Data-driven clustering differentiates subtypes of major depressive disorder with distinct brain connectivity and symptom features

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Background
Major depressive disorder (MDD) is a clinically and biologically heterogeneous syndrome. Identifying discrete subtypes of illness with distinguishing neurobiological substrates and clinical features is a promising strategy for guiding personalised therapeutics.

Aims
This study aimed to identify depression subtypes with correlated patterns of functional network connectivity and clinical symptoms by clustering patients according to a weighted linear combination of both features in a relatively large, medication-naive depression sample.

Method
We recruited 115 medication-naive adults with MDD and 129 matched healthy controls, and evaluated all participants with magnetic resonance imaging. We used regularised canonical correlation analysis to identify component mapping relationships between functional network connectivity and symptom profiles, and K-means clustering was used to define distinct subtypes of patients.

Results
Two subtypes of MDD were identified: insomnia-dominated subtype 1 and anhedonia-dominated subtype 2. Subtype 1 was characterised by abnormal hyperconnectivity within the ventral attention network and sleep maintenance insomnia. Subtype 2 was characterised by abnormal hypoconnectivity in the subcortical and dorsal attention networks, and prominent anhedonia symptoms.

Conclusions
Our study identified two distinct subtypes of patients with specific neurobiological and clinical symptom profiles. These findings advance understanding of the biological and clinical heterogeneity of MDD, offering a pathway for defining categorical subtypes of illness via consideration of both biological and clinical features.

Keywords
Depression; anxiety; heterogeneity; functional network connectivity; magnetic resonance imaging.

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Major depressive disorder (MDD) is a heterogeneous syndrome varying considerably in clinical presentation, progression, treatment response and neurobiology.1 Failure to effectively resolve this heterogeneity, especially in biological terms, limits drug discovery and personalised treatment approaches for patients with MDD. Most data-driven MDD subtyping efforts have focused on discriminating different symptom profiles. In general, these approaches have not proven to be clinically useful or helpful for treatment development, as similar particular symptom profiles could be attributed to different biological mechanisms.2 Also, symptom-based subtyping has low stability over time.3 Most importantly, symptom-based clustering has not been useful in illuminating heterogeneity at the biological level.

Neuroimaging-based subtyping
Although there are multiple biological approaches that might prove useful for subtyping patients with MDD, considerable attention has been paid to psychoradiological substrates for depression.4 Some neuroimaging studies typically have identified MDD subtypes based on a single modality of neuroimaging data, such as functional activity or structural patterns.5 Partly because of the focus on high-dimensional neurobiological data, cluster results have had weak correspondence to clinical symptoms, which limits understanding of the clinical relevance of brain-based subtyping.

Intergrating analysis with cluster technique
This concern can be addressed by using analytic approaches designed to identify MDD subtypes based on both neurobiology and clinical phenomenology.6 Recent resting-state magnetic resonance imaging (MRI) studies7 have identified neurophysiological subtypes of MDD by applying clustering analysis, together with some dimensionality reduction via canonical correlation analysis (CCA) or principal component analysis, with functional connectivity data, clinical ratings and gene expression data. These approaches not only enhance the quality of clustering, but also facilitate interpretation of clustering results.6

Although previous efforts in this area have made significant advances, they have been limited by factors including small samples, multisite imaging with different scanners and acquisition protocols, limited or absent approaches for data reduction before clustering to reduce concerns with overfitting and validation, separate consideration of symptoms and biology in cluster analyses, and complex effects of medications and illness course on both symptoms and brain function on input parameters for cluster analysis.5 An
approach that defines robust cluster results that considers MRI and symptom features simultaneously during clustering, in a large sample of medication-naïve patients with MDD, could address these concerns and advance progress in this area. To this end, we conducted a data-driven decomposition of functional MRI-defined functional brain connectivity, and defined subtypes by clustering medication-naïve patients with MDD according to a multivariate correspondence between functional network connectivity (FNC) and clinical symptoms, using a regularised canonical correlation analysis (rCCA). We then examined symptom severity and functional brain connectivity differences between identified subtypes. Finally, we applied multivariate regression modelling, to detect associations between clinical profiles and abnormal FNC in each identified subtype.

