The Relationship Between Mental Health, Disease Severity, and Genetic Risk for Depression in Early Rheumatoid Arthritis

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ABSTRACT

Objective: Reduced mental health (MH) is prevalent in rheumatoid arthritis (RA). Although longitudinal studies are limited, there is evidence that depression is associated with worse disease outcomes. We evaluated reciprocal relationships between MH, RA severity, and genetic risks for depression for 2 years in a well-characterized cohort of RA patients.

Methods: We evaluated 520 early RA patients previously enrolled to two clinical trials. MH was measured using the short form-36 MH domain and mental component summary scores (MCS). MCS/MH associations over 2 years with disease activity (disease activity score on a 28-joint count), disability (health assessment questionnaire), pain visual analog scale scores, and a weighted genetic risk score for depression were tested using linear mixed-effects and regression models.

Results: Poorer MH was associated with worse RA outcomes. Lower MCS scores (indicating worse MH) were seen in patients with a greater genetic risk for depression (weighted genetic risk score: coefficient = −1.21, p = .013). Lower baseline MCS was associated with lower 2-year improvements in disease activity score on a 28-joint count (coefficient = −0.02, p < .001), pain (coefficient = −0.33, p < .001), and health assessment questionnaire (coefficient = −0.01, p = .006). Baseline MCS was associated with changes in the swollen joint count (coefficient = −0.09, p < .001) and patient global assessment (coefficient = −0.28, p < .001) but not the tender joint count (p = .983) and erythrocyte sedimentation rate (p = .973). Only baseline pain visual analog scale (coefficient = −0.07, p = .002) was associated with 2-year changes in MCS.

Conclusions: Reduced baseline MH was associated with lower improvements in disease activity, disability, and pain for 2 years, supporting current national guidelines recommending screening for depression in RA. Pain had a bidirectional relationship with MH. Depression genetic risk had a significant association with MH.

Key words: disability, disease activity, genetics, mental health, pain, rheumatoid arthritis.

INTRODUCTION

Reduced mental health (MH) is prevalent in rheumatoid arthritis (RA), with major depression present in 16.8% of patients (1). The cause of this excess burden of MH impairment is uncertain. Comorbid depression also seems to have a detrimental impact on the disease course of RA, being associated with increased healthcare use and costs (2) and representing an independent risk factor for non-suicide-related mortality (3). Determining the cause and effect of depression in RA is, therefore, a key research goal. Research in this area has mainly involved cross-sectional studies in patients with long-standing RA. These identified associations between depression and pain (4), disability (5), and arthritis disease activity (6). Their cross-sectional nature, however, made...
it impossible to infer causality. Although longitudinal studies are limited, there is some evidence for a bidirectional effect with pain in patients with musculoskeletal disorders, whereby depression influences pain and vice versa (7,8). There is also some evidence that depression predicts the subsequent disease activity of RA, with an analysis of established RA patients finding a slower rate of decline in disease activity over time in patients with a history of depression (9).

Depression also has a substantial genetic component (10), with several variants associated with depression identified in a genome-wide association study (GWAS) (11). This is consistent with previous twin studies of the heritability of major depressive disorder (MDD), which found that the disorder has a heritability of 48% to 75%, depending on assumptions made on the prevalence of MDD in the general population (12). The role of these in determining MH in RA has not previously been evaluated.

The aim of our study was to evaluate the relationship between MH and disease activity, disability, pain, and genetic risk for depression for 2 years in a well-characterized clinical trial cohort of patients with early RA. The direction of any associations was tested by examining the impact of baseline MH on changes in disease activity, disability and pain, and vice versa.

