

Testing the Predictive Value of Peripheral Gene Expression for Nonremission Following Citalopram Treatment for Major Depression

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Major depressive disorder (MDD) in general, and anxious-depression in particular, are characterized by poor rates of remission with first-line treatments, contributing to the chronic illness burden suffered by many patients. Prospective research is needed to identify the biomarkers predicting nonremission prior to treatment initiation. We collected blood samples from a discovery cohort of 34 adult MDD patients with co-occurring anxiety and 33 matched, nondepressed controls at baseline and after 12 weeks (of citalopram plus psychotherapy treatment for the depressed cohort). Samples were processed on gene arrays and group differences in gene expression were investigated. Exploratory analyses suggest that at pretreatment baseline, nonremitting patients differ from controls with gene function and transcription factor analyses potentially related to elevated inflammation and immune activation. In a second phase, we applied an unbiased machine learning prediction model and corrected for model-selection bias. Results show that baseline gene expression predicted nonremission with 79.4% corrected accuracy with a 13-gene model. The same gene-only model predicted nonremission after 8 weeks of citalopram treatment with 76% corrected accuracy in an independent validation cohort of 63 MDD patients treated with citalopram at another institution. Together, these results demonstrate the potential, but also the limitations, of baseline peripheral blood-based gene expression to predict nonremission after citalopram treatment. These results not only support their use in future prediction tools but also suggest that increased accuracy may be obtained with the inclusion of additional predictors (eg, genetics and clinical scales).

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INTRODUCTION

Major depressive disorder (MDD) is a common psychiatric disorder, affecting an estimated 19 million people each year in the United States alone (Kessler *et al*, 2003, 2005; Murray and Lopez, 1997). For patients with MDD, the lifetime mortality due to suicide approaches 15% (Mann, 2003), and the annual cost associated with MDD-related disability and death ranks in the tens of billions of dollars in the US (Kupfer *et al*, 2012; Murray and Lopez, 1997). A critical barrier to

devise optimal treatment and achieve sustained remission is the lack of predictive tests for disease state and treatment remission. The uncomplicated, noninvasive, and cost-effective availability of blood samples make blood-based biomarkers leading candidates for large-scale development.

The potential of blood transcriptome (the panel of detectable gene transcripts) for monitoring health and disease states is being investigated for peripheral organ and brain diseases (for review, see (Mohr and Liew, 2007)). Convergent functional genomic approaches have proposed biomarkers for mood states in bipolar depression (Le Niculescu *et al*, 2008). Padmos *et al* (2008) have reported a profile of proinflammatory gene transcripts that discriminated bipolar-depressed patients from controls. Pajer *et al* (2012) differentiated between MDD and control subjects, and between MDD with and without anxiety in adolescents, using candidate blood and brain gene expression from animal models of depressive-like behaviors. Using inflammatory, stress and neuroplasticity markers, Papakostas *et al* (2011) developed a serum-based protein assay that robustly

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discriminated MDD from control patients, and that replicated in an independent validation cohort. Similarly, leukocyte expression of proinflammatory, neuroendocrine stress axis and neuroplasticity markers discriminated between MDD and control subjects; and baseline proinflammatory genes were differentially expressed between responders and nonresponders to two antidepressant treatments (Nortriptyline or escitalopram) (Cattaneo *et al*, 2012). Using a microarray approach, Mamdani *et al* (2011) found that changes in the expression of 32 probesets predicted changes in Hamilton Rating Scale for Depression (HRSD) scores following citalopram treatment. Belzeaux *et al* (2012) reported baseline changes between responder and nonresponder MDD subjects and identified four gene transcripts with predictive value for treatment response.

In this study, we tested the potential of gene expression to predict nonremission prior to treatment initiation. We focused on nonremission, as it significantly contributes to the chronic illness burden suffered by many patients. Specifically, we hypothesize that MDD patients who do not achieve symptom remission after 12 weeks of treatment (MDD-NR, MDD-nonremitter) differ in baseline gene expression from the subjects who will remit (MDD-R, MDD-remitter). If so, baseline blood gene expression should predict remission after treatment with citalopram, a common antidepressant. We measured large-scale blood transcriptome changes in 34 MDD and 33 healthy control subjects (initial cohort), as part of an ongoing study of anxious-depression treated with a 12-week course of combined citalopram and weekly psychotherapy treatment. In the first phase, we performed an exploratory analysis of differential expression to seek evidence for group differences, namely MDD-NR and MDD-R *vs* controls, being cognizant of the high false positive rate associated with these approaches. In the second and independent phase, we used the same dataset to construct a predictive model with a machine learning method with support vector machines and corrected the results for the bias of reporting the 'best' prediction rates. We then sought to confirm the prediction results using the existing clinical and blood gene expression data in a second cohort of MDD subjects with similar, although not identical characteristics, and who were treated for 8 weeks with citalopram alone (validation cohort) (Mamdani *et al*, 2011). We also had two sets of secondary predictions; first, we predicted that baseline gene expression profiles of MDD nonremitters would be characterized by an upregulation of genes associated with immune activation and inflammation. Second, we predicted that the inclusion of clinical scales would improve prediction of nonremission.

MATERIALS AND METHODS

A short summary is provided here. Details and technical information on the cohorts and analyses are in the online supplements. The initial cohort included 34 anxious-depressed adults recruited for an outpatient trial of combined medication and psychotherapy treatment, and an age- and gender-matched sample of 33 nondepressed controls (Table 1). The patient sample met criteria for a nonpsychotic MDD episode of sufficient severity (documented by DSM-IV SCID interview and score ≥ 15

on the 25-item HRSD (Hamilton, 1960)) and elevated symptoms of panic or anxiety (score ≥ 7 on the past-month panic and agoraphobic spectrum self-report (Cassano *et al*, 1997; Shear *et al*, 2001)). Demographic and clinical scales are listed in Supplementary Table S1. Nonpatient controls did not meet criteria for any mood or anxiety disorder. Individuals with a history of schizophrenia, bipolar disorder or antisocial personality disorder; drug or alcohol abuse/dependence in the past 3 months; severe, uncontrolled medical illness; history of nonresponse to citalopram; and women who were pregnant or planning to become pregnant were excluded from participation. Subjects were free of antidepressant treatment prior to the start of the study. All study procedures were approved by the University of Pittsburgh Institutional Review Board. Treatment for the MDD cohort included citalopram (average dose at week 12, 37.5 ± 10.5 mg) and weekly psychotherapy with either (randomly assigned) interpersonal psychotherapy for depression with panic and anxiety symptoms (IPT-PS) or brief supportive psychotherapy (BSP) (Hellerstein *et al*, 1998). Analyses were conducted on an intent-to-treat basis, with depression remission defined as achieving three consecutive weeks of average HRSD-17 scores ≤ 7 . The trial is registered at ClinicalTrials.gov under NCT00930293.

Blood was collected in PAXgene tubes at study entry (T0) and after 12 weeks of treatment (T12) and processed onto Illumina HT12-v4.0 gene array. Gene functional analysis was performed using Ingenuity Pathway Analysis (<http://www.ingenuity.com>). Analysis of promoter transcription factor enrichment was performed with Enrichr (Chen *et al*, 2013). Real-time quantitative polymerase chain reaction (qPCR) was performed on a Mastercycler real-time PCR machine (Eppendorf, Hamburg, Germany) using universal PCR conditions (Sibille *et al*, 2009). Differential expression was analyzed by random intercept model with variable selection using Bayesian information criteria and class label permutation (Wang *et al*, 2012). Prediction analyses for nonremission were performed using standard machine learning method with support vector machines (with linear kernel) protocol. To correct for the selection bias of reporting the best model, we added a nested cross-validation step (Tibshirani, 2009; Varma and Simon, 2006).

For the validation cohort, we used clinical and transcriptome data from a published study (Table 1; Mamdani *et al* (2011) for details), in which MDD patients were treated with citalopram (10–60 mg, titration based on response and tolerability) for 8 weeks, with depression remission defined as HRSD-17 score ≤ 7 at week 8 assessment. Blood samples were collected at T0 and T8 in PAXgene tubes and processed onto U133 + 2.0 Affymetrix GeneChips. The top genes identified in the initial cohort predictions were used to construct the prediction model for response/remission in the validation cohort.