**Method**

**Participants**

We recruited 115 first-episode and medication-naïve patients with MDD at the West China Hospital of Sichuan University. Diagnoses of MDD were confirmed with the Structured Clinical Interview for DSM-IV Disorders (SCID), determined by consensus of two experienced clinical psychiatrists. All patients had not received any lifetime psychotherapy or psychiatric medications before MRI scanning, and none had a significant history of systemic or neurological illness. They received different antidepressant treatments later according to clinical guidelines. We also recruited 129 healthy controls, matched for age, gender, handedness and education, from the local area by poster advertisement. Healthy controls were screened with the SCID Non-Patient Version to confirm the lifetime absence of non-affective psychosis, eating disorders and mood and anxiety disorders. See Supplementary Table 1 available at https://doi.org/10.1192/bjp.2021.103 for detailed inclusion and exclusion criteria. Two experienced clinical psychiatrists rated clinical symptoms of patients with MDD by using the 17-item Hamilton Rating Scale for Depression (HRSD)9 and the 14-item Hamilton Rating Scale for Anxiety (HRSA).10 All participants provided written informed consent and the protocol was approved by the Research Ethics Committee of West China Hospital, Sichuan University (study ethical approval number: 2018(610)).

**MRI acquisition and preprocessing**

Neuroimaging data were acquired with a whole-body 3.0 T MRI scanner (Siemens Trio, Erlangen, Germany), using a 12-channel head coil, including T1-weighted anatomic imaging and resting-state functional MRI. For functional studies, participants were instructed to relax with their eyes closed, and each scan lasted 6 min and 40 s (i.e. 200 volumes). See Supplementary material for the detailed information on MRI sequences and protocols.

MRI data preprocessing was done with MATLAB version R2016b for Windows (MathWorks, Inc, USA; see https://www.mathworks.com/products/matlab.html), and MRI data analyses were done with SPM12 (Wellcome Centre for Human Neuroimaging, UK; http://www.fil.ion.ucl.ac.uk/spm) and CONN version 2019b for Windows (The Gabrieli Lab. McGovern Institute for Brain Research, MIT, USA; see http://www.nitrc.org/projects/conn). Briefly, resting-state images of each participant were motion corrected, corrected for slice timing, normalised to Montreal Neurological Institute space and smoothed with a Gaussian kernel of 8 mm full width at half maximum. Denoising (e.g. physiological aliasing) and head motion correction procedures are detailed in the Supplementary material.

**Group independent component analysis**

Independent component analysis (ICA) is data-driven approach for constructing a whole-brain FNC within the CONN toolbox, and has been well validated. Unlike region of interest-based functional connectivity analyses, ICA can delineate distinct, coherent resting-state networks.11 We performed a participant-wise group ICA to uncover ordered components across all participants. Group ICA was performed with 30 components and a dimensionality reduction of 64, which are the default settings of CONN. This approach provides an optimised determination of network spatial locations for the sample, image acquisition and analysis pipeline. It also allows identification of discrete subregions/networks within networks identified by Yeo et al.12 Spatial ICA was performed on the preprocessed images to decompose the data into spatial independent components. We then examined the similarity of extracted independent components with canonical resting-state networks (Yeo et al’s seven networks), using spatial correlation analysis with confirmation by visual inspection. Independent components were assigned into different networks, including the visual network, sensorimotor network (SMN), dorsal attention network (DAN), ventral attention network (VAN), affective/limbic network, frontoparietal network (FPN) and default mode network (DMN). We additionally defined subcortical and cerebellar networks, which have been implicated in MDD pathophysiology (see Supplementary Fig. 1(a)).13 A detailed description of group ICA procedures is presented in the Supplementary material.

