groups. An advanced statistical approach called multiple-group confirmatory factor analysis (CFA) can evaluate an instrument's psychometric performance across groups through measurement invariance testing. Measurement invariance is the statistical property of measurement that signifies the same underlying construct is being measured across groups. If measurement invariance is established, it would demonstrate that PHQ-9 assesses the same construct across U.S. sociodemographic groups and that observed differences in PHQ-9 scores among these groups reflect true group differences in depressive symptoms. Thus, we could conclude that the PHQ-9 is acceptable to use in major U.S. sociodemographic groups and that the PHQ-9 allows for meaningful comparisons in depressive symptoms across these groups. However, if measurement invariance is not established, it would raise serious concerns regarding the validity and utility of the PHQ-9 for population-level screening in the U.S. Widespread use of a depression screener shown not to possess measurement invariance would likely result in under-detection or overdetection of depression in certain groups. Under-detection would lead to under-treatment of depression, which could increase sociodemographic disparities in depression care and outcomes (Kim, 2014; Simpson, Krishnan, Kunik, & Ruiz, 2007), whereas over-detection would lead to the wasting of limited treatment resources.

Although a few investigations have examined the measurement invariance of the PHO-9. these studies have been limited in two ways. One, they have not examined a consistent factor structure, with some claiming the PHQ-9 has one factor (Baas et al., 2011; Cameron, Crawford, Lawton, & Reid, 2013; Crane et al., 2010; Galenkamp, Stronks, Snijder, & Derks, 2017; González-Blanch et al., 2018; Huang, Chung, Kroenke, Delucchi, & Spitzer, 2006; Keum, Miller, & Inkelas, 2018; Merz, Malcarne, Roesch, Riley, & Sadler, 2011) and others claiming two factors (Petersen et al., 2015), making the generalizability of findings confusing. Two, they have utilized non-representative samples – namely, primary care patients (Huang et al., 2006), college students (Keum et al., 2018), people with HIV (Crane et al., 2010), Latina women (Merz et al., 2011), and non-U.S. samples (Baas et al., 2011; Cameron et al., 2013; Galenkamp et al., 2017; González-Blanch et al., 2018; Petersen et al., 2015). Consequently, it is not known which PHQ-9 factor structure provides the best fit and justification across major U.S. sociodemographic groups and whether the PHQ-9 can be used in these groups without bias. To fill these gaps, we examined a large, diverse sample representative of the U.S. adult population and used a state-of-the-art analytic approach to determine the factor structure and measurement invariance of the PHQ-9 across sex, race/ ethnicity, and education level.

#### **Methods**

#### Study Design and Sample

The continuous National Health and Nutrition Examination Survey (NHANES) is a cross-sectional, population-based study designed to assess the health and nutritional status of the U.S. population. Using stratified multistage probability sampling, NHANES enrolls a nationally representative sample of ~5,000 non-institutionalized civilians each year. Those

selected to participate are initially interviewed in their homes by trained personnel, who administer questionnaires using computer-assisted technology. One to two weeks after the household interview, respondents are asked to visit a Mobile Examination Center (MEC) to complete additional interviews, examinations, and laboratory assessments. See the NHANES website (<a href="https://www.cdc.gov/nchs/nhanes.htm">www.cdc.gov/nchs/nhanes.htm</a>) for further details.

We examined NHANES data from all survey years in which the PHQ-9 was administered (2005–2016). From the total sample (N=60,936), we selected all respondents aged 18+ years who had complete PHQ-9 data (n = 31,366; Table 1). This sample was used to determine the PHQ-9 factor structure and measurement invariance across sex. For our analyses across race/ethnicity, our sample was 30,179, as we excluded the 1,187 respondents in the other/multiracial group (because this highly heterogeneous group would cloud result interpretation). For our analyses across education level, our sample was 31,344, as we excluded the 22 respondents with missing data. The study was approved by the local institutional review board.