METHODS

Participants
We studied patients in the Combination Anti-Rheumatic Drugs in Early RA (CARDERA) genetics cohort. It has been described in detail previously (13). In brief, it comprises European ancestry RA patients enrolled to two multicenter randomized controlled trials (RCTs): CARDERA-1 and CARDERA-2 (14,15). Both recruited patients with early RA (<2-year duration) and active disease defined as three of three or more swollen joints, six or more tender joints, 45-minute or more morning stiffness, or erythrocyte sedimentation rate (ESR) of 28 mm/h or greater. CARDERA-1 recruited patients between 2000 and 2002; CARDERA-2 recruited patients between 2003 and 2010. CARDERA-1 randomized patients to receive either (1) methotrexate; (2) methotrexate and ciclosporin; (3) methotrexate and prednisolone; or (4) methotrexate, ciclosporin, and prednisolone. CARDERA-2 randomized patients to receive either (1) methotrexate or (2) methotrexate and anakinra. Because the original aim of the CARDERA studies was to investigate the performance of combination therapy with reference to monotherapy, a placebo group was not assigned. Rheumatoid factor (RF), a biomarker providing clinical information on the antibody composition of patient serum, was assayed as described previously (16). Follow-up was 2 years. The current analysis is restricted to the 520 patients with baseline MH data available.

Disease Outcomes
The following disease outcomes were captured. First, disease activity (how active a patient's arthritis is) was recorded using the disease activity score on a 28-joint count (DAS28). This composite score combines information on the number of swollen and tender joints (assessed by a clinician from 28 joint counts), the patient global assessment (PGA) of disease activity (which involves a patient rating his or her overall disease activity on a 100-mm visual analog scale [VAS]), and the ESR in a mathematical formula to give an assessment of RA activity ranging from 0 to 10. Lower scores indicate less active disease, with scores of higher than 5.1, lower than 3.2, and lower than 2.6 indicating high disease activity, low disease activity, and remission, respectively. Second, disability was recorded using the health assessment questionnaire (HAQ), a patient-completed questionnaire giving a score of function ranging from 0 to 3. HAQ scores of lower than 1, 1 to 2, and higher than 2 indicate mild, moderate, and severe disability, respectively. Third, pain was recorded using a 100-mm patient-completed pain VAS, a method for quantifying the severity of self-reported pain (17). Fourth, health-related quality of life (HRQoL) was recorded using the short-form-36 (SF-36), which is described in detail hereinafter. In CARDERA-1, the aforementioned outcomes were captured every 6 months. In CARDERA-2, they were captured at 0, 6, 12, and 24 months.

Mental Health
The SF-36 is a generic measure of health status, capturing HRQoL across eight domains (4 physical and 4 mental) (18). These domains are scored from 0 to 100, with higher scores indicating better HRQoL. They can be normalized, z-transformed, and combined into mental component summary (MCS) and physical component summary scores providing summary measures of overall mental and physical health, relative to a population mean score of 50 (SD = 10) (19).

We used the MH domain score and MCS as measures of MH in our analysis. Both have been used to screen for depression, with an MCS cutoff of 42 having a sensitivity and specificity of 74% and 81%, respectively, for detecting depressive disorder (20). They also associate with depression severity, both cross-sectionally and over time (21).

Genotyping
CARDERA patients were genotyped on the Illumina ImmunoChip array (described in detail previously (13)). Single nucleotide polymorphism (SNP) markers were removed that had more than 5% missingness, were duplicates, were not in Hardy-Weinberg equilibrium (HWE, p < .00001), and had a minor allele frequency (MAF) of less than 0.01. From 196,524 pre-identification control markers, 138,373 were available in the final data set. Imputation was subsequently performed using IMPUTE2 (22) and the 1000 Genomes Phase 1 integrated variant version 3 (March 2012) reference panel (variants filtered with a European MAF <0.01). Postimputation SNPs were removed with low INFO scores (<0.7), MAF (<0.05), HWE (p < .000001), and genotyping rate (<0.1), resulting in 429,193 available markers.

Genetic Risk for Depression
The Psychiatric Genomics Consortium MDD GWAS mega-analysis failed to find a locus of genome-wide significance, likely reflecting limited power caused by the genetic architecture of MDD (small effect sizes of individual genetic variants) and the high prevalence of MDD, which increases the difficulty in recruiting large samples of screened, low-risk controls (23). We therefore tested a weighted genetic risk score (wGRS), combining loci of genetic variants) and the high prevalence of MDD, which increases the difficulty in recruiting large samples of screened, low-risk controls (23). We therefore tested a weighted genetic risk score (wGRS), combining loci of genetic variants with MDD for an association with MH in CARDERA. This approach is commonly used in studies of polygenic disorders, whose genetic architecture comprises thousands of very small effect common alleles (24,25). We used a c value threshold (c) for SNPs to include in the wGRS of 0.05 (representing nominal association with MDD). A continuous wGRS based on MDD GWAS results has been shown to predict depression in independent cohorts, with a c value of .05 demonstrating the wGRS that most strongly predicts MDD risk (26). After linkage disequilibrium pruning, 3010 SNPs were included in the wGRS. The wGRS was generated for each individual in CARDERA by calculating the number of nominally associated risk alleles they carried, weighted by the log odds ratio from the MDD mega-analysis, summed across SNPs.