**Functional connectivity comparison between patients and healthy controls**

First-level correlation maps were produced by extracting the residual blood oxygen level-dependent time course from each independent component, and Pearson correlations were performed between each pair of independent components to index FNC. Correlation coefficients of each participant were converted to normally distributed z-scores, using the Fisher transformation for second-level generalised linear model analyses. In the second-level analysis, connectivity maps from all participants were entered into GLM to compute differences in functional connectivity between patients and healthy controls, with age, gender and head motion as covariates. Statistical significance for all comparisons was thresholded at connection-wise \( P < 0.001 \) and cluster-level false discovery rate (FDR)-corrected \( P < 0.05 \).14 Correlation coefficients of each participant were extracted for the subsequent rCCA.

**rCCA and K-means clustering**

High-dimensional neuroimaging data invariably includes features not relevant to clustering results. We reasoned that more reliable and meaningful depression subtypes would be marked by a low-dimensional representation of the significant correlation between neuroimaging features and clinical symptoms. Therefore, to reduce dimensionality, in a first-step preliminary screen we first identified all significant computed Spearman’s correlations \( (P < 0.005) \) between connectivity features and each item of HRSD and HRSA psychopathology ratings. Next, we used rCCA to define a low-dimensional projection from these associations for use in clustering. Then, to identify potential MDD subtypes, the two-dimensional variates (representing connectivity features and clinical symptoms, respectively) of the first component derived from the significant rCCA models were subjected to K-means clustering. The first component represents canonical correlations that describes the maximum covariation between the two multidimensional variables. The significance of the rCCA model was determined by permutation tests. Finally, the optimal cluster number and

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the validity of the cluster solution were determined with the ‘Nbclust’ package in R version 4.0.3 for Windows (R Foundation for Statistical Computing, Vienna, Austria; see https://cran.r-project.org/mirrors.html). Clustering stability was examined with the Jaccard coefficient and a bootstrap technique \((n = 1000)\) in the ‘fpc’ package in R. Methodological details regarding rCCA and \(k\)-means clustering algorithms are described in the Supplementary material.

**Functional connectivity comparisons between subtypes and healthy controls**

To assess whether dysfunctional FNC differed by MDD subtypes, or between subtypes and healthy controls, we performed a one-way ANOVA analysis for subtype 1, subtype 2 and healthy controls by using a GLM in CONN, with age, gender and head motion as covariates. Statistical significance for all comparisons was thresholded at connection-wise \(P < 0.001\) and cluster level FDR-corrected \(P < 0.05\). Post hoc \(t\)-tests were performed to compare connectivity patterns across each MDD subtype and the healthy controls group. The FDR correction maintained a corrected \(P < 0.05\) for these comparisons.

**Partial least square regression analysis**

We applied partial least square regression (PLSR) to create a low-dimensional representation that relates symptoms and network connectivity. PLSR involves projecting a predictor space (\(X\)) and a response space (\(Y\)) onto a latent space, and then finding the variance in \(X\)-space that explains the maximum variance in \(Y\)-space. This statistical approach is suited for the current problem because it can work with multivariate response variables as well as multicollinearity among the predictors. We constructed a PLSR model for both subtype 1 and subtype 2 (the two subtypes defined by the cluster analysis), to examine relations between subtype-differentiated connectivity features and subtype-differentiated symptoms. In addition, to assess the association of connectivity alterations and symptoms in patients with MDD as a whole, we also conducted a PLSR model for the whole MDD group. Details of this analysis are presented in the Supplementary material. A graphic representation of the analysis pipeline is presented in Fig. 1.

**Results**

**Group ICA**

The spatial matching across all participants between extracted independent components and Yeo et al’s seven networks implemented in CONN are shown in Supplementary Fig. 1. Among the 30 independent components, four were identified as noise components, leaving 26 for subsequent connectivity analyses. The selected 26 independent components included four independent components in the visual network, four independent components in the SMN, three independent components in the DAN, two independent components in the VAN, two independent components in the limbic network (LIM), four independent components in the FPN, four independent components in the DMN, two independent components in the subcortical network and one independent component representing the cerebellar network (see Fig. 2(a)). More details including the labels and peak coordinates of these independent components are presented in Supplementary Table 2.