#### **Measures**

**Depressive Symptoms**—The PHQ-9 was administered during the face-to-face MEC interview to assess depressive symptoms over the last two weeks (Kroenke et al., 2001). Respondents indicated, on a 0–3 scale, the frequency with which they experienced the following symptoms: (1) anhedonia, (2) depressed mood, (3) sleep disturbance, (4) fatigue, (5) appetite changes, (6) low self-esteem, (7) concentration problems, (8) psychomotor disturbances, and (9) suicidal ideation. Total scores range from 0 to 27, with scores 10 representing clinically significant depressive symptoms (Kroenke & Spitzer, 2002). Furthermore, the PHQ-9 is validated as a depressive symptom severity measure (total score 1–4: minimal depression, 5–9: mild depression, 10–14: moderate depression, 15–19: moderately severe depression, and 20–27: severe depression; Kroenke et al., 2001). The PHQ-9 demonstrates high internal consistency and good sensitivity and specificity for identifying cases of MDD (Hirschtritt & Kroenke, 2017; Kroenke et al., 2001; Mitchell, Yadegarfar, Gill, & Stubbs, 2016; Moriarty, Gilbody, McMillan, & Manea, 2015; Zuithoff et al., 2010).

Sociodemographic Factors—Sex, race/ethnicity, and education level data were collected by NHANES personnel during the household interview. Sex was coded by NHANES personnel as either male or female. Race/ethnicity was assessed by two questions: (1) "Do you consider yourself to be Hispanic, Latino, or Spanish origin?", and (2) "What race do you consider yourself to be?" Using responses to these questions, NHANES personnel classified respondents into five groups (non-Hispanic White, non-Hispanic Black, Mexican American, other Hispanic, other/multi-racial) for the 2005–2010 years and six groups (non-Hispanic Asian was added) for the 2011–2016 years.

Education level was assessed by the question: "What is the highest grade or level of school you have completed or the highest degree you have received?" Using responses to this question, NHANES personnel classified respondents into the following groups: those aged 20+ years – less than 9<sup>th</sup> grade, 9<sup>th</sup>–12<sup>th</sup> grade with no diploma, high school graduate/GED

or equivalent, some college or associate degree, or college graduate or above; those aged 18–19 years – never attended/kindergarten only, grade level ranging from 1<sup>st</sup>-12<sup>th</sup> grade with no diploma, high school graduate, GED or equivalent, or more than high school. We used the categories for respondents aged 20+ years to reclassify respondents aged 18–19 years.

#### **Data Analysis**

We performed CFAs using MPlus software (Muthén & Muthén, 2015). Consistent with current recommendations (Rhemtulla, Brosseau-Liard, & Savalei, 2012), we used the means and variance adjusted weighted least squares (WLSMV) estimation method due to the ordinal scale. To determine the factor structure, we conducted five single-group CFAs on our full sample (n = 31,366), each of which examined a plausible model that has received support (Model 1, Cameron, Crawford, Lawton, & Reid, 2008; Dum, Pickren, Sobell, & Sobell, 2008; Galenkamp et al., 2017; González-Blanch et al., 2018; Huang et al., 2006; Keum et al., 2018; Kocalevent, Hinz, & Brähler, 2013; Model 2, Chilcot et al., 2013; Elhai et al., 2012; Krause, Bombardier, & Carter, 2008; Petersen et al., 2015; Model 3, Baas et al., 2011; Petersen et al., 2015; Model 4, Elhai et al., 2012; Petersen et al., 2015; Model 5 Kalpakjian et al., 2009; Krause, Reed, & McArdle, 2010; Richardson & Richards, 2008; see Table 2). In justifying our baseline model, we considered fit indices (root mean square error of approximation [RMSEA], Tucker-Lewis index [TLI], and comparative fit index [CFI]) and current depression theory. To assess model fit, we used the following guidelines: For absolute fit indices (RMSEA), exact fit = 0.00, close fit = 0.01–0.05, acceptable fit = 0.05– 0.08, mediocre fit = 0.08-0.10, and poor fit = greater than 0.10; for relative fit indices (TLI and CFI), exact fit = 1.00, close fit = 0.95-0.99, acceptable fit = 0.90-0.95, mediocre fit = 0.85-0.90, and poor fit = less than 0.85 (Hu & Bentler, 1999).