Statistical Analysis
 Associations with MH
Two different modeling approaches were used to evaluate the relationship between MH, RA severity measures, and genetic risk for depression. The first approach established whether MH was associated with either RA severity measures or the wGRS for depressive disorder over time. This used a linear mixed-effects model, which incorporated either MCS or MH measured at each time point as the response variable regressed on the corresponding predictors (wGRS, DAS28 and its components, HAQ, or pain VAS) from each time point. Effect size estimates (coefficients) for predictor variables provided information on the average differences in the MCS or
MH score for the 2-year period relative to the average predictor variable score. We have included a level 1 random effect of individual, fitting random intercepts for each individual. These models have a level 2 random effect for time point, modeling deviation from the overall effect of time point on outcome within each individual as random slopes. We specify this correlation structure using the lm4 package in R (27). The following variables were tested for their associations with MCS: age, disease duration, and RF status. Of these, only sex improved the model fit—as determined using a stepwise selection approach, with the optimal model determined using the Akaike information criterion—and thus, only this variable was included as a covariate (Supplementary Tables A.1 and A.2, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380). The wGRS was standardized to a z score to provide interpretable β values. Examination of residuals from a model containing time, sex, and treatment only confirmed a good model fit (Supplementary Figure A.1, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380), and variance inflation factors calculated for each predictor-ensured multicollinearity between RA outcomes and DAS28 components were not an issue (Supplementary Table A.3, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380).

The second approach evaluated the direction of associations between MH and RA severity by testing whether MH at study baseline was associated with 2-year changes in RA outcomes over time or vice versa. This used linear regression models to look at the association between (a) baseline MCS or MH and 2-year changes in RA severity measures and (b) baseline RA severity measures and 2-year changes in MH scores. These models included the baseline response variable score, treatment, and sex as covariates. Examination of model residuals confirmed good model fits (Supplementary Figure A.1, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380).

**Missing Data Imputation**

In the original CARDERA-1 trial, missing data at each time point had previously been imputed using last observation carried forward (LOCF) analysis for study end points (DAS28, HAQ, and SF-36). LOCF is a commonly used procedure to address missing data in clinical trials with repeated measures over multiple time points. For each individual, missing values at a time point are replaced by the last observed value of that variable (14,28). In the original CARDERA-2 trial, missing data were not imputed (15). For consistency in the current analysis, we imputed missing, previously nonimputed CARDERA-1 data (swollen joint count [SJC], tender joint count [TJC], ESR, PGA, pain VAS) and missing CARDERA-2 data using LOCF. The largest amount of missing data was seen for pain VAS (11% of observations missing across all time points). We repeated our analysis with nonimputed data only; this excluded a significant impact of the LOCF assumption (Supplementary Table A.4, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380).

**Statistical Software**

Analyses were performed in the statistical environment R (R Foundation for Statistical Computing, Vienna, Austria), PRSice (Version 1.2) (29), IMPUTE2 (22), and PLINK (Version 1.9) (30).

**Ethics, Consent, and Permissions**

CARDERA-1 (South Thames Multicentre Research Ethics Committee (REC) reference: MREC (1) 99/04) and CARDERA-2 (South East REC reference: MREC 02/1/089) were ethically approved. Approval was obtained to genotyping archived DNA (National Research Ethics Service Committee East of England-Essex REC reference: 11/EE/0544). All patients provided consent.

**RESULTS**

**Patient Baseline Characteristics**

Most patients were female (69%, Table 1) and possessed RF in their serum (67%). Baseline disease activity was high (mean DAS28 = 5.88), moderate disability (mean HAQ = 1.56), and short disease duration (mean duration = 3.3 months). Baseline MH was reduced relative to the general population (mean MCS score = 40.6, which is 9.4 U lower than the general population mean).