**rCCA and \(k\)-means clustering**

Results of rCCA analysis indicated that the first component has the highest canonical correlation coefficient \((r = 0.86)\) between FNC and clinical symptoms, by permutation test (see Supplementary Fig. 2). Therefore, the first component, including two canonical variates of FNC and clinical features, was used as input for the cluster analysis.

Participants with MDD were subdivided into two subtypes by \(k\)-means clustering analysis of the two-dimensional variates. In terms of the cluster distribution, 59 and 56 patients were partitioned into subtypes 1 and 2, respectively. Validity and stability analyses showed that the two cluster solution had the highest silhouette index (0.56) and Jaccard similarity (0.82) (Supplementary Fig. 3).

**Demographic and clinical profiles in MDD subtypes**

Demographic and clinical data for each subtype are shown in Table 1. HRSD and HRSA total scores did not differ between subtypes, indicating that the discrimination was not by illness severity (Fig. 3(c)).

We used the Kruskal–Wallis ANOVA to test subtype differences in scores for each item of the HRSD and HRSA. Eleven symptom measures showed significant differences, with anhedonia symptoms prominent in subtype 1 and sleep maintenance insomnia prominent in subtype 2 (Fig. 3(a) and 3(b)).

**Group comparison of FNC**

Comparisons of FNC between the combined MDD group and healthy controls showed abnormal connectivity patterns, including hypoconnectivity within the FPN, visual network and SMN, LIM and SMN, and LIM and DAN, and hyperconnectivity between the FPN and LIM in the MDD group (Fig. 2(c)).

Distinct patterns of abnormal FNC between networks were observed in the two MDD subtypes. Compared with healthy controls, subtype 1 showed hyperconnectivity between the VAN (left central opercular gyrus) and DAN (right lateral occipital gyrus) and within the VAN (between left supramarginal gyrus and left central opercular gyrus). By contrast, subtype 2 showed hyperconnectivity between the DMN (left praecuneus) and VAN (left operculum), FPN (right frontal pole) and visual network (right occipital gyrus), and subcortical network (bilateral thalamus) and DAN (bilateral lateral occipital gyrus), and hyperconnectivity within the VAN (between left supramarginal gyrus and left operculum) (Fig. 2(e)).

In addition, there were common (‘shared’) connectivity abnormalities across the two patient groups versus healthy controls (subtype 1 versus healthy controls and subtype 2 versus healthy controls), which involved hyperconnectivity between the visual network (bilateral occipital gyrus) and SMN (right superior temporal gyrus), affective network (frontal orbital/medial gyrus) and SMN (right superior temporal gyrus), and affective network (frontal orbital/medial gyrus) and DAN (right middle temporal gyrus), and hyperconnectivity between the affective network (left amygdala/hippocampus/temporal pole) and FPN (left cerebellum-crus II) (Fig. 2(d) and 2(e)).

**Association between FNC and MDD symptoms**

Significance of exploratory pairwise correlations in PLSR model was determined by permutation test \((P < 0.05, n = 1000)\). Hyperconnectivity within the VAN was only negatively correlated with sleep maintenance insomnia in subtype 1. Anhedonia was only positively correlated to hypoconnectivity between the subcortical network and DAN in subtype 2. To illustrate relations between FNC and symptoms, feature loadings are shown in
Main findings
The considerable heterogeneity of MDD presents a major obstacle for the development of new treatment and personalised medicine. Many previous studies have worked to parse this heterogeneity in either purely clinical or biological terms — strategies that have had limited utility and an unclear relationship to clinical practice. In the present study, we evaluated a cohort of 115 patients with first-episode MDD, and used an analytic approach that considered neuroimaging and clinical ratings simultaneously. Importantly, the patients with MDD were treatment-naïve, so that medication effects on functional connectivity patterns symptoms and symptoms did not confound the data analysis. We identified two specific MDD subtypes with different symptom domains (insomnia dominated and anhedonia dominated) that had distinct patterns of FNC alteration. Our findings provide insight into MDD heterogeneity in biological and clinical terms, and represent a promising step toward categorical subtype classification and, potentially, the development of personalised treatments for MDD.