To determine the PHQ-9's measurement invariance, we carried out the four steps described by Gregorich (2006) and Sass (2014) using single and multiple-group CFAs. In these steps, four models were tested sequentially, each representing a specific level of measurement invariance, going from the least to most restrictive level of invariance. First, we evaluated dimensional invariance (equivalence in the number of latent factors across groups) by separately fitting the selected baseline model to each sex, race/ethnicity, and education level group using a single-group CFA approach. Second, we evaluated configural invariance (the latent factors are indicated by the same items across groups) by simultaneously fitting the selected model to the groups within each sociodemographic factor (e.g., the model was simultaneously fit to men and women) using a multiple-group CFA approach. Third, we evaluated scalar invariance (equivalence in the meaning of the latent factors and in potential item response biases unrelated to the latent factors across groups) by imposing equality constraints on the factor loadings (correlation between the item and factor) and item thresholds (the ordinal equivalent of an item intercept) of the configural invariance model. Fourth, we evaluated strict invariance (equivalence in the item error estimates unexplained by the latent factors across groups) by further imposing equality constraints on item residual variances of the scalar invariance model.

Consistent with current recommendations, we used a < -0.004 change in CFI ( CFI), to determine whether measurement invariance held at each step (Rutkowski & Svetina, 2017).

If CFI was <-0.004 from one step to the next (e.g., from configural to scalar invariance), indicating that adding more equality constraints did not substantially decrease model fit, we concluded that the latter model (e.g., scalar invariance) is not significantly worse. Conversely, if CFI was -0.004, we selected the former model (e.g., configural invariance). As is also recommended, we used RMSEA <0.010, in conjunction with CFI, for invariance testing (Rutkowski & Svetina, 2017).

#### Results

#### **Depressive Symptoms and Sociodemographic Factors**

The mean PHQ-9 total score was 3.20 (SD=4.27), falling in the minimal depression range. Even so, 9% of respondents had a PHQ-9 total score 10, which is indicative of clinically significant depressive symptoms (Kroenke & Spitzer, 2002). As is presented in Table 1, the mean PHQ-9 total score for each group (2.23-3.92) also fell in the minimal depression range, and the percentage of respondents with clinically significant depressive symptoms ranged from 3.8-13.2%. The mean age was 47.5 years (SD=18.8). About half of the sample were women and non-White, and there was good representation across the education levels (see Table 1). Finally, all groups demonstrated a similar correlation between the somatic and cognitive/affective constructs.

#### **PHQ-9 Factor Structure**

We conducted single-group CFAs, assessing the factor structure for the PHQ-9. All five models demonstrated close model-data fit, as the RMSEAs fell within the 0.01–0.05 range and the TLIs and CFIs fell within the 0.95–0.99 range (Table 2). Based on the fit indices alone, all models were plausible; however, we advocate for a two-factor model. Because depression is a multifaceted disorder, the high correlation of the factors and the items most likely suggest a higher order factor of depression (Byrne, 2005), rather than a more simplistic one-factor solution. In addition, a two-factor model has research utility. Specifically, studies examining depressive symptoms clusters have found that they are differentially associated with various health-related outcomes (Case & Stewart, 2014; Holzapfel et al., 2008; Roest et al., 2013; Roest et al., 2011; Smolderen et al., 2009; Vrany, Berntson, Khambaty, & Stewart, 2015).

Because all two-factor models had similar model-data fit, we turned to current depression theory to guide our selection of the best model. Models 2–5 differ with respect to on which factor the psychomotor disturbances and concentration difficulties items load. Concentration difficulties refer to depression-related issues with attention (a domain of cognitive functioning) and are thus conceptualized as a cognitive symptom of depression (De Jonge, Mangano, & Whooley, 2007; Duivis, Vogelzangs, Kupper, de Jonge, & Penninx, 2013; Hoen et al., 2010; McIntyre et al., 2015). For this theoretical reason, we conclude it is more reasonable that the concentration difficulties item fall under the cognitive/affective factor, which rules out Model 4. Model 5, in which the psychomotor disturbances item loads on both factors, revealed that this item was a stronger indicator of the cognitive/affective factor (factor loading = 0.6) than the somatic factor (factor loading = 0.2). This finding is consistent with current views on the neurobiological underpinnings of depression-related