RESULTS

Table 1 provides the baseline demographic and clinical features of the two cohorts used in the report. A total of 34 MDD and 33 nondepressed control subjects comprised the initial cohort. Six MDD subjects dropped out of treatment prior to the week 12 (T12) blood draw. Only one of these six

Table 1 Clinical and Demographic Factors of the Initial Cohort

	Combined cohort						Test cohort				
	T0			T12							
	MDD	CTRL	Statistic	P-value	df	MDD	CTRL	Statistic	P-value	df	
Age	30.27 (10.31)	34.01 (11.15)	t = -1.42	0.16	64.2						
Sex	27 (F)/7 (M)	27 (F)/6 (M)	$\chi^2 = 0$	1.00	1						
Race	28 ^W /3 ^B /2 ^{BW} /1 ^{PI}	28 ^W /5 ^B	$\chi^2 = 3.49$	0.32	3						
Marital	22 ^A /6 ^B /6 ^D	20 ^A /9 ^B /2 ^C /2 ^D	$\chi^2 = 4.68$	0.20	3						
Education	8 ^α /13 ^β /13 ^γ	7 ^α /5 ^β /21 ^γ	$\chi^2 = 5.49$	0.06	4						
HRSD-17	18.47 (2.88)	1.36 (1.60)	t = 30.15	1.58E-34	51.8	8.61 (4.81)	2.44 (3.73)	t = 5.54	1.09E-06	50.3	
HRS-A	16.33 (4.65)	1.13 (1.36)	t = 17.97	4.33E-20	37.7	7.75 (4.37)	1.91 (2.45)	t = 6.30	1.68E-07	40.6	
QIDS-16	14.29 (3.70)	1.54 (1.20)	t = 19.06	1.08E-21	40.1						
MDD-R cohort	MDD-R	CTRL (R)	Statistic	P-value	df	MDD-R	CTRL (R)	Statistic	P-value	df	
Age	28.66 (9.64)	31.72 (9.92)	t = -0.94	0.36	34.0						
Sex	13 (F)/5 (M)	13 (F)/5 (M)	$\chi^2 = 0$	1	1						
Race	15 ^W /2 ^{BW} /1 ^{PI}	16 ^W /2 ^B	$\chi^2 = 5.03$	0.17	3						
Marital	11 ^A /5 ^B /2 ^D	14 ^A /4 ^B /0 ^D	$\chi^2 = 2.47$	0.29	2						
Education	4 ^α /8 ^β /6 ^γ	4 ^α /3 ^β /11 ^γ	$\chi^2 = 3.74$	0.15	4						
HRSD-17	17.50 (2.81)	1.11 (1.08)	t = 23.08	7.39E-17	21.9	5.83 (2.31)	2.67 (2.91)	t = 3.62	0.001	32.3	
HRS-A	15.76 (4.38)	1.06 (1.23)	t = 13.29	5.77E-11	18.7	5.50 (2.68)	2.00 (1.49)	t = 4.83	4.94E-05	26.6	
QIDS-16	11.94 (3.00)	1.56 (1.20)	t = 13.64	2.64E-12	22.3						
MDD-NR cohort	MDD-NR	CTRL (NR)	Statistic	P-value		MDD-NR	CTRL (NR)	Statistic	P-value		
Age	32.08 (11.12)	35.87 (12.21)	t = -0.86	0.40	25.8						
Sex	13 (F)/1 (M)	13 (F)/1 (M)	$\chi^2 = 0$	1	1						
Race	11 ^W /3 ^B	11 ^W /3 ^B	$\chi^2 = 0$	1	1						
Marital	9 ^A /1 ^B /4 ^D	6 ^A /4 ^B /2 ^C /2 ^D	$\chi^2 = 5.07$	0.17	3						
Education	4 ^α /4 ^β /6 ^γ	3 ^α /2 ^β /9 ^γ	$\chi^2 = 1.41$	0.49	3						
HRSD-17	19.64 (2.71)	1.71 (2.13)	t = 19.49	1.77E-16	24.6	13.60 (4.03)	1.50 (2.91)	t = 7.69	7.92E-07	16.4	
HRS-A	16.93 (5.00)	1.07 (1.49)	t = 11.37	7.27E-09	15.3	11.80 (3.91)	1.50 (3.06)	t = 6.56	4.85E-06	17.0	
QIDS-16	17.29 (2.40)	1.43 (1.22)	t = 22.03	3.73E-15	19.3						
Validation cohort											
T0					T8						
Age	38.68 (11.65)										
Sex	35 (F)/28 (M)										
Education	12.4 ^δ										
HRSD-17	30.06 (6.47)					10.35 (7.28)					

Abbreviations: HRS-A, Hamilton Rating Scale for Anxiety; HRSD-17, 17-item Hamilton Rating Scale for Depression; QIDS-16, Quick Inventory of Depressive Symptomatology.

Values are mean (± SD). Group codes are as following Sex: F, female; M, male; Self-declared race: ^Wwhite, ^Bblack, ^{BW}black,white, ^{PI}pacific islander; Marital status: ^Anever married; ^Bmarried; ^Cliving together as married; ^Dseparated/divorces. Education: ^α<high school, high school, GED, technical school; ^βsome college; ^γcollege; ^δaverage years of education.

met remission criteria prior to drop out, and was thus classified as a remitter; the remaining five were classified as nonremitters in intent-to-treat prediction analyses, in line with CONSORT clinical trial reporting standards (<http://www.consort-statement.org/>). Of those who remained at

T12, 18 were classified as remitters (MDD-R) and 10 as nonremitters (MDD-NR). Thus, 19 of 34 subjects (55.9%) were classified as treatment remitters in intent-to-treat analyses. At baseline assessment, MDD-R and MDD-NR patients did not significantly differ in age, race, education,

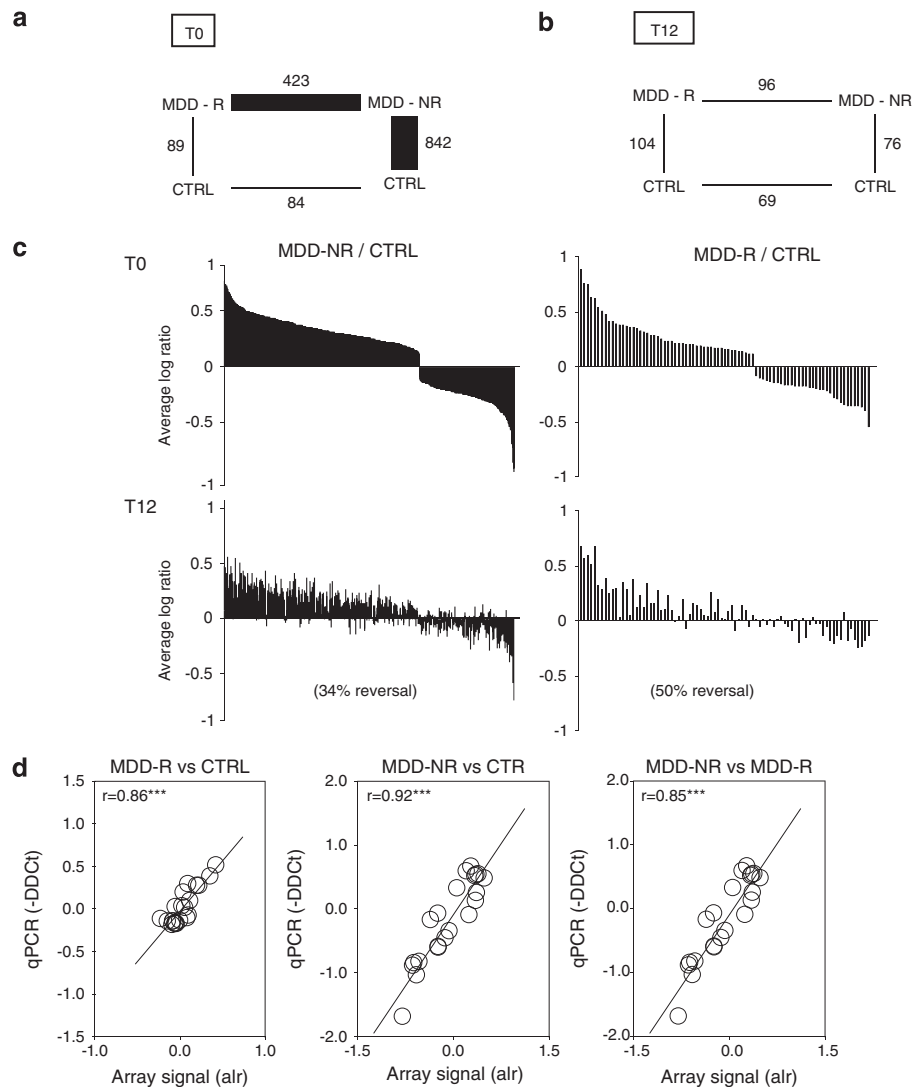


Figure 1 Peripheral gene expression in MDD. (a) At T0, MDD subjects who will not remit (MDD-NR) displayed a greater number of differentially expressed genes compared with controls (CTRL) and to MDD subjects who will remit (MDD-R). The line thickness indicates weighted numbers of differentially expressed genes between groups. The two control groups (CTRL) correspond to subjects matched to the respective MDD groups (Table 1). (b) At T12, the number of differentially expressed genes was reduced between MDD-NR and other groups, and was at the level observed between control groups, suggesting that it reflected background gene variability. (c) The T0 profiles of changes in gene transcript levels for MDD-related genes are reversed by antidepressant treatments. T0, upper panels: MDD-related genes are organized by the extent of their changes in expression along the x axis ($p < 0.01$, $FC > 20\%$). Vertical bars indicate the extent of changes [average \log_2 of MDD-NR/CTRL (Left) or MDD-R/CTRL (Right) expression ratios]. T12, lower panels: vertical bars indicate the relative transcript levels for the same MDD-related genes after citalopram/psychotherapy treatments. Genes are sorted in the same order as in upper panels. The T12 profiles correspond to a 34% reversal in MDD-NR and to a 50% reversal in MDD-R subjects (both $p < 0.0001$). (d) Validation of array results by independent qPCR measurements in MDD-R and MDD-NR subjects. See Supplementary Tables S7-8 for details. Alr: average log ratio.

anxiety (HRSA scores), or gender distribution (all p -values > 0.10). However, the MDD-NR group did display higher baseline clinician-rated and self-reported depression scores as compared with the MDD-R group (HRSD-17 = 19.6 vs 17.5, $p = 0.02$; Quick Inventory of Depressive Symptom scores, QIDS = 17.3 vs 11.9, $p < 0.01$; Table 1).