**Disease Severity Associations With MH**

In a sex- and treatment-adjusted linear mixed-effects model, DAS28 (p < .001), HAQ (p < .001), and pain VAS (p < .001) were significantly associated with MCS (Table 2). On average, for 2 years, MCS scores were 2.22, 6.07, and 0.14 units lower per unit increase in DAS28, HAQ, and pain VAS scores, respectively. This indicates that the higher a patient's disease activity, disability, and pain levels, the worse their MH. In multivariate models, the following three disease severity measures retained a highly significant association with MCS (Table 2): HAQ (coefficient = −3.38, p < .001), DAS28 (coefficient = −0.91, p < .001), and pain VAS (coefficient = −0.05, p < .001). Similar associations were seen with the MH domain.

**Major Depressive Disorder Genetic Risk Score Associations With MH**

A significant association was seen between the wGRS for depression and MCS (p = .013) and MH (p = .041) (Table 2). The association with MCS (p = .033) but not MH (p = .080) was retained in multivariate models including DAS28, HAQ, and pain VAS as covariates. Higher wGRS scores, which indicate a greater genetic risk for depression, were associated with worse MH (lower MCS and MH scores) (MCS: coefficient = −1.21; MH: coefficient = −1.37). Repeating the analysis with a linear mixed-effects model that incorporated a wGRS-time interaction term provided some evidence that genetic risk for depression also predicted the rate at which MH improved across time points, with a significant association seen between the wGRS-time term and MH (p = .039, coefficient = −0.83) but not MCS (p = .330, coefficient = −0.30).

**TABLE 1.** CARDERA Genetics Cohort Baseline Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Summary Statistic</th>
</tr>
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<tbody>
<tr>
<td>Demographics</td>
<td></td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>358 (69)</td>
</tr>
<tr>
<td>Age, M (SD), y</td>
<td>54.7 (12.6)</td>
</tr>
<tr>
<td>RA specific</td>
<td></td>
</tr>
<tr>
<td>RA duration, M (SD), mo</td>
<td>3.3 (4.9)</td>
</tr>
<tr>
<td>RF positive, n (%)</td>
<td>350 (67)</td>
</tr>
<tr>
<td>DAS28, M (SD)</td>
<td>5.88 (1.29)</td>
</tr>
<tr>
<td>HAQ, M (SD)</td>
<td>1.56 (0.70)</td>
</tr>
<tr>
<td>Mental health</td>
<td></td>
</tr>
<tr>
<td>MCS, M (SD)</td>
<td>40.6 (14.1)</td>
</tr>
<tr>
<td>MH, M (SD)</td>
<td>61.0 (18.0)</td>
</tr>
<tr>
<td>Treatment</td>
<td></td>
</tr>
<tr>
<td>Receiving MTX, n (%)</td>
<td>159 (31)</td>
</tr>
<tr>
<td>Receiving MTX and CIC, n (%)</td>
<td>108 (21)</td>
</tr>
<tr>
<td>Receiving MTX and pred, n (%)</td>
<td>102 (20)</td>
</tr>
<tr>
<td>Receiving triple therapy, n (%)</td>
<td>107 (21)</td>
</tr>
<tr>
<td>Receiving MTX and anakintra, n (%)</td>
<td>44 (8)</td>
</tr>
</tbody>
</table>

M = mean; SD = standard deviation; RA = rheumatoid arthritis; RF = rheumatoid factor; DAS28 = disease activity score on a 28-joint count; HAQ = health assessment questionnaire; MCS = mental component summary score; MH = mental health; MTX = methotrexate; CIC = ciclosporin; pred = prednisolone; triple therapy = MTX, CIC, and pred.