psychomotor disturbances (Drevets, 2001). Specifically, psychomotor disturbances are thought to arise from dysfunction in the same brain reward pathways that are thought to underlie anhedonia (Drevets, 2001), an affective symptom of depression (Treadway & Zald, 2011). Thus, the psychomotor disturbances item theoretically justified under the PHQ-9 cognitive/affective factor, ruling out Models 3 and 5. Therefore, we selected Model 2 (see Figure 1), which had factor loadings above 0.70, for subsequent testing.

#### PHQ-9 Measurement Invariance across Sex, Race/Ethnicity, and Education Level

To evaluate *dimensional invariance*, we separately fit Model 2 to each group using a single-group CFA approach. All models demonstrated close fit (Table 3); therefore, this model served as a baseline for subsequent invariance testing.

To evaluate *configural invariance*, we simultaneously fit Model 2 to the groups within each sociodemographic factor by running three multiple-group CFAs. The models for sex, race/ethnicity, and education level all demonstrated close fit (Table 4), and the factor loadings across the groups were similar (Table 5). These results demonstrate that the number of factors are indicated by the same pattern of item loadings across groups, meaning that Model 2 exists across sex, race/ethnicity, and education level groups.

To evaluate *scalar invariance*, we further equated the factor loadings and item thresholds across sociodemographic groups from the configural model. The models for sex, race/ethnicity, and education level all demonstrated close fit (Table 4). The CFI (<-0.004) and RMSEA (<0.010) criteria were met for all three models, signifying that scalar invariance was established. These results demonstrate that there is equivalence in both the meaning of the latent factors and the systematic influences on item responses unrelated to the latent factors across groups.

To evaluate *strict invariance*, we equated the error variances of the three multiple group CFAs from the scalar model. Yet again, the models of sex, race/ethnicity, and education level all demonstrated close fit (Table 4). The CFI (< -0.004) and RMSEA (< 0.010) criteria were met for all three models, indicating that strict invariance was established. These results demonstrate that there is equivalence in the item error variances across groups.

Altogether, measurement invariance tests yield three conclusions. One, the PHQ-9 cognitive/ affective and somatic factors as specified in Figure 1 carry the same meaning across sex, race/ethnicity, and education level groups in U.S. adults. Two, it is defensible to compare PHQ-9 observed means and variances/covariances across sex, race/ethnicity, and education level groups (Gregorich, 2006). In other words, the PHQ-9 allows for meaningful comparisons in depressive symptoms across these major U.S. sociodemographic groups with minimal risk of bias. Three, our use of a two-factor solution demonstrates that it is defensible to compare PHQ-9 cognitive/affective and somatic subscale scores across these groups.

## **Discussion**

Our study examined the factor structure and measurement invariance of the PHQ-9 across major U.S. sociodemographic groups. First, we justified a two-factor solution for the PHQ-9 consisting of cognitive/affective and somatic factors. A subscale score for the *cognitive/affective factor* can be computed by summing the responses to items 1 (anhedonia), 2 (depressed mood), 6 (low self-esteem), 7 (concentration difficulties), 8 (psychomotor disturbances), and 9 (suicidal ideation). A subscale score for the *somatic factor* can be computed by summing items 3 (sleep disturbance), 4 (fatigue), and 5 (appetite changes). Future studies should consider standardizing subscale scores (i.e., *z* scoring) when comparing them.

