# The Genetic Basis of Major Depression

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The Genetic Basis of Major Depression

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Abstract:

Major depressive disorder (MDD) is a common, debilitating, phenotypically heterogeneous disorder with heritability ranges from 30% to 50%. Compared to other psychiatric disorders, its high prevalence, moderate heritability, and strong polygenicity have posed major challenges for gene-mapping in MDD. Studies of common genetic variation in MDD, driven by large international collaborations such as the Psychiatric Genomics Consortium, have confirmed the highly polygenic nature of the disorder and implicated over 100 genetic risk loci to date. Rare copy number variants associated with MDD risk were also recently identified. The goal of this review is to present a broad picture of our current understanding of the epidemiology, genetic epidemiology, molecular genetics, and gene-environment interplay in MDD. Insights into the impact of genetic factors on the aetiology of this complex disorder hold great promise for improving clinical care.
Introduction

Major depressive disorder (MDD) is a common mood disorder that substantially and negatively impacts both quality and length of life (Kessler and Bromet, 2013), and represents the leading cause of disability worldwide (GBD 2017 DALYs and HALE Collaborators, 2018). This review summarises our existing knowledge about the genetic basis of depression. Specifically, it details how research in this area has informed our understanding of the biology underlying depression and how this knowledge may be of use to improve patient care in the future.

Definition

The term ‘depression’ most often refers to the clinical diagnosis of MDD, as defined by the International Classification of Diseases (ICD) and the Diagnostic and Statistical Manual of Mental Disorders (DSM) diagnostic criteria. It is characterised by the core symptoms of i) depressed mood (e.g., feelings of sadness, irritability, or emptiness), and ii) anhedonia (e.g., decreased ability to feel pleasure or to experience interest in usual activities). Individuals with depression also experience a wide range of other symptoms which can be grouped into cognitive (e.g., concentration difficulties and indecisiveness), emotional (e.g. feelings of worthlessness and hopelessness), and vegetative symptoms (e.g., significant appetite or weight changes, insomnia or hypersomnia) (Malhi and Mann, 2018).

MDD is a heterogeneous disorder, which has important implications for clinical practice and research (Flint and Kendler, 2014; Cai et al., 2020a). A clinical diagnosis of MDD may be reached in many different ways using established criteria (Fried et al., 2020). This phenotypic heterogeneity in clinical practice is mirrored in the broad range of case definition strategies for genetic research. Genetic studies have adopted multiple methods of classifying depression -
from self-reported clinical diagnosis or electronic health record (EHR) diagnosis codes to structured diagnostic interviews (Smith \textit{et al.}, 2013; Howard \textit{et al.}, 2019). Despite the shortcomings of dichotomizing a heterogeneous disorder like MDD, a categorical definition of depression has been widely used in research. Apart from such a diagnosis-based definition, the severity of depressive symptoms is often studied as a quantitative trait, so the full variation of depressive symptoms can be taken into account. These phenotypic definitions can have a considerable impact on genetic findings (Cai \textit{et al.}, 2020b) and their potential interpretation and translation to the clinic. In line with previous studies (Wray \textit{et al.}, 2018), we will use ‘MDD’ where the study subjects met the clinical criteria for major depressive disorder, and ‘major depression’ where the subjects were ascertained using alternative strategies, but nevertheless likely to have met the diagnostic criteria for MDD.

Epidemiology

The 12-month prevalence of MDD is estimated to be \(~6\%\) and the lifetime prevalence \(~20\%\), although these figures vary considerably across countries (e.g., the 12-month prevalence ranged from 2.2\% in Japan to 10.4\% in Brazil) (Bromet \textit{et al.}, 2011). Established epidemiological risk factors for MDD include age, female sex, separated/divorced marital status, physical health problems, and childhood trauma (Moskvina \textit{et al.}, 2007; Bromet \textit{et al.}, 2011; Van Assche \textit{et al.}, 2017). The peak age of onset of MDD extends from mid to late adolescence to the early 40s, with the median in the mid-20s (Kessler and Bromet, 2013). Women are twice as likely as men to be diagnosed with MDD (Bromet \textit{et al.}, 2011). Sex-specific differences in clinical presentation (Rice \textit{et al.}, 2015), environmental risk factors, and differences in the brain’s exposure to sex hormones have been suggested as possible explanations. MDD is associated with chronic physical health problems including cardiovascular diseases, chronic pain disorders, and cancers.
(Kessler and Bromet, 2013). The nature of the association between MDD and physical ill health is complex. There is evidence for a bidirectional causal association, as well as common antecedents such as social deprivation increasing the risk of both MDD and physical ill health (Kessler and Bromet, 2013).

Genetic epidemiology

Early evidence from family-based studies clearly demonstrated a familial aggregation of MDD, with an increased risk in first-degree relatives of MDD patients (odds ratio, OR=2.84, 95% CI =2.31-3.49) (Sullivan et al., 2000). However, families share both genetic and environmental factors and quantifying their relative contributions to MDD liability has been central to genetic epidemiological studies.

Classic quantitative genetic modelling partitions phenotypic variance into genetic and environmental components. The pedigree heritability is then estimated as the proportion of phenotypic variance attributable to additive genetic effects ("narrow-sense heritability") (Lynch and Walsh, 1998; Visscher et al., 2008). Such modelling is typically performed in twin or adoption studies. Twin studies, comparing monozygotic (MZ) and dizygotic (DZ) twins, have consistently demonstrated that MDD is moderately heritable (Table 1) and an early meta-analysis estimated the heritability at 37% (95% CI = 31-42%) (Sullivan et al., 2000). Subsequent twin studies showed similar estimates, ranging between 29-49%, with higher estimates among treated or recurrent MDD cases – potentially due to greater clinical severity in the study samples (Kendler et al., 2006; Polderman et al., 2015; Kendler et al., 2018a). These studies also suggested that the liability variance attributable to the shared family environment is low, ranging from 0 to 11% (Sullivan et al., 2000; Kendler et al., 2006; Polderman et al., 2015;
Kendler et al., 2018a). Adoption studies, comparing the transmission patterns among adoptive and intact families, are better able to separate shared genetics from other aspects of a shared family environment than twin studies (Kendler et al., 2018b). Early adoption studies in MDD, however, were based on small samples and produced conflicting results (Sullivan et al., 2000). Contrary to findings from twin studies, a recent population-based adoption study reported a much lower heritability (approximately 16%) and a higher contribution of the family environment (Kendler et al., 2018b). In addition, the overestimation of the MDD heritability in twin studies has been raised as a potential reason for the “missing heritability” – i.e. the discrepancy between the pedigree heritability and the much lower phenotypic variance explained by genomic loci identified in genome-wide association studies (GWAS) (Manolio et al., 2009). Recent studies utilised national registers and EHR to reconstruct extended familial relationships, and therefore provide estimates of pedigree heritability beyond classic twin design (Wray and Gottesman, 2012; Kendler et al., 2018a; Polubriaginof et al., 2018). These estimates, however, were very similar to those from twin studies (Table 1), suggesting that the heritability of MDD has not been overestimated.

One of the most intensively studied questions regarding the heritability of MDD is whether sex differences in the genetic architecture of the disorder exist. A higher heritability in women than in men might contribute to the higher rates of MDD observed in women. Early meta-analyses failed to detect either quantitative or qualitative evidence for sex differences in MDD heritability (Sullivan et al., 2000). However, more recent studies demonstrated a significantly higher MDD heritability in women than in men (Table 1), although the genetic correlation between the sexes was high (0.63-0.89) (Kendler et al., 2006, 2018a). These results suggest that
most, but not all, genetic risk is shared between the sexes (Kendler et al., 2018a), consistent with the highly polygenic nature of MDD.

Research efforts have attempted to identify genetically more homogeneous MDD subtypes (Flint and Kendler, 2014). Parents with early-onset and recurrent MDD confer the highest risk for MDD in offspring (Birmaher et al., 1996; Sullivan et al., 2000). Other clinical features, including greater clinical severity, comorbid anxiety disorder, and the use of antidepressant and electroconvulsive therapy, also confer higher MDD risk for relatives (Kendler et al., 2018a). Similarly, the Psychiatric Genomics Consortium (PGC) MDD GWAS demonstrated that individuals from many of these subtypes carry higher polygenic risk scores (PRS) for MDD (Power et al., 2017; Wray et al., 2018). Studies examining the heritability of these subtypes are, however, sparse. The heritability among recurrent MDD cases (41%) may be significantly higher than in those who had single-episode (28%) (Fernandez-Pujals et al., 2015), and postpartum depression has been shown to have a higher heritability (44-54%) than non-postpartum depression (32%) (Viktorin et al., 2016). Ascertaining these more heritable forms of MDD has led to some of early GWAS findings (CONVERGE consortium, 2015).

Candidate gene studies

Candidate gene studies of MDD, first performed more than 40 years ago (Beckman et al., 1978), focused on genes thought to be involved in the pathophysiology of the disorder and in drug mechanisms – as reviewed in (Shadrina et al., 2018). More than 1500 publications have assessed variants in over 200 candidate genes with often conflicting results (Flint and Kendler, 2014). Most of these studies were vastly underpowered, lacked correction for population stratification, and the reported significance levels did not exceed what would be expected by
chance (Flint and Kendler, 2014). Based on the observed OR and minor allele frequencies (MAF), all published candidate gene variants should be detectable in the current GWAS at genome-wide significance (Flint and Kendler, 2014). Yet, this has not been the case (Bosker et al., 2011; Wray et al., 2012, 2018; Howard et al., 2019). In large-scale data from the UK Biobank (UKBB) and PGC (n=62,138-443,264 per subsample), systematic analyses of the most commonly studied polymorphisms mapping to 18 candidate genes found no evidence for significant associations with major depression, except at the gene-level for DRD2 (dopamine receptor D2) (Border et al., 2019a).

Candidate gene studies were the mainstay of genetic research in MDD until the emergence of GWAS and some are still among the most highly cited works in the field (McIntosh et al., 2019). However, a lack of robust association has led to the strong recommendation that candidate gene studies should no longer be performed (Duncan and Keller, 2011; Farrell et al., 2015; Border et al., 2019a, 2019b).

The role of rare genetic variants

Historical linkage studies in families with MDD failed to identify causal variants but implicated regions on chromosomes 2, 3, 11, and 15 in disorder risk (Zubenko et al., 2003; Holmans et al., 2004, 2007; Camp et al., 2005, p. 2005; Levinson et al., 2007; Middeldorp et al., 2008; Scholg Gelok et al., 2010; Breen et al., 2011). A highly polygenic architecture, as is the case for MDD, can generate significant linkage study results which could be erroneously interpreted as large-effect loci (Hemani et al., 2013). Indeed, no significant overlap was observed between the reported linkage regions and association loci from current well-powered GWAS (McIntosh et
A major contribution of single highly penetrant variants within these regions to the genetic architecture of MDD is therefore unlikely.

In the past decade, association testing of rare and low-frequency variants has progressed to whole-exome (WES) and whole-genome sequencing (WGS) studies. Rare (MAF <0.5%) or low-frequency (MAF range >0.5% and <5%) variants are expected to contribute to the genetic basis of MDD (Lee et al., 2014; Yu et al., 2018). However, the largest WGS study of MDD to date, by the CONVERGE consortium (5,303 cases and 5,337 controls), failed to find evidence for the association of individual rare or low-frequency variants with MDD (CONVERGE consortium, 2015). Using the same data set, it was subsequently shown that individuals with MDD had a significantly higher overall burden of singleton exonic variants predicted to be deleterious (Peterson et al., 2017). This suggests that larger sample sizes are required to identify associated rare variants with moderate effect sizes (Lee et al., 2014; Border et al., 2019b). An approach to mitigate against the need for vast sample sizes is to examine MDD-related subphenotypes (summarised in Table 2). The majority of these studies identified single rare variants or gene-level burden scores as associated with at least one of the assessed subphenotypes. In most cases, these results await replication in independent and well-powered datasets.

Rare copy number variants (CNVs), deletions or duplications of >1kb of DNA, play a well-established role in risk of neurodevelopmental and psychiatric conditions such as autism spectrum disorder, schizophrenia, and intellectual disability (International Schizophrenia Consortium, 2008; Stefansson et al., 2008; Cooper et al., 2011; Coe et al., 2014), but have received less attention in MDD. Until recently, studies of CNVs in depression have been underpowered and yielded inconsistent and sometimes conflicting results (Glessner et al.,
Two recent large-scale efforts have assessed the contribution of rare CNVs to the genetic architecture of MDD. In a meta-analysis of four cohorts (5,780 cases, 6,626 controls), short deletions <100 kb were significantly enriched in individuals with MDD (Zhang et al., 2019). The enriched deletions mapped primarily to intergenic regions and, within those, to enhancers, suggesting that rare CNVs may increase risk of MDD by altering gene expression. A study of 407,074 individuals from the UKBB (23,979 cases) implicated 53 rare CNVs associated with neurodevelopmental conditions in risk of major depression (OR=1.3–1.5) (Kendall et al., 2019). Three CNV loci—duplications at 1q21.1, the Prader-Willi Syndrome locus at 15q11-13, and at 16p11.2—were individually associated with self-reported depression. Outside of this group, there was no evidence for a residual burden of rare CNVs >100 kb in major depression. However, smaller CNVs were not tested in the study by Kendall and colleagues. Rare CNVs therefore likely contribute to MDD risk, but the exact nature of the association remains unclear.

**Genome-wide association studies**

Many MDD GWAS have been conducted during the past decade, predominantly on subjects of European ancestry (Table 3). Therefore, European-ancestry studies are described first, followed by a review of GWAS in more diverse populations.

Initial MDD GWAS of common variants (i.e., MAF ≥1%) were limited by small sample sizes and failed to identify genome-wide significant loci (p-value threshold <5×10⁻⁸) (Table 3). However, the first suggestive GWAS result implicated the gene encoding the presynaptic active zone
component Piccolo (*PCLO*) (Sullivan *et al.*, 2009) which has been confirmed as a risk gene in subsequent studies (Hek *et al.*, 2010; Wray *et al.*, 2018; Howard *et al.*, 2019). In 2011, the first genome-wide significant locus was reported - *SLC6A15* (a neuron-specific neutral amino acid transporter), although this was in a recessive model (Kohli *et al.*, 2011). This association did not reach genome-wide significance with additive models in larger follow-up GWAS (Wray *et al.*, 2018; Howard *et al.*, 2019). Recognising the importance of combining data across studies to increase statistical power, the PGC published its first MDD GWAS meta-analysis in 2013, although no significant locus was identified (Ripke *et al.*, 2013).

Since then, large-scale GWAS of major depression have been conducted with data from the direct-to-consumer company 23andMe and large-scale biobank efforts such as UKBB (*Table 3*). In 2016, a GWAS by 23andMe using self-reported depression diagnosis identified 15 genome-wide significant loci (Hyde *et al.*, 2016), confirming the estimate that >75,000 cases would be required for genome-wide significant results (Levinson *et al.*, 2014). A second large GWAS meta-analysis from the PGC described 44 associated genetic loci, pooling cohorts with different phenotype definitions (Wray *et al.*, 2018). The three datasets used in the latter studies were eventually combined in an analysis yielding 102 genome-wide significant variants, 87 of which replicated (Howard *et al.*, 2019). The latest GWAS to date meta-analysed these datasets with data from the Million Veteran Program (MVP), yielding 178 associated loci (Levey *et al.*, 2020). Collectively, these GWAS confirmed that major depression is highly polygenic, with ORs of individual variants under 1.05 (Wray *et al.*, 2018; Howard *et al.*, 2019), as suggested previously (Nishino *et al.*, 2018). The two loci with the overall most robust support for an association map to the genes *NEGR1* and *OLFM4*, both with an OR of 1.04 and a MAF of ~40% (Wray *et al.*, 2018;
The neuronal growth regulator 1 (NEGR1) cell adhesion molecule modulates neurite outgrowth, synapse formation, and synaptic function (Wray et al., 2018; Noh et al., 2019). The secreted glycoprotein Olfactomedin 4 (OLFM4) has an established function in inflammation, innate immunity, and cancer (Liu and Rodgers, 2016; Wray et al., 2018). Several other associated loci have functions in neurite outgrowth, synaptic function and plasticity, as well as in immunity and inflammation (Wray et al., 2018). To establish the functions of the proposed risk SNPs (single nucleotide polymorphisms) and their links to specific genes, analyses of cell-type-specific gene expression and epigenetic data are necessary (Zhong et al., 2019; Arloth et al., 2020; Chan et al., 2020; Li et al., 2020; Wang et al., 2020).

Given the small effect sizes of single SNPs, gene-level and pathway analyses that integrate the effects of many variants across genes and pathways may be more suitable for studying the aetiology of MDD. Gene-level association analyses have implicated genes encoding pre- and post-synaptic proteins, especially receptor units, e.g. voltage-gated calcium channels (VGCC), the dopamine receptor DRD2, and glutamate receptors (Wray et al., 2018). Identified by both SNP- and gene-level analyses, the splicing regulator RBFOX1 may play an important role in the aetiology of major depression (Wray et al., 2018; Noh et al., 2019). Gene-set analyses implicated pathways with roles in neuronal differentiation and projection and, most prominently, synaptic formation and function, including gene sets linked to VGCC and synaptic RNA-binding proteins like FMRP and CELF4 (Wray et al., 2018; Noh et al., 2019; Levey et al., 2020).

Despite a large number of identified genomic loci, the variance explained by major depression PRS is still very low (Nagelkerke’s pseudo-$R^2$ 1.5-3.2% on the liability scale) (Howard et al.,
This prediction accuracy was lower than achieved for, e.g. schizophrenia, where Nagelkerke’s pseudo-$R^2$ is 7.7% on the liability scale (Schizophrenia Working Group of the PGC et al., 2020). At much smaller sample sizes, the GWAS on schizophrenia also identified many more associated variants (Figure 1). The identification of MDD risk variants has been more challenging than for schizophrenia because of its higher prevalence, lower heritability, and more heterogeneous samples (Wray et al., 2012; Levinson et al., 2014; Ormel et al., 2019). Schizophrenia GWAS saw a critical inflection point at approximately 15,000 cases and are currently discovering a new locus for, on average, every 230 new cases (Levinson et al., 2014). Such an inflection point was observed for depression GWAS only at approximately 75,000 cases, as predicted (Wray et al., 2012; Levinson et al., 2014). GWAS using liberal phenotype definitions of major depression now identify a new locus for every 1500 cases (Figure 1).

Importantly, phenotype definition matters. Given the strong genetic correlations across cohorts, recent GWAS meta-analyses combined cohorts with different ascertainment strategies to maximize the overall sample size. However, a comparison of genetic analyses using different phenotype definitions in UKBB revealed that GWAS of broad phenotype definitions might identify associated loci that are not specific to MDD, but, more likely, shared with other psychiatric disorders (Cai et al., 2020b). In order to identify robustly associated MDD variants, a balanced strategy of maximising sample size while focusing on more stringent phenotype definitions has been suggested (Davies et al., 2019; Schwabe et al., 2019; Byrne et al., 2020).

Quantitative definitions have also been analysed in GWAS, and a continuum between depressive symptoms and MDD was suggested (Direk et al., 2017). Several variants were reported to be associated with depressive symptoms (Table 3), with limited replication success. Of these symptom-associated variants, rs62100776 (Okbay et al., 2016) maps to the gene DCC,
coding for a neuronal cell adhesion molecule, which was a genome-wide significant locus in the 2018 PGC case-control GWAS (Wray et al., 2018). Further noteworthy GWAS were conducted on MDD stratified by the age at onset (Power et al., 2017), sex (Hall et al., 2018), or exposure to adversity (Peterson et al., 2018), or combined GWAS of different depression phenotypes (Amare et al., 2020).

The studies discussed so far focused on patients of European ancestry. However, complex disorders show population-specific landscapes of genetic risk (Martin et al., 2017). Accordingly, GWAS in non-European populations have also been conducted (Table 3). The CONVERGE study on female Han Chinese recurrent in-patient MDD cases identified two significant loci near the SIRT1 and in the LHPP gene, and replicated them in the same population (CONVERGE consortium, 2015). Neither variant was significantly associated in GWAS on European samples (Wray et al., 2018; Howard et al., 2019), possibly due to large differences in allele frequency across ancestries – both variants exhibit markedly lower allele frequencies in Europeans. Follow-up studies estimated the genetic correlation between East Asian and European MDD cohorts at 0.33-0.41, while the average genetic correlations between European cohorts was 0.76 (Bigdeli et al., 2017; Wray et al., 2018), and identified trans-ancestral risk variants (Li et al., 2018). In a study of Hispanic/Latino individuals, two out of 22 European depression GWAS loci were replicated (Dunn et al., 2018). The MVP study identified no genome-wide significant loci in over 20,000 African-ancestry patients with major depression (Table 3), and only one of 206 analysed European risk SNPs formally replicated in the African sample (Levey et al., 2020). However, a trans-ancestry meta-analysis conducted within the same study increased the number of risk loci from 178 in Europeans alone to 183. These findings highlight the need for
greater geographical, ethnic, and economic diversity in depression genetic studies (McIntosh et al., 2019).

Genetic overlap between MDD and other traits

Several lines of evidence, including cross-disorder analyses, support the notion that MDD shares genetic underpinnings with many other psychiatric phenotypes. A genetic overlap has also been demonstrated between MDD and non-psychiatric phenotypes. Common techniques for studying cross-disorder genetic overlap include genetic correlation ($r_g$) estimation and PRS analyses (Grotzinger et al., 2019).

Genetic correlation quantifies the genetic relationship between two traits, reflecting their underlying modes of pleiotropy – i.e. the same genetic variant influencing both traits (van Rheenen et al., 2019; Baselmans et al., 2020). Mathematically, it is derived as the genetic covariance between the two traits scaled by the square roots of their genetic variances (van Rheenen et al., 2019). Two methods are commonly used to estimate $r_g$ – i) bivariate restricted maximum likelihood (REML) which uses individual-level genotype data as the input (Lee et al., 2012), and ii) linkage disequilibrium score regression (LDSC) (Bulik-Sullivan et al., 2015) and its recent extension, high-definition likelihood (HDL) method (Ning et al., 2020), which uses GWAS summary statistics. $r_g$ ranges between −1 and 1, with 1 meaning that shared liability is caused by the same risk SNPs and −1 meaning that identical SNPs increase the risk for one trait while decreasing risk for the other. It is important to note that $r_g$ estimates cannot infer causality.

Many studies have examined $r_g$ between MDD and both psychiatric and non-psychiatric disorders or traits. The strongest $r_g$ detected for MDD are with depressive symptoms measured
as a continuous trait (0.63-1.0) (Direk et al., 2017; Turley et al., 2018), confirming the concept of a continuum between subthreshold depressive symptoms and clinical diagnoses of MDD.

Similarly, strong $r_g$ have been found for personality traits such as neuroticism ($r_g = 0.70-0.75$) and subjective well-being (-0.75) (Okbay et al., 2016; Howard et al., 2019). Major depression was moderately correlated with several other psychiatric disorders, e.g. autism spectrum disorder (0.45), attention-deficit/hyperactivity disorder (0.44), bipolar disorder (BD) (0.36), and schizophrenia (0.34) (Cross-Disorder Group of the PGC, 2019). Further studies reported a weaker $r_g$ of major depression with type-I (0.26-0.31) than with type-II BD (0.61-0.69) (Stahl et al., 2019; Coleman et al., 2020). Moreover, a positive $r_g$ was reported between the body-mass index (BMI) and atypical depression with increased appetite/weight, which was not present for those without changes in appetite (Milaneschi et al., 2017). An investigation of genetic correlations between neurological and psychiatric disorders found little evidence of shared genetic underpinnings, except for a significant but small positive $r_g$ between MDD and migraine (The Brainstorm Consortium et al., 2018). Negative genetic correlations have been detected for major depression with educational attainment and cognitive performance (each $\sim$-0.15) (Howard et al., 2019).

Analyses of PRS is another method used to assess the polygenic overlap between MDD and other traits. PRS use SNP effect sizes from a training GWAS to compute the cumulative genetic risk burden of each individual in a target sample. PRS are calculated as the weighted sum of the allele count multiplied by its effect size across all selected SNPs (Andlauer and Nöthen, 2020; Lewis and Vassos, 2020). Therefore, they capture the combined effects of many SNPs and are used to examine the polygenic liability for a disorder or trait or to examine the genetic overlap across traits (Wray et al., 2014). Although the generation of PRS requires large training GWAS, it
can be applied to much smaller target sets. For examples, a MDD PRS was predictive of population-based obsessive-compulsive symptoms (Zilhão et al., 2018). Clinically, PRS may be used for diagnosis, prognosis, and treatment prediction (Schijven et al., 2020). However, PRS for schizophrenia have, so far, shown greater potential for such applications, given their higher variance explained (Schizophrenia Working Group of the PGC et al., 2020). Importantly, PRS were used to demonstrate that MDD subtypes may diverge in their genetic overlaps with other disorders. For example, early-onset MDD cases show a stronger association with PRS for schizophrenia and BD than late-onset MDD patients do (Power et al., 2017). In the future, PRS might also be used to identify aetiologically more homogeneous MDD subtypes. For example, subgrouping on the basis of treatments such as electroconvulsive therapy might be informative (Foo et al., 2019) and provide further insights into the underlying genetic differences of the severity of the disorder. Finally, MDD PRS may also be used to study PRS-environment (PRS×E) interactions, to support treatment choice, and to refine the penetrance of high-risk variants (Lewis and Vassos, 2020).

**Gene environment interaction**

In addition to the heritable component of MDD, environmental factors such as stressful life events (e.g. childhood trauma) play a major role in the aetiology of MDD, both independently and in combination with genetic factors. One active area of research has been the study of gene-environment interactions (G×E). Early G×E efforts examined specific candidate genes with putative biological significance for MDD, e.g. the 5-HTTLPR polymorphism of the serotonin transporter gene, and found evidence of interaction with stressful life events (Caspi et al., 2003). However, these initial reports showed inconsistent replicability (Duncan and Keller, 2011), and recent meta-analytic evidence indicated no robust evidence of candidate G×E
effects on MDD (Van der Auwera et al., 2018; Border et al., 2019a). Studies using genome-wide data have reported varied G×E results. First, stratified GWAS results from the CONVERGE study have suggested that MDD may have a stronger genetic basis when it occurs in the absence of environmental adversity, with genome-wide significant hits for MDD identified only among women not exposed to adversity (Peterson et al., 2018). Data from the UKBB, however, have suggested the opposite, with a higher SNP-based heritability for major depression observed among individuals reporting lifetime trauma exposure (Coleman et al., 2020). Second, PRS have been used to aggregate genome-wide effects for MDD and examine their interactions with stressful life events. Again, this revealed discrepant findings where MDD PRS were more strongly associated with MDD among individuals both exposed (Peyrot et al., 2014) and unexposed (Mullins et al., 2016) to childhood trauma. A later but still modestly powered meta-analysis (n=5,765) identified limited evidence for the interaction of PRS and childhood trauma on MDD (Peyrot et al., 2018), which was echoed in epidemiological and clinical cohorts (n=184-1,450) (Halldorsdottir et al., 2019). However, large-scale studies like the UKBB have provided the opportunity to re-examine PRS×E effects with greater power. One study showed stronger PRS effects on major depression among individuals exposed to childhood trauma and socioeconomic adversity (Shen et al., 2020), though not other types of exposures such as adulthood stressful life events. These findings support the notion that genetic influences could vary by specific environments. Third, genome-wide G×E analyses have identified few variants with genome-wide significant G×E effects (Dunn et al., 2016; Arnau-Soler et al., 2019), although these studies have been largely underpowered and results have not been further replicated. Together, current evidence suggests that genome-wide influences on MDD may vary by stressful life exposures but that the detection of G×E may be sensitive to the research design (e.g. how depression and adversity are defined).
The role of stressful life events, specifically childhood trauma, was the predominant focus of G×E in MDD, but other environmental exposures have also received attention. For example, studies examining social support in combination with MDD PRS in vulnerable populations have consistently shown that social support appears to reduce risk of depression. This effect remained even after accounting for genetic vulnerability for depression, but did not vary by or modify the genetic effects (Choi et al., 2020a; Stringa et al., 2020). Similar observations of independent effects have been made for physical activity and major depression, suggesting that even among individuals at high genetic vulnerability for depression, physical activity levels may still be protective (Choi et al., 2020c). Another genetically informed approach to estimating environmental effects on MDD is Mendelian randomisation (MR). MR methods have emerged as an alternative strategy to examine causal inference, obviating typical sources of confounding (Smith and Ebrahim, 2003). Here, strongly associated genetic variants are selected as statistical instruments — or proxies — to estimate the effect of an exposure of interest (e.g., substance use) on an outcome (e.g., MDD) under specific assumptions. MR studies have supported the role of educational attainment in reducing the depression risk and of BMI in increasing the risk (Wray et al., 2018). Studies have also suggested a potential role of lifestyle exposures in shaping the risk of major depression, such as social connections and sedentary media use (Choi et al., 2020b). Establishing stronger causal evidence for different environmental factors with updated methods may help to clarify inconsistencies across G×E findings (Ni et al., 2019). An additional source of potential bias and inconsistency in G×E research are underlying gene-environment correlations, as the presence of gene-environment correlations may lead to false GxE (Purcell and Sham, 2002; Ni et al., 2019).
Discussion

In summary, MDD is a common, debilitating, phenotypically heterogeneous disorder. Its heritability ranges between 30-50%. MDD with early-onset, recurrence, comorbid anxiety, greater severity, and postpartum depression may be more heritable subtypes. Compared to other psychiatric disorders, its high prevalence, moderate heritability, and strong polygenicity have posed major challenges for gene-mapping. Recently, rare CNVs have been implicated in risk of MDD. Studies of common variation in MDD, driven forward by large international collaborations such as the PGC, confirmed the highly polygenic nature of the disorder and implicated over 100 loci in disorder risk. These studies identified genes involved in neuronal growth, synaptic function, and inflammation in the pathophysiology of the disorder. Finally, gene-environment interaction and Mendelian randomisation studies have implicated childhood trauma, socioeconomic adversity, social support, physical activity, and their interactions with genetic factors, in risk of MDD.

Evidence on the genetic basis of MDD has multiple potential applications including the development of pharmacological treatments, informing discussions regarding disorder risk, and treatment choice. A greater understanding of the biological processes underlying MDD may facilitate the development of new, potentially more effective medications. MDD GWAS have implicated biological processes underlying the disorder including neurotransmission and synaptic structure as well as genetically informed pharmacological modes of action (Howard et al., 2019). MDD is a very heterogeneous disorder and genetic research will likely benefit from the study of more homogeneous disorder subtypes. For example, research on patients with atypical features of increased appetite or weight suggested that this patient subgroup might
benefit from treatments specifically targeting immunometabolic pathways (Milaneschi et al., 2017).

An individual’s genetic make-up likely impacts the disease course – from prevention to diagnosis, treatment, and prognosis (Lewis and Vassos, 2020). Whilst the application of MDD PRS to predict individual risk is currently limited by its low discriminative accuracy, in the future PRS might be used for risk stratification at the population level (Andlauer and Nöthen, 2020). This is supported by recent evidence that MDD PRS – well-established to predict prevalent depression – also predict incident cases in the general population (Musliner et al., 2019). For other complex disorders like coronary artery disease and type-2 diabetes, PRS-based risk stratification is already powerful (Khera et al., 2018). Further development of PRS to allow the identification of individuals at heightened genetic risk, coupled with targeted prevention and intervention measures, holds potential to mitigate depression risk (McIntosh et al., 2019).

Knowledge of the genetic architecture of MDD and its links with other psychiatric and non-psychiatric disorders might inform genetic counselling and clinical care. For example, identified genetic correlations with physical health phenotypes might facilitate the development of clinical guidelines for physical health monitoring in individuals with MDD.

MDD is a multifactorial disorder. Other levels of genetic variation, such as DNA methylation (Barbu et al., 2020; Van Assche et al., 2017b) and subsequent gene-expression differences (Arloth et al., 2015; Schiweck et al., 2020), and also the telomere length (Lin et al., 2016), influence depression risk and severity. Importantly, epigenetic mechanisms serve as an interface between the individual and its environment (Provencal and Binder, 2015). A better understanding of the role of epigenetic variation and an improved integration of multi-omics
data may enhance our understanding of the aetiology and treatment of MDD. In addition, this review focuses on genetic factors influencing the risk or severity of depression. We did not discuss how genetic variants affect response to treatment, especially the metabolisation of antidepressant drugs via cytochrome P450 genes. This important topic was recently reviewed by Bousman and colleagues (Bousman et al. 2021).

Here, we highlighted the state of the art, the promises, and the pitfalls of genetic research in MDD. The field would benefit from further exploration of the stratification properties of genetics – both for the understanding of its aetiology, including the role of environmental factors, and for the development of targeted treatment opportunities. Rich datasets with high-quality and deep phenotyping are crucial for studying heterogeneous disorders like MDD. Combining deep, harmonized phenotypes with molecular data will greatly aid the genetic stratification of MDD. Improving our insights into the impact of genetic factors on MDD holds great promise for improving clinical care, and this intense and fast-developing field has a bright future.

Acknowledgments

We thank Andrew McIntosh, Cathryn Lewis, and Naomi Wray for critical comments on our manuscript.
# Tables and Figures

## Table 1. MDD heritability estimates from selected publications

<table>
<thead>
<tr>
<th>Key paper</th>
<th>Type</th>
<th>$h^2$ (95% CI)</th>
<th>$c^2$ (95% CI)</th>
<th>$e^2$ (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Twin data</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Sullivan et al., 2000)</td>
<td>Meta-analysis</td>
<td>0.37 (0.31-0.42)</td>
<td>0 (0-0.05)</td>
<td>0.63 (0.58-0.67)</td>
</tr>
<tr>
<td>(Kendler et al., 2006)</td>
<td>Largest twin study (15.5K pairs)</td>
<td>male: 0.29 (0.19-0.38)</td>
<td>NA$^a$</td>
<td>0.71 (0.62-0.81)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.42 (0.36-0.47)</td>
<td></td>
<td>0.58 (0.53-0.64)</td>
</tr>
<tr>
<td>(Polderman et al., 2015)</td>
<td>Meta-analysis of 50 years of twin studies</td>
<td>depressive episode: 0.34 (0.31-0.38)</td>
<td>0.11 (0.08-0.13)</td>
<td>0.55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recurrent MDD: 0.45 (0.36-0.55)</td>
<td>0.03 (0-0.08)</td>
<td>0.52</td>
</tr>
<tr>
<td>(Kendler et al., 2018a)</td>
<td>Twins identified from SWE register; Treated cases</td>
<td>male: 0.41 (0.19-0.49)</td>
<td>0 (0-0.13)</td>
<td>0.59 (0.51-0.66)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.49 (0.31-0.56)</td>
<td>0.02 (0-0.17)</td>
<td>0.49 (0.44-0.55)</td>
</tr>
<tr>
<td><strong>EHR or register-based data</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Wray and Gottesman, 2012)</td>
<td>DEN register, parent-offspring</td>
<td>0.32 (0.30-0.34)</td>
<td>NA$^b$</td>
<td>NA$^b$</td>
</tr>
<tr>
<td>(Kendler et al., 2018a)</td>
<td>SWE register, siblings (full vs half, reared together vs apart)</td>
<td>male: 0.36 (0.31-0.38)</td>
<td>0.05 (0.05-0.05)</td>
<td>0.59 (0.58-0.61)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.51 (0.51 -0.53)</td>
<td>0.02 (0.02-0.02)</td>
<td>0.47 (0.46-0.49)</td>
</tr>
<tr>
<td>(Polubriaginof et al., 2018)</td>
<td>US EHR, inferred extended pedigree</td>
<td>depressive episode: 0.25 (0.17-0.30)</td>
<td>NA$^a$</td>
<td>0.75 (0.70-0.83)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recurrent MDD: 0.36 (0.21-0.51)</td>
<td>NA$^a$</td>
<td>0.64 (0.49-0.79)</td>
</tr>
</tbody>
</table>

$h^2$ narrow-sense heritability; $c^2$ shared environment; $e^2$ unique environment. SWE: Sweden; DEN: Denmark. EHR: Electronic Health Records.

- a. Missing estimate of $c^2$ as a result of AE models (i.e., quantitative genetic models with only additive genetics and unique environment components) in the original study.
- b. Only the heritability was derived from the summary level data.
Table 2. Whole-exome sequencing and exome chip studies assessing the role of rare variants in subphenotypes of depression.

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Method</th>
<th>Sample size</th>
<th>Ancestry</th>
<th>Implicated gene or variant</th>
<th>Implicated pathway</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population isolate</td>
<td>WES, exome chip</td>
<td>1999 (discovery) 1604 (replication)</td>
<td>European</td>
<td>NKPD1</td>
<td>de novo synthesis of sphingolipids</td>
<td>(Amin et al., 2017a)</td>
</tr>
<tr>
<td>Depressive symptoms in families &amp; population isolate</td>
<td>Linkage, haplotype analysis, WES, exome chip</td>
<td>1336 WES 1527 exome chip</td>
<td>European</td>
<td>RCLI p.Leu186Phe (rs115482041)</td>
<td>interlaminar astrocytes</td>
<td>(Amin et al., 2018)</td>
</tr>
<tr>
<td>Familial &amp; early-onset MDD</td>
<td>targeted resequencing of 1742 synaptic genes</td>
<td>259 cases 334 controls</td>
<td>European</td>
<td>Cav2-adaptor gene set</td>
<td>calcium signaling, actin polymerisation, spine formation</td>
<td>(Pirooznia et al., 2016)</td>
</tr>
<tr>
<td>Familial &amp; early-onset MDD</td>
<td>WES</td>
<td>12</td>
<td>European</td>
<td>ZNF34</td>
<td>brain TFs</td>
<td>(Subaran et al., 2016)</td>
</tr>
<tr>
<td>Sex differences in MDD</td>
<td>WES and WGS</td>
<td>1000 cases (70.7% female) 72 controls</td>
<td>Asian</td>
<td>PDE4A (rs201432982), FDX1L (rs62640397, rs79442975), MYO15B (rs820182, rs820148)</td>
<td>more frequent in female cases</td>
<td>none</td>
</tr>
<tr>
<td>Suicide in MDD</td>
<td>WES specifically with 5’/3’-UTRs and promoters</td>
<td>23 cases 21 controls (post-mortem brain samples)</td>
<td>European</td>
<td>COL6A6 variants in 17% of suicide completers; 3 calcium channel genes highlighted (CACNA1B, -1C, -2D4) 86 non-synonymous variants in 42 genes found in at least two suicide victims but not controls</td>
<td>TGF-β pathway</td>
<td>(Tomáč et al., 2017)</td>
</tr>
<tr>
<td>MRI grey matter volume changes in MDD</td>
<td>WES</td>
<td>77 cases 245 controls</td>
<td>Asian</td>
<td>CSMD1, CNTNAP5</td>
<td>neuroactive ligand receptor interacting pathway</td>
<td>(Zhang et al., 2020)</td>
</tr>
<tr>
<td>Depressive symptoms, general population</td>
<td>WES</td>
<td>3612</td>
<td>European</td>
<td>LIPG p.Asns396Ser (rs77960347)</td>
<td>steroid and cholesterol biosynthesis</td>
<td>(Amin et al., 2017b)</td>
</tr>
<tr>
<td>----------------------------------------</td>
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<td>-------------------------------------</td>
<td>---------------------</td>
</tr>
<tr>
<td>Depressive symptoms in context of environmental stress</td>
<td>exome chip, WGS</td>
<td>203 cases, 196 controls (discovery) 473 cases, 497 controls (replication)</td>
<td>Hispanic (discovery), European (replication)</td>
<td>PHF21B</td>
<td>9 pathways including innate immune response, glutamate signalling, sensory perception</td>
<td>(Wong et al., 2017)</td>
</tr>
<tr>
<td>Treatment resistance</td>
<td>GWAS, WES, exome chip</td>
<td>1209 cases (discovery) 1128 cases (replication)</td>
<td>European</td>
<td>response predictor in resistance vs. response to treatment with serotonergic antidepressants using rare variants</td>
<td>cell survival and proliferation, neurodegeneration, immune response</td>
<td>(Fabbri et al., 2020)</td>
</tr>
</tbody>
</table>

MDD=major depressive disorder; WES=whole-exome sequencing; WGS=whole-genome sequencing; GWAS=genome-wide association study; TFs=transcription factors; UTR=untranslated region; TGF-β=Transforming growth factor beta; MRI=magnetic resonance imaging
Table 3. Published genome-wide association studies on depression.

<table>
<thead>
<tr>
<th>Year</th>
<th>Discovery: Nca/Nco or Ntot</th>
<th>Discovery: Loci</th>
<th>Replication: Cases/Controls</th>
<th>Replicated loci</th>
<th>Definition of depression</th>
<th>Ancestry</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>2008</td>
<td>1738/1802</td>
<td>0</td>
<td>6079/5893</td>
<td>0*1</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Sullivan et al., 2009)‡</td>
</tr>
<tr>
<td>2008</td>
<td>1359/1782</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Muglia et al., 2010)‡</td>
</tr>
<tr>
<td>2009</td>
<td>3957/3428</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Shyn et al., 2011)‡</td>
</tr>
<tr>
<td>2010</td>
<td>1636/1594</td>
<td>0</td>
<td>1418/1918</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Lewis et al., 2010)‡</td>
</tr>
<tr>
<td>2010</td>
<td>604/1364</td>
<td>0</td>
<td>409/541</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Rietschel et al., 2010)‡</td>
</tr>
<tr>
<td>2010</td>
<td>1020/1636</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Shi et al., 2011)‡</td>
</tr>
<tr>
<td>2010</td>
<td>2431/3673</td>
<td>0</td>
<td>3332/3228</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Wray et al., 2012)‡</td>
</tr>
<tr>
<td>2011</td>
<td>353/366</td>
<td>0</td>
<td>3735/10635</td>
<td>1*2</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Kohli et al., 2011)‡</td>
</tr>
<tr>
<td>2012</td>
<td>9240/9519</td>
<td>0</td>
<td>6783/50695</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Ripke et al., 2013)‡</td>
</tr>
<tr>
<td>2013</td>
<td>34549</td>
<td>0</td>
<td>16709</td>
<td>1*3</td>
<td>Depressive symptoms</td>
<td>European</td>
<td>(Hek et al., 2013)</td>
</tr>
<tr>
<td>2015</td>
<td>5303/5337</td>
<td>2</td>
<td>3231/3186</td>
<td>2</td>
<td>Structured interview: Women with recurrent MDD</td>
<td>Han Chinese</td>
<td>(CONVERGE consortium, 2015)</td>
</tr>
<tr>
<td>2015</td>
<td>16948</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Depressive symptoms</td>
<td>Trans-ancestry</td>
<td>(Ware et al., 2015)</td>
</tr>
<tr>
<td>Year</td>
<td>Sample Size</td>
<td>n</td>
<td>Age (Mean)</td>
<td>Year of Diagnosis</td>
<td>Method of Diagnosis</td>
<td>Population</td>
<td>Source</td>
</tr>
<tr>
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</tr>
<tr>
<td>2015</td>
<td>1443</td>
<td>1*4</td>
<td>NA</td>
<td>NA</td>
<td>Depressive symptoms</td>
<td>Hispanic American</td>
<td>(Ware et al., 2015)</td>
</tr>
<tr>
<td>2016</td>
<td>32528</td>
<td>1</td>
<td>6813</td>
<td>0</td>
<td>Depressive symptoms</td>
<td>European</td>
<td>(Demirkan et al., 2016)</td>
</tr>
<tr>
<td>2016</td>
<td>7179</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Depressive symptoms</td>
<td>African American</td>
<td>(Dunn et al., 2016)</td>
</tr>
<tr>
<td>2016</td>
<td>3138</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Depressive symptoms</td>
<td>Hispanic/ Latino</td>
<td>(Dunn et al., 2016)</td>
</tr>
<tr>
<td>2016</td>
<td>161460</td>
<td>2</td>
<td>368890</td>
<td>2</td>
<td>Depressive symptoms</td>
<td>European</td>
<td>(Okbay et al., 2016)</td>
</tr>
<tr>
<td>2016</td>
<td>3869/9519</td>
<td>1</td>
<td>6107/124230</td>
<td>1</td>
<td>Structured interview: Adult-onset MDD</td>
<td>European</td>
<td>(Power et al., 2017)</td>
</tr>
<tr>
<td>2016</td>
<td>75607/231747</td>
<td>7 and 4*5</td>
<td>45773/106354</td>
<td>15*6</td>
<td>Self-reported diagnosis</td>
<td>European</td>
<td>(Hyde et al., 2016)</td>
</tr>
<tr>
<td>2016</td>
<td>70017</td>
<td>1</td>
<td>38328</td>
<td>1</td>
<td>Mixed (including depressive symptoms)</td>
<td>European</td>
<td>(Direk et al., 2017)</td>
</tr>
<tr>
<td>2017</td>
<td>14729/14435</td>
<td>1</td>
<td>4504/7007</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European &amp; Han Chinese</td>
<td>(Bigdeli et al., 2017)</td>
</tr>
<tr>
<td>2017</td>
<td>5347/1871</td>
<td>1</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD with decreased vs increased appetite/ weight</td>
<td>European</td>
<td>(Milaneschi et al., 2017)</td>
</tr>
<tr>
<td>2018</td>
<td>12310</td>
<td>0</td>
<td>7948</td>
<td>0</td>
<td>Depressive symptoms</td>
<td>Hispanic/ Latino</td>
<td>(Dunn et al., 2018)</td>
</tr>
<tr>
<td>Year</td>
<td>Study ID</td>
<td>N</td>
<td>Cases</td>
<td>Follow-up</td>
<td>Condition</td>
<td>Ethnicity</td>
<td>Reference</td>
</tr>
<tr>
<td>------</td>
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<td>-----------</td>
<td>-----------</td>
<td>----------</td>
<td>-----------</td>
</tr>
<tr>
<td>2018</td>
<td>90150/246603</td>
<td>10</td>
<td>2659/17237</td>
<td>6*7</td>
<td>Mixed</td>
<td>European &amp; Han Chinese</td>
<td>(Li et al., 2018)</td>
</tr>
<tr>
<td>2018</td>
<td>10851/32211</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Mixed</td>
<td>European</td>
<td>(Hall et al., 2018)‡</td>
</tr>
<tr>
<td>2018</td>
<td>3852/16034</td>
<td>1</td>
<td>NA</td>
<td>NA</td>
<td>Mixed: Men only</td>
<td>European</td>
<td>(Hall et al., 2018)</td>
</tr>
<tr>
<td>2018</td>
<td>3139/3832</td>
<td>3</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: Women without exposure to adversity</td>
<td>Han Chinese</td>
<td>(Peterson et al., 2018)</td>
</tr>
<tr>
<td>2018</td>
<td>113769/208811</td>
<td>14</td>
<td>75607/231747</td>
<td>14*3</td>
<td>Broad depression</td>
<td>European</td>
<td>(Howard et al., 2018)‡</td>
</tr>
<tr>
<td>2018</td>
<td>30603/143916</td>
<td>2</td>
<td>75607/231747</td>
<td>2*3</td>
<td>Probable MDD</td>
<td>European</td>
<td>(Howard et al., 2018)‡</td>
</tr>
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<td>2019</td>
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<td>3</td>
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</tr>
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<td>(Levey et al., 2020)</td>
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* Year indicates the year in which the study was first published (including online ahead of print and, were no published version as available, preprint), which may deviate from the year the article was published in print. Nca/Nco for number of cases and controls in case-control design; Ntot as total sample size for quantitative trait.
* ‡ Studies used for Figure 1.
* 1 The top locus replicated in subsequent studies (Hek et al. 2010, Wray et al. 2018, Howard et al. 2019).
* 2 Significant in the meta-analysis of discovery and replication cohorts in a recessive model. This association did not replicate in subsequent GWAS in additive models. Replication has, to our knowledge, not been attempted using a recessive model.
* 3 Significant in the meta-analysis of discovery and replication cohorts.
* 4 This association did not replicate in the follow-up Hispanic ancestry GWAS meta-analysis Dunn et al. 2018.
* 4 Meta-analysis of 23andMe and PGC MDD cohorts.
* 5 Replication of top results from the meta-analysis of 23andMe and PGC MDD cohorts.
* 6 Only SNPs not reaching genome-wide significance in the discovery-stage were analysed in the replication phase. Six loci (of which two were not significant in the discovery GWAS) covered by these selected SNPs reached significance in the meta-analysis of discovery and replication cohorts.
* 7 From a meta-analysis of overlapping subjects (PGC and UKBB), case and control numbers may thus be inflated.
The study identified 102 independent variants at 101 loci.

Figure 1: Identified GWAS loci in relation to the number of cases.
The relationship of identified loci to the number of cases based on published studies on schizophrenia, bipolar disorder (BD), and major depression (MD). To simplify the presentation, studies with different depression definitions have been combined into the MD category. The plot shows the number of cases and independent significant loci in the discovery-stage analyses (without replication). Schizophrenia GWAS saw a critical inflection point at approximately 15,000 cases and are currently discovering a new locus for, on average, every 230 new cases (Levinson et al., 2014). BD currently discovers a new locus for every 400 new cases. Such an inflection point was observed for MD GWAS only at approximately 75,000 cases, as predicted (Wray et al., 2012; Levinson et al., 2014), and MD GWAS using liberal phenotype definitions now identify a new locus every 1500 cases. Slopes starting at the inflection point were estimated using linear regression. The MD studies used for the plot are provided in Table 3.
References


The Genetic Basis of Major Depression

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Current word count: 55795138
Abstract:

Major depression depressive disorder (MDD) is a common, debilitating, phenotypically heterogeneous disorder with a heritability of ranges from 30% to 50%. Compared to other psychiatric disorders, its high prevalence, relatively low/moderate heritability, and strong polygenicity have posed major challenges for gene-mapping in MMD. Studies of common genetic variation in MMD, driven by large international collaborations such as the Psychiatric Genomics Consortium, have confirmed the highly polygenic nature of the disorder and implicated over 100 genetic risk loci to date. Rare copy number variants associated with MDD risk were also recently identified. The goal of this review is to present a broad picture of our current understanding of the epidemiology, genetic epidemiology, molecular genetics, and gene-environment interplay in MMD. Insights into the impact of genetic factors on the aetiology of this complex disorder hold great promise for improving clinical care.
Introduction

**Depression**-Major depressive disorder (MDD) is a common mood disorder that substantially and negatively impacts both quality and length of life (Kessler and Bromet, 2013), and represents the leading cause of disability worldwide (GBD 2017 DALYs and HALE Collaborators, 2018). This review summarises our existing knowledge about the genetic basis of depression. Specifically, it details how research in this area has informed our understanding of the biology underlying depression and how this knowledge may be used to improve patient care in the future.

Definition

The term ‘depression’ most often refers to the clinical diagnosis of MDD, as defined by the **International Classification of Diseases (ICD)** and the **Diagnostic and Statistical Manual of Mental Disorders (DSM)** diagnostic criteria. Depression is characterised by the core symptoms of i) depressed mood (e.g., feelings of sadness, irritability, or emptiness), and ii) anhedonia (e.g., decreased ability to feel pleasure or to experience interest in usual activities). Individuals with depression also experience a wide range of other symptoms which can be grouped into cognitive (e.g., concentration difficulties and indecisiveness), emotional (e.g. feelings of worthlessness and hopelessness), and somatic domainsvegetative symptoms (e.g., significant appetite or weight changes, insomnia or hypersomnia) (Malhi and Mann, 2018).

**Depression-MDD** is a heterogeneous disorder, which has important implications for clinical practice and research (Flint and Kendler, 2014; Cai *et al.*, 2020a). Even within one classification system, multiple combinations of symptoms can lead to a diagnosis. A study of depression symptoms in participants of the STAR*D trial found over a thousand unique symptom profiles with the most common only being reported by 1.8% of the sample (Fried and Nesse, 2015).
Furthermore, the term depression is used inconsistently, and its clinical diagnosis of MDD may be reached in many different ways using established criteria (Fried et al., 2020). This phenotypic heterogeneity in clinical practice is mirrored in the broad range of case definition strategies for genetic research. Genetic studies have adopted multiple methods of classifying depression - from self-reported clinical diagnosis or electronic health record (EHR) diagnosis codes to structured diagnostic interviews (Smith et al., 2013; Howard et al., 2019). Despite the shortcomings of dichotomizing a heterogeneous disorder like MDD, a categorical definition of depression has been widely used in research. Apart from such a diagnosis-based definition, the severity of depressive symptoms is often studied as a quantitative trait, so the full variation of depressive symptoms can be taken into account. These phenotypic definitions can have a considerable impact on genetic findings (Cai et al., 2020b) and their potential interpretation and translation to the clinic. In line with previous studies (Wray et al., 2018) The Psychiatric Genomics Consortium (PGC) has adopted the term Major Depression (MD) to reflect the degree of uncertainty present when analysing data from a combination of studies with varying classification methods. Throughout this review, we will use ‘MDD’ where the study subjects met the clinical criteria for major depressive disorder, and ‘major depression’ where the subjects were ascertained using alternative strategies, but nevertheless likely to have met the diagnostic criteria for MDD.

Epidemiology

The 12-month prevalence of MD-MDD is estimated to be ~6% and the lifetime prevalence ~20%, although these figures vary considerably across countries (e.g., the 12-month prevalence ranged from 2.2% in Japan to 10.4% in Brazil) (Bromet et al., 2011). Established epidemiological risk factors for MD-MDD include age, female sex, separated/divorced marital status, physical
health problems, and childhood trauma (Moskvina et al., 2007; Bromet et al., 2011; Van Assche et al., 2017). The peak age of onset of MD-MDD extends from mid to late adolescence to the early 40s, with the median in the mid-20s (Kessler and Bromet, 2013). Women are twice as likely as men to be diagnosed with MD-MDD (Bromet et al., 2011). Sex-specific differences in clinical presentation (Rice et al., 2015), environmental risk factors, and differences in the brain’s exposure to sex hormones have been suggested as possible explanations. MD-MDD is associated with chronic physical health problems including cardiovascular diseases, chronic pain disorders, and cancers (Kessler and Bromet, 2013). The nature of the association between MD-MDD and physical ill health is complex. There is evidence for a bidirectional causal association, as well as common antecedents such as social deprivation increasing the risk of both MD-MDD and physical ill health (Kessler and Bromet, 2013).

Genetic epidemiology

Early evidence from family-based studies clearly demonstrated a familial aggregation of MD-MDD, with an increased risk in first-degree relatives of MD-MDD patients (odds ratio, OR=2.84, 95% CI =2.31-3.49) (Sullivan et al., 2000). However, families share both genetic and environmental factors and quantifying their relative contributions to MD-MDD liability has been central to genetic epidemiological studies.

Classic quantitative genetic modelling partitions phenotypic variance into genetic and environmental components. The pedigree heritability is then estimated as the proportion of phenotypic variance attributable to additive genetic effects (“narrow-sense heritability”) (Lynch and Walsh, 1998; Visscher et al., 2008). Such modelling is typically performed in twin or adoption studies. Twin studies, comparing monozygotic (MZ) and dizygotic (DZ) twins, have
consistently demonstrated that MD-MDD is moderately heritable (Table 1) and an early meta-
analysis estimated the heritability at 37% (95% CI = 31-42%) (Sullivan et al., 2000). Subsequent
twin studies showed similar estimates, ranging between 29-49%, with higher estimates among
treated or recurrent MD-MDD cases – potentially due to greater clinical severity in the study
samples (Kendler et al., 2006; Polderman et al., 2015; Kendler et al., 2018a). These studies also
suggested that the liability variance attributable to the shared family environment is low,
ranging from 0 to 11% (Sullivan et al., 2000; Kendler et al., 2006; Polderman et al., 2015;
Kendler et al., 2018a). Adoption studies, which comparing the transmission patterns among
adoptive and intact families, are better able to separate shared genetics from other aspects of a
shared family environment than twin studies (Kendler et al., 2018b). Early adoption studies in
MD-MDD, however, were based on small samples and produced conflicting results (Sullivan et
al., 2000). Contrary to findings from twin studies, a recent population-based adoption study
reported a much lower heritability (approximately 16%) and a higher contribution of the family
environment (Kendler et al., 2018b). In addition, the overestimation of the MD-MDD heritability
in twin studies has been raised as a potential reason for the “missing heritability” – i.e. the
discrepancy between the pedigree heritability and the much lower phenotypic variance
explained by genomic loci identified in genome-wide association studies (GWAS) (Manolio et
al., 2009). Recent studies utilised national registers and EHR to reconstruct extended familial
relationships, and therefore provide estimates of pedigree heritability beyond classic twin
design (Wray and Gottesman, 2012; Kendler et al., 2018a; Polubriaginof et al., 2018). These
estimates, however, were very similar to those from twin studies (Table 1), suggesting that the
heritability of MD-MDD has not been overestimated.
One of the most intensively studied questions regarding the heritability of MD-MDD is whether sex differences in the genetic architecture of the disorder exist. A higher heritability in women than in men might contribute to the higher rates of MD-MDD observed in women. Early meta-analyses failed to detect either quantitative or qualitative evidence for sex differences in MD-MDD heritability (Sullivan et al., 2000). However, more recent studies demonstrated a significantly higher MD-MDD heritability in women than in men (Table 1), although the genetic correlation between the sexes was high (0.63-0.89) (Kendler et al., 2006, 2018a). These results suggest that most, but not all, genetic risk is shared between the sexes (Kendler et al., 2018a), consistent with the highly polygenic nature of MD-MDD.

Research efforts have attempted to identify genetically more homogeneous MD-MDD subtypes (Flint and Kendler, 2014). Parents with early-onset and recurrent MD-MDD confer the highest risk for MD-MDD in offspring (Birmaher et al., 1996; Sullivan et al., 2000). Other clinical features, including greater clinical severity, comorbid anxiety disorder, and the use of antidepressant and electroconvulsive therapy, also confer higher MD-MDD risk for relatives (Kendler et al., 2018a). Similarly, the Psychiatric Genomics Consortium (PGC) MD-MDD GWAS have demonstrated that individuals from many of these subtypes carry higher polygenic risk scores (PRS) for MD-MDD (Power et al., 2017; Wray et al., 2018). Studies examining the heritability of these subtypes are, however, sparse. The heritability among recurrent MD-MDD cases (41%) may be significantly higher than in those who had single-episode MD (28%) (Fernandez-Pujals et al., 2015), and postpartum depression has been shown to have a higher heritability (44-54%) than non-postpartum depression (32%) (Viktorin et al., 2016). Ascertaining these more heritable forms of MD-MDD has led to some of early GWAS findings (CONVERGE consortium, 2015).
Candidate gene studies

Candidate gene studies of MDMDD, first performed more than 40 years ago (Beckman et al., 1978), focused on genes thought to be involved in the pathophysiology of the disorder and in drug mechanisms – as reviewed in (Shadrina et al., 2018). More than 1500 publications have assessed variants in over 200 candidate genes with often conflicting results (Flint and Kendler, 2014). Most of these studies were vastly underpowered, lacked correction for population stratification, and the reported significance levels did not exceed what would be expected by chance (Flint and Kendler, 2014). Based on the observed odds ratios (OR) and minor allele frequencies (MAF), all published candidate gene variants should be detectable in the current GWAS at genome-wide significance (Flint and Kendler, 2014). Yet, this has not been the case (Bosker et al., 2011; Wray et al., 2012, 2018; Howard et al., 2019). In large-scale data from the UK Biobank (UKBB) and PGC (n=62,138-443,264 per subsample), systematic analyses of the most commonly studied polymorphisms mapping to 18 candidate genes found no evidence for significant associations with MDMmajor depression, except at the gene-level for DRD2 (dopamine receptor D2) (Border et al., 2019a).

Candidate gene studies were the mainstay of genetic research in MDM-MDD until the emergence of GWAS and some are still among the most highly cited works in the field (McIntosh et al., 2019). However, a lack of robust association has led to the strong recommendation that candidate gene studies should no longer be performed (Duncan and Keller, 2011; Farrell et al., 2015; Border et al., 2019a, 2019b).
The role of rare genetic variants

Historical linkage studies in families with MDD failed to identify causal variants but implicated regions on chromosomes 2, 3, 11, and 15 in disorder risk (Zubenko et al., 2003; Holmans et al., 2004, 2007; Camp et al., 2005, p. 2005; Levinson et al., 2007; Middeldorp et al., 2008; Schol-Gelok et al., 2010; Breen et al., 2011). A highly polygenic architecture, as is the case for MDD, can generate significant linkage study results which could be erroneously interpreted as large-effect loci (Hemani et al., 2013). Indeed, no significant overlap was observed between the reported linkage regions and association loci from current well-powered GWAS (McIntosh et al., 2019). A major contribution of single highly penetrant variants within these regions to the genetic architecture of MDD is therefore unlikely.

In the past decade, association testing of rare and low-frequency variants has progressed to whole-exome (WES) and whole-genome sequencing (WGS) studies. Rare (MAF <0.5%) or low-frequency (MAF range >0.5% and <5%) variants are expected to contribute to the genetic basis of MDD (Lee et al., 2014; Yu et al., 2018). However, the largest WGS study of MDD to date, by the CONVERGE consortium (5,303 cases and 5,337 controls), failed to find evidence for the association of individual rare or low-frequency variants with MDD (CONVERGE consortium, 2015). Using the same data set, it was subsequently shown that individuals with MDD had a significantly higher overall burden of singleton exonic variants predicted to be deleterious (Peterson et al., 2017). This suggests that larger sample sizes are required to identify associated rare variants with moderate effect sizes (Lee et al., 2014; Border et al., 2019b). An approach to mitigate against the need for vast sample sizes is to examine MDD-related subphenotypes (summarised in Table 2). The majority of these studies identified single rare variants or gene-level burden scores as associated with at least one of the assessed...
subphenotypes. In most cases, these results await replication in independent and well-powered datasets.

Rare copy number variants (CNVs), deletions or duplications of >1kb of DNA, play a well-established role in risk of neurodevelopmental and psychiatric conditions such as autism spectrum disorder, schizophrenia, and intellectual disability (International Schizophrenia Consortium, 2008; Stefansson et al., 2008; Cooper et al., 2011; Coe et al., 2014), but have received less attention in MDMDD. Until recently, studies of CNVs