Cohort size used in analysis is composed of 520 patients. A DAS28 of 5.88 indicates highly active disease. A HAQ of 1.56 indicates moderate disability. An MCS of 40.6 is 9.4 units lower than that observed in the normal population (which has a mean score of 50.0 units).
TABLE 2. Longitudinal Associations Between MH, RA Outcomes, and Depression Genetic Risk Score

<table>
<thead>
<tr>
<th>RA Outcomes</th>
<th>MCS Sex- and Treatment-Adjusted Model</th>
<th>MH Domain Score Sex- and Treatment-Adjusted Model</th>
<th>Multivariate Model*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>coefficient (SE)</td>
<td>p</td>
<td>coefficient (SE)</td>
</tr>
<tr>
<td>DAS28</td>
<td>-2.22 (0.16)</td>
<td>&lt;.001</td>
<td>-0.91 (0.21)</td>
</tr>
<tr>
<td>HAQ</td>
<td>-6.07 (0.40)</td>
<td>&lt;.001</td>
<td>-3.88 (0.48)</td>
</tr>
<tr>
<td>Pain VAS</td>
<td>-0.14 (0.01)</td>
<td>&lt;.001</td>
<td>-0.05 (0.01)</td>
</tr>
</tbody>
</table>

Genetic risk

| MDD wGRS   | -1.21 (0.48)      | .013     | -0.92 (0.43)      | .033     | -1.37 (0.67)      | .041     | -1.05 (0.60) | .080  |

DAS28 components

| SJC        | -0.32 (0.04)      | <.001    | -0.12 (0.04)      | .003     | -0.47 (0.05)      | <.001    | -0.25 (0.05) | <.001 |
| ESR        | -0.07 (0.01)      | <.001    | -0.04 (0.01)      | .003     | -0.11 (0.02)      | <.001    | -0.06 (0.02) | <.001 |
| PGA        | -0.13 (0.01)      | <.001    | -0.11 (0.01)      | <.001    | -0.16 (0.01)      | <.001    | -0.11 (0.01) | <.001 |
| TJC        | -0.27 (0.04)      | <.001    | -0.03 (0.04)      | .461     | -0.34 (0.05)      | <.001    | -0.03 (0.05) | .519  |

MCS = mental component summary score; MH = mental health; SE = standard error; RA = rheumatoid arthritis; DAS28 = disease activity score on a 28-joint count; HAQ = health assessment questionnaire; pain VAS = pain visual analog scale score; MDD wGRS = weighted genetic risk score for major depressive disorder; SJC = swollen joint count; ESR = erythrocyte sedimentation rate; PGA = patient global assessment of disease activity; TJC = tender joint count.

All linear mixed-effects models include sex, treatment, and time as explanatory variables; multivariate model for DAS28 components also includes SJC, ESR, PGA, and TJC as explanatory variables.

* Multivariate model for RA outcomes and genetic risk also includes DAS28, HAQ, pain VAS, and MDD wGRS as explanatory variables.

Disease Activity Score on a 28-Joint Count Component Associations With MH

In a sex- and treatment-adjusted linear mixed-effects model, all four DAS28 components—SJC, TJC, ESR, and PGA—were associated with MCS and MH scores when tested individually (Table 2). Higher scores in each DAS28 component were associated with lower MCS and MH scores; this indicates that more active disease is linked with poorer MH. On average, for 2 years, MCS scores were lower in the highest SJC octile (group 2) and the lowest SJC octile (group 8) (Table 3). Lower baseline MCS and MH scores (indicating poorer MH) were associated with lesser improvements in DAS28, pain VAS, and HAQ scores. The effect sizes were, however, modest: per 10 unit increase in baseline MCS, the 2-year reductions in DAS28, HAQ, and pain VAS were 0.20, 0.17, and 13.91 units, respectively, in the lowest MCS octile (group 1) and 8.02, and 11.98 units, respectively, in the highest MCS octile (group 8).

Dividing patients into octiles based on their baseline MCS and plotting the mean disease severity measure in each octile demonstrated the effect of baseline MCS on RA outcomes (Figure 1). Trends toward (a) worse disease outcomes at each time point and (b) lower improvements in disease outcomes for 2 years across increasing baseline MCS octiles were seen (Figure 1). For 2 years, the mean DAS28, HAQ, and pain VAS scores changed by −1.14, −0.23, and −8.11 U, respectively, in the lowest MCS octile (group 1) and −1.94, −0.49, and −18.49 units, respectively, in the highest MCS octile (group 8).