Blood Gene Expression of MDD-NR Subjects Differ from MDD-R and Control Subjects at Baseline

In the initial cohort, results from exploratory analyses of differential gene expression at baseline (pretreatment; T0)

between MDD and control subjects are summarized in Figure 1a. Few changes were observed between MDD-R and matched control subjects, compared with between MDD-NR and matched control subjects. Gene changes in MDD-R subjects did not correlate with effects in MDD-NR subjects (Pearson correlation factor, $R = -0.05$), and a direct comparison between MDD-R and MDD-NR revealed intermediate number of differentially expressed genes. False positives and/or normal background variability in gene expression was assessed as differences observed between subgroups of healthy (nondepressed) control subjects. Results from these exploratory analyses suggest that at T0,

Table 2 Biological Functions Over-represented in Differentially-expressed Gene Groups

Group comparisons	Numbers of biological groups at $q < 0.01$	Biological groups
(T0) MDD-NR/CTRL (Details in Supplementary Table S3)	154	Neutrophils: function, homing, chemotaxis, cell movement. Myeloid cells: response, cell movement, phagocytosis. Granulocytes: homing, chemotaxis, cell movement. Leukocytes: function, cell movement, migration, chemotaxis, quantity, proliferation. Phagocytes: phagocytosis, immune response, function, engulfment, homing, cell movement, response of Lymphocytes: proliferation. Immune Cells: proliferation, adhesion. Cells: chemotaxis, homing, immune response, phagocytosis Viral Infection, function of blood cells. Apoptosis of lymphoma cell lines, Inflammatory response.
(T0) MDD-NR/MDD-R (Details in Supplementary Table S4)	12	Viral Infection, replication of virus, replication of RNA virus, function of leukocytes, cell rolling of phagocytes, replication of Influenza A virus, function of neutrophils, cell death of myeloid cells, apoptosis of myeloid cells, infection of mammalia, immune response of cells, invasion of cells.
(T0) MDD-R/CTRL	0	None
(T0) CTRL(NR)/CTRL(R)	0	None
(T12) MDD-NR/CTRL	0	None
(T12) MDD-NR/MDD-R	0	None
(T12) MDD-R/CTRL	0	None
(T12) CTRL(NR)/CTRL(R)	0	None

Significant biological groups defined using an ingenuity pathway analysis were regrouped and summarized by cell types. See Supplementary Table S4 for details.

(a) untreated MDD subjects differ from control subjects, (b) MDD differences are mostly driven by MDD-NR subjects, and (c) MDD-NR subjects differ from MDD-R subjects (Figure 1a). See Supplementary Table S2 for details on individual genes.

An analysis of cell-specific markers between MDD-NR and control subjects did not suggest that these findings were supported by changes in cell-type densities (Supplementary Results and Supplementary Table S3). Biological functions identified as over-represented in MDD-NR patients at T0 compared with controls corresponded almost exclusively to increased immune function, inflammation, and white blood cell recruitment (Table 2; details in Supplementary Table S4). Moreover, transcription factor binding sites enriched in promoter regions of differentially expressed genes were similarly linked to immune and inflammatory responses, including NF- κ B-related signaling and cell cycle regulation, and/or mediation of tumor growth/suppression (Supplementary Table S6).

qPCR was performed for 21 genes with T0 differences (ADSL, ARHGEF11, CD3D, ECGF1 (TYMP), GBA, GZMA, IFITM3, IL17RA, MBOAT1, MPST, MSH6, NAP1L4, OSCAR, PRDM4, RPL4, RPL5, RPL17, RPL24, RSPH3, TBXAS1, and TIMM23). The qPCR results correlated with array data for the MDD-R vs control (Pearson correlation $r = 0.86$, $p = 5e^{-7}$), MDD-NR vs control ($r = 0.92$, $p < 1e^{-7}$) and MDD-NR vs MDD-R ($r = 0.85$, $p = 5e^{-7}$) (Figure 1d), together supporting the technical reliability of the array data, although not all individual assays reached significance. Results from individual genes are in Supplementary Tables S7-8. Results from post-treatment (T12) blood gene expression and treatment effects are summarized in Figure 1b. All T12 groups differences were at the level of background gene variability, as measured between control subjects matching MDD-R or MDD-NR subjects. Comparing T0 and T12, most genes reverted back

toward control expression levels after treatment (ie, closer to x axis in Figure 1c).

Together, the results of those exploratory approaches suggest that nonremitter subjects are biologically different from remitter subjects, hence providing a rationale for performing an independent prediction analysis for non-remission.

Prediction of Nonremission After Treatment

To predict depression remission at T12 based on T0 transcriptome data, we constructed a predictive model with a machine learning method with support vector machines. As standard practice for machine learning protocols, we started with the T0 dataset, ranked all genes based on statistical significance, and filtered the data based on effect sizes in the training set. We then systematically constructed prediction models using the top sets of genes, ranging from 2–30 genes, and applied these models to the test set (Figure 2). This process is iterated until each sample is treated as test set once. Typically, the prediction accuracy is reported for an optimal gene set; however, to address the common bias that is introduced by reporting the ‘best result’, we added an additional step to correct the results for model selection, using a nested cross-validation approach (Tibshirani, 2009; Varma and Simon, 2006) (Figure 2). The results from these studies indicate an average cross-validated accuracy (ie, model selection bias corrected) of 79.4% in predicting remission status, with the 13-gene model displaying the highest individual noncorrected prediction value (88%) (Tables 3 and 4 and Supplementary Table S9). An analysis of correlation of expression with cell-specific markers suggests that the 13-gene set is potentially, although not exclusively, enriched in genes expressed in lymphocytes (Supplementary Results and Supplementary

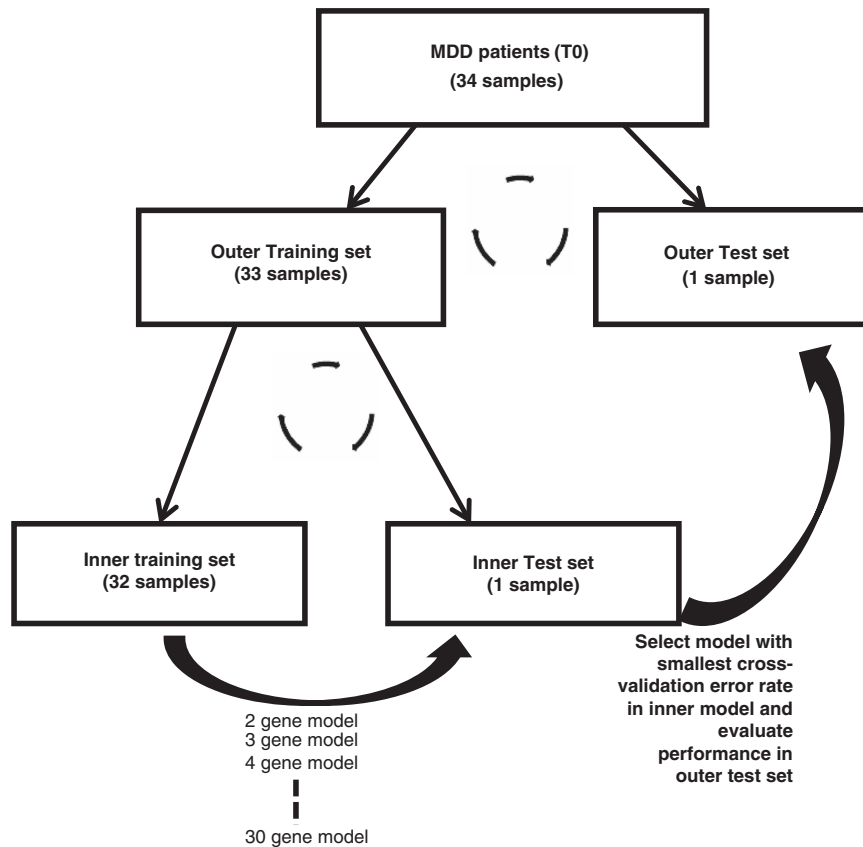


Figure 2 Flowchart of blood biomarker prediction performance assessment, including correction of model selection bias. To search for the optimal number of features (2–30 genes) in the prediction model, we applied a nested cross-validation (CV) with two nested CV loops. The dataset is initially divided into two: one sample as the test set, and the second as the training set. Then a leave-one-out cross validation (LOOCV) method is applied on the training set using all classifiers (top 2–30 genes). The classifiers with the smallest error rates are selected and used to build the model within the training set. The model is then evaluated on the left-out test sample. This procedure is repeated until all samples have been left out once. This means that the left-out test sample is independent from the model selection stage, including the selection of the model with the minimum error rate. Together, this procedure guarantees an unbiased error estimate.

Table 3 Prediction of Nonremission in MDD Patients at Baseline

Features used in prediction model	Initial cohort (genes only)	Validation cohort (genes only)
# features used in model	13	6
Corrected accuracy (%)	79.4	76.2
Sensitivity (%)	66.7	86.1
Specificity (%)	89.5	59.3

Accuracy represents the percentage of correctly identified samples. Sensitivity represents the ability to predict nonremission. Specificity represents the ability to predict remission. The corrected accuracy was obtained by nested cross-validation approach as described in Figure 2.

