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Comparing sensitivity to change using the 6-item versus the 17-item Hamilton depression rating scale in the GUIDED randomized controlled trial

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Abstract

Background: Previous research suggests that the 17-item Hamilton Depression Rating Scale (HAM-D17) is less sensitive in detecting differences between active treatment and placebo for major depressive disorder (MDD) than is the HAM-D6 scale, which focuses on six core depression symptoms. Whether HAM-D6 shows greater sensitivity when comparing two active MDD treatment arms is unknown.

Methods: This post hoc analysis used data from the intent-to-treat (ITT) cohort (N = 1541) of the Genomics Used to Improve DEpression Decisions (GUIDED) trial, a rater- and patient-blinded randomized controlled trial. GUIDED compared combinatorial pharmacogenomics-guided care with treatment as usual (TAU) in patients with MDD. Percent of symptom improvement, response rate and remission rate from baseline to week 8 were evaluated using both scales. Analyses were performed for the full cohort and for the subset of patients who at baseline were taking medications predicted by the test to have moderate or significant gene-drug interactions. A Mokken scale analysis was conducted to compare the homogeneity of HAM-D17 with that of HAM-D6.

Results: At week 8, the guided-care arm demonstrated statistically significant benefit over TAU when the HAM-D6 (Δ = 4.4%, p = 0.023) was used as the continuous measure of symptom improvement, but not when using the HAM-D17 (Δ = 3.2%, p = 0.069). Response rates increased significantly for guided-care compared with TAU when evaluated using both HAM-D6 (Δ = 7.0%, p = 0.004) and HAM-D17 (Δ = 6.3%, p = 0.007). Remission rates also were significantly greater for guided-care versus TAU using both measures (HAM-D6 Δ = 4.6%, p = 0.031; HAM-D17 Δ = 5.5%, p = 0.005). Patients in the guided-care arm who at baseline were taking medications predicted to have gene-drug interactions showed further increased benefit over TAU at week 8 for symptom improvement (Δ = 7.3%, p = 0.004) response (Δ = 10.0%, p = 0.001) and remission (Δ = 7.9%, p = 0.005) using HAM-D6. All outcomes showed continued improvement through week 24. Mokken scale analysis demonstrated the homogeneity and unidimensionality of HAM-D6, but not of HAM-D17, across treatment arms.

Conclusions: The HAM-D6 scale identified a statistically significant difference in symptom improvement between combinatorial pharmacogenomics-guided care and TAU, whereas the HAM-D17 did not. The demonstrated utility of pharmacogenomics-guided treatment over TAU as detected by the HAM-D6 highlights its value for future biomarker-guided trials comparing active treatment arms.

(Continued on next page)
Background
Roughly half of patients with major depressive disorder (MDD) fail to respond to treatment with an antidepressant medication, and approximately two-thirds fail to achieve remission [1]. These inadequate outcomes have sparked great interest in exploring biological subtypes of depression that correlate with variability in medication response [2]. Pairing clearly defined subtypes with validated biomarkers such as genetic and epigenetic, proteomic, metabonomic, inflammation, neuroimaging, and electroencephalography measures might enable more precise treatment selection and response monitoring.

Genetic variation is an important biological contributor to both MDD development [3, 4] and to treatment response [5, 6]. On their own, individual gene variants explain little of the variance in disease risk or outcomes; rather, clinical manifestation of MDD and treatment response appear to result from the combined effects of many genes, along with other clinical and environmental factors. Combinatorial pharmacogenomic tests, which evaluate the weighted effects of genetic variants to predict which medications may be impacted by gene-drug interactions, hold promise for aiding patient-specific treatment selection [7]. Recently, the Genomics Used to Improve DEpression Decisions (GUIDED) randomized controlled trial (RCT) reported on the efficacy of using a combinatorial pharmacogenomic test in medication selection (guided-care), compared with treatment as usual (TAU), for patients with treatment non-responsive MDD [8]. This trial differed from traditional drug studies in that patients in both arms received active treatment. GUIDED approached but did not achieve a statistically significant difference between guided-care versus TAU for its primary outcome, percent symptom improvement at week 8 ($p = 0.069$; intent-to-treat [ITT] cohort), as assessed by the Hamilton Depression Rating Scale, 17-item (HAM-D17). However, significantly more patients achieved the secondary outcomes, response ($p = 0.007$) and remission ($p = 0.005$) at week 8, measured using HAM-D17, when they received pharmacogenomics-guided care.