Examining individual DAS28 components revealed that baseline MCS and MH scores had significant inverse associations with 2-year changes in the SJC (MCS and MH, p < .001) and PGA (MCS and MH, p < .001) but not the TJC (MCS, p = .983; MH, p = .226) and ESR (MCS, p = .973; MH, p = .355) (Table 3). This differential impact on DAS28 components is shown in Figure 2. For 2 years, the mean SJC, PGA, TJC, and ESR levels changed by −0.17, −13.91, −8.02, and −11.98 units, respectively, in the lowest MCS octile (group 1) and −4.69, −20.46, −5.97, and −11.03, respectively, in the highest baseline MCS octile (group 8).

Direction of Association Between RA Outcomes and MH

Association Between Baseline Disease Severity and Changes in MH

The only baseline RA severity measure that had a significant association with 2-year changes in both MCS and MH scores was pain VAS (Table 3). Higher baseline pain VAS scores (indicating greater levels of pain) were associated with lesser increases in MCS and MH scores (indicating lower improvements in MH). The increase in MCS was 0.07 units less per 1-mm increase in baseline pain VAS. A significant association between the baseline TJC and 2-year changes in MH domain scores was also seen (p = .023), although this variable was not significantly associated with 2-year changes in MCS scores (p = .122).

Association Between Baseline MH and Changes in RA Outcomes

Baseline MCS and MH domain scores had significant inverse associations with 2-year changes in DAS28 (MCS and MH, p < .001), pain VAS (MCS and MH, p < .001), and HAQ (MCS, p = .006; MH, p = .008) (Table 3). Lower baseline MCS and MH scores (indicating poorer MH) were associated with lesser improvements in DAS28, pain VAS, and HAQ scores. The effect sizes were, however, modest: per 10 unit increase in baseline MCS, the 2-year reductions in DAS28, HAQ, and pain VAS were 0.20, 0.10, and 3.30 units greater, respectively (Table 3).

Dividing patients into octiles based on their baseline MCS and plotting the mean disease severity measure in each octile demonstrated the effect of baseline MCS on RA outcomes (Figure 1). Trends toward (a) worse disease outcomes at each time point and (b) lower improvements in disease outcomes for 2 years across increasing baseline MCS octiles were seen (Figure 1). For 2 years, the mean DAS28, HAQ, and pain VAS scores changed by −1.14, −0.23, and −8.11 U, respectively, in the lowest MCS octile (group 1) and −1.94, −0.49, and −18.49 units, respectively, in the highest MCS octile (group 8).

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DISCUSSION

Our study evaluated the relationship between MH and disease activity, disability, pain, and genetic risk for depression for 2 years in a well-characterized clinical trial cohort of patients with early RA. It has three key findings. The first, and most clinically important, is
that low MH was associated with poorer disease outcomes. In a repeated measures analysis, lower MCS and MH scores had significant associations with more active disease, increased disability, and greater pain for 2 years; because MCS and MH scores increased over time, DAS28, HAQ, and pain levels fell. Lower baseline MCS and MH scores (indicating worse MH) were associated with a reduced improvement in disease activity and disability, suggesting that depression predicts the degree to which RA improves over time. The relationship between pain and MH seemed bidirectional, with baseline pain associating with lower improvement in MCS and MH domain scores and vice versa; this is in keeping with existing studies of musculoskeletal disorders (7,8).

The second finding was that swollen, but not TJC, had a significant association with reduced MH. In a multivariate model incorporating all four DAS28 components, the tender joint component of the DAS28 (TJC) failed to retain a significant association with MCS and MH scores. In established RA patients attending routine clinics, the opposite relationship seems true, with an analysis of the CORRONA registry reporting that a lifetime depression history was associated with slower improvements in the TJC but not the SJC (9). One explanation for the lack of association between MCS/MH scores and the TJC in CARDERA is that the short disease duration of patients means the pain pathway sensitization characterizing fibromyalgic RA—which could be particularly