Interpretation of our findings
Of the two subtypes we defined, subtype 1 had more prominent sleep maintenance insomnia and genital symptoms. In contrast, subtype 2 demonstrated more prominent depressed mood, feelings of guilt, suicide attempts and lack of enjoyment (anhedonia). Importantly, total scores of depression and anxiety severity did not differ between subtypes, indicating that our subtype delineation did not merely separate mild and more severely ill patients. Heterogeneity regarding insomnia and anhedonia features have long been considered in depression research, and our findings advance those clinical conceptualisations by linking them to different regional alterations in functional brain systems.

Fig. 3(d). There were no other significant associations in the PLSR analyses.

Discussion

Fig. 1  Overview of data processing and analysis pipeline. Group independent component analysis (ICA) was used to calculate and extract the average signal time series from brain parcellations. Following data reduction steps, connectivity networks and clinical symptoms were used to perform a regularised canonical correlation analysis (CCA). Regarding the discovery of categorical biotypes, K-means clustering was used to test identify subtypes defined by associations between functional network connectivity and symptoms patterns. Finally, the Wilcoxon rank-sum tests were used to test for differences in the severity of clinical symptom scores between subtypes. The generalised linear model analyses were performed to examine common and distinct network connectivity patterns across groups. The associations between specific symptoms and distinct network connectivity patterns were examined with partial least squares regression. DAN, dorsal attention network; DMN, default mode network; FPN, frontoparietal network; HRSA, Hamilton Rating Scale for Anxiety; HRSD, Hamilton Rating Scale for Depression; LIM, limbic network; MDD, major depressive disorder; SMN, sensorimotor network; VAN, ventral attention network.

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Fig. 2  The functional network connectivity (FNC) differences between MDD, MDD subtypes and healthy controls. (a) Nine functional networks identified by group independent component analysis. (b) Whole-brain 64 × 64 averaged functional connectivity matrix between independent component pairs was computed over all participants. The value in the correlation matrix represents the Fisher’s z-transformed Pearson correlation coefficient. (c) Heat map and circular plot showing features that were significantly abnormal in patients with MDD relative to healthy controls. (d) Heat map showing significant results of ANOVA in FNC among MDD subtypes and healthy controls. (e) Circular plots showing significant FNC patterns between subtype 1, subtype 2 and healthy controls, using post hoc analyses. Results of statistical comparisons were thresholded at connection-wise $P < 0.001$ with false discovery rate-corrected $P < 0.05$. DAN, dorsal attention network; DMN, default mode network; FPN, frontoparietal network; LIM, limbic network; MDD, major depressive disorder; SMN, sensorimotor network; VAN, ventral attention network.
world, were more characteristic of subtype 2, consistent with their association with anhedonia symptoms. These findings indicate that distinct patterns of FNC were associated with hyperarousal features of MDD and with the deficits of social motivation and hedonic tone. Importantly, not only were there different symptom-brain correlations, but the two patterns were evident in different subgroups of patients with MDD. Thus, these observations may be helpful for understanding the complex pathophysiological mechanisms of MDD and its clinical heterogeneity. Further, they may suggest the need to target different neural systems in patients with MDD, depending on their pattern of symptom manifestation.

In addition to these distinguishing subtype features, some abnormal network patterns were shared across the two patient subgroups. Overlapping abnormalities included hypoconnectivity among the visual network and SMN, and affective network and SMN, networks involved in emotion and sensorimotor processing, and hyperconnectivity between the FPN and affective network, networks involved in visceral/emotional monitoring. These findings highlight that although patients with MDD as a group share some traits, distinguishing features of subtypes of patients can still be identified. From a broader perspective, our findings indicate that combining clinical and neurobiological features in subtype identification may permit delineation of distinct and