The results observed in the GUIDED trial highlight the challenges in detecting clinically and statistically significant differences in randomized trials when patients in all study arms receive active treatment. This is especially true in psychiatry, where several well-powered randomized trials comparing active MDD treatments have failed to show differences in efficacy, including the Sequenced Treatment Alternatives in Depression (STAR*D) trial [9], the Genome-Based Therapeutic Drugs for Depression (GENDEP) trial [10], and the Combining Medications to Enhance Depression Outcomes COMED trial [11]. Lack of significant differences in efficacy extends even to large trials that compare psychotherapy, antidepressant medications, or their combination [12, 13]. Such equivalent outcomes, despite the treatments’ distinct mechanisms, raise the possibility that the assessment metrics used are flawed [14].

The Hamilton Depression Rating Scale (HAM-D) is the most widely used outcome measure in MDD clinical trials, with the 17-item version (HAM-D17) originally published in 1960, serving as the standard [15, 16]. Over the past four decades, however, researchers have raised concerns about the ability of the HAM-D17 scale to assess accurately the severity of and change in depression symptoms [17–19]. Factor analyses of HAM-D17 have determined that the scale is not a unidimensional measure of depression severity but rather consists of two to eight symptom factors [20]. Although multidimensionality in a scale is useful for detecting a broad array of clinical features, a multidimensional (or multifactorial) scale may reduce the ability to detect change over time, because some factors may not adequately distinguish groups when valid differences exist [21]. The ability to scale appropriately with illness severity is a fundamental aspect of construct validity. Medication side effects may affect some factors on multidimensional scales more than others, potentially producing total score changes that do not align with changes in core depressive symptoms [22, 23]. In studies such as GUIDED that allow concomitant treatments (e.g., sedative hypnotics for insomnia and anxiety in conjunction with antidepressant medication), assessing efficacy with HAMD-17 becomes even more problematic, as the uncontrolled additional medications can result in score changes unrelated to antidepressant treatment.

To address these shortcomings, researchers developed abbreviated, more focused versions of HAM-D17 [24]. Of these, the most widely used is the six-item subscale of HAM-D17, known as the HAM-D6 or melancholia subscale [23, 25]. The HAM-D6 scale is specific to the core depressive symptoms of depressed mood, guilt, work and activities, psychomotor retardation, psychic anxiety, and general somatic symptoms (energy and physical pain), and...
it is unidimensional [26]. HAM-D17 symptoms omitted from the HAM-D6 scale include suicidal thoughts, initial insomnia, middle insomnia, late insomnia, psychomotor agitation, somatic anxiety, gastrointestinal symptoms [appetite], sexual disturbances, hypochondriasis [somatization], insight, and weight disturbances. The HAM-D6 scale correlates better with the Clinical Global Impressions Scale-Severity than does the HAM-D17 scale, particularly among more severely ill patients [21]. It has repeatedly demonstrated greater effect sizes for second-generation antidepressants than has HAM-D17, as well as similar effect sizes for medications that have sedating side effects, such as TCAs and mirtazapine [27–29].

This post hoc analysis of GUIDED trial data evaluated whether the HAM-D6 scale showed significant differences in outcomes between patients whose treatment was guided by combinatorial pharmacogenomic testing versus TAU. We hypothesized that the more sensitive and unidimensional HAM-D6 would detect a statistically significant difference in symptom improvement between the guided-care and TAU arms, whereas the difference approached but did not achieve significance ($p = 0.069$) using HAM-D17. We also examined whether the statistically significantly higher rates of response and remission observed using the HAM-D17 scale would be replicated using HAM-D6.

Methods

Pharmacogenomic testing
All enrolled patients were tested with a combinatorial pharmacogenomic test (GeneSight Psychotropic, Assurex Health, Inc., now Myriad Neuroscience, Mason, OH). At the time of the study, the test evaluated genotypes for 59 alleles and variants across eight genes (CYP1A2, CYP2C9, CYP2C19, CYP3A4, CYP2B6, CYP2D6, HTR2A, and SLC6A4) [30]. Using a proprietary algorithm that weighted the combined influences of individual genotypes on each of 38 medications, a report was generated that categorized the medications into three levels of gene-drug interaction: ‘use as directed’ (no detected gene-drug interactions); ‘use with caution’ (moderate gene-drug interactions, i.e., medications may be effective with dose modification); and ‘use with increased caution and with more frequent monitoring’ (significant gene-drug interactions that may significantly impact drug safety and/or efficacy) [31].

Study description
The GUIDED trial was a 24-week blinded, randomized, controlled trial that evaluated the utility of combinatorial pharmacogenomic testing in medication selection (guided-care) compared with TAU for adults with MDD. Unlike traditional drug studies, patients in both study arms received active treatment. The study was performed in primary care and psychiatry specialty clinics across 60 U.S. community and academic sites.