Our two-factor solution differs from previous studies advocating for a one-factor solution (Dum et al., 2008; Galenkamp et al., 2017; González-Blanch et al., 2018; Huang et al., 2006; Keum et al., 2018; Kocalevent et al., 2013). Three of these studies (Dum et al., 2008; Huang et al., 2006; Kocalevent et al., 2013) used principal component analysis, which is mathematically inappropriate (Brown, 2014). The studies that used CFA did not consider the research utility of a two-factor solution. Specifically, investigations using the PHQ-9 have found that depressive symptom clusters are differentially associated with various health-related outcomes (Case & Stewart, 2014; Holzapfel et al., 2008; Roest et al., 2013; Roest et al., 2011; Smolderen et al., 2009; Vrany et al., 2015). Consistent with CFA recommendations (Brown, 2014), our study takes into consideration empirical evidence, current theory, and practical utility. Nonetheless, the PHQ-9 total score currently has high clinical utility, and future research is needed to compare the utility of the total and subscale scores in different clinical scenarios (e.g., screening versus monitoring).

Second, we established strict measurement invariance for the PHQ-9 across major U.S. sociodemographic groups. Therefore, the observed differences in the PHQ-9 total, cognitive/affective, and somatic scores among these groups reflect true group differences in these depressive symptoms. The findings we report support the conclusions that (a) the PHQ-9 is acceptable to use in major sociodemographic groups in the U.S. and that (b) the PHQ-9 allows for meaningful comparisons in total, cognitive/affective, and somatic depressive symptoms across these groups with minimal risk of bias. Although we tested measurement invariance using a two-factor solution, it is still appropriate to compute and use the single PHQ-9 total score for research, clinical, or other purposes across the examined groups, as the high correlation among the factors and items implies a higher order factor of depression (Byrne, 2005). However, it may be useful in future research to explicitly test this assumption.

As with all studies, ours is not without limitations. First, although we use a large U.S. representative sample, there were missing values which could limit the generalizability of findings. However, missingness in the sample was below the 10% guideline (Little, Jorgensen, Lang, & Moore, 2013). Second, due to the cross-sectional nature of the NHANES data, we were not able to assess measurement invariance overtime, as done by González-Blanch et al. (2018). Future studies are needed to determine whether measurement invariance of the selected two-factor solution for the PHQ-9 holds overtime. Third, age was

beyond the scope of this study, future investigations are needed to determine the factor structure and measurement invariance of the PHQ-9 across various categorizations for age groups.

#### Conclusion

We extend previous research validating the PHQ-9 as a depression screener and severity measure by determining its factor structure in U.S. adults and its measurement invariance across sex, race/ethnicity, and education level groups. Our findings demonstrate that the PHQ-9 is acceptable to use in major sociodemographic groups in the U.S. and allows for meaningful comparisons in total, cognitive/affective, and somatic depressive symptoms across these groups, extending the PHQ-9's use from the clinic to the community. This knowledge is especially useful and timely as medicine moves toward wider adoption of alternative payment models emphasizing high-quality and cost-efficient care, which will likely incentivize behavioral health integration and population health management. Finally, we also provide a consistent, evidence-based approach for computing PHQ-9 cognitive/ affective and somatic subscale scores, which should facilitate comparisons of results across studies and future meta-analytic efforts.

### Acknowledgments

Conflicts of Interest and Source of Funding: A portion of Dr. Stewart's time was supported by the National Heart, Lung, and Blood Institute under Award Number R01HL122245. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

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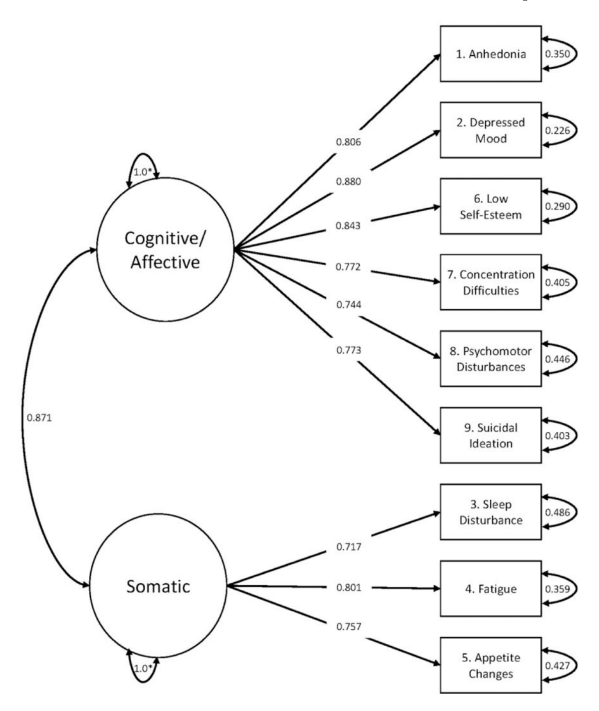