Table S10), highlighting the value of using whole blood transcriptome for predictive purposes.

Validation of Gene-based Prediction of Nonremission in an Independent Cohort

Given the small size and unique features of the initial cohort and the potential bias in model selection, we sought to

confirm the prediction model in an independent cohort recruited and tested at another institution. We obtained an archival dataset drawn from a study that evaluated gene expression in blood from 63 MDD subjects who were subsequently treated for 8 weeks with similar doses of citalopram (see Table 1 and Mamdani *et al* (2011) for details). Thus, this study represented a close, but not identical, match to our initial cohort. We built a new prediction model in the validation cohort using the same 13 genes identified in the initial cohort, and found through another round of leave-one-out cross validation that a 6-gene model achieved the highest accuracy (76.2%) in the validation cohort (Table 4 and Supplementary Table S11).

Exploratory Analyses

Predictive models including clinical data. We explored the potential of clinical features to augment the gene-based prediction models. These analyses were performed in the initial cohort only, because of the availability of multiple clinical scales in that study (Supplementary Table S1). We first applied machine learning prediction methods using clinical scale data only, considering models with 2–30

Table 4 Top Genes Included in the Prediction Models

Gene symbol	Gene name	Frequency of use in initial cohort	Frequency of use in validation cohort
CD3D	CD3d molecule, delta (CD3-TCR complex)	34	46
CD97	CD97 molecule	34	
IFITM3	Interferon induced transmembrane protein 3	34	63
RPL5	Ribosomal protein L5	34	
GZMA	Granzyme A	33	63
TAGLN2	Transgelin 2	33	
TIMP1	TIMP metalloproteinase inhibitor 1	33	
RPL24	Ribosomal protein L24	29	63
PSMA4	Proteasome (macropain) subunit, alpha type, 4	25	
MATR3	Matrin 3	18	63
RPL9	Ribosomal protein L9	15	
PSMA6	Proteasome (macropain) subunit, alpha type, 6	12	
RPL17	Ribosomal protein L17	7	58

Genes in bold were included in the independent qPCR validation (Figure 1). Consistent with the previous functional and transcription factor analyses, this 13-gene model included genes associated with immune/inflammatory activation (CD3D, CD97, IFITM3, and GZMA) and mediation of cell proliferation (GZMA and TIMP1), although not exclusively.

clinical variables. The following three baseline clinical variables provided optimal accuracy in predicting T12 remission status, with corrected accuracy of 70.6% as determined by nested cross validation (as in Figure 2): clinician-rated depressive symptoms (HRSD-17 scores), patient-reported depressive symptoms (QIDS (Rush *et al*, 2003)), and perceived stress scores (4-item Perceived Stress Scale (Cohen *et al*, 1983)). Next we tested models including both clinical and gene expression data and show that T12 depression nonremission could be predicted with a corrected accuracy of 97% based on T0 gene expression and clinical data (Supplementary Tables S12-13), in a model including just three features: two genes (IFITM3 and TIMP1) and one clinical feature (QIDS). Notably the type of adjunctive psychotherapy treatment provided in that cohort in addition to citalopram (IPT-PS or BSP) did not contribute to any of the prediction models.

Treatment response trajectory. Because MDD-NR and MDD-R patients differed significantly in baseline depression scores (Table 1), we explored whether observed differences in posttreatment remission status represented differences in treatment trajectory. Figure 3 shows a large overlap in baseline depression scores between the MDD-NR and MDD-R groups, and significant group differences in slopes between T0 and T12 in the initial cohort ($p < 0.00137$) and T0 and T8 in the validation cohort ($p < 1.25e^{-6}$), hence confirming the presence of a different trajectory of treatment response in MDD-NR subjects. Consistent with the overlap in values (Figure 3), baseline HRSD-17 values provided moderate prediction for nonremission (67.6% in initial and 57.1% in validation cohort).

DISCUSSION

Results from the first and exploratory phase of this study suggest that at pretreatment assessment, the gene

expression profiles obtained from blood samples of MDD subjects who will not attain remission after treatment (MDD-NR group) differ from nondepressed controls and also from MDD patients who will remit with treatment (MDD-R) (Figure 1). The nature of genes affected in MDD-NR subjects suggests a state of inflammation and immune system activation in those patients, although this was not confirmed by independent approaches (Table 2). In the second phase of the study, using a machine learning prediction model and a leave-one-out cross validation to correct for multiple model testing, we show that pretreatment baseline gene expression predicted nonremission after treatment with 79.4% accuracy (Table 3), based on a 13-gene panel (Table 4). A strength of this study is that we confirmed our results in an independent validation cohort. Specifically, we showed that 6 out of 13 genes identified in the initial cohort could predict remission in an independent cohort ($n = 63$ MDD patients) with 76.2% accuracy (Table 3). Together, this report demonstrates the potential, but also the limitations, of pretreatment peripheral gene expression profiles to predict nonremission following an 8- to 12-week course of citalopram treatment.

Although the sample size of the initial cohort was small, the gene expression prediction results replicated in an independent cohort, despite differences in experimental design (8- vs 12-week drug exposure, no psychotherapy) and inclusion of nonanxious MDD patients in the validation cohort. It is worth noting that we did not apply the optimized 13-gene model from our cohort directly to the validation cohort study, but instead used those top 13 candidate markers to perform a new model selection in the independent cohort with feature selection and leave-one-out cross validation to assess the prediction accuracy. This approach was designed to facilitate the analysis and mitigate differences across the two cohorts and platforms, and it may explain why genes differentially expressed in specific cohorts (see introduction) may not necessarily be included in the final common set of predictors. For future

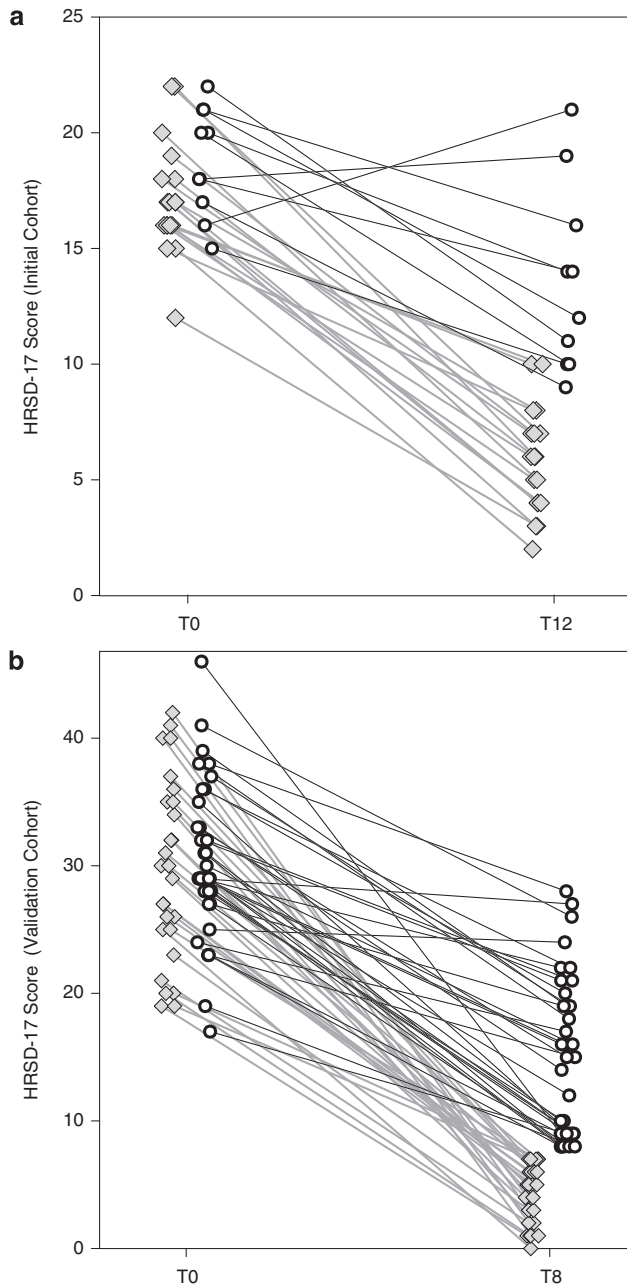


Figure 3 Trajectories of HRSD-17 scores between MDD-NR and MDD-R subjects in the initial and validation cohorts. Grey squares indicate MDD-R subjects. Empty black circles indicate MDD-NR subjects.

translational and clinical utility, a randomized prospective clinical trial using a preestablished predictive model is the ultimate validation. To date, however, direct application of predictive models to different test studies has been difficult due to cross-laboratory variability in probe designs from different array platforms and different experimental protocols. Such issues could be addressed in the future through the use of carefully standardized experimental protocols or commercialized products. For example, prognostic prediction of breast cancer treatment (Cheng *et al*, 2009; Shen *et al*, 2004; Xu *et al*, 2008) has been greatly facilitated by the use of MammaPrint (van't Veer *et al*, 2002) and Oncotype

DX (Paik *et al*, 2006), two successful microarray and qPCR examples with cross-laboratory standardization.