Patients and raters were blinded to study arm. Physicians in TAU were blinded to pharmacogenomic test results. The study protocol was approved by the Copernicus Group independent review board (INC1-14-012) and conducted in accordance with the principles of the Declaration of Helsinki and its amendments. All patients provided written informed consent for participation. Detailed methods and primary analyses for the GUIDED trial have been described previously [8]. Methods relevant to the current analysis are summarized here.

Prior to the baseline visit, patients were randomized 1:1 to the guided-care or TAU arm. Active treatment was provided to patients in both arms, with medications selected based on clinician judgment, informed by the pharmacogenomic test report for the guided-care arm, and “standard” clinician judgment in the non-guided arm. Clinicians for patients in the guided-care arm were not required to adhere to the test results in making medication decisions, and no medications were prohibited.

Patient assessments were performed at week 0 (baseline) and at the end of weeks 4, 8, 12 and 24. Patients and raters in both arms were blinded to study arm and pharmacogenomic test results. Clinicians for patients in the TAU arm were blinded to test results until after completion of the week 8 visit. Blinding of patients, sites, and physicians was maintained through week 8. Sites were instructed to unblind patients to their randomization assignment following the week 12 assessment. Because patient unblinding may have occurred before week 12 assessments were performed, however, only data collected through the week 8 assessment were considered blinded.

Participants
Patients were enrolled if they were diagnosed with DSM-IV-TR-defined MDD, confirmed by both the self-rated and site-rated 16-item Quick Inventory of Depressive Symptomology (QIDS-SR16 and QIDS-C16 $\geq 11$) at screening and baseline, and if they reported an inadequate response within the current depressive episode to at least one medication included on the pharmacogenomic test report. Key exclusion criteria included significant short-term suicide risk, bipolar disorder, current delirium or neurocognitive disorder, psychotic disorder or psychotic symptoms during the current or a previous depressive episode, a current substance use disorder, or a significant unstable medical condition.

Statistical analysis
Analyses described herein were conducted using the ITT cohort, which included all patients who met eligibility criteria, were randomized to a study arm, and had at least one post-baseline visit. Outcomes analyses were
performed for the ITT cohort and separately for the subset of patients who at baseline were taking medications predicted to have moderate or significant gene-drug interactions (those in the ‘use with caution’ and ‘use with increased caution and more frequent monitoring’ report categories). This subset excluded patients who were taking only medications in the ‘use as directed’ category.

The protocol-defined primary efficacy measure for GUIDED was the HAM-D17 scale, administered by blinded central raters (MedAvante-ProPhase Inc., Hamilton, NJ). For this post hoc scale comparison, HAM-D6 scores were derived from the HAM-D17 assessments. These included: item 1, depressed mood; item 2, guilt feelings; item 7, work and activities; item 8, psychomotor retardation; item 10, psychic anxiety; and item 13, general somatic symptoms. Items 1, 2, 7, 8, and 10 each were scored from 0 to 4, and item 13 was scored from 0 to 2, for a maximum possible HAM-D6 score of 22. For HAM-D17, the maximum possible score was 52.

The primary endpoint was percent symptom improvement from baseline to week 8, and secondary endpoints were response and remission rates at week 8. Response was defined as a ≥50% decrease in score at week 8 from baseline and was assessed for both HAM-D17 and HAM-D6. Remission was defined as having a score of ≤7 for HAM-D17 [32] and ≤4 for HAM-D6 [21, 33]. The durability of pharmacogenomic testing utility was evaluated in the guided-care arm through outcome assessments at weeks 4, 8, 12, and 24.

Identical statistical methods were used for the primary HAM-D17 analyses and the post hoc HAM-D6 analyses. A mixed model for repeated measures was used to assess percent change in symptoms from baseline to week 8. A generalized linear mixed model was used for separate analyses of response and remission at week 8. Because the response and remission outcomes were measured at both week 4 and week 8, a generalized linear mixed model (GLMM) was used to account for both within-subject and between-subject variability over time. Both the mixed model for repeated measures and the GLMM included treatment, week, treatment-by-week interaction, baseline HAM-D6 score, and baseline HAM-D6 score-by-week interaction as fixed effects. Binomial distribution with a log-link function was used for the GLMM model. The pairwise comparisons between the two treatment arms at week 8 were tested at a significance level of 0.05 (2-sided). Missing values were handled using maximum likelihood method via mixed models for repeated measures for both symptom improvement and via generalized linear mixed model for categorical variables – response and remission. Analyses were performed with SAS software (version 9.4) or JMP 14 (SAS Institute).