### Table 1: Demographic features of the two subtypes of patients with major depressive disorder and healthy controls

<table>
<thead>
<tr>
<th>Subtype</th>
<th>Mean (SD)</th>
<th>Statistics</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subtype 1</td>
<td>Subtype 2</td>
<td>Healthy controls</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>33.63 (11.74)</td>
<td>30.43 (11.26)</td>
<td>32.95 (10.39)</td>
</tr>
<tr>
<td>Gender, male/female</td>
<td>21/38</td>
<td>19/37</td>
<td>57/72</td>
</tr>
<tr>
<td>Education level</td>
<td>1.81 (1.04)</td>
<td>1.50 (0.79)</td>
<td>1.52 (0.69)</td>
</tr>
<tr>
<td>Age at onset, years</td>
<td>33.18 (11.82)</td>
<td>29.90 (11.33)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Illness duration, weeks</td>
<td>23.37 (24.91)</td>
<td>27.68 (28.15)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Handedness, right/left</td>
<td>56/37</td>
<td>49.7</td>
<td>129.7</td>
</tr>
<tr>
<td>Depression symptoms</td>
<td>HRSD score 25.03 (5.16)</td>
<td>HRSD score 24.46 (9.67)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Anxiety symptoms</td>
<td>HRSA score 24.46 (9.67)</td>
<td>HRSA score 24.34 (8.83)</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

All data are shown as mean (s.d.) or ratios. HRSD, Hamilton Rating Scale for Depression; HRSA, Hamilton Rating Scale for Anxiety.

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**Identifying MDD subtypes with data-driven clustering**

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### Fig. 3: Subtype-specific clinical profiles for (a) depression symptoms and (b) anxiety symptoms that varied most significantly by cluster (P < 0.05, Wilcoxon rank-sum tests, false discovery rate corrected). Each item value was z-transformed with respect to the mean for all patients in each subtype. (c) Boxplot of subtype differences in overall depression and anxiety severity (total HRSD/HRSA score). (d) Loadings of the distinct functional network connectivity (FNC) and clinical symptoms from partial least squares regression (PLSR) in each of the two identified subtypes. The left graph shows loadings of PLSR relating distinct FNC (within the VAN) to insomnia-middle (DS) in subtype 1; the right graph shows loadings of PLSR relating distinct FNC (between the subcortical network and DAN) to anhedonia symptoms (D7) in subtype 2. Items in HRSA and HRSD rating instruments are presented on the y-axis (e.g. D7 is the seventh item of HRSD). The asterisk indicates significant difference from mean symptom severity rating for all patients, P < 0.05; error bars depict s.e.m. **P < 0.05, ***P < 0.005, ****P < 0.001.**
clinically relevant subtypes of patients with MDD, to advance inte-
gration of neurobiology with clinically relevant behavioural features for refining disease classification and treatment for depression
syndromes.23

Heterogeneity is a common issue not only for MDD but across psychiatric syndromes. Identification of discrete homogeneous sub-
types is becoming recognised as an important path for the develop-
ment of more personalised diagnostics and treatments.2 Working to
resolve heterogeneity by using both clinical and neurobiological information together represents a promising strategy in this
regard. Although our findings represent a significant step forward in
this direction, they to be replicated and their clinical utility
needs to be established in clinical trials.8 Further, given the com-
plexity of the problem, group clustering efforts such as ours are
beginning with single or a small domain of biological phenotypes.
In the longer term, combining wider ranges of phenotypes in
predictive/classification models likely will be needed to optimize subtype characterisation for nosology refinement and identifi-
yng targets for subtype-specific therapies. Comparison of different
methodological approaches for identifying subtypes will also be
needed. In the present study, we used a regularised version of
CCA to incorporate latent relations between neuroimaging and
clinical data-sets, which tends to avoid overfitting and unstable
solutions that can result from use of classical CCA,24 but alternative
approaches may have benefits as well.

Limitations
Out study has limitations that need to be considered. First, although
we used the bootstrapping method to establish the stability of clus-
tering, we did not have an external validation data-set to replicate
the clustering results. Second, our study included treatment-naïve
patients with first-episode MDD. This group is useful for identify-
ing disease-related features independent of drug treatment effects, but it is unclear whether our findings would be applicable to MDD
groups with multiple prior episodes and active antidepressant therapy, and across the lifespan, to paediatric and elderly patients. Third, more
precise estimates of functional connectivity may be obtained using
longer data acquisitions, which can increase the reliability of intrin-
sic connectivity estimates compared with 5–7 min of resting-state
data.25