The combination of clinical scales and gene expression data enhanced the accuracy of the nonremission prediction to 97% in the initial cohort (Table 3 and Supplementary Table S11). Notably, of all clinical scales included in the combined machine learning models (including those assessing current and lifetime mood and anxiety symptoms, childhood trauma, and social/interpersonal function; see supplements and Supplementary Table S1), the only scale retained in the combined prediction model was the QIDS-16; a patient-reported indicator of depressive symptom severity. This finding may highlight the importance of assessing subjective levels of patient-reported distress which may, moreover, be more closely linked to over-activation of pro-inflammatory pathways (Cole, 2010; Miller *et al*, 2008; Pace and Miller, 2009).

What could we learn from the exploratory analyses of differential gene expression? We are cognizant of the high false positive rate associated with the uncorrected statistical approaches used here, so we briefly discuss results at the gene group level rather than for independent genes. As hypothesized, biological functions represented by genes showing baseline expression differences largely suggest pretreatment elevations in inflammation and immune activation in MDD-NR subjects (Table 2). This may, in part, relate to baseline elevations in symptoms of depression and of perceived stress in the MDD-NR group. In this regard, MDD-NR subjects also displayed heightened expression of numerous transcripts with response elements for NF- κ B, a proinflammatory transcription factor associated with the experience of chronic stress (Cole, 2010; Miller *et al*, 2008; Pace and Miller, 2009). The fact that the prevalence of these predictive gene markers was reduced after treatment despite elevated HRSD scores (Table 2) suggests that while inflammation and immune-related pathways could have contributed to MDD onset, their presence may not be necessary for the maintenance of MDD in those subjects. Alternatively, putative remodeling of brain circuitry and other molecular/neurochemical changes (such as changes in glucocorticoid receptor sensitivity) may sustain the depressive state despite the absence of continued peripheral immune and inflammatory gene markers. Finally, it is worth noting that the extent of correlation of expression between the predictive genes and cell-specific markers suggests the contribution of various cell types. Future studies may address the contribution of changes within and across specific leukocyte cell populations, but the current results suggest that using total blood gene transcripts may be appropriate for predictive purpose.

The finding that MDD-NR subjects showed differential gene expression and trajectories of reduction in HRSD-17 scores following treatment (Figure 3) supports the intrinsic difference of this group compared with other MDD subjects. Future studies will need to characterize those differences on longer timescales and to examine the association between the observed pretreatment gene expression profiles and functional indicators of systemic inflammation or stress-induced immune/inflammatory reactivity (Maes, 1995; Zunszain *et al*, 2013). Conversely, subjects who remitted after treatment (MDD-R) had pretreatment gene expression profiles similar to background variability in control

subjects, demonstrating that diagnostic MDD status is not necessarily accompanied by peripheral gene changes (at least with the technical sensitivity of the gene arrays used here). This observation highlights the underlying biological heterogeneity in MDD subjects, and underscores the need for future mental health biomarker research to move beyond traditional diagnostic categories, as articulated in the NIMH Research Domain Criteria initiative (Insel *et al*, 2010).

Together, the current and prior studies (see introduction) not only support the biomarker potential of peripheral gene expression in predicting MDD treatment outcome but also highlight the need for additional studies. Future studies will need to be performed in larger cohorts and may need to combine measures of gene expression with selected clinical scales, specifically related to mood symptoms and chronic life stress. The inclusion of genetic information, measures of peripheral metabolites and treatment history may further increase the predictive value of those assays. For instance, Gudayol-Ferre *et al* (2013) showed that a combination of clinical, genetic (val/Met COMT polymorphism) and neuropsychological variables were associated with a pattern of remission in MDD subjects. The fact that in our study baseline peripheral gene expression did not differ between MDD subjects who remitted and healthy controls also suggests that different sets of biomarkers may be needed to capture the heterogeneity of MDD subjects. The findings from this study show the potential of blood gene expression to identify prior to treatment—with high sensitivity and specificity—a population of individuals that would benefit from treatment augmentation. Future studies will need to assess means of treatment augmentation in the context of prediction of nonremission. Whether gene expression biomarkers will provide insight into disease mechanisms for a rational choice of drug augmentation is currently unknown, so the added value of those assays will have to be determined based on optimized design and better knowledge of successful augmenting factors, but the levels of accuracy and confirmation obtained under the current conditions are promising. Finally, future treatment paradigms will benefit from determining whether other antidepressants are successful in treating the nonremitting patients identified in this study.

Notes and Limitations

Several limitations to the current study are noteworthy. First, the initial cohort sample size was small and thus potentially associated with high false positive rates for both differential expression and prediction analyses. Second, with the exception of the Mamdani *et al* (2011) study, direct comparisons of prediction results with prior reports (see Introduction) were not performed due to differences in study goals (eg, disease status *vs* outcome prediction here). Third, the lack of available anxiety measures in the validation cohort may have limited the analysis in that cohort. Fourth, we did not attempt to separate blood cells and instead investigated transcriptome profiles at the time of collection, so that expression of putative markers would not be influenced by sample manipulation. Instead, using prior results, we ruled out the possible interference of blood cell proportions on differential expression of genes. Fourth, the

biological validity of the inflammation and immune activation was not confirmed by other biological tests and is thus speculative at this time. Fifth, the study was not designed to assess later time points regarding delayed remission and potential biomarkers for future relapse.

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DISCLAIMER

The content is solely the responsibility of the authors and does not necessarily represent the official views of the NIMH or the National Institutes of Health.

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-- Supplement Material --

**Peripheral gene expression predicts non-remission following
citalopram treatment for major depression**

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DETAILED MATERIAL AND METHODS

Initial cohort: Subjects and treatment

Subjects. (**Table 1**) 34 anxious-depressed adults were recruited for an outpatient trial of combined medication and psychotherapy treatment (NIH-MH085874, J. Cyranowski, PI), at the Western Psychiatric Institute and Clinic (WPIC) Depression and Manic-Depression Treatment Program (Pittsburgh, PA). The patient sample met current criteria for a non-psychotic, MDD episode of sufficient severity, as documented by the Structured Clinical Interview for Axis I, DSM-IV Disorders (SCID) and by a rating of ≥ 15 on the 25-item HRSD (Cyranowski *et al*, 2005; Hamilton, 1960). Patients were also required to report significant co-occurring panic or anxiety symptoms, as determined by a score ≥ 7 on the past-month Panic and Agoraphobic Spectrum Self-Report (PAS-SR) (Cassano *et al*, 1997c; Markowitz *et al*, 2005; Rush *et al*, 2003; Shear *et al*, 2001a). By contrast, non-patient controls (n=33) assessed with the same scales did not meet criteria for any SCID-diagnosed mood or anxiety disorder. Individuals with a history of schizophrenia, bipolar disorder, or antisocial personality disorder; primary diagnosis of anorexia or bulimia nervosa; drug or alcohol abuse/dependence in the past three months; severe, uncontrolled medical illness; a history of nonresponse to an adequate trial of citalopram; and women who were pregnant or planning to become pregnant were excluded from study participation. All subjects were free of antidepressant treatment prior to study start, including 4 subjects with wash-out periods (3 weeks, 2 weeks and 8 days (2 subjects)). Study procedures were approved by the University of Pittsburgh Institutional Review Board (IRB). All subjects provided written, informed consent after receiving a complete description of the study and prior to study participation.

Treatment and blood sampling. Citalopram was administered using a flexible titration schedule, starting at 10 mg per day and, if tolerated, increased daily by 10 mg increments until stabilization was achieved or until a daily dose of 60 mg was reached. At 12-weeks, the average

citalopram dose was 37.5 +/- 10.5 mg. In addition to medication, patients received weekly 50-minute sessions with one of two randomly-assigned manualized psychotherapies: Interpersonal Psychotherapy for Depression with Panic and Anxiety Symptoms (IPT-PS) (Cyranowski *et al*, 2005) a version of IPT for depression modified to explicitly target both depression and co-occurring anxiety symptoms, or a standardized depression psychotherapy control, Brief Supportive Psychotherapy (BSP) (Hellerstein *et al*, 1998; Markowitz *et al*, 2005). Whole blood (2.5 ml) was collected in PAXgene tubes at entry (T0) and after 12 weeks of citalopram treatment (T12), and stored at -80°C.

Initial cohort: Measures

Mood and Anxiety Symptoms. Clinician ratings of patients' current (past-week) symptoms of depression and anxiety were evaluated with the 25-item HRSD (which allowed for calculation of traditional 17-item HRSD scores, more atypical presentations of depression, and factor-derived HRSD subscales for core mood, anxiety and sleep items(Hamilton, 1960; Shafer, 2006)) as well as the 14-item Hamilton Rating Scale for Anxiety, (HRSA) (Shear *et al*, 2001c). Treatment progress was evaluated weekly with 17-item HRSD scores (Hamilton, 1960) with remission defined as a mean HRSD-17 score ≤ 7 over three consecutive weeks. Self-reported assessment of mood and anxiety symptoms were also obtained with the Quick Inventory of Depressive Symptoms–Self Report (QIDS-SR) (Rush *et al*, 2003), the Mood and Anxiety Symptom Questionnaire (MASQ) (Watson *et al*, 1995a; Watson *et al*, 1995b), and the Anxiety Sensitivity Scale (ASI) (Reiss *et al*, 1986).