Figure 1:
Two-Factor Measurement Model of the Patient Health Questionnaire-9 (PHQ-9).On the right, the boxes represent the PHQ-9 items (indicator variables). Circular arrows that point back to the indicator variables represent item error variances. Moving to the left, unidirectional linear arrows pointing from circles to boxes represent standardized factor loadings. The circles represent latent factors. Circular arrows that point back to the latent

factors represents latent variances (fixed to 1.0 for identification purposes, as demonstrated by the asterisk). The bidirectional arrow between latent factors represents a correlation

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Table 1

Characteristics of NHANES Respondents

	Frequency (%)	PHQ-9 Total Score mean (SD)	PHQ-9 Total Score 10 (%)	PHQ-9 Cognitive/Affective Score (range: 0-18) mean (SD)	PHQ-9 Somatic Score (range 0-9) mean (SD)	Correlation between Cognitive/Affective and Somatic Scores
Sex $(n = 31,366)$						
Women	15,935 (50.8)	3.72 (4.57)	11.0	1.67 (2.80)	2.05 (2.23)	0.65
Men	15,431 (49.2)	2.67 (3.87)	6.5	1.24 (2.40)	1.43 (1.90)	0.61
Race/Ethnicity $(n = 30,179)$						
Non-Hispanic White	13,455 (42.9)	3.19 (4.19)	8.4	1.38 (2.55)	1.81 (2.09)	0.63
Non-Hispanic Black	6,793 (21.7)	3.21 (4.35)	9.1	1.48 (2.65)	1.73 (2.16)	0.63
Non-Hispanic Asian	1,745 (5.6)	2.23 (3.07)	3.3	1.00 (1.90)	1.22 (1.58)	0.55
Mexican American	5,205 (8.6)	3.15 (4.20)	8.6	1.51 (2.59)	1.64 (2.05)	0.64
Other Hispanic	2,981 (12.2)	3.80 (4.93)	12.2	1.88 (3.11)	1.92 (2.24)	0.69
Education Level $(n = 31,344)$						
Less than 9th grade	3,156 (10.1)	3.77 (5.03)	13.2	1.95 (3.17)	1.82 (2.31)	0.68
9th to 12th grade (no diploma)	5,017 (16.0)	3.92 (4.79)	12.4	1.92 (3.01)	2.00 (2.27)	0.64
High school graduate/GED equivalent	7,410 (23.6)	3.31 (4.32)	0.6	1.51 (2.65)	1.81 (2.14)	0.63
Some college or associate degree	9,101 (29.0)	3.21 (4.22)	8.6	1.41 (2.54)	1.81 (2.12)	0.64
College graduate or above	6,660 (21.2)	2.26 (3.19)	3.8	0.89 (1.88)	1.37 (1.70)	0.59

Note. The cognitive/affective score was computed as the sum of items 1, 2, 6, 7, 8, and 9, and the somatic score was computed as the sum items 3, 4, and 5.NHANES = National Health and Nutrition Examination Survey. PHQ-9 = Patient Health Quesionnaire-9

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Table 2

Single-Group Confirmatory Factor Analysis Models and Fit Indices Evaluating Factor Structure of the Patient Health Questionnaire-9 (PHQ-9) in U.S. Adults