Practical implications
The aim of the present study was to use the association between clin-
cal profiles and functional brain imaging features in an integrated
way for the purpose of defining MDD subtypes. We applied a
rCCA and clustering-based method, considering both neuroima-
ging and clinical data, to define two subtypes of MDD that differen-
tiated both in their symptom profiles (insomnia dominated and
anhedonia dominated) and distinct patterns of altered brain func-
tion. Although the two subtypes were associated with different
MDD symptom features, they cannot be simply distinguished in
overall depression severity or duration, as has been a challenge in
some previous efforts at subgroup identification. Further, we note
that neither subtype was a rare entity, with the two subtypes
being relatively equally represented in our sample. Our specific
findings regarding the neurobiological-clinical framework for defining
subtypes need to be examined in terms of its utility for personalising
MDD treatments, and for guiding treatment development efforts.
Success in such efforts would accelerate personalised treatment for
depression, and speed the integration of neuroscience-based
evaluations and clinical/behavioural evaluation in providing optimal
treatment for patients with MDD.

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center of MDD treatments, and for guiding treatment development efforts.
Success in such efforts would accelerate personalised treatment for
depression, and speed the integration of neuroscience-based
evaluations and clinical/behavioural evaluation in providing optimal
treatment for patients with MDD.

Supplementary material
To view supplementary material for this article, please visit https://doi.org/10.1192/bjp.2021.103

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Drs. Q. Gong and X. Huang contributed equally to playing the role of corresponding and senior
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Data availability
The data that support the findings of this study are available from the corresponding author,
X. Huang, upon reasonable request.

Author contributions
Y.W., S.T. and Q.G. contributed to the conception and design of the study. Y.W. and S.T. per-
formed the analyses and drafted the manuscript. L.J., X.B., J.A.S. and L. participated in discus-
sion of statistical analyses and imaging analyses. H.L., Y.G., X.H., Z.J. and W.K. recruited participants and collected the data. J.A.S., Q.G. and X.Huang reviewed and revised the manu-
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Declaration of interest
J.A.S. consults for VerdGt. All other authors report no conflicts of interest.

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Saturn: star of melancholy

George Ikkos

Celebrated ‘Saturn and Melancholy: Studies in the History of Natural Philosophy, Religion and Art’, by Klibansky et al, was published in English by the Warburg Institute in London in 1964. The Institute started life in Hamburg in 1909 as the Warburg-Bibliothek für Kulturwissenschaft, a library for art studies established by Aby Warburg (1866–1929), scion of the city’s renowned Jewish banking family. He had ceded his inheritance to his brother, in exchange for a promise to buy him any book he requested. In 1926 the Library became an Institute, later affiliating with Hamburg University. Under Nazi threat, in 1933 it relocated to London and in 1944 became associated with London University. Established in 1798, M. M. Warburg & Co, still in Hamburg, is the longest surviving bank in the world.

Saturn is our solar system’s furthest planet visible to the naked eye, hence the darkest. The first direct reference to it as relating to melancholia occurs in Arab astrology during the 9th century. Attributed to it were what initially appear to be bewildering temperaments and illness. It also carries it forward in history to Renaissance literature, coloured by Christianity. In the Western pictorial tradition Melancholia was most famously depicted in 1514 at the threshold of Enlightenment in Albrecht Dürer’s Melencolia I. The work is rich in metaphors and symbolism: the dog represents the antithesis of the rationalizing spirit in man, the cornucopia symbolizes the human capacity to use nature, the scythe is a reminder of the inevitability of death, and the Blocks are a reference to the Roman games. The work has been interpreted as a criticism of the rationalist spirit in the human mind, and as a reflection of the artist’s own inner turmoil.

Saturn is also associated with melancholia in Renaissance literature, specifically in the work of the Italian poet Giorgio Vasari. In his Lives of the Artists, Vasari describes Saturn as the planet of melancholy and coldness, and as a symbol of the human soul. He also mentions that Saturn was associated with the classical Greek goddess Mnemosyne, the goddess of memory, who is often depicted with a cornucopia.


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