Baseline Social / Role Function. Patients also provided self-report data regarding baseline levels of social role function including, for example, measures of life enjoyment and satisfaction (14-item Quality of Life Enjoyment and Satisfaction Questionnaire (QLESQ) (Endicott *et al*, 1993)), social support (12-item Interpersonal Support Evaluation List (ISEL) (Cohen S *et al*,

1985)), perceived stress (4-item Perceived Stress Scale (PSS) (Cohen *et al*, 1983)), and perceived mastery (7-item Perceived Sense of Mastery scale (Pudrovska *et al*, 2005a; Pudrovska *et al*, 2005b)). Interpersonal difficulties or dysfunction were also assessed with the 5-item Negative Interaction Scale (NIS) (Krause, 1995) and a 25-item short-form of the Inventory of Interpersonal Problems (IIP) (Horowitz *et al*, 1989).

Childhood Trauma or Abuse. Patient-reported levels of abuse or neglect experienced prior to age 18 were assessed with subscales of the 28-item Childhood Trauma Questionnaire-Short Form (CTQ) (Bernstein *et al*, 2003).

Lifetime Mood and Anxiety Symptoms. Lifetime mood and anxiety symptom loadings (many of which have been associated with poorer depression treatment outcomes in previous clinical trials (Frank *et al*, 2011; Frank *et al*, 2000)) were assessed using four self-report instruments: the Mood Spectrum scale (MOODS-SR) (Dell'Osso *et al*, 2002a) the Panic-Agoraphobic Spectrum scale (PAS-SR) (Cassano *et al*, 1997a, b; Shear *et al*, 2001a, b), the Obsessive-Compulsive Spectrum scale (OBS-SR) (Dell'Osso *et al*, 2002b) and the Social Phobia spectrum scale (SHY-SR) (Dell'Osso *et al*, 2002b; Fournier *et al*, 2012). All self-report and assessment scales are summarized in **Table S1**.

Validation cohort

Clinical and transcriptome data from a previous study were used (**Table 1**). See Mamdani *et al* (Mamdani *et al*, 2011) for details. Briefly, subjects were recruited from a community outpatient clinic at the Douglas Mental Health University Institute, Montreal, Canada. 63 patients were analyzed, consisting of males and females with a diagnosis of MDD without psychotic features, according to the SCID. Subjects did not previously use citalopram and were not treated for their current episode if they had a recurrent form of MDD. Patients were treated with citalopram (initial dose of 10 mg per day, titrated up to a maximum of 60mg per day) for 8 weeks. Assessments of depression severity were carried out at pretreatment (T0), one month after (T4)

and post-treatment (T8) using the HRSD-17 rating scale. Blood samples were collected at T0 and T8 in PAXgene blood RNA tubes. All subjects provided informed consent and the project was approved by the internal review board for the Douglas Mental Health University Institute.

Microarray samples

Initial cohort. Whole blood RNA was extracted using the PreAnalytix kit (Qiagen), verified for quality (A260/A280 ratio >2) and integrity (RNA integrity number >7; Agilent Technologies, Palo Alto, CA, USA). A GlobinClear (Invitrogen, Carlsbad, USA) treatment was applied to deplete globin mRNA transcripts from total RNA samples. Samples were processed for hybridization onto Illumina HT12 human array v4.0, allowing for the analysis of 47,231 probes.

Validation cohort. Whole blood RNA was processed for hybridization onto Affymetrix GeneChip Human Genome U133 Plus 2.0 array (Affymetrix, Santa Clara, CA, USA), consisting of 54,000 probes. Quality control for all array data included normalization for potential batch effects and cross-correlation and principal component analyses for the putative presence of outlier samples. All samples passed the quality control and were included in the analyses. See details in Mamdani et al (Mamdani *et al*, 2011).

Gene functional analysis

Gene lists were analyzed using Ingenuity Pathway Analysis (IPA), using Holm-Bonferroni corrected p-value <0.01 for over-representation of biological functions within gene groups.

Real-time quantitative polymerase chain reaction (qPCR)

500ng total RNA was converted into cDNA using the qScript kit (Quanta, Gaithersburg, MD). Small PCR products (70-160 base-pairs) were amplified in quadruplets on a Mastercycler real-time PCR machine (Eppendorf, Hamburg, Germany), using universal PCR conditions (Sibille *et al*, 2009). Results were calculated as the geometric mean of relative intensities compared with

two validated internal controls (actin and glyceraldehyde-3-phosphate dehydrogenase).

Statistical analysis

Antidepressant reversal. Antidepressant reversal of MDD effect represents the extent by which transcripts measured at T0 were brought back to control level at T12. Percentage of reversal was calculated using “0%” as no change in expression over time compared to control, and “100%” meaning that gene transcript levels were back to control levels at T12. Reversal was capped at 100% for genes whose antidepressant-related changes were in opposite directions. Gene-wise T0 and T12 values were compared by paired t-tests in the MDD-NR and MDD-R groups.

Differential analysis to detect candidate marker genes. Subjects with T0 and T12 expression data were included for analyses of differential expression to compare pre- to post-treatment (n=28/group). After filtering for low expression and variance, 11,797 probes were retained. A random intercept model with variable selection was applied to detect candidate marker genes for all two-group comparisons. We described this method in Wang et al (Wang *et al*, 2012). Specifically, a random intercept model was used to accommodate disease-control paired information and covariates (age, gender and dosage) to detect differential effects. All possible RIM models that included at most two (i.e. 0, 1 or 2) clinical variables were computed and compared. The model with the smallest Bayesian Information Criterion (BIC) value was selected. This additional variable selection avoided the inclusion of more than 2 clinical variables in the model and allowed the inclusion of different sets of covariates for each gene, which offers more relevant biological conclusions and interpretations. The obtained p-values from the best BIC model were, however, not corrected p-values for differential expression since they were biased by the variable selection procedure and the type I error control was voided. As a result, a permutation analysis that randomly shuffled the disease labels within each pair was performed

to generate a null distribution for p-value assessment (B=500). Subsequently, the permutation-corrected and unbiased p-values were further adjusted by Benjamini-Hochberg procedure for multiple comparisons within each study to control the false discovery rate (FDR).

Prediction analysis-model building. To investigate the prediction power of genomics features on predicting MDD status and non-remission status in 12 weeks, support vector machine (SVM) with feature selection was applied to construct the models. SVM has been shown to give excellent performance in comparative study (Lee *et al*, 2005). We use SVM as the classification algorithm with a linear kernel (implemented in R package e1041). Feature selection is applied on the 11797 probes with 4 criterions: (i) p-value calculated from standard t test between two class in the training samples, (ii) effect size (difference of average of log₂ expression between two groups), (iii) the background probe variability, as identified by differentially-expressed between CNR and CR at both time 0 and time 12; (iv) the probes should correspond to a matched gene symbol and should also have a matched gene in the independent cohort. In selecting the genes to construct model, probes were first ranked by p-value, then the log₂ fold changes of less than 0.5 was filtered away as well as the probes associated with background variation. This is because small log₂ fold change could result in false positives (Spijker *et al*, 2010). Probes without corresponding gene symbol or matched genes in the independent cohort were further filtered. The background variation genes were defined as probes between CNR and CR at time 0 and time 12 with p value smaller than 0.05. In this study, 766 probes were obtained as background variation probes. We attempted different models with features from 2 to 30.

Prediction analysis-performance assessment. N-fold cross validation is commonly used to evaluate the performance of individual classifier. In this study, we evaluated different models with leave-one-out cross validation (LOOCV). However, when multiple models are considered in the study, the model selection bias is often ignored by simply reporting the model with the

highest CV accuracy. In this study, we not only assessed the performance of each classifier with leave-one-out cross validation but we also report the performance by correcting the potential model selection bias by applying nested-cross validation (Varma and Simon, 2006). For each iteration of nested-cross validation, 33 of the 34 samples are treated as training samples and the rest 1 sample is treated as the independent test sample. The models are only evaluated within the 33 training samples with another round of leave-one-out cross validation. The model with the best CV accuracy is then selected and applied to the independent test sample. The whole process was iterated 34 times and the average performance is reported. The performance evaluation workflow is illustrated in **Table S8**. The 9 and 13 genes features can achieve 88.2% accuracy; however, the accuracy could be biased upward because multiple classifiers are attempted. The corrected accuracy with nested CV is 79.4%. If 47 baseline clinical variables are included together with the genomic features for model selection, a 3-feature model is most frequently selected in nested CV, with a corrected accuracy of 97%, the same as the performance of 3-feature model without nested CV correction.

Validation in an independent cohort with the 13 top genes. In the prediction analysis of the initial cohort, we concluded that the 13 gene model provided the best performance. Therefore, we applied the top 13 genes selected in the initial cohort to an independent cohort of 63 samples from Mamdani et al (Mamdani *et al*, 2011) publication. The original study was designed to examine the difference of responder versus non-responder groups, instead of remission status. We obtained the HAMD 17 item score and divided the samples into 36 remitted and 27 non-remitted samples by the HAMD score 7 cutoff which is consistent with our definition. We applied the same feature selection criteria in the prediction model construction: (i) p value, (ii) effect size and (iii) background variation genes. For assessing the predictability of the 13 genes on the new cohort, we performed an additional round of feature selection due to the larger cohort and different platform of the gene expression arrays (Illumina HT12-v4.0 in initial cohort and

Affymetrix U133 plus in validation cohort). In the end, we selected 6 out of 13 genes to achieve an optimal 76.2% accuracy (with nested CV correction, see **Table S9**) and we showed that the effect sizes of the top frequent features involved in the 6-gene model in both the initial and validation cohorts are highly consistent and correlated.