An analysis of scalability was performed using the non-parametric item response theory model developed by Mokken [34]. Using this framework, the deviation of either the HAM-D17 scale or the HAM-D6 scale from a perfectly homogeneous structure was expressed using Loevinger’s scalability coefficient (H) [35], a measure of the extent to which the scale items represented a single dimension. Loevinger’s coefficient was interpreted as follows: ≥0.5, strong scale homogeneity; 0.40–0.49, moderate but acceptable homogeneity; 0.30–0.39, doubtful homogeneity; <0.30, no homogeneity.

Results
Cohort description
At baseline, the ITT cohort included 1541 patients (guided-care, n = 760; TAU, n = 781). Baseline clinical characteristics of the cohort are presented in Table 1. There were no meaningful differences between the two treatment arms in depression characteristics, HAM-D17 scores or HAM-D6 scores at baseline. At the week 8 time point, the ITT cohort included 1298 patients (guided-care, n = 621; TAU, n = 677).

Symptom improvement, response and remission: HAM-D6 versus HAM-D17
At week 8, there was a 28.3% decrease in HAM-D6 scores from baseline in the guided-care arm, compared with a 23.9% decrease in the TAU arm (Fig. 1). This difference in mean percent symptom improvement between arms was statistically significant (Δ = 4.4%, p = 0.023) compared to that reported previously using the HAM-D17 scale (Δ = 3.2%, p = 0.069). The response rate at 8 weeks among patients in the guided-care arm (29.6%) similarly showed a significant increase over TAU (22.5%) using HAM-D6 (Δ = 7.0%, p = 0.004) (Fig. 1). The percent difference between study arms was also statistically significant for HAM-D17 (Δ = 6.3%, p = 0.007). Remission rates at week 8 favored pharmacogenomics-guided treatment (20.8%) versus TAU (16.2%) at week 8 using HAM-D6 (Fig. 1), and the percent difference between study arms was statistically significant for HAM-D6 (Δ = 4.6%, p = 0.031). The remission rate in the guided-care versus TAU arms was significant using the HAM-D17 scale (Δ = 5.4%, p = 0.005). Overall, the results for response rate and remission rate were similar for both scales.

Patients entering on medications with predicted gene-drug interactions
To examine the impact of guided-care versus TAU more specifically for patients who stand to benefit most from pharmacogenomic testing, HAM-D6 outcomes were assessed in the subset of patients who at baseline were prescribed medications predicted by the patient’s test results to have gene-drug interactions (Fig. 2). At week 8, the mean percent symptom improvement in the guided-care arm (28.6%) was significantly greater than that measured in TAU (21.3%) (Δ = 7.3%, p = 0.004). Response rate in the
guided-care arm (29.5%) also was significantly improved over TAU (19.5%) ($\Delta = 10.0\%$, $p = 0.001$). Finally, remission rate was improved for guided-care (22.2%) versus TAU (14.3%) in these patients ($\Delta = 7.9\%$, $p = 0.005$). Compared with the outcomes assessed using the HAM-D17 scale in this subset of patients (Fig. 2) [36], the HAM-D6 scale showed equal or greater sensitivity to detect differences between guided-care and TAU for all three depression outcomes. In addition, the percent differences between guided-care and TAU across all three outcomes were substantially higher in patients predicted to be most impacted by gene-drug interactions than were those observed in the full patient cohort using either HAM-D17 or HAM-D6 (Fig. 1).

**Scale homogeneity**

To assess the dimensionality of the HAM-D17 and HAM-D6 assessments in the GUIDED ITT cohort, a Mokken scale analysis was performed. Table 2 shows the Loevinger’s coefficient of homogeneity (H) at week 8 for each assessment scale. For the combined treatment arms, HAM-D17 had a coefficient of 0.30, indicating that the scale is heterogeneous and multidimensional. In contrast, HAM-D6 had a coefficient of 0.53 for the combined arms, indicating that the scale is homogeneous and unidimensional. Similar results were observed for individual treatment arms.

**Durability of response**

To evaluate the durability of the guided-care treatment results, patient HAM-D6 scores in the guided-care arm were evaluated at time points extending through the end of the 24-week trial period (Fig. 3). Consistent increases were observed for all three measured outcomes from baseline through weeks 4, 8, 12, and 24.