	Model 1	Model 2	Model 3	Model 4	Model 5
1. Anhedonia	Depression	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective Cognitive/Affective	Cognitive/Affective
2. Depressed Mood	Depression	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective
3. Sleep Disturbance	Depression	Somatic	Somatic	Somatic	Somatic
4. Fatigue	Depression	Somatic	Somatic	Somatic	Somatic
5. Appetite Changes	Depression	Somatic	Somatic	Somatic	Somatic
6. Low Self-Esteem	Depression	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective
7. Concentration Difficulties	Depression	Cognitive/Affective	Cognitive/Affective	Somatic	Cognitive/Affective
8. Psychomotor Disturbances	Depression	Cognitive/Affective	Somatic	Somatic	Both
9. Suicidal Ideation	Depression	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective	Cognitive/Affective
Chi-Square ( $\chi 2$ )	1783.11	941.82	1182.37	994.99	910.98
Degrees of Freedom	27	26	26	26	25
p-value	<0.001	<0.001	<0.001	<0.001	<0.001
RMSEA	0.046	0.034	0.038	0.034	0.034
RMSEA 90% Confidence Interval	0.044 - 0.047	0.032-0.035	0.036-0.040	0.033-0.036	0.032-0.036
TLI	0.972	0.985	0.981	0.984	0.985
CFI	0.979	0.989	0.986	0.988	0.989

Note. N=31,366. For absolute fit indices (RMSEA), exact fit = 0.00, close fit = 0.01-0.05, acceptable fit = 0.05-0.08, mediocre fit = 0.08-0.10, and poor fit = greater than 0.10. For relative fit indices (TLI and CFI), exact fit = 1.00, close fit = 0.95-0.99, acceptable fit = 0.90-0.95, mediocre fit = 0.85-0.90, and poor fit = less than 0.85. RMSEA = root mean square error of approximation. TLI = Tucker-Lewis index. CFI = comparative fit index

Table 3

Confirmatory Factor Analysis Models and Fit Indices Evaluating Dimensional Invariance of the Patient Health Questionnaire-9 (PHQ-9) across Sex, Race/Ethnicity, and Education Level in U.S. Adults

		$\chi^2$	Df	p-value	RMSEA	RMSEA 90% CI	TLI	CFI
9. 21.266	Women	638.62	26	0.000	0.038	0.036-0.041	0.984	0.988
Sex $n = 31,366$	Men	427.30	26	0.000	0.032	0.029-0.034	0.986	0.990
	Non-Hispanic White	556.47	26	0.000	0.039	0.036-0.042	0.983	0.988
	Non-Hispanic Black	247.90	26	0.000	0.035	0.031-0.040	0.986	0.990
Race/Ethnicity $n = 30,179$	Non-Hispanic Asian	79.24	26	0.000	0.034	0.026-0.043	0.982	0.987
	Mexican American	177.50	26	0.000	0.033	0.029-0.038	0.987	0.990
	Other Hispanic	113.44	26	0.000	0.034	0.027-0.040	0.988	0.991
	Less than 9th grade	98.54	26	0.000	0.030	0.024-0.036	0.990	0.993
	9th to 12th grade (no diploma)	213.14	26	0.000	0.038	0.033-0.043	0.981	0.986
Education Level $n = 31,344$	High school graduate/GED equivalent	266.67	26	0.000	0.035	0.032-0.039	0.986	0.990
	Some college or associate degree	350.29	26	0.000	0.037	0.034-0.041	0.984	0.989
	College graduate or above	159.61	26	0.000	0.028	0.024-0.032	0.986	0.990

Note. We used the two-factor Model 2 shown in Figure 1. For absolute fit indices (RMSEA), exact fit = 0.00, close fit = 0.01-0.05, acceptable fit = 0.05-0.08, mediocre fit = 0.08-0.10, and poor fit = greater than 0.10. For relative fit indices (TLI and CFI), exact fit = 1.00, close fit = 0.95-0.99, acceptable fit = 0.90-0.95, mediocre fit = 0.85-0.90, and poor fit = less than 0.85. RMSEA = root mean square error of approximation. CI = confidence interval. TLI = Tucker-Lewis index. CFI = comparative fit index

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Multiple-Group Confirmatory Factor Analysis Models and Fit Indices Evaluating Measurement Invariance of the Patient Health Questionnaire-9 (PHQ-9) across Sex, Race/Ethnicity, and Education Level in U.S. Adults