Supplemental Results

Inference of blood cell-type proportions

MDD-related genes: To address the possibility that our results may reflect differences in the cell pool rather than differences in per-cell gene expression, we estimated the distribution per blood cell compartment of our 842 differentially expressed genes between MDD-NR subjects and CTRLS (at T0). Percent of genes identified as enriched in various cell types were assessed and compared with previous findings (Abbas *et al*, 2005). Results show statistical differences ($\chi^2=41.1$, $p<0.0001$, **Table S3**), although the percentage changes are modest (10% at best). Importantly for our analysis, over 50% of the genes appear to be expressed in various blood cell types, and 45% of genes are expressed across different cell groups.

Biomarkers of non-remission (13 genes list): Due to the smaller group size we used an alternative approach and assessed putative cellular origin of transcripts for the 14 genes using Pearson correlation with known cellular markers. Based on a (Palmer *et al*, 2006) reported list of genes selectively enriched in specific cell types, we measured the proportion of genes within a blood-cell population that displayed correlated patterns with each of the 13 predictive genes using data from T0 and T12. 9 out of the 13 genes (CD3D, GZMA, MATR3, PSMA4, PSMA6, RPL17, RPL24, RPL5, RPL9, **Table S9**) displayed correlated expression with lymphoid-related genes. However this was not specific, as the same genes also moderately correlated with expression of genes belonging to other cell types. Moreover 4 of the 13 genes (CD97, IFITM3, TAGLN2, TIMP1, **Table S9**) showed correlated patterns with granulocytes-related genes, together confirming that the genes used in our model do not belong to a specific cell compartment.

SUPPLEMENTARY TABLES

Table S1. Self-report and assessment scales

Demographic Variables	
AGE	
SEX	
RACE	
MARITAL	
EDUCATION	
BODY MASS INDEX	
INCOME	
Clinical Variables	Assessment Scales
Clinician-rated mood symptoms	17-item Hamilton Depression Rating Scale (HRSD-17)
	HRSD - core mood symptoms
	HRSD - core anxiety symptoms
	HRSD - core sleep symptoms
Clinician-rated anxiety symptoms	HRSA (Hamilton Rating Scale for Anxiety)
Self-reported depression	Quick Inventory of Depressive Symptomatology (QIDS-SR)
Self-reported anxiety sensitivity	Anxiety Sensitivity Scale (ASI)
Mood and Anxiety Symptom Questionnaire (MASQ) self-report subscales	MASQ General Distress-Anxiety
	MASQ General Distress-Depression
	MASQ Anxious Arousal
	MASQ Anhedonic Depression
Lifetime panic symptoms	Panic-Agoraphobic Spectrum Scale (PAS-SR) - total
Lifetime mood symptoms	Mood Spectrum Scale (MOOD-SR) - total
Lifetime OCD symptoms	Obsessive-Compulsive Spectrum Scale (OC-SR) - total
Lifetime social phobia symptoms	Social Phobia Spectrum Scale (SHY-SR) - total
Baseline Social / Role Function	
Self-reported social/role function	Negative Interaction Scale (NIS)
	Interpersonal Support Evaluation List (ISEL)
	Quality of Life Enjoyment and Satisfaction (QLESQ) scale
	Perceived Stress Scale (4-item PSS)
	Perceived Sense of Mastery (PMAST) scale
Inventory of Interpersonal Problems (IIP) subscales	IIP Interpersonal Sensitivity
	IIP Interpersonal Ambivalence
	IIP Aggression
	IIP Need for Social Approval
	IIP Lack of Sociability
	IIP Personality Disorder screener
Childhood Trauma or Abuse	
Childhood Trauma Questionnaire (CTQ) subscales	CTQ Emotional Abuse
	CTQ Physical Abuse
	CTQ Sexual Abuse
	CTQ Emotional Neglect
	CTQ Physical Neglect

Additional information included in the prediction models included: Treatment type, number of lifetime depressive episodes, age at first depressive episode, duration (in weeks) duration of current episode.

Table S2. Differential expression of genes between MDD, control, MDD-NR, and MDD-R at T0 and T12.

See separate excel tabl

Note that probesets are indicated in the supplemental tables, which were reduced to gene numbers in the manuscript.

- **Worksheet a)** Differential expression of genes between MDD and control, MDD-NR and control, MDD-R and control and MDD-NR and MDD-R at **T0**;
- **Worksheet b)** Differential expression of genes between MDD and control, MDD-NR and control, MDD-R and control and MDD-NR and MDD-R at **T12**
- **Worksheet c)** Differential expression of genes between MDD, MDD-NR and MDD-R at **T12 vs T0**.

Table S3: Blood cell group distribution of genes differentially expressed at T0 between MDD-Non remitter (NR) versus controls

	B-cells	Dendritic cells	Lymphoids	Monocyte	Multiple	Myeloid	Neutrophils	NK-cells	T-cells
Number of genes in whole blood (%)	93 (5.2)	74 (4.1)	241 (13.4)	84 (4.7)	821 (45.8)	343 (19.1)	45 (2.5)	16 (0.9)	77 (4.3)
Number of genes in our dataset (%)	0 (0)	2 (1.6)	3 (2.4)	7 (5.6)	67 (53.6)	37 (29.6)	8 (6.4)	0 (0)	1 (0.8)

Table S4: Functional analysis in MDD-Non remitter (NR) versus controls

See separate excel table

Table S5: Functional analysis in MDD-Remitter (R) versus controls

[See separate excel table](#)

Table S6. Transcription factor analysis. p-value are from fisher's exact test using the Enrich-R software (Chen *et al*, 2013).

Term	P-value	Adjusted P-value
PPARG	1.08E-17	3.16E-15
E2F1	3.28E-17	4.82E-15
RELA	1.67E-15	1.23E-13
TP53	8.26E-15	4.86E-13
FOXC1	6.40E-13	2.69E-11
PCBP1	2.37E-12	7.75E-11
RUNX1	3.01E-13	1.48E-11
CREB1	1.08E-12	3.99E-11
NR1H3	3.43E-11	8.41E-10
WT1	1.09E-11	2.92E-10
NR5A2	7.58E-12	2.23E-10
SP3	2.23E-10	4.69E-09
NFKB1	7.64E-11	1.73E-09
SP1	3.79E-10	7.44E-09
HIF1A	4.58E-10	7.92E-09
TCFAP2A	1.52E-09	2.13E-08
RARA	9.31E-10	1.44E-08
E2F6	1.17E-09	1.71E-08
GATA2	5.29E-09	5.98E-08
ZBTB7A	1.81E-09	2.31E-08
SRF	8.12E-10	1.33E-08
NR1I2	6.13E-09	6.67E-08
SREBF1	5.03E-09	5.92E-08
ETS1	1.62E-08	1.70E-07
HINFP	4.89E-09	5.92E-08
ATF2	3.94E-08	3.73E-07
NFE2	3.13E-08	3.11E-07
RELB	1.13E-07	1.00E-06
PITX2	3.17E-08	3.11E-07
ELK1	1.49E-07	1.29E-06
MTF1	2.42E-07	1.98E-06
JUN	9.23E-08	8.48E-07
KLF5	2.73E-07	2.11E-06
TEAD4	2.72E-07	2.11E-06
SPI1	3.28E-07	2.41E-06
MIR138	2.86E-07	2.16E-06
RBPJ	5.28E-07	3.70E-06
AHR	6.63E-07	4.53E-06
CEBPD	8.64E-07	5.29E-06
MAX	7.86E-07	4.91E-06
IRF8	7.37E-07	4.71E-06
NFYA	3.56E-07	2.56E-06
TFAP2A	9.84E-07	5.90E-06
RXRA	1.35E-06	7.93E-06
SNAI2	3.05E-06	1.50E-05
TCF3	3.05E-06	1.50E-05
SNAI1	3.05E-06	1.50E-05
GF11	2.99E-06	1.50E-05
LTF	1.43E-06	8.26E-06
NRF1	2.46E-06	1.36E-05

Table S7: Validation of microarray results by qPCR (with the primer sequences). Effect size are in average \log_2 ratio of (MDD-R/control), (MDD-NR/control) and (MDD-NR/MDD-R).

Red shadow indicates up-regulation. Green shadow indicates down-regulation. Grey shadow indicates p<0.05.