**Discussion**

This comparative, post hoc analysis of the HAM-D6 and HAM-D17 depression scales in the GUIDED trial for MDD treatment found greater sensitivity to differences in treatment effects with the abbreviated version of the scale. This result likely is due to the narrower focus of the HAM-D6 scale, as compared with HAM-D17, for the core symptoms of depression. Furthermore, although both versions of the scale achieved statistically significant differences for the response and remission outcomes, the greater differences for symptom improvement seen with the HAM-D6 scale suggest that HAM-D6 provided a more precise measure for MDD outcome assessment. This is supported further by the observation of the high sensitivity of the HAM-D6 scale in the subset of patients who entered the trial on medications predicted by the pharmacogenomic test to have gene-drug interactions. Mokken scale analysis further supported the increased homogeneity of HAM-D6 relative to HAM-D17. Altogether, these

### Table 1 Clinical features of the GUIDED intent-to-treat study population at baseline (week 0)

| Characteristic                          | Treatment | Guided-Care (N = 760) | Total (N = 1541) |
|----------------------------------------|-----------|-----------------------|------------------|
|                                        | TAU (N = 781) |                       |                  |
| HAM-D17                                |           |                       |                  |
| Mean (SD)                              | 20.74 (4.86) | 20.49 (4.84)          | 20.62 (4.85)     |
| Min, Max                               | 6.0, 35.0 | 4.0, 37.0             | 4.0, 37.0        |
| Depression Category, n (%)             |           |                       |                  |
| None (0–7)                             | 5 (0.6) | 6 (0.8)                | 11 (0.7)         |
| Mild (8–13)                            | 46 (5.9) | 45 (5.9)              | 91 (5.9)         |
| Moderate (14–18)                       | 189 (24.2) | 209 (27.5)            | 398 (25.8)       |
| Severe (19–22)                         | 270 (34.6) | 241 (31.7)            | 511 (33.2)       |
| Very Severe (≥ 23)                     | 271 (34.7) | 259 (34.1)            | 530 (34.4)       |
| HAM-D6                                 |           |                       |                  |
| Mean (SD)                              | 10.90 (2.16) | 10.99 (2.11)          | 10.94 (2.13)     |
| Min, Max                               | 1.0, 17.0 | 3.0, 18.0             | 1.0, 18.0        |
| Failed Medication Trials               |           |                       |                  |
| Mean (SD)                              | 3.54 (3.01) | 3.45 (3.02)           | 3.49 (3.01)      |
| Min, Max                               | 1, 34 | 1, 25                  | 1, 34            |
| Psychiatric Comorbidities, n (%)       |           |                       |                  |
| General anxiety disorder               | 104 (13.4) | 127 (16.7)            | 231 (15.0)       |
| Panic disorders/social phobia          | 112 (14.4) | 119 (15.7)            | 231 (15.0)       |
| Post-traumatic stress disorder         | 35 (4.5) | 41 (5.4)              | 76 (4.9)         |

SD standard deviation, TAU treatment as usual
results mirror those seen in many placebo-controlled pharmacological trials, wherein the HAM-D17 scale failed to identify an antidepressant effect, while the HAM-D6 scale did [27].

Although the percent differences in response and remission rates were generally similar for HAM-D17 and HAM-D6, the slightly lower difference in remission rate between arms (0.8%) as assessed by the HAM-D6 (Δ = 4.6%) versus the HAM-D17 (Δ = 5.4%) is of interest. A concern in the field is that the standard HAM-D17 remission threshold of ≤7 may be high, capturing many patients who continue to experience impairment or distress from persisting symptoms [35, 36]. Thus, low levels of core symptoms, as determined by the standard HAM-D6 remission threshold (≤4), might comprise a more valid measure for defining the state of clinical remission.

The question of whether the HAM-D6 or the HAM-D17 remission threshold better predicts restoration of function and long-term wellness should be a focus of future work.

The importance of maximizing signal detection through use of the most sensitive scale to detect treatment effects is of particularly great importance for comparative effectiveness studies and for biomarker-based clinical trials, both of which provide active treatment to all patients [37]. Over the past several decades, adequately powered MDD trials comparing active treatments, be they medications or psychotherapies, have found no difference between treatment arms [9–11]. Notably, all of these large trials have used either the HAM-D17 scale, the Montgomery-Åsberg Depression Rating Scale (MADRS), or the Quick Inventory of Depressive Symptomatology-Self-Report (QIDS-SR) as the efficacy measure, each of which contains numerous items unrelated to core depression symptoms captured by the HAM-D6 scale. Trials such as GUIDED that allow concomitant medications for specific symptoms, such as sedative hypnotics for anxiety or insomnia, can further diminish the ability to identify a difference between treatment arms when the outcome measure includes non-core depressive symptoms [38]. Consequently, future randomized trials applying biomarker-based approaches to treatment selection...
for MDD may benefit from using the HAM-D6 or a similar, more focused symptom scale.

The greater discriminative ability of the HAM-D6 scale also allows for smaller sample sizes to test hypotheses about efficacy [39–41]. Given the greater precision and numerous advantages of HAM-D6, it is difficult to justify continued use of the full HAM-D17 scale as the sole primary outcome measure in MDD treatment trials. In the future, HAM-D17 could be used to enable historic comparisons of baseline severity among trials, but new study protocols should consider specifying the HAM-D6 or a similarly more precise assessment of core symptoms [42, 43] as the primary efficacy variable for analysis. Administering the shorter version can have the added benefit of reducing time burdens on clinical trial participants.

