Familial risk for major depression: differential white matter alterations in healthy and depressed participants

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Abstract

Background. Major depressive disorder (MDD) has been associated with alterations in brain white matter (WM) microstructure. However, diffusion tensor imaging studies in biological relatives have presented contradicting results on WM alterations and their potential as biomarkers for vulnerability or resilience. To shed more light on associations between WM microstructure and resilience to familial risk, analyses including both healthy and depressed relatives of MDD patients are needed.

Methods. In a 2 (MDD v. healthy controls, HC) × 2 (familial risk yes v. no) design, we investigated fractional anisotropy (FA) via tract-based spatial statistics in a large well-characterised adult sample (N = 528), with additional controls for childhood maltreatment, a potentially confounding proxy for environmental risk.

Results. Analyses revealed a significant main effect of diagnosis on FA in the forceps minor and the left superior longitudinal fasciculus (pFWE = 0.009). Furthermore, a significant interaction of diagnosis with familial risk emerged (pFWE = 0.036). Post-hoc pairwise comparisons showed significantly higher FA, mainly in the forceps minor and right inferior fronto-occipital fasciculus, in HC with as compared to HC without familial risk (pFWE < 0.001), whereas familial risk played no role in MDD patients (pFWE = 0.797). Adding childhood maltreatment as a covariate, the interaction effect remained stable.

Conclusions. We found widespread increased FA in HC with familial risk for MDD as compared to a HC low-risk sample. The significant effect of risk on FA was present only in HC, but not in the MDD sample. These alterations might reflect compensatory neural mechanisms in healthy adults at risk for MDD potentially associated with resilience.

Introduction

Major depressive disorder (MDD) is a debilitating, life-quality diminishing, and often recurring mental disorder with a lifetime prevalence ranging from 10% to 29.9% (Kessler, Petukhova, Sampson, Zaslavsky, & Wittchen, 2012; for a meta-analysis, see Lim et al. 2018). Besides examining the course and symptomatology of the disorder itself, it is of high clinical relevance to identify risk factors contributing to the development of depressive psychopathology. Family history of MDD is a consistently replicated risk factor: Genetic approaches involving twin studies estimate a heritability of 30–40% (Kendler, Gatz, Gardner, & Pedersen, 2006; Sullivan, Michael Neale, & Kendler, 2000) and one of the largest genome-wide association studies has recently highlighted the complex genetic architecture of depression by identifying more than 200 genes associated with MDD (Howard et al. 2019). Moreover, parental depression increases the risk to develop MDD threefold, and was associated with a higher probability of recurring episodes, mortality and overall poorer functioning as compared to offspring of non-depressed parents (Hammen, Burge, Burney, & Adrian, 1990; Weissman et al. 2016a, 2016b). The period between 15 and 25 years of age is most critical to develop MDD for individuals with familial risk, but also without familial risk, as revealed in the 30-year
follow-up study (Weissman et al. 2016b). Not only offspring, but first-degree relatives of MDD patients in general exhibit elevated rates of MDD as compared to individuals without a familial pre-disposition (Klein, Lewinsohn, Seeley, & Rohde, 2001).

Importantly, this familial risk entails environmental stressors, as genetic risk and in some cases, the emotionally and socially strenuous environment among depressed relatives, jointly contribute to psychopathology in offspring (Goodman & Gotlib, 1999) by, for instance, emotionally neglectful upbringing and other forms of childhood maltreatment (Lovejoy, Graczyk, O’Hare, & Neuman, 2000; Pawlby, Hay, Sharp, Waters, & Pariente, 2011).

Yet, not all healthy individuals exposed to depression in their familial surrounding manifest clinically relevant depressive symptoms, but are resilient to this risk. Resilience is understood as the dynamic ability to maintain well-being and mental health by activating protective resources, even when facing adversity, i.e., suffering, discomfort or a potentially traumatic event in life (Sisto et al. 2019; Windle, 2011). However, little is known regarding neurobiological differences between resilient individuals exposed to familial risk, and exposed individuals who developed MDD themselves. On the one hand, it could be speculated that familial risk exposure shapes brain structure and function towards alterations found in patient samples, suggesting risk-associated markers of vulnerability. On the other hand, the opposite might be the case, suggesting (over-)compensatory mechanisms related to resilience in exposed persons.

While some neuroimaging studies rather point towards a structural and functional neurobiological similarity of at-risk individuals and MDD patients, e.g., regarding reduced hippocampal volume (Amico et al. 2011; Peterson et al. 2009) and alterations in emotion-processing circuits (e.g. Gotlib, Joormann, & Folan-Ross, 2014; Opel et al. 2017), others have provided evidence for correlates of resilience by increased grey matter volume in the dorsolateral prefrontal cortex (Brosch et al. 2021). Despite the soaring awareness that differences in white matter (WM) fibre structure seem to play a role in the pathophysiology of MDD (Murphy & Frodl, 2011; van Velzen et al. 2020), examining WM microstructure alterations in participants with familial risk has been neglected. The superior longitudinal fasciculus (SLF) is a major WM tract connecting the frontal lobe with posterior regions (Makris et al. 2005). While its function has not been fully established yet, it has been associated with depression (Murphy & Frodl, 2011), emotion processing (Koshiyama et al. 2020), and working memory (Karlgodt et al. 2008). Other tracts which have been associated with depression in a large meta-analysis are the corpus callosum, which connects the two hemispheres, and the corona radiata, a segment of the limbic-thalamo-cortical circuitry involved in emotion regulation (van Velzen et al. 2020). However, the authors point towards a structural disconnectivity in depressed individuals as the revealed alterations in WM microstructure in adults with MDD are rather widespread. Alterations in these tracts can be observed through magnetic resonance diffusion tensor imaging (DTI), which presents a non-invasive technique investigating WM architecture in vivo based on the tissue water diffusion rate. Most frequently studied is fractional anisotropy (FA), which is regarded as a quantitative index of WM coherence. FA quantifies directional diffusion from zero (= isotropic) to one (= anisotropic/constrained along one axis). High FA values are interpreted as revealing a highly organised and normally myelinated axon structure. In contrast, decreased FA might represent reduced coherence in the main preferred diffusion direction and accordingly reflect WM dysfunction.

To the best of our knowledge, only few DTI studies, with small sample sizes, examined participants with familial risk. On the one hand, there is evidence of decreased FA in at-risk participants, pointing towards family history as a risk factor (Brachat, Linden, & Keedwell, 2015; Huang, Fan, Williamson, & Rao, 2011; Keedwell et al. 2012), implying that healthy at-risk groups resemble diagnosed patients on a neurobiological level. On the other hand, Frodl et al. (2012) provided evidence for greater FA in healthy controls (HC) at-risk as compared to low-risk HC. This study did not include MDD patients, but the results suggest a compensation mechanism which might be further used to distinguish healthy at-risk participants from already-affected at-risk MDD patients.

In sum, there are contradictory reports on WM alterations associated with familial risk, and a lack of DTI studies which analyse (1) healthy as well as depressed individuals (2) with and without familial risk for MDD, (3) including adult, not only adolescent participants, (4) while controlling for the closely related, and therefore potentially confounding factor environmental risk, as previous studies have found distinct WM correlates of childhood maltreatment (Frodl et al. 2012; Huang, Gundapuneedi, & Rao, 2012; Meintert et al. 2019). Therefore, the objective of the present study was to investigate the familial risk for MDD and its associations with WM microstructure in a large well-characterised sample of HC and MDD with additional consideration of childhood maltreatment. Based on multiple previous findings (Huang et al. 2012; Meintert et al. 2019; Repple et al. 2020; van Velzen et al. 2020), we expect a main effect of diagnosis, more specifically, reduced FA in MDD compared with HC, specifically in the left SLF. We further expect that healthy relatives of MDD patients might show adaptation on a neural level, i.e., increased FA, despite the familial risk as they have already entered adulthood and passed the critical age range in adolescence according to findings by Weissman et al. (2016b). Lastly, we explore the effects of childhood maltreatment as an environmental component of familial risk by correcting our findings for the degree of self-reported maltreatment experiences.

Materials and methods

Participants

The present sample was drawn from the bicentric Marburg-Münster Affective Cohort Study (MACS/FOR2107-cohort) which has been previously described in more detail elsewhere (Kircher et al. 2019; see Vogelbacher et al. 2018, for the MRI quality assurance protocol). It consisted of N = 528 participants (N = 401 female, Mage = 31.26, S.D.age = 11.72), aged from 18 to 65 years with West-European ancestry; N = 262 HC compared with N = 266 patients with a lifetime diagnosis of MDD. Furthermore, we divided these two groups into two similarly-sized risk and low-risk groups, respectively; N = 129 HC (HCr) and N = 132 MDD with familial risk for MDD (MDDfr), and N = 133 HC (HClr) and N = 134 MDD without MDDfr (see online Supplement S1 for further information on sample selection). We operationalised familial risk as reporting at least one first-degree relative (biological mother, father, siblings and/or children) with a known history of past or current psychological treatment due to a diagnosis of MDD, while no other primary psychiatric diagnoses were allowed in these relatives. In contrast, we only included participants in the low-risk groups...
Association, 2000) for a lifetime diagnosis of MDD. The Beck Mental Disorders IV (DSM-IV-TR; American Psychiatric Association, 1994) was employed by trained personnel to assess whether participants fulfilled standardised criteria defined by the Diagnostic and Statistical Manual of Mental Disorders IV (DSM-IV-TR; American Psychiatric Association, 2000) for a lifetime diagnosis of MDD. The Beck Depression Inventory (BDI; Beck, Ward, Mendelson, Mock, & Steer, 1988) was administered additionally to assess the presence and severity of current (subclinical) depressive symptomatology, while all HCs with BDI score \( \geq 10 \), considering the cut-off for none or minimal depression, were excluded (Beck, Steer, & Carbin, 1988). HCr v. HCrl as well as MDDrl v. MDDlr did not differ significantly in BDI scores (Table 1).

For an overview of antidepressant medication intake, we calculated a Medication Load Index (MedIndex) for every MDD participant, i.e., the sum of absent medication \( (=0) \), equal or lower than the average dose \( (=1) \), or higher than the average dose \( (=2) \) for each psychopharmacological agent. This is an established method (Hassel et al., 2008; Redlich et al., 2014) considering the active ingredient and the daily dose intake recommended by the Physician’s Desk Reference (Reynolds, 2008).

Since age of onset of MDD (van Velzen et al., 2020) and lifetime cumulative duration of depressive episodes (De Diego-Adelino et al. 2014) have been shown to be significantly associated with disruption of WM, it is important to note that the two MDD groups did not differ significantly with respect to these variables. Furthermore, MedIndex, remission status, and recurrence of episodes were evenly distributed across the two groups (Table 2).

Concerning childhood maltreatment, HCr v. HCrl as well as MDDlr v. MDDlr, respectively, did not differ significantly (all \( p > 0.05 \)) in the total score of the German version of the Childhood Trauma Questionnaire (CTQ; Wingenfeld et al., 2010) (Table 1).

The study protocol for this cohort was approved by the ethics committee of the Medical Faculties, University of Marburg (AZ: 2014-422-b-S) and University of Münster (AZ: 2014-422-b-S). The procedure was performed in accordance with the ethical guidelines and regulations. All participants provided written informed consent and received financial compensation for participation.

### DTI data acquisition

All participants were examined on a 3 T whole body MRI scanner (Marburg: Tim Trio, Siemens, Erlangen, Germany; Münster: Prisma fit, Siemens, Erlangen, Germany) with a GRAPPA acceleration factor of 2 and identical sequence parameters for both sites: Fifty-six axial slices, 2.5-mm thick with no gap, were measured with cubic voxels of 2.5 mm edge length (TE = 90 ms, TR = 7300 ms). Five non-diusion-weighted (DW) images \( (b_0 = 0) \) and 2 × 30 DW images with a b-value of 1000 s/mm² for spatial directions were acquired. These scan parameters were consistent over the two sites. GRE field mapping: 2:01; DTI 1 & 2 each

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>HC (n = 262)</th>
<th>MDD (n = 266)</th>
<th>Across all groups</th>
<th>Test statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HCrl (n = 133)</td>
<td>HCr (n = 129)</td>
<td>MDDlr (n = 134)</td>
<td>MDDlr (n = 132)</td>
</tr>
<tr>
<td>Sex, m/f</td>
<td>35/98</td>
<td>33/96</td>
<td>29/105</td>
<td>30/102</td>
</tr>
<tr>
<td>Site: Marburg/ Münster</td>
<td>64/69</td>
<td>64/65</td>
<td>53/81</td>
<td>67/65</td>
</tr>
<tr>
<td>Age</td>
<td>31.03 ± 11.74</td>
<td>31.78 ± 12.51</td>
<td>29.97 ± 10.83</td>
<td>32.27 ± 11.83</td>
</tr>
<tr>
<td>Education, in years</td>
<td>14.05 ± 2.40</td>
<td>13.84 ± 2.20</td>
<td>13.62 ± 2.39</td>
<td>13.67 ± 2.63</td>
</tr>
<tr>
<td>CTQ</td>
<td>31.32 ± 8.54</td>
<td>32.22 ± 7.90</td>
<td>41.77 ± 16.09</td>
<td>44.67 ± 14.05</td>
</tr>
<tr>
<td>BDI( ^d )</td>
<td>2.61 ± 2.18</td>
<td>3.00 ± 2.35</td>
<td>17.67 ± 10.94</td>
<td>16.56 ± 11.56</td>
</tr>
<tr>
<td>SIGH-ADS( ^f )</td>
<td>1.6 ± 2.36</td>
<td>2.1 ± 2.38</td>
<td>12.43 ± 8.74</td>
<td>11.59 ± 9.46</td>
</tr>
</tbody>
</table>

Abbreviations: HC, healthy controls; MDD, major depressive disorder; HCrl, low-risk HC; HCr, at-risk HC; MDDlr, low-risk MDD; MDDlr, at-risk MDD; CTQ, Childhood Trauma Questionnaire; BDI, Beck Depression Inventory; SIGH-ADS, Structured Interview Guide for the Hamilton Depression Rating Scale with Atypical Depression online Supplement.

\( ^a \) One-way analysis of variance (ANOVA) assuming equal variance.

\( ^b \) Test for independent samples, assuming unequal variance.

\( ^c \) Test for independent samples, assuming equal variance.

\( ^d \) Data not available for one MDD participant.

\( ^f \) Data not available for 14 participants.

Note. Numbers represent respective n, or Mean ± Standard Deviation.

** ** p < 0.001, two-tailed.
Table 2. Clinical characteristics and medication in the MDD sample (n = 266)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>MDDr (n = 132)</th>
<th>MDDlr (n = 134)</th>
<th>Test statistic</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lifetime psychiatric comorbidity</td>
<td>56</td>
<td>64</td>
<td>$\chi^2(1) = 7.65$</td>
<td>0.382*</td>
</tr>
<tr>
<td>Recurrent episodes</td>
<td>87</td>
<td>73</td>
<td>$\chi^2(1) = 3.63$</td>
<td>0.057*</td>
</tr>
<tr>
<td>Remission status: Acute/partially/fully remitted</td>
<td>50/31/51</td>
<td>70/25/39</td>
<td>$\chi^2(2) = 5.56$</td>
<td>0.062*</td>
</tr>
<tr>
<td>Lifetime cumulative duration of depressive episodes (months)$^b$</td>
<td>41.67 ± 72.62</td>
<td>29.32 ± 42.01</td>
<td>(227) = 2.85</td>
<td>0.144*</td>
</tr>
<tr>
<td>Number of lifetime depressive episodes$^d$</td>
<td>3.46 ± 6.75</td>
<td>2.59 ± 2.70</td>
<td>(253) = 2.47</td>
<td>189*</td>
</tr>
<tr>
<td>Duration of current episode (months)$^c$</td>
<td>17.54 ± 26.34</td>
<td>13.34 ± 19.52</td>
<td>(156) = 1.72</td>
<td>0.252*</td>
</tr>
<tr>
<td>Age of onset$^e$</td>
<td>21.6 ± 9.52</td>
<td>22.78 ± 9.89</td>
<td>t(263) = 1.00</td>
<td>0.320*</td>
</tr>
<tr>
<td>Medication</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agomelatine</td>
<td>6</td>
<td>9</td>
<td>$\chi^2(1) = 0.59$</td>
<td>0.443*</td>
</tr>
<tr>
<td>Tricyclic Antidepressant</td>
<td>9</td>
<td>5</td>
<td>$\chi^2(1) = 1.27$</td>
<td>0.260*</td>
</tr>
<tr>
<td>Noradrenaline dopamine reuptake inhibitor</td>
<td>5</td>
<td>4</td>
<td>$\chi^2(1) = 0.13$</td>
<td>0.717*</td>
</tr>
<tr>
<td>Selective Serotonin Reuptake Inhibitor</td>
<td>33</td>
<td>32</td>
<td>$\chi^2(1) = 0.05$</td>
<td>0.832*</td>
</tr>
<tr>
<td>Serotonin-Norepinephrine Reuptake Inhibitor</td>
<td>33</td>
<td>30</td>
<td>$\chi^2(1) = 0.25$</td>
<td>0.616*</td>
</tr>
<tr>
<td>Noradrenergic and specific serotonergic antidepressant</td>
<td>9</td>
<td>7</td>
<td>$\chi^2(1) = 0.30$</td>
<td>0.585*</td>
</tr>
<tr>
<td>Hypericum perforatum</td>
<td>1</td>
<td>2</td>
<td>$\chi^2(1) = 0.32$</td>
<td>0.570*</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>0</td>
<td>2</td>
<td>$\chi^2(1) = 1.99$</td>
<td>0.159*</td>
</tr>
<tr>
<td>No antidepressant medication</td>
<td>52</td>
<td>60</td>
<td>$\chi^2(1) = 0.79$</td>
<td>0.374*</td>
</tr>
<tr>
<td>MedIndex</td>
<td>1.16 ± 1.18</td>
<td>1.22 ± 1.41</td>
<td>t(264) = 0.41</td>
<td>0.685*</td>
</tr>
</tbody>
</table>

Abbreviations: MDD, major depressive disorder; MDDlr, low-risk MDD; MDDR, at-risk MDD; MedIndex, Medication Load Index.

$^a$t-test (two-tailed).
$^b$Data available for n = 229.
$^c$t-test for independent samples (two-tailed), assuming equal variance.
$^d$Data available for n = 255.
$^e$Only available for patients in their acute stage, n = 158.
$^f$Data missing for one participant; $^t$ t test for independent samples (two-tailed), assuming unequal variance.

Note. Numbers represent respective n, or Mean ± Standard Deviation.

4:10; DTI b0 3x 0:31; reverse phase encoding 0:53; in total, the DTI acquisition lasted 10 min and 46 s.

Image processing

Preprocessing and analyses were conducted with FSL6.0.1 [http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/, FMRIB, Oxford Center for Functional MRI of the Brain, University of Oxford, Department of Clinical Neurology, John Radcliffe Hospital, Oxford, United Kingdom (Jenkinson, Beckmann, Behrens, Woolrich, & Smith, 2012; Smith et al. 2004; Woolrich et al. 2009)]. DTI image processing methods and quality controls were already described in Meinert et al. (Meinert et al. 2019). Briefly, the DW images were corrected for motion artefacts and eddy currents with FSL’s ‘eddy’ (Andersson & Sotiropoulos, 2016), with a subsequent rotation of b-vectors. As reference for alignment, the first b0 was used after automated skull stripping. For this purpose, FMRIB’s brain extraction tool (Smith, 2002) was applied prior to fitting a diffusion tensor model at each voxel using ‘DTIFIT’ within FMRIB’s Diffusion Toolbox (FDT) (Behrens et al. 2003). For quality assurance of our data, the open-source software DTIPrep (Oguz et al. 2014) was used with default options with an automatic pipeline. Gradients failing checks for intensity-related artefacts are deleted. Intensity artefacts are gradients with a large deviation from the mean of all gradients and were subsequently deleted. As a result, affected images up until a threshold of $\leq 20\%$ were deleted, in which case the participant was excluded from further analyses. This resulted in a number of 64.32 images on average for our sample [s.d. = 1.31, range: (54–65)].

FA is the most frequently reported measure of diffusion in DTI studies besides mean diffusivity (MD), radial diffusivity (RD), and axial diffusivity (AD). We focus on FA with the intention of making our results comparable to previous studies in this area. Our analyses on MD, RD, and AD are available in more detail in the online Supplements. These additional measures contribute to findings of FA providing information about tissue microstructure alterations. MD is a parameter of overall diffusion, averaged across the three eigenvalues of the diffusion tensor. Frequently denoted as the apparent diffusion coefficient. RD reflects diffusion radial to the axes, whereas AD is associated with diffusion parallel to the axes (Alexander, Lee, Lazar, & Field, 2007; Feldman, Yeatman, Lee, Barde, & Gaman-Bean, 2010).

Analyses

Demographic and clinical data were analysed with the Statistical Package for Social sciences (IBM SPSS Statistic 27; SPSS Inc., Chicago, IL, USA). Tract-based spatial statistics (TBSS) (Smith et al. 2006) were applied for the reduction of partial volume effects and registration misalignments. We corrected for multiple
comparisons with Threshold-Free Cluster Enhancement (TFCE) with 5000 permutations. Additionally, estimated cluster sizes were corrected for the family-wise error (FWE) at $p < 0.05$. Since data were acquired with two different MRI scanners, and due to a body coil exchange in Marburg, two dummy-coded variables (Marburg pre body-coil: yes/no, Marburg post body-coil: yes/no) with Münster as reference category were created (Vogelbacher et al. 2018).

In a first step, analyses of covariance (ANCOVAs) were conducted in FSL with FA (as well as AD, RD and MD, respectively) as dependent variables, and diagnosis as well as state of familial risk as independent variables to estimate the main effect of diagnosis and familial risk as well as their interaction. Nuisance variables were age, sex, total intracranial volume (TIV), Marburg pre body-coil, and Marburg post body-coil. We investigated the main effect of diagnosis, and the main effect of risk as well as the interaction effect of diagnosis $\times$ risk ($F$ tests). In case of significant results, post-hoc pairwise $t$ tests were calculated to investigate individual group differences and the direction of the effects. For analyses in FSL, effect sizes were calculated based on the $t$ value of the peak voxel provided by FSL and respective sample sizes according to Cooper, Hedges, & Valentine (2009).

Furthermore, we conducted accessory analyses and robustness checks: In two sub-analyses, we excluded (a) participants under the age of 26, aiming to confirm results in a sample which has presumably passed the critical age range for developing MDD according to Weissman et al. (2016b), and (b) participants of or above the age of 60 to rule out effects of age on FA (Salat et al. 2005; Sexton et al. 2014; Wallhovd et al. 2005; Westlye et al. 2010). Next, we conducted control analyses for potential effects of outliers using SPSS by extracting individual mean FA values of the significant cluster from the main analysis. In the total sample, we examined outliers with Cook’s distance $> 3$ S.D. and recalculated the identical general linear model (GLM) in SPSS. Furthermore, we investigated the influence of childhood maltreatment including CTQsum as a covariate. For analyses in SPSS, given effect sizes in $\eta^2_p$ were converted into Cohen’s $d$ for comparability. In addition, we conducted an accessory sensitivity analysis in which we only included participants in the risk groups whose parents had been treated for MDD (online Supplement S5).

**Results**

**Main analysis: ANCOVA with main effect of diagnosis and risk, and their interaction**

A significant main effect of diagnosis was found ($p_{\text{tfce-FWE}} = 0.021$, total $k = 1774$ voxels in 3 clusters, peak voxel of a largest cluster: $x = 27, y = -23, z = 16$). A post-hoc $t$ test revealed that individuals from both HC groups had higher FA compared with both MDD groups in the forceps minor (FM) and the left SLF with a moderate effect size ($d = 0.40$, $p_{\text{tfce-FWE}} = 0.009$, total $k = 19,792$ voxels in 4 clusters, peak voxel of largest cluster: $x = 24, y = -20, z = 6$, Fig. 1). Significant effects were also found on RD ($p_{\text{tfce-FWE}} = 0.033$) but not on MD ($p_{\text{tfce-FWE}} = 0.314$), with increased RD in MDD patients (online Supplement S2). The main effect of familial risk was not significant neither for FA ($p_{\text{tfce-FWE}} > 0.265$), nor RD ($p_{\text{tfce-FWE}} > 0.129$), nor MD ($p_{\text{tfce-FWE}} > 0.125$). However, the interaction of diagnosis $\times$ familial risk yielded significant results ($p_{\text{tfce-FWE}} = 0.036$, total $k = 9297$ voxels in 5 clusters, peak voxel of largest cluster: $x = 26, y = -21, z = 28$). Clusters predominantly in the left corticospinal tract (CST) and the bilateral SLF were affected by altered FA (Fig. 2a). The effect was also significant for MD ($p_{\text{tfce-FWE}} = 0.042$) and RD ($p_{\text{tfce-FWE}} = 0.008$) (online Supplement S2). No significant main or interaction effects were found for AD (all $p_{\text{tfce-FWE}} > 0.115$).

Post-hoc pairwise $t$ tests revealed a significant difference between the two HC groups, with widespread increased FA values in the HCr group ($p_{\text{tfce-FWE}} < 0.001$, total $k = 41,494$ voxels in one cluster, peak voxel: $x = 37, y = -37, z = 12$), mainly in the FM and right inferior fronto-occipital fasciculus (IFOF), with a moderate

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**Fig. 1.** Affected white matter tracts by effect of diagnosis (HC > MDD).

Note. Increased FA in healthy controls as compared to patients with major depressive disorder, mainly in the forceps minor and superior longitudinal fasciculus, $p_{\text{tfce-FWE}} = 0.009$. In order to illustrate the effects on the FMRIB58 template (visualised in green) for all three sectional views in the Montreal Neurological Institute (MNI) Atlas coordinate system, the mean FA value was obtained from FA values of all significant voxels ($p_{\text{tfce-FWE}} < 0.05$). Red-yellow areas represent voxels in significant clusters, using the ‘fill’ command in FSL. The colour bar indicates the probability of a voxel being a member of the different labelled regions within the JHU-atlas, averaged over all the voxels in the significant mask ($p_{\text{tfce-FWE}} < 0.05$). MNI coordinates of selected plane: $x = 26, y = -19, z = 10$. FM, forceps minor; SLF, superior longitudinal fasciculus.
Effect size ($d = 0.59$) (Fig. 2b). Furthermore, HCr also had significantly higher FA values compared to the MDDr and MDDlr group (MDDr: $p_{FWE} < 0.001$, total $k = 42,419$ voxels in one cluster, peak voxel: $x = 28, y = -18, z = 21$, $d = 0.67$; MDDlr: $p_{FWE} < 0.005$, total $k = 34,734$ voxels in one cluster, peak voxel: $x = -15, y = -9, z = -6$, $d = 0.61$). For location and size of all significant clusters as well as affected tracts see online Supplementary Tables S2 and S3. All other effects were not significant (all $p_{FWE} > 0.332$; online Supplementary Table S3).

### Accessory analyses and robustness checks

Analysing only participants with $\geq 26$ years of age ($N = 284$) the post-hoc differences between healthy risk and low-risk participants found in the original sample could be replicated with a nominally higher effect size ($p_{FWE} = 0.009$, $d = 0.75$) (online Supplement S3). Similarly, including participants with < 60 years of age yielded the same results (online Supplement S4). The results from our first analyses remained also unchanged when excluding participants whose children or siblings were affected (online Supplement S5).

Control analyses in SPSS confirmed our main results. Even after excluding 14 outliers from the original sample, the main effect of diagnosis and interaction effect remained significant (online Supplement S4). A weak but significantly negative correlation between CTQsum and FA scores ($r(526) = -0.094$, $p = 0.031$) did not abolish the effect either: when including CTQsum as a covariate in the general linear model in SPSS, the main effect ($F(1,527) = 4.28, p = 0.039, d = 0.18$) and interaction effect ($F(1,527) = 36.18, p < 0.001, d = 0.53$) were still significant with no significant impact of CTQ ($F(1,527) = 0.325, p = 0.569$).

### Discussion

In the present study, we investigated cross-sectional correlates of familial risk for MDD and WM microstructure in a large sample of HC and patients with MDD. We were (a) able to replicate a previously reported significant main effect of diagnosis and (b) found a significant interaction effect of familial risk with diagnosis. HC had increased FA as compared to MDD participants in the FM and left SLF. The interaction effect was driven by widespread increased FA in healthy participants with a first-degree relative with MDD as compared to a low-risk healthy sample, mainly in the FM and right IFOF. The additional covariance of self-reported retrospective childhood maltreatment scores did not change the significant results, suggesting the reported WM alterations reflect rather genetic than environmental risk which has also been associated with distinct disruptions in WM microstructure (Frodl et al. 2012; Huang et al. 2012; Meinert et al. 2019), effects on cognition (Goltermann et al. 2021) and grey matter anomalies (Opel et al. 2016). No significant differences were found between MDDr and MDDlr concerning all four analysed diffusion indices while recurrence of episodes and state of remission did not differ between the MDD samples.

We were able to replicate results by Murphy & Frodl (2011) indicating decreased FA in MDD participants across different states of remission in the left SLF as compared to HC. Connecting the frontal cortex with the parietal, temporal and occipital cortex, the SLF is a major bidirectional association tract which is involved in specific behavioural and cognitive functions in healthy adults such as verbal memory, but also non-verbal cognition, working memory, and visuo-spatial functions (Koshiyama et al. 2020), as well as recognition of emotional
faces (Ioannucci, George, Friedrich, Cerlani, & Thiebaut de Schotten, 2020). A meta-analysis across emotional disorders identified reduction in FA in the SLF, as compared to HC, as the most replicable finding and concludes that the functions associated with the SLF are in line with impaired perception of and attention to emotional information in these disorders (Jenkins et al. 2016). Consolidation of WM fibre microstructure in the SLF might be positively associated with emotional perception and cognitive control in HC as compared to MDD in general.

In contrast to previous studies, no significant reduction in FA in our healthy risk v. low-risk participants were found in the bilateral cingulum bundles (Bracht et al. 2015; Huang et al. 2011; Keedwell et al. 2012). Instead, the present increases in FA in HC r as compared to HClr are partly in contrast to previous studies with a similar design, revealing decreases in FA. Nevertheless, our results are in line with Frodl et al. (2012) who also reported increased FA in the right IFOF, among other tracts, in healthy relatives of MDD patients as compared to HC without familial risk. These structural differences could result in differences in the neural processing of emotion. The IFOF is involved in emotional visual function, building an association between the vision-related ventro-medial occipital to the emotion-related infero- and dorsolateral regions of the frontal lobe (Catani, Howard, Pajevic, & Jones, 2002). Altered emotional visual perception, i.e., altered transmission of valenced signals linked to reduced FA in IFOF has been observed in depression (Kieseppä et al. 2010) and female adolescent HCr (Joormann, Cooney, Henry, & Gotlib, 2012). We also found increased FA in the FM in HCr v. HClr. The FM connects the two hemispheres, more specifically the dorsolateral prefrontal cortices (DLPFC). The DLPFC is involved in attention, executive functions and internally guided behaviour during goal-oriented and working memory tasks (Kane & Engle, 2002). Furthermore, it is activated during emotion regulation processes (Dixon, Thiruchselvam, Todd, & Christoff, 2017; Versace et al. 2018), more specifically mindfulness-based negative emotion regulation strategies (Opialla et al. 2015). Enhanced WM fibres in the IFOF and FM might point towards enhanced negative emotion regulation and activation of cognitive strategies in HCr which prevent depressive symptomatology and promote mental health.

A reason why we found results contradicting previous studies might be a different age range of participants. Importantly, Huang et al. (2011) included adolescents from age 12 to 20 only, whereas our sample consisted of individuals between 18 and 65, with the majority of participants in an age range from 25 to 30. Weissman et al. (2016b) found that the risk for first onset of major depression in high-risk and low-risk individuals was highest between ages 15 and 25. In our reported sub-analysis excluding participants from our sample aged under 26 years, we were indeed able to replicate the findings of increased FA in HCr v. HClr with a higher effect size than in the original analysis. Our participants might therefore have adapted to the increased risk: Increased FA in HCr might reflect overcompensation, a neurobiological mechanism allocating more, yet protective, resources in tracts connecting areas which are involved in emotion-processing, emotion regulation and executional tasks. Extending the results of Frodl et al. (2012), we revealed an interaction between diagnosis and familial risk, suggesting that our HCr group is resilient in contrast to MDDr. However, this should be addressed in future studies with measurements of resilience, e.g., questionnaires.

### Limitations

Our study's cross-sectional design does not allow disentangling between precursor and consequence. Longitudinal studies are needed to account for putative causal relationships between familial risk factors and alterations in WM microstructure.

Even though reported in the follow-up study by Weissman et al. (2016a, 2016b), studies have exposed contradictory findings on the age of onset for MDD (Kessler et al. 2005; Solmi et al. 2021). However, we based our decision to conduct an accessory analysis with an older age group with this specific cut-off on data in high-risk and low-risk individuals which were investigated over a time period of 30 years.

Secondly, we relied on participants’ self-report on whether a first-degree relative suffered from MDD. In order to increase the level of confidence regarding diagnosis, we only included participants who confirmed that the affected relative received treatment related to the diagnosis, but however, we cannot exclude that participants categorised as low-risk might have been unaware of affected relatives. Including only participants whose relatives were treated for depression might also indicate more severe cases of depression, and might also be influenced by the social and financial ability to seek and receive treatment. Moreover, we did not control for the number of affected relatives. Relatedly, family history and personal depression are independent. On the one hand, the experience of having a first-degree relative with MDD might lead to more attention in the individual towards this disorder. The individual potentially reports symptoms earlier and seeks treatment faster, as they are aware of the increased risk. On the other hand, self-experienced stigma and shame might lead to not mentioning affected family members.

Furthermore, our operationalisation of familial risk does not only represent an estimation of genetic, but also of a stressful familial environment. Future studies should investigate this association between genetic markers and WM microstructure. However, childhood maltreatment scores as measured with the CTQSum did not differ significantly between HCr and HClr. Additionally, the interaction effect remained stable when including CTQSum as a covariate in our analyses, making strong influences of the environmental component of familial risk rather unlikely. However, disentangling the specific mechanisms of neglect and abuse as mediators on developing depression needs to be addressed in future studies with specific study designs including groups recruited for presence/absence of childhood maltreatment.

Notably, there were more females than males in our sample. Further studies on the interaction of sex and familial risk on FA would be of interest.

### Conclusion

The main finding of our study shows that resilient individuals with familial risk for MDD exhibit widespread increased FA, with an emphasis on the forceps minor and right IFOF as compared to healthy individuals without such risk. This challenges previous results that HC at risk have decreased FA in the cingulum bundle. The effects remained significant after the exclusion of participants under the age of 26, and when correcting for a measure of environmental risk, i.e., self-reported childhood maltreatment. Even though we were able to replicate results indicating increased FA in HC in the left SLF as compared to MDD participants, the significant interaction of familial risk and diagnosis suggests adaptation processes on a neural level in a healthy, but
not in an already affected-at-risk sample, possibly reflecting resilience. Future longitudinal studies are needed to disentangle precursor and consequence of these WM alterations in resilient individuals and to which degree they can help understand the biological basis of depression, particularly taking the rather modest effect sizes into account.

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