Gene Symbol	MDDR vs control t0				MDDNR vs control t0				MDDNR vs MDDR t0			
	microarray		qPCR		microarray		qPCR		microarray		qPCR	
	alr	p-value	alr	t-test paired	alr	p-value	alr	t-test paired	alr	p-value	alr	t-test unpaired
ADSL	0.05	0.427	0.02	0.496	-0.24	0.002	-0.60	0.001	-0.33	0.0001	-0.48	0.002
ARHGEF11	-0.06	0.237	-0.17	0.066	0.35	0.0001	0.54	0.021	0.21	0.001	0.37	0.006
CD3D	0.22	0.144	0.28	0.229	-0.54	0.037	-0.82	0.023	-0.59	0.004	-0.76	0.009
ECGF1 (TYMP)	-0.09	0.459	-0.14	0.258	0.20	0.152	0.60	0.142	0.44	0.001	0.80	0.025
GBA	-0.01	0.941	-0.13	0.139	0.35	0.002	0.14	0.164	0.32	0.001	0.15	0.153
GZMA	0.35	0.006	0.39	0.242	-0.58	0.022	-1.03	0.035	-0.64	0.002	-0.65	0.037
IFITM3	-0.23	0.476	-0.11	0.510	0.36	0.499	0.26	0.307	1.28	0.000	0.51	0.034
IL17RA	-0.09	0.486	-0.14	0.136	0.39	0.005	0.55	0.002	0.46	0.003	0.37	0.004
MBOAT1	-0.05	0.737	-0.17	0.079	0.24	0.001	-0.09	0.191	0.31	0.0001	0.09	0.233
MPST	-0.06	0.444	-0.16	0.342	0.27	0.009	0.67	0.149	0.28	0.002	0.67	0.042
MSH6	0.02	0.719	0.04	0.441	-0.13	0.105	-0.45	0.016	0.19	0.007	-0.37	0.002
NAP1L4	0.04	0.558	0.20	0.453	-0.25	0.003	-0.06	0.253	-0.27	0.001	0.03	0.365
OSCAR	-0.06	0.512	0.03	0.421	0.48	0.002	0.49	0.006	0.41	0.001	0.25	0.015
PRDM4	0.08	0.136	-0.10	0.105	-0.07	0.389	-0.34	0.001	-0.19	0.001	-0.26	0.013
RPL17	0.42	0.121	0.52	0.224	-0.80	0.022	-1.68	0.006	-0.79	0.013	-1.16	0.005
RPL24	0.09	0.514	0.30	0.270	-0.62	0.012	-0.84	0.054	-0.59	0.006	-0.96	0.012
RPL4	0.20	0.112	0.28	0.313	-0.24	0.045	-0.59	0.003	-0.46	0.001	-0.60	0.008
RPL5	0.11	0.444	0.10	0.422	-0.64	0.01	-0.88	0.007	-0.73	0.002	-0.76	0.027
RSPH3	-0.15	0.005	-0.14	0.094	0.05	0.333	0.33	0.132	0.24	0.001	0.57	0.002
TBXAS1	-0.10	0.302	-0.18	0.238	0.34	0.026	0.52	0.021	0.34	0.001	0.56	0.01
TIMM23	0.10	0.117	-0.07	0.186	-0.36	0.003	-0.16	0.036	-0.33	0.001	-0.03	0.097

Table S8. Primer sets used in RT-qPCR experiments.

Gene Name	Forward Primer Sequence	Reverse Primer Sequence
ACTIN	ACGGTGAAGGTGACAGCA	TTAGGATGGCAAGGGACTTC
ADSL	TCCCATTCACTCCCAGTT	TCACCTTCATCACGCTTTC
ARHGEF11	TCCCTGGACTTCTCATCTTC	GCTCCGTACCTGCTTCAC
CD3D	TGAGGACAGAGTGTGGTGAA	TCCCAGGTCCAGTCTTGT
GAPDH	TGCACCACCAACTGCTTAGC	GGCATGGACTGTGGTCATG
GBA	ACATCACCAAGGACACGTT	CACAACAGCAGAGCCATC
GZMA	GTGAAACCAGGAACCATGT	TATCGGACCAAGATGCACTAT
IFITM3	GGACAGGAAGATGGTTGG	CCCAGATGTTCCAGGCACT
IL17RA	GTGCAGCGGTCTGGTTAT	CATGCAATTTATCTTGGTATCC
MBOAT	ATTGGAATCAGAAAGTCTCCT	TGAGGCATGAACATTTGG
MSH6	TCTCCCAGAGGAAGTTATTCA	TCAAAGTCAGCAATTTATGGA
MSPT	CCTACGAGGACATCAAGGA	CTCAGGGCTCTTCTCCAG
NAP1L4	AGCGTCAAGTCTGCTTT	TAAGGTTAGGCCGAGCAC
OSCAR	TGTGCTGTAGCTTCTTTCCA	CACGTCTCACTACTACCTTTCTGT
PRDM4	CACATCCAGCGTTTCTCA	CTACTCCAGCCCAGAGGTT
RPL17	CCTAAACCAGAAGAGGAGGTT	ATTTACTCCCGTGCCATAAG
RPL24	CTGGTGCATCTCTTGCTG	GCAGCCCTGATAGCTTGT
RPL4	ACCCACTGAAAACTTGAGAAT	CTTTCCTTTCTTACCTACCACAG
RPL5	TGCTATACGAGAGAATCCAGTC	AGCTACCCGATCCTTCTTC
RSPH3	GGGGAGGAAGAATTGTCATA	CCCAGTCCATTACACAAAAA
TBXAS	GTCTAGGGCTGCTTGAGG	AGCGGGATACGATCTTGA
TIMM23	CATTAGTGAGTTGAAGCCAAA	TTAAATGAATGGACTTGGTGATA
TYMP	CACCTTGATAAGCTGGAG	CTGACCCACGATACAGCA

Table S9: Prediction of non-remission in MDD patients at baseline with gene information only. The accuracy shown is from single loop cross validation and is overly optimistic from selection bias. The corrected accuracy of the 13 gene model by nested cross validation is 79.4%. The threshold was selected to balance sensitivity and specificity such that Youden index (sensitivity+specificity-1) was maximized.

Model (gene number)	Accuracy (using single loop cross validation)	Sensitivity	Specificity
2	0.85	0.73	0.95
3	0.85	0.80	0.89
4	0.82	0.73	0.89
5	0.79	0.73	0.84
6	0.76	0.60	0.89
7	0.82	0.67	0.95
8	0.79	0.60	0.95
9	0.88	0.80	0.95
10	0.85	0.73	0.95
11	0.79	0.67	0.89
12	0.82	0.80	0.84
13	0.88	0.80	0.95
14	0.79	0.67	0.89
15	0.79	0.67	0.89
16	0.79	0.67	0.89
17	0.82	0.73	0.89
18	0.82	0.73	0.89
19	0.85	0.80	0.89
20	0.82	0.80	0.84
21	0.82	0.80	0.84
22	0.85	0.80	0.89
23	0.82	0.73	0.89
24	0.82	0.73	0.89
25	0.79	0.73	0.84
26	0.76	0.73	0.79
27	0.74	0.67	0.79
28	0.74	0.67	0.79
29	0.76	0.67	0.84
30	0.71	0.67	0.74

Table S10: Percentage of blood-cell specific genes showing positive correlation of expression changes between groups with genes belonging to the 13-genes model list.

	B-Cells	T-cells	CD8+ T-cells	GRANS	LYMPHS	
<i>Number of genes enriched in the population (from Palmer et al., 2006)</i>						
	428	222	23	411	67	
<i>Number of cell-specific genes detected in our dataset</i>						
	172	124	14	228	26	
<i>Cell-specific genes displaying similar directional effect across group comparisons (in %)</i>						
13 genes model list	CD3D	33.1	25.8	35.7	5.3	80.8
	CD97	24.4	27.4	7.1	62.3	3.8
	GZMA	30.2	24.2	35.7	6.6	76.9
	IFITM3	16.9	19.4	0.0	50.0	3.8
	MATR3	30.2	23.4	14.3	5.7	76.9
	PSMA4	32.6	24.2	28.6	5.3	80.8
	PSMA6	32.6	25.8	28.6	5.7	80.8
	RPL17	32.6	25.8	28.6	5.7	80.8
	RPL24	34.9	26.6	28.6	5.7	80.8
	RPL5	31.4	24.2	14.3	5.3	80.8
	RPL9	34.3	25.0	35.7	5.3	80.8
	TAGLN2	26.2	26.6	0.0	61.4	3.8
	TIMP1	15.7	20.2	0.0	47.8	3.8

GRANS : granulocytes cells

LYMPHS: lymphoid cells

Table S11. Prediction of non-remission in the independent validation cohort with top 13 gene features from our cohort. Accuracy represents the percentage of correctly identified samples. Sensitivity represents the ability to predict non-remission. Specificity represents the ability to predict remission.

	Accuracy	Sensitivity	Specificity
2	0.63	0.72	0.52
3	0.65	0.83	0.41
4	0.71	0.83	0.56
5	0.75	0.86	0.59
6	0.76	0.86	0.63
7	0.73	0.83	0.59
8	0.75	0.86	0.59
9	0.68	0.81	0.52
10	0.70	0.83	0.52
11	0.70	0.81	0.56
12	0.65	0.75	0.52
13	0.68	0.81	0.52

Table S12: Prediction of non-remission in MDD patients at baseline with gene information plus clinical information: Top genes frequency for the three features model. IDS16TOT: Quick Inventory of Depressive Symptomatology.

probe_definition	matched_geneSymbol	freq
Homo sapiens interferon induced transmembrane protein 3 (1-8U) (IFITM3), mRNA.	IFITM3	34
Homo sapiens TIMP metalloproteinase inhibitor 1 (TIMP1), mRNA.	TIMP1	33
IDS16TOT	<i>Clinical variable</i>	34

Table S13: Prediction of non-remission in MDD patients at baseline with gene information plus clinical information.

Features used in prediction model	Initial cohort (genes & clinical information)
# features used in model	3
Corrected Accuracy (%)	97.1
Sensitivity (%)	1
Specificity (%)	94.7

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