### TABLE 3. Direction of Associations Between MH and RA Outcomes

<table>
<thead>
<tr>
<th></th>
<th>Model 1: Baseline MCS Predicting 2-Year Changes in RA Outcomes</th>
<th>Model 2: Baseline RA Outcomes Predicting 2-Year Change in MCS</th>
<th>Model 1: Baseline MH Predicting 2-Year Changes in RA Outcomes</th>
<th>Model 2: Baseline RA Outcomes Predicting 2-Year Change in MH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>coefficient (SE)</td>
<td>p</td>
<td>coefficient (SE)</td>
<td>p</td>
</tr>
<tr>
<td>DAS28</td>
<td>−0.02 (0.01)</td>
<td>&lt;.001</td>
<td>−0.31 (0.45)</td>
<td>.492</td>
</tr>
<tr>
<td>SJC</td>
<td>−0.09 (0.02)</td>
<td>&lt;.001</td>
<td>0.04 (0.09)</td>
<td>.625</td>
</tr>
<tr>
<td>ESR</td>
<td>0.00 (0.07)</td>
<td>.973</td>
<td>0.03 (0.02)</td>
<td>.203</td>
</tr>
<tr>
<td>PGA</td>
<td>−0.28 (0.08)</td>
<td>&lt;.001</td>
<td>−0.03 (0.02)</td>
<td>.144</td>
</tr>
<tr>
<td>TJC</td>
<td>0.00 (0.02)</td>
<td>.983</td>
<td>−0.12 (0.07)</td>
<td>.122</td>
</tr>
<tr>
<td>HAQ</td>
<td>−0.01 (0.00)</td>
<td>.006</td>
<td>0.50 (0.85)</td>
<td>.554</td>
</tr>
<tr>
<td>Pain VAS</td>
<td>−0.33 (0.08)</td>
<td>&lt;.001</td>
<td>−0.07 (0.02)</td>
<td>.002</td>
</tr>
</tbody>
</table>

MCS = mental component summary score; MH = mental health; RA = rheumatoid arthritis; SE = standard error; DAS28 = disease activity score on a 28-joint count; SJC = swollen joint count; ESR = erythrocyte sedimentation rate; PGA = patient global assessment of disease activity; TJC = tender joint count; HAQ = health assessment questionnaire; pain VAS = pain visual analog scale score.

All linear regression models include sex, treatment, and baseline measure of response variable tested as covariates.

**FIGURE 1.** Mean DAS28, HAQ, and pain VAS stratified by baseline MCS octile. MCS divided into octiles (8 quantiles); mean scores with SE bars for octiles 1, 4, 5, and 8 plotted at each time point; to facilitate visual interpretation, octiles 2, 3, 6, and 7 are not plotted, although the same trends are observed (Supplementary Figure A.2, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A380). Color image is available only in online version (www.psychosomaticmedicine.org).
influenced by MH—is yet to occur. An explanation for the association observed between MCS/MH scores and the SJC is that overlapping proinflammatory cytokines, which are present in high levels in early active RA, play important roles in mediating both reduced MH and RA activity. Although this hypothesis is supported by evidence that administering interleukin 1β and tumor necrosis factor α induces depressive behavior in mice (31) and that these cytokines are elevated in the serum of depressed patients (32,33), it fails to explain why baseline MH scores did not predict changes in ESR levels. There is an extensive body of literature investigating the pathophysiology of inflammation-related depression (34), with one proposed mechanism being the activation of the enzyme indoleamine 2,3-dioxygenase by inflammatory cytokines, which catalyzes tryptophan leading to a downstream depletion in serotonin (35)—indeed, inflammation-related depression seems to be dependent on the activation of indoleamine 2,3-dioxygenase (36). In light of this, anti-inflammatory medication has been proposed as a treatment for inflammation-related depression; however, its efficacy is still contested (37); therefore, further research is required in other early active RA cohorts to confirm the generalizability of our results.

Our third finding was that genetic risk for MDD was a significant predictor of MH. We tested a wGRS combining 3010 SNPs of nominal association with MDD in the publicly available Psychiatric Genomics Consortium GWAS for its association with MH in CARDERA. Although a significant association with lower MCS and MH scores was observed, the comparatively large standard error (SE) of the wGRS variable makes any conclusions on its relative importance challenging. The significant interaction term for wGRS-time predicting MH, indicating slower improvements in MH among individuals with high MDD genetic risk, is consistent with previous work indicating that depression genetic risk increases an individual’s sensitivity to adverse environmental effects (38,39). Taken together, these findings support the notion that depression is a complex disorder with a modest, albeit important, genetic contribution comprising thousands of alleles of a small effect size.