This analysis had several strengths that were inherent to the GUIDED primary analysis. First, the diversity of the study cohort mirrors that seen across varied clinical scenarios for MDD treatment, including clinicians in both psychiatric specialty and primary care clinics. Second, the study’s two active treatment arms reflect real-world clinical practice and provide a relevant evaluation of clinical utility. The limitations of the primary GUIDED analysis also apply to this study [8]. Specifically, the treating clinician was not blinded to study arm, though this limitation was mitigated somewhat by using blinded central raters, and by keeping the site raters and patients blinded to study arm until after week 8. The impact of polypharmacy is another intrinsic limitation; however, as was discussed in the primary analysis,

Table 2 Mokken scale analysis of homogeneity of HAM-D17 and HAM-D6 scores at week 8

| Treatment Arm | N  | HAM-D17 (H) | HAM-D6 (H) |
|---------------|----|-------------|------------|
| Combined      | 1298 | 0.30        | 0.53       |
| Guided-Care   | 621  | 0.34        | 0.56       |
| TAU           | 677  | 0.27        | 0.51       |

Loevinger’s coefficient (H) values were determined for combined treatment arms, guided-care and treatment as usual. HAM-D6, Hamilton depression rating scale, 6-item; HAM-D17, Hamilton depression rating scale, 17-item; TAU treatment as usual.
confounding effects likely would be equivalent between study arms. A specific limitation of using the HAM-D6 scale is that it does not assess for some important depressive symptoms, including physical symptoms [24] and suicide. The routine use of separate, more comprehensive suicide assessments in modern clinical trials of MDD treatments reduces concern about this limitation.

**Conclusion**
The results of this analysis are consistent with a substantial body of published evidence showing that HAM-D6, which is focused more precisely on core depressive symptoms, is more sensitive than HAM-D17 in assessing depression symptom improvement in patients with MDD. The demonstrated utility of pharmacogenomics-guided treatment over TAU as detected by the HAM-D6 in the GUIDED trial highlights its value for future biomarker-guided trials comparing multiple active treatment arms.

**Abbreviations**
GUIDED: Genomics Used to Improve DEpression Decisions trial; HAM-D: Hamilton Depression Rating Scale; MADRS: Montgomery-Åsberg Depression Rating Scale; MDD: major depressive disorder; QIDS-SR: Quick Inventory of Depressive Symptomatology-Self-Report; TAU: treatment as usual

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**Authors’ contributions**
BWD, SVP and JFG contributed to the conception and design of the work and the acquisition and interpretation of data. AJR, MET, CD, CRC, BPF, FMM, RCS, and MM contributed to the acquisition and interpretation of data. PT and JL contributed to the interpretation of data. BWD wrote the first draft, and JL, HY, SVP, and JFG substantively revised the work. All authors read and approved the final manuscript.

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**Availability of data and materials**
All data generated or analyzed during this study are included in this published article.

**Ethics approval and consent to participate**
The GUIDED study protocol was approved by the Copernicus Group independent review board (INC1-14-012) and conducted in accordance with the principles of the Declaration of Helsinki and its amendments. All patients provided written informed consent for participation.