Table 4

Configural 1048.15 Sex $n = 31,366$ Scalar 877.68				,	MINISER	At praise amosea ministrative of the		7.7	CFI	MAISEA
	1048.15	52	!	<0.001	0.035	0.033-0.037 0.985	0.985	0.989	!	
	89.77.88	84	117.93	<0.001	0.025	0.023-0.026	0.992	0.991	0.002	-0.010
Strict	907.43	75	54.46	<0.001	0.027	0.025-0.028	0.991	0.991	0.000	-0.008
Configural	<b>Configural</b> 1204.45 166	166	1	<0.001	0.032	0.030-0.034 0.988	0.988	0.989	1	
Race/Ethnicity $n = 30,179$ Scalar	1629.07	262	411.65	<0.001	0.029	0.028-0.031 0.990	0.990	0.986	-0.003	-0.003
Strict	<b>Strict</b> 1419.80 235	235	287.26	<0.001	0.029	0.027-0.030 0.990	0.990	0.987	0.001	-0.003
Configural	<b>Configural</b> 1092.39 130	130	1	<0.001	0.034	0.032-0.036	0.985	0.989	1	
Education Level $n = 31,344$ Scalar	1507.24	258	700.51	<0.001	0.028	0.026-0.029	0.990	0.986	-0.003	-0.006
Strict	<b>Strict</b> 1088.25 222		486.91	<0.001	0.025	0.023-0.026 0.992	0.992	0.990	0.004	-0.009

Note. We used the two-factor Model 2 shown in Figure 1. For absolute fit indices (RMSEA), exact fit = 0.00, close fit = 0.01-0.05, acceptable fit = 0.05-0.08, mediocre fit = 0.08-0.10, and poor fit = greater than 0.10. For relative fit indices (TLI and CFI), exact fit = 1.00, close fit = 0.95-0.99, acceptable fit = 0.90-0.95, mediocre fit = 0.85-0.90, and poor fit = less than 0.85. For nested model testing, a change in CFI -0.004 or change in RMSEA 0.010 signifies that measurement invariance was not established for that step. RMSEA = root mean square error of approximation. CI = confidence interval. TLI = Tucker-Lewis index. CFI = comparative fit index. Page 19

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Table 5

Factor Loadings from the Multiple-Group Confirmatory Factor Analyses Evaluating Configural Invariance

			Sex			Race/Ethnicity	ty			4	Education Level	el	
Construct	PHQ-9 Item	Men	Women	Non- Hispanic White	Non- Hispanic Black	Non- Hispanic Asian	Mexican American	Other Hispanic	Less than 9th grade	9th to 12th grade (no diploma)	High school graduate/ GED equivalent	Some college or associate degree	College graduate or above
	1. Anhedonia	0.79	0.82	0.84	0.74	0.70	0.74	0.80	0.75	0.75	7.00	0.83	0.86
	2. Depressed Mood	0.89	0.88	0.89	0.87	0.85	98.0	0.87	0.89	0.85	0.88	0.89	0.89
	6. Low Self-Esteem	0.84	0.84	0.85	0.83	0.77	0.83	0.83	0.84	0.83	0.85	0.85	0.81
Cognitive/Affective	7. Concentration Difficulties	0.75	0.79	0.77	0.78	0.71	0.75	0.83	08.0	0.73	0.79	0.77	0.75
	8. Psychomotor Disturbances	0.74	0.74	0.72	0.78	0.65	0.76	0.82	0.81	0.77	0.73	0.73	99.0
	9. Suicidal Ideation	0.79	0.76	0.77	0.81	0.68	0.74	0.79	0.70	0.73	0.78	08.0	0.80
	3. Sleep Disturbance	0.74	69:0	0.70	0.78	0.67	0.75	0.77	0.76	0.74	0.72	0.73	0.66
Somatic	4. Fatigue	0.79	0.80	0.81	0.79	0.74	0.81	0.79	08.0	0.79	0.81	0.81	0.80
	5. Appetite Changes	0.73	0.77	0.76	0.76	0.72	0.72	0.76	0.74	0.75	0.75	0.77	0.73