Background. Clinical depression is associated with reductions in white-matter integrity in several long tracts of the brain. The extent to which these findings are localized or related to depressive symptoms or personality traits linked to disease risk remains unclear.

Method. Members of the Lothian Birth Cohort 1936 (LBC936) were assessed in two waves at mean ages of 70 and 73 years. At wave 1, they underwent assessments of depressive symptoms and the personality traits of neuroticism and extraversion. Brain diffusion magnetic resonance imaging (MRI) data were obtained at the second wave and mood assessments were repeated. We tested whether depressive symptoms were related to reduced white-matter tract fractional anisotropy (FA), a measure of integrity, and then examined whether high neuroticism or low extraversion mediated this relationship.

Results. Six hundred and sixty-eight participants provided useable data. Bilateral uncinate fasciculus FA was significantly negatively associated with depressive symptoms at both waves (standardized $\beta = 0.12–0.16$). Higher neuroticism and lower extraversion were also significantly associated with lower uncinate FA bilaterally (standardized $\beta = 0.09–0.15$) and significantly mediated the relationship between FA and depressive symptoms.

Conclusions. Trait liability to depression and depressive symptoms are associated with reduced structural connectivity in tracts connecting the prefrontal cortex with the amygdala and anterior temporal cortex. These effects suggest that frontotemporal disconnection is linked to the etiology of depression, in part through personality trait differences.

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Key words: Depression, DTI, extraversion, MRI, neuroticism.

Introduction

Clinical depression is a common disorder that affects approximately 15% of the population at some time in their lives and shows a tendency to become recurrent or persistent (Thornicroft & Sartorius, 1993). The disorder is predicted to become the leading cause of worldwide disability (Murray & Lopez, 1997) and has significant costs for affected individuals, their families and society at large. Despite the considerable burden, the neurobiology of depression is poorly understood although genetic factors and stressful life events both play a part in its etiology (Levinson, 2006).

Brain imaging has provided important insights into the biological basis of depression, with evidence of modest gray-matter reductions in frontotemporal regions using both region of interest and voxel-based methods (Drevets et al. 1997; Videbech, 1997; Kumar et al. 1998; Videbech & Ravnkilde, 2004). Depression has not, however, typically been associated with a single cortical or subcortical region (Arnone et al. 2012) and this could suggest that the disorder is caused by abnormal interactions between different brain regions due to disrupted connectivity. In support of this idea, some studies have found evidence of reduced white-matter integrity in the prefrontal and temporal lobes, and in several other regions (Taylor et al. 2007;
Cullen et al., 2010; Dalby et al., 2010). Many of these white-matter tracts, for example the uncinate fasciculus and cingulum, connect prefrontal areas with the medial temporal lobe. Disrupted connectivity of these emotion-related areas may therefore lead to emotional dysregulation and possibly an increased risk of depression. The extent to which these findings are secondary to the effects of clinical psychiatric illness or its treatment is unclear.

To overcome the issue of confounding by treatment-related factors, an alternative approach to studying clinical depression directly is to examine the relationship of white-mater tract integrity to depressive symptoms and known risk factors in a non-clinical population cohort. Trait liability to depression has long been associated with neuroticism (Kendler et al., 1993a, 1994, 2002; Matthews et al., 2009), a heritable and relatively stable personality trait. Family and twin studies indicate that there is a high genetic correlation between neuroticism, clinical depression (Kendler et al., 2006) and depressive symptoms in non-clinical population samples (Ivkovic et al., 2007), with an overlap of more than 50% (Kendler et al., 1993b; Levinson, 2006) in their underlying architecture.

Lower extraversion has also been proposed as a risk factor for depression, although the literature (Barnett et al., 2011) implicating this dimension of personality in the etiology of depression is sparse.

In the current study we first examined the relationship of white-matter tract fractional anisotropy (FA), derived from diffusion tensor imaging (DTI) as a proxy for tract integrity, with depressive symptoms in individuals from a population-based sample, the Lothian Birth Cohort 1936 (LBC1936). Second, we examined whether trait liability to depression was also associated with reductions in white-matter tract FA in the same tracts using the personality traits of (higher) neuroticism and (lower) extraversion. Third, because we hypothesized that reduced frontotemporal white-matter tract FA confers greater risk of depressive symptoms through effects on trait liability, we conducted a mediation analysis specifically to test this directional hypothesis.

Method

Subjects

The LBC1936 comprises 1091 community-dwelling relatively healthy individuals without dementia, residing around the city of Edinburgh, Scotland, most of whom had participated in the Scottish Mental Survey 1947 (SMS1947). The LBC1936 undertook personality, medical and cognitive testing at a mean age of about 70 years (for full details on participant recruitment and cognitive testing, see Deary et al., 2007). Subjects were then reassessed approximately 3.2 years later at which time they received additional investigations, including detailed structural brain magnetic resonance imaging (MRI) (Wardlaw et al., 2011), and retook the medical and cognitive assessments, including the Mini Mental State Examination (MMSE) of cognitive function (Folstein et al., 1975). Participants lived independently in the community around the city of Edinburgh, Scotland, and were able to travel on both occasions to the Wellcome Trust Clinical Research Facility in Edinburgh for testing. The study was approved by the Lothian (REC 07/MRE00/58) and Scottish Multicentre (MREC/01/0/56) Research Ethics Committees and all subjects gave written informed consent.

Depressive symptoms and personality traits

Hospital Anxiety and Depression Scale (HADS; Zigmond & Snaith, 1983) scores were available from the assessments performed at 70 and 73 years of age. These assessments were available separately as depression (HADS-D) and anxiety (HADS-A) measures with seven self-report items for each subscale. Neuroticism and extraversion were measured at the first assessment using the Neuroticism, Extraversion and Openness Five Factor Inventory (NEO-FFI; Costa & McCrae, 1989), a self-report scale consisting of 12 items for each NEO-FFI personality trait. The other three traits (Openness, Agreeableness and Conscientiousness) were not included in the analyses in the current study.

Diffusion MRI

Diffusion MRI was performed on a GE Signa HDx 1.5-T clinical scanner (General Electric, USA) using a self-shielding gradient set with maximum gradient strength of 33 mT m⁻¹, and a manufacturer-supplied eight-channel phased-array head coil. Single-shot, spin-echo echo-planar diffusion-weighted volumes (b=1000 s mm⁻²) were acquired in 64 non-collinear directions, along with seven T²-weighted volumes (b=0 s mm⁻²). Seventy-two contiguous axial slices of thickness 2 mm were acquired with a field of view of 256 x 256 mm and matrix size of 128 x 128, giving a resolution of 2 x 2 x 2 mm. The repetition time was 16.5 s and echo time 95.5 ms, producing a total scan time of approximately 20 min (see Wardlaw et al., 2011 for full details of the imaging protocol).

White-matter tractography

Datasets were converted into NIFTI-1 (http://nifti.nimh.nih.gov/nifti-1) format and preprocessed using
FSL tools (FMRIB, Oxford, UK; www.fmrib.ox.ac.uk/) to extract the brain, remove bulk motion and eddy current-induced artifacts and estimate water diffusion tensor parameters. Brain connectivity data were created using the BEDPOSTX/ProbTrackX tractography algorithm with a two-fiber model and 5000 streamlines to reconstruct tracts of interest. An automatic tract selection method with good reproducibility (Clayden et al. 2009a), based on a model of tract topology (Clayden et al. 2007), was used to generate equivalent tracts of interest in each subject. This technique, termed probabilistic neighborhood tractography (PNT) and implemented in the TractoR package (http://github.com/jonclayden/tractor/wiki/), optimizes the choice of seed point for tractography by estimating the best matching tract from a series of candidate tracts generated from a neighborhood of voxels placed around a voxel transferred from standard space against a reference tract that was derived from a digital human white-matter atlas (Mori & van Zijl, 2007). The topological tract model was also used to reject any false positive connections (Clayden et al. 2009b), thereby significantly improving tract segmentation (Maniega et al. 2008). As displayed in Fig. 1, 12 white-matter tracts were segmented, namely the genu and splenium of corpus callosum, cingulum cingulate gyri (CCG) (Clayden et al. 2007), anterior thalamic radiations (ATR), uncinate, arcuate and inferior longitudinal fasciculi (ILF). For each subject, the seed point that produced the best match tract to the reference for each of the 12 pathways was determined, with the resulting tractography mask applied to each subject's FA volume. Tract-averaged FA values were calculated from these masks and used in all subsequent analyses. To ensure that the segmented tracts were anatomically plausible representations of the fasciculi of interest, a researcher (S.M.M.) visually inspected all masks blind to the other study variables and excluded tracts with aberrant or truncated pathways.

Statistical analyses

Statistical analyses were conducted in the R statistical software package (www.r-project.org/) using linear regression. To meet underlying statistical assumptions, we applied a square root transformation to the HADS symptom measures. Initially, we entered the transformed HADS scores as the dependent variable and age at symptom assessment, sex and white-matter tract FA as the independent variables (analysis 1). To correct for multiple testing, we applied a false discovery rate (FDR; Benjamini & Hochberg, 1995) correction to these initial associations. Subsequently, we examined whether higher neuroticism or lower extraversion were also related to white-matter tract FA using the same model, substituting these personality measures as the dependent variables (analysis 2). We then examined whether previous antidepressant treatment or a history of stroke confounded any significant relationships between FA and depression or personality measures by conducting a series of planned additional analyses. We then adjusted for the effects of antidepressant treatment, current or past cigarette use or stroke by including these as covariates in the statistical models. Finally, to assess whether neuroticism/extraversion could account for any associations between depression and white-matter tract FA, we examined the influence of these variables on analysis 1. Specifically, we noted the
attenuation in the effect of white-matter tract FA after neuroticism/extraversion was included in the model, and conducted a formal mediation analysis using the Sobel test (Baron & Kenny, 1986).

**Results**

A total of 668 individuals from the LBC1936 provided useable tractography data from the 740 who presented for MRI. As indicated in Table 1, the average age of the subjects was 72.7 (s.d. = 0.7) years at the time of the imaging examination and 356 (53%) were male. The rates of antidepressant prescribing were lower than in the general Scottish population (see www.isdscotlandarchive.scot.nhs.uk/isd/6517.html), approximately 10% of the sample reported a diagnosis of diabetes and 48.4% reported a diagnosis of hypertension. MMSE scores (Folstein et al. 1975) showed that only one subject met a specific threshold for organic brain disorder whereas 6.4% of the sample met a sensitive threshold. At both assessments, the HADS-D score correlated moderately and positively with neuroticism (Pearson’s r = 0.35 at age 69.5, and r = 0.35 at age 72.7) and negatively with extraversion (Pearson’s r = −0.38 at age 69.5, and r = −0.34 at age 72.7). Table 1 also shows that the number of individual tracts that did not meet quality criteria (truncation or failed to follow expected path) ranged from 0.6% for the right inferior longitudinal fasciculus to 16.8% for the left anterior thalamic radiation.

We first examined the relationship of the HADS-D subscale scores with FA in each tract (Table 2). Depressive symptoms measured at the time of the imaging assessment were significantly and negatively associated with FA in the left and right uncinate fasciculus (standardized β = −0.12, p = 0.0038 in the left hemisphere; standardized β = −0.13, p = 0.0011 in the right hemisphere). Depressive symptoms measured at the first assessment 3 years earlier were also significantly and negatively associated with bilateral uncinate FA, with similar effect sizes (standardized β = −0.16, p = 0.0002 in the left hemisphere).

<table>
<thead>
<tr>
<th>Male sex, n (%)</th>
<th>356 (53)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at assessment 1 (years), mean (s.d.)</td>
<td>69.5 (0.8)</td>
</tr>
<tr>
<td>Age at assessment 2 (imaging) (years), mean (s.d.)</td>
<td>72.7 (0.7)</td>
</tr>
<tr>
<td>Antidepressants at assessment 1, n (%)</td>
<td>23 (3.4)</td>
</tr>
<tr>
<td>Antidepressants at assessment 2, n (%)</td>
<td>30 (4.5)</td>
</tr>
<tr>
<td>Diabetes, n (%)</td>
<td>72 (10.8)</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>323 (48.4)</td>
</tr>
<tr>
<td>MMSE score &lt; 22, n (%)</td>
<td>1 (0.1)</td>
</tr>
<tr>
<td>MMSE score &lt; 27, n (%)</td>
<td>43 (6.4)</td>
</tr>
<tr>
<td>MMSE score, median (IQR)</td>
<td>29 (28–30)</td>
</tr>
<tr>
<td>HADS Depression – assessment 1, mean (s.d.)</td>
<td>2.6 (2.2)</td>
</tr>
<tr>
<td>HADS Depression – assessment 2, mean (s.d.)</td>
<td>2.6 (2.2)</td>
</tr>
<tr>
<td>HADS Anxiety – assessment 1, mean (s.d.)</td>
<td>4.8 (3.1)</td>
</tr>
<tr>
<td>HADS Anxiety – assessment 2, mean (s.d.)</td>
<td>4.4 (3.2)</td>
</tr>
<tr>
<td>NEO-FFI Neuroticism – assessment 1, mean (s.d.)</td>
<td>16.6 (7.6)</td>
</tr>
<tr>
<td>NEO-FFI Extraversion – assessment 1, mean (s.d.)</td>
<td>27.3 (5.9)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tract</th>
<th>Complete n (%)</th>
<th>Mean FA (s.d.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genu of corpus callosum</td>
<td>646 (96.7)</td>
<td>0.41 (0.046)</td>
</tr>
<tr>
<td>Splenium of corpus callosum</td>
<td>663 (99.3)</td>
<td>0.49 (0.069)</td>
</tr>
<tr>
<td>Left arcuate fasciculus</td>
<td>639 (95.7)</td>
<td>0.45 (0.042)</td>
</tr>
<tr>
<td>Right arcuate fasciculus</td>
<td>580 (86.8)</td>
<td>0.43 (0.043)</td>
</tr>
<tr>
<td>Left anterior thalamic radiation</td>
<td>556 (83.2)</td>
<td>0.32 (0.033)</td>
</tr>
<tr>
<td>Right anterior thalamic radiation</td>
<td>643 (96.3)</td>
<td>0.33 (0.047)</td>
</tr>
<tr>
<td>Left uncinate fasciculus</td>
<td>567 (84.9)</td>
<td>0.33 (0.030)</td>
</tr>
<tr>
<td>Right uncinate fasciculus</td>
<td>628 (94.0)</td>
<td>0.33 (0.032)</td>
</tr>
<tr>
<td>Left inferior longitudinal fasciculus</td>
<td>663 (99.3)</td>
<td>0.40 (0.048)</td>
</tr>
<tr>
<td>Right inferior longitudinal fasciculus</td>
<td>664 (99.4)</td>
<td>0.38 (0.048)</td>
</tr>
<tr>
<td>Left cingulum cingulate gyrus</td>
<td>641 (96.0)</td>
<td>0.44 (0.047)</td>
</tr>
<tr>
<td>Right cingulum cingulate gyrus</td>
<td>650 (97.3)</td>
<td>0.39 (0.044)</td>
</tr>
</tbody>
</table>

**Table 1. Demographic and historical information**

MMSE, Mini Mental State Examination; HADS, Hospital Anxiety and Depression Scale; NEO-FFI, Neuroticism, Extraversion and Openness Five Factor Inventory; FA, fractional anisotropy; s.d., standard deviation; IQR, interquartile range.
We then adjusted for the effects of antidepressants prescribed at time 1 or time 2, the presence of a self-reported history of stroke, current alcohol use at the time of the scan, and whether the individual reported a history of past or current tobacco use. In each case the relationship between uncinate white-matter integrity remained in the same direction as reported in the whole sample (see online Supplementary material). However, the relationship between right uncinate white-matter integrity and neuroticism became non-significant.

We then assessed the effect of adding neuroticism or extraversion to a statistical model where tract FA and the independent variable also improved the model fit when neuroticism was the dependent variable and right uncinate FA was the independent variable. The purpose of this analysis was to examine whether the addition of neuroticism/extraversion attenuated the effect of uncinate tract FA on depressive symptoms, indicating that neuroticism was more closely associated with depression and potentially mediating the association of tract FA with depressive symptoms. The addition of neuroticism led to a significantly improved model fit in both the left (see online Supplementary Table S2) and right hemispheres. The regression coefficient estimates for the effect of uncinate tract FA on depressive symptoms were somewhat attenuated in both the left hemispheres after the inclusion of neuroticism in the model (see Supplementary Table S2). Similarly, the addition of extraversion to a model where depressive symptoms was the dependent variable and right uncinate FA was the independent variable also improved the model fit whereas the effect of tract FA on depressive symptoms remained significant after FDR adjustment across all 12 tracts assessed.

We then reanalyzed the data after the graphical detection and subsequent removal of a single outlying observation with an FA value of <0.2, with no effect on any of our significant findings, all of which remained significant.
was also somewhat attenuated (see supplementary Table S2).

Mediation analysis supported our hypothesis that neuroticism and extraversion partly mediate the relationship between tract FA and depression. Specifically, Sobel tests showed a significant mediating effect of neuroticism between lower uncinate fasciculus FA and depressive symptoms at the first wave of assessment in both the left (test for mediation: Z = -2.38, p = 0.017) and right (test for mediation: Z = -2.03, p = 0.042) hemispheres. Extraversion also significantly partly mediated the relationship between right uncinate fasciculus FA and depressive symptoms in the right hemisphere (test for mediation: Z = -2.90, p = 0.004) but not the left (test for mediation: Z = -1.91, p = 0.056). In each case, the mediating effect of neuroticism was partial, as the (albeit attenuated) effect of white-matter integrity on depressive symptoms remained significant in each case.

Discussion

Using data from the LBC1936 we found that there were significant negative associations between depressive symptoms and white-matter integrity in the bilateral uncinate fasciculus. This relationship was highly significant when the depressive symptoms were rated either at the time of the imaging assessment or 3 years earlier. Subsequently we found that the personality traits of neuroticism and extraversion also have significant associations with white-matter integrity in the uncinate, consistent with their risk-conferring effects on depression. The association between depressive symptoms and white-matter integrity was slightly and significantly attenuated after the inclusion of neuroticism or extraversion in the model. Sobel analyses confirmed that neuroticism and extraversion partially mediated the relationship between white-matter integrity and depression. These associations were significant after adjustment for age and sex, after correction for multiple testing, and were robust to planned sensitivity analyses that took into account previous antidepressant treatment or stroke. These findings suggest that the effect of reduced uncinate white-matter integrity on depressive symptoms is conferred partially through effects on extraversion and neuroticism that have been shown to increase liability to depression in other studies.

No significant associations were found between depressive symptoms and white-matter integrity in any other tract, nor were there any associations between anxiety rating scale scores and white-matter integrity. By contrast, there was a significant negative association (surviving FDR correction) between left cingulum white-matter integrity and neuroticism. Similarly, there was a significant positive association between left ILF integrity and extraversion. To the best of our knowledge, neither of these associations has been reported previously in the literature. The absence of a relationship with anxiety is superficially surprising, as anxiety frequently accompanies depressive symptoms in the clinical disorder. The absence of a relationship between white-matter integrity and anxiety implies a more specific relationship with

Table 3. Association between white-matter tract FA, neuroticism and extraversion

<table>
<thead>
<tr>
<th>Neuroticism</th>
<th>Extraversion</th>
</tr>
</thead>
<tbody>
<tr>
<td>β</td>
<td>s.e.(β)</td>
</tr>
<tr>
<td>Genu</td>
<td>-4.98</td>
</tr>
<tr>
<td>Splenium</td>
<td>-6.64</td>
</tr>
<tr>
<td>LArc</td>
<td>-6.26</td>
</tr>
<tr>
<td>RArc</td>
<td>-1.27</td>
</tr>
<tr>
<td>LATR</td>
<td>-19.93</td>
</tr>
<tr>
<td>RATR</td>
<td>-18.59</td>
</tr>
<tr>
<td>LCCG</td>
<td>-21.78</td>
</tr>
<tr>
<td>RCCG</td>
<td>-17.41</td>
</tr>
<tr>
<td>LUnc</td>
<td>-28.61</td>
</tr>
<tr>
<td>RUnc</td>
<td>-21.04</td>
</tr>
<tr>
<td>LILF</td>
<td>-7.09</td>
</tr>
<tr>
<td>RILF</td>
<td>-7.75</td>
</tr>
</tbody>
</table>

FA, Fractional anisotropy; L, left; R, right; Arc, arcuate fasciculus; ATR, anterior thalamic radiation; CCG, cingulum cingulate gyrus; Unc, uncinate fasciculus; ILF, inferior longitudinal fasciculus.

Bold text used to highlight effects on Unc.

* False discovery rate (FDR) < 0.05.
Depressive symptoms, and suggests that anxiety symptoms are associated with a smaller detectable signal or that their neural correlates are more detectable as changes in gray-matter structure or brain function.

The uncinate fasciculus connects the prefrontal cortex to limbic and emotional regions of the temporal lobe, such as the amygdala and anterior temporal cortex. These areas are implicated in the encoding of rewarding or punishing stimuli and in normal emotional processing and regulation in both the animal (Cardinal et al. 2002) and human (Phillips et al. 2003) literature. Abnormalities in these tracts have been found in studies of unipolar depression (Taylor et al. 2007; Cullen et al. 2010; Dalby et al. 2010) and people at high familial risk of major mental illness (Huang et al. 2011). There is also a growing body of literature showing abnormal frontotemporal activation and connectivity in depression (Anand et al. 2005a,b) and in a variety of other mental disorders (Lawrie et al. 2002; Chepenik et al. 2010) for which neuroticism is an identified risk factor (Maier et al. 1994; van Os & Jones, 2001; Angst et al. 2003; Barnett et al. 2011).

Few studies have directly examined the relationship of white-matter tract FA to neuroticism, although recently a negative association between harm avoidance and measures of tract integrity was reported in several long white-matter tracts (Westlye et al. 2011). Structural and functional abnormalities of frontotemporal brain regions have, however, been linked to specific genetic risk factors for depression and other mental illnesses. These studies include several examining the role of the 5-HTTLPR polymorphism (Ball et al. 1997), in which the s allele confers reduced frontotemporal functional connectivity (Pezawas et al. 2005), increased neuroticism (Barnett et al. 2011) and an increased risk of depression (Caspi et al. 2003). The current findings extend that literature to imply that the personality traits of neuroticism and extraversion may mediate the effects of reduced frontotemporal connectivity on the risk of depressive symptoms.

Although no relationship between left ILF FA and depressive symptoms was found in the current study, left ILF FA was significantly and positively related to extraversion. The ILF connects occipital brain areas with the temporal lobe whereas the cingulum connects the medial prefrontal cortex to the hippocampus. There is evidence that the integrity of these tracts is also associated with liability to depression (Sheline et al. 2008), bipolar disorder (Chaddock et al. 2009; Zanetti et al. 2009; Sprooten et al. 2011) and schizophrenia. It is also possible that reductions in tract integrity in these regions may have important consequences for cognitive function in non-demented adults (Kantarci et al. 2011), a hypothesis that could be addressed in the current cohort at a later imaging assessment.

The current study is the largest investigation to date of depressive symptoms and white-matter tract FA as a proxy for integrity in any non-clinical sample. It should be noted that <10 people in the current study met the specific HADS-D threshold for clinical depression. These thresholds have been largely determined in healthcare-seeking populations, so the positive predictive value is likely to be higher in the current sample where the prevalence of depression is likely to be lower. The association of depressive symptoms with white-matter FA in a non-clinical cohort is likely to have broader applicability to clinical depression for several reasons. First, the symptoms of clinical depression are identical to those captured by the items of most depression-rating scales, albeit without the usual 2-week duration criterion needed to make a clinical diagnosis. However, the choice of a 2-week duration criterion for depression is arbitrary and has limited validity, with shorter duration of symptoms predicting depression in both affected subjects and co-twins (Kendler & Gardner, 1998). Depressive symptoms measured at a single episode also show similar heritability estimates to clinically defined syndromal depression (McGue & Christensen, 1997) and higher estimates of heritability when measured repeatedly over time (McGue & Christensen, 2003). Furthermore, there is evidence that major and subthreshold ‘minor’ depression share a common neural basis (Kumar et al. 1998). Overall, these findings suggest that scale-measured depressive symptoms are associated with trait liability to depression and are not merely measures of current mental state.

A further limitation of this study is the relatively small effect sizes that were found for the relationships between white-matter integrity, depression and the personality traits of neuroticism and depression. All scales for personality and depressive symptoms have strengths and limitations and each only approximates the measurement of the underlying latent variables. The estimates given here may therefore underestimate the true relationships, which cannot be measured in an error-free way. Finally, in the present sample, neuroticism and extraversion were correlated with one another (Pearson’s $r = -0.37$) and it is possible that both measures may be measuring a common underlying trait. Future studies should explore the facets of each personality measure that are independently associated with white-matter integrity, and also perhaps mediate the association with symptoms of depression.

The current study is the first large-scale investigation showing an association between frontotemporal white-matter tract FA, depression and trait
liability to clinical mood disorder. These findings are not due to differences in age, sex or the secondary effects of antidepressant treatment, and suggest that variation in frontotemporal interactions may underlie depressive symptoms at a population level, and may also apply to clinical populations.

Supplementary material

For supplementary material accompanying this paper visit http://dx.doi.org/10.1017/S003329171200150X.

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Declaration of Interest

A.M.M. and S.M.L. report having received research support from Pfizer Pharmaceuticals and consultancy fees from Roche Pharmaceuticals. J.H. reports having received research funding from Pfizer Pharmaceuticals.

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