Our study replicates existing research that depression and pain have a bidirectional relationship. In CARDERA, baseline MCS and MH scores predicted 2-year changes in pain VAS and vice versa. This finding has been documented in psoriatic arthritis; for example, Husted et al. (7) identified a small bidirectional relationship between MCS and pain in 394 patients followed up for a mean of 7.5 years. It has also been reported in patients with persistent back, hip, or knee pain (8), back pain (40,41), and pain from a variety of disorders (42). The complex nature of pain makes it difficult to discern mechanisms by which this pain-depression bidirectional relationship could occur. Possible mechanisms include the following: (1) low mood could affect pain through promoting maladaptive coping strategies, especially catastrophizing (perceiving a situation to be worse than it is) (43); (2) pain could affect MH through reducing daily activities (44) and reducing social activities (45); and (3) imbalances in shared neurotransmitters (serotonin and norepinephrine) in affective and nociceptive pathways could contribute to both mood and pain (44). Further research is required to better characterize the mechanisms underlying this complex relationship.

Supporting our finding that MH predicts disease outcomes across time points, other studies have reported a detrimental impact...
of reduced MH on patients’ responsiveness to anti-inflammatory medication—specifically anti–tumor necrosis factor—as defined by DAS28 change (46,47). This effect is highly relevant to stratified medicine in RA. Although in CARDERA, the impact of baseline MCS on improvements in disease outcomes for 2 years was modest, if considered alongside other poor prognostic markers, such as anti–citrullinated protein antibody status (48), human leukocyte antigen variants (49), smoking, and sex (50), it could provide clinically useful prognostic information, guiding decisions on treatment intensity and facilitating a stratified approach to managing early RA patients.

Our study has several strengths. These include its large size, recruitment from multiple centers spanning two clinical trials, the measurement of multiple disease outcomes in a highly standardized manner, and the short disease duration of RA (mean = 3.3 months), leaving it well placed to examine the effects of MH in very early disease. It also has several weaknesses. As a secondary post hoc analysis of existing RCTs, it did not test a prespecified hypothesis according to a predetermined analysis plan. It evaluated a clinical trial cohort of severe RA patients, limiting its generalizability to patients seen in routine clinical practice. In addition, we only evaluated European ancestry individuals; the relevance of our findings to other ethnic populations is uncertain.

Current National Institute for Health and Care Excellence guidelines for RA management recognize the importance of assessing for comorbid depression, recommending this as part of an annual review process (51). Our findings strongly support this recommendation in early RA. One unresolved issue is the impact of treating depression on the disease course of RA. Although we did not evaluate the impact of MH therapies on RA outcomes, there is some evidence that psychological interventions (such as cognitive behavioral therapy, disclosure therapy, and biofeedback) are useful adjunctive management tools in RA patients. Two systematic literature reviews have evaluated the evidence base for this. Astin et al. (52) reported significant pooled effect sizes for psychological interventions at reducing postinterventional pain, disability, and psychological status across 25 trials. Similarly, Dissanyake and Bertouch (53) found evidence for the efficacy of disclosure therapy and cognitive behavioral therapy with maintenance therapy across four and five studies, respectively. The evidence base is, however, limited with both reviews noting that available trials had methodological limitations. Further research is required to better define the impact of specific psychological interventions at improving disease outcomes in large, well-conducted clinical trials of RA patients.

CONCLUSIONS
In this cohort of 520 early active RA patients, reduced MH (captured using the SF-36) was associated with worse disease outcomes. Lower MCS and MH scores (indicating poorer MH) were significantly associated with more active disease, increased disability, and greater pain for 2 years. Worse baseline MH was associated with lesser improvements in RA outcome measures, suggesting that depression predicts the rate at which RA improves over time. A bidirectional relationship was observed between MH and pain, replicating existing work in musculoskeletal disorders. Depression genetic risk had a significant, albeit modest, impact on MH. Our findings support the current National Institute for Health and Care Excellence RA management guidelines recommending the annual screening of RA patients for comorbid depression. Further research is needed to establish the impact of specific MH management strategies on improving RA outcomes.

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