**Consent for publication**
Not applicable.

**Competing interests**
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References

1. Trivedi MH, Rush AJ, Wisniewski SR, Nierenberg AA, Warden D, Ritz L, Norquist G, Howland RH, Lebowitz B, McGrath PJ, et al. Evaluation of outcomes with citalopram for depression using measurement-based care in STAR*D: implications for clinical practice. Am J Psychiatry. 2006;163(1):28–40.
2. Haddler G, Drevets WC, Manji HK, Charney DS. Discovering endophenotypes for major depression. Neuropsychopharmacology. 2004;29(10):1765–81.
3. Wray NR, Ripke S, Mattheisen M, Traskoukoski M, Byrne EM, Abdollaau A, Adams MJ, Agerbo E, Air TM, Andlauer TMF, et al. Genome-wide association studies identify 44 risk variants and refine the genetic architecture of major depression. Nat Genet. 2015;47(5):668–81.
4. Howard DM, Adams MJ, Clarke TK, Hafferty JD, Gibson J, Shirali M, Coleman BJ, Hagenaaars SP, Ward J, Wigmore EM, et al. Genome-wide meta-analysis of depression identifies 102 independent variants and highlights the importance of the prefrontal brain regions. Nat Neurosci. 2019;22(3):343–52.
5. Conrado DJ, Rogers HL, Zineh I, Pacanowski MA. Consistency of drug-diff and gene-drug interaction information in US FDA-approved drug labels. Pharmacogenomics. 2013;14(2):215–23.
6. Hicks JR, Bishop JR, Sangkkuhl K, Muller DJ, Ji Y, Leckband SG, Leedger JS, Graham RL, Chiulli DL, Llerena A, et al. Clinical pharmacogenetics implementation consortium (CPC) guideline for CYP2D6 and CYP2C19 genotypes and dosing of selective serotonin reuptake inhibitors. Clin Pharmacol Ther. 2015;98(2):127–34.
7. Bousman CA, Arandjelovic K, Mancuso SG, Eyre HA, Dunlop BW. Pharmacogenetic tests and depressive symptom remission: a meta-analysis of randomized controlled trials. Pharmacogenomics. 2019;20(1):37–47.
8. Greden JF, Parkih SV, Rothchild AJ, Thase ME, Dunlop BW, Defattista C, Conway CR, Forester B, Mondimore FM, Shelton RF, et al. Impact of pharmacogenomics on clinical outcomes in major depressive disorder in the GUIDED trial: a large, patient- and rater-blinded, randomized, controlled study. J Psychiatr Res. 2019;115:59–67.
9. Rush AJ, Trivedi MH, Wisniewski SR, Stewart JW, Nierenberg AA, Thase ME, Ritz L, Biggs MM, Warden D, Luther JF, et al. Bupropion-SR, sertraline, or venlafaxine-XR after failure of SSRIs for depression. N Engl J Med. 2006;354(12):1231–42.
10. Uher R, Maier W, Hauser J, Marusic A, Schmael C, Mors O, Henigsberg N, Souery D, Placentino A, Rietzch M, et al. Differential efficacy of escitalopram and norifinpyline on dimensional measures of depression. Br J Psychiatry. 2009;194(3):252–9.
11. Rush AJ, Trivedi MH, Stewart JW, Nierenberg AA, Fava M, Kusnir BT, Warden D, Morris DW, Luther JF, Husain MM, et al. Combining medications to enhance depression outcomes (CO-MED): acute and long-term outcomes of a single-blind randomized study. Am J Psychiatry. 2011;168(7):689–701.
12. Cipriani A, Furkaeva TA, Salant G, Chaimara J, Atkinson LZ, Ogawa Y, Leucht S, Ruhe HG, Turner HE, Higgins JPT, et al. Comparative efficacy and acceptability of 21 antidepressant drugs for the acute treatment of adults with major depressive disorder: a systematic review and network meta-analysis. Lancet. 2018;391(10128):1357–66.

Dunlop BW. Evidence-based applications of combination psychotherapy and pharmacotherapy for depression. Focus (Am Psychiatr Publ). 2016;14(1):156–73.

Fried EI, Nesse RM. Depression sum-scores don’t add up: why analyzing specific depression symptoms is essential. BMC Med. 2015;13:72.

Hamilton M. A rating scale for depression. J Neurol Neurosurg Psychiatry. 1962;256:6–7.

Hamilton M. Development of a rating scale for primary depressive illness. Br J Clin Psychol. 1967;6(4):278–96.

Bech P, Allerup P, Gram LF, Reisby N, Rosenberg R, Jacobsen O, Nagy A. The Hamilton depression scale. Evaluation of objectivity using logistic models. Acta Psychiatr Scand. 1981;63(3):290–9.

Santor DA, Coyne JC. Examining symptom expression as a function of symptom severity: item performance on the Hamilton rating scale for depression. Psychol Assess. 2001;13(1):127–39.

Ostergaard SD, Bech P, Trivedi MH, Wisniewski SR, Rush AJ, Fava M, Brief unidimensional melancholia rating scales are highly sensitive to the effect of citalopram and may have biological validity: implications for the research domain criteria (RDoC). J Affect Disord. 2014;163:18–24.

Bagby RM, Ryder AG, Schuller DR, Marshall MB. The Hamilton depression rating scale: has the gold standard become a lead weight? Am J Psychiatry. 2004;161(12):2163–77.
21. Ruhe HG, Dekker JJ, Peen J, Holman R, de Jonghe F. Clinical use of the Hamilton depression rating scale: is increased efficiency possible? A post hoc comparison of Hamilton depression rating scale, Maier and Bech subscales, clinical global impression, and symptom Checklist-90 scores. Compr Psychiatry. 2005;46(6):417–27.

22. Moller HJ. Methodological aspects in the assessment of severity of depression by the Hamilton depression scale. Eur Arch Psychiatry Clin Neurosci. 2001;251(Suppl 2):113-20.

23. Licht RW, Qvitzaa S, Alereup P, Bech P. Validation of the Bech-Rafaelsen melancholia scale and the Hamilton depression scale in patients with major depression; is the total score a valid measure of illness severity? Acta Psychiatr Scand. 2005;111(2):144–9.

24. Bech P. The ABC profile of the HAM-D17. Revista brasileira de psiquiatria (Sao Paulo, Brazil). 1999. 2011;33(2):109–10.

25. Bech P, Gram LF, Dein E, Jacobsen O, Vitger J, Bolwig TG. Quantitative rating of depressive states. Acta Psychiatr Scand. 1975;51(3):161–70.

26. Lercker Y, Bech P. The Ham D(6) is more homogenous and as sensitive as the Ham D(17). Eur Psychiatry. 2007;22(4):252–5.

27. Timmerby N, Andersen JH, Sondegaard S, Ostergaard SD, Bech P. A systematic review of the clinimetric properties of the 6-item version of the Hamilton depression rating scale (HAM-D6). Psychother Psychosom. 2017;86(3):141–9.

28. O’Sullivan RL, Fava M, Agustin C, Baer L, Rosenbaum JF. Sensitivity of the six-item Hamilton depression rating scale. Acta Psychiatr Scand. 1997;95(5):379–84.

29. Hooper CL, Bakshi D. An examination of the sensitivity of the six-item Hamilton rating scale for depression in a sample of patients suffering from major depressive disorder. J Psychiatry Neurosci. 2000;25(2):178–84.

30. Jablonski M, King N, Wang Y, Winner JG, Watterson LR, Gurselman S, Dechairo BM. Analytical validation of a psychiatric pharmacogenomic test. Personal Med. 2018;15(3):189–97.

31. Hall-Flavin DK, Winner JG, Allen JD, Jordan JU, Nesheim RS, Snyder KA, Drews MS, Esterhold LL, Biennack JM, Mazeek DA. Using a pharmacogenic algorithm to guide the treatment of depression. Transl Psychiatry. 2012;2:e172.

32. Frank E, Pien RF, Jarrett RB, Keller MB, Kupper DJ, Lavori PW, Rush AJ, Weissman MM. Conceptualization and rationale for consensus definitions of terms in major depressive disorder: Remission, recovery, relapse, and recurrence. Arch Gen Psychiatry. 1991;48(9):851–5.

33. Kyle PR, Lemming OM, Timmerby N, Sondergaard S, Andreasson K, Bech P. The validity of the different versions of the Hamilton depression scale in separating remission rates of placebo and antidepressants in clinical trials of major depression. J Clin Psychopharmacol. 2016;36(5):453–6.

34. Thase ME, Parikh SV, Rothschild AJ, Dunlop BW, DeBattista C, Conway CR, Forester BP, Mondimore FM, Shelton RC, Macaluso M, et al. Impact of pharmacogenomics on clinical outcomes for patients taking medications with gene-drug interactions in a randomized, controlled trial. J Clin Psychiatry. 2019;80(6).

35. Dunlop BW, Rapaport MH. When should a patient be declared recovered from a major depressive episode? J Clin Psychiatry. 2016;77(8):e1026–8.

36. Zimmerman M, Martinez J, Attuliah N, Friedman M, Toba C, Boreescu DA, Rahjeb M. Further evidence that the cutoff to define remission on the 17-item Hamilton depression rating scale should be lowered. Depress Anxiety. 2012;29(2):69–65.

37. Cumming G. Understanding the new statistics: effect sizes, confidence intervals, and metaanalysis. London: Routledge; 2012.

38. Dunlop BW, Davis PG. Combination treatment with benzodiazepines and SSRIs for comorbid anxiety and depression: a review. Prim Care Companion J Clin Psychiatry. 2008;10(3):222–8.

39. Entsuah R, Shaffer M, Zhang J. A critical examination of the sensitivity of unidimensional subscales derived from the Hamilton depression rating scale to antidepressant drug effects. J Psychiatr Res. 2002;36(6):437–48.

40. Ostergaard SD, Bech P, Mikowsiak KW. Fewer study participants needed to demonstrate superior antidepressant efficacy when using the Hamilton melancholia subscale (HAM-D(6)) as outcome measure. J Affect Disord. 2016;190:842–5.

41. Leon AC, Marzuk PM, Portera L. More reliable outcome measures can reduce sample size requirements. Arch Gen Psychiatry. 1995;52(10):867–71.

42. Moller HJ. Improving the assessment of severity of depressive states: a reduction of the Hamilton depression scale. Pharmacopsychiatry. 1985;18:114–5.

43. Cleary PJ. Problems of internal consistency and scaling in life event schedules. J Psychiatr Res. 1981;25(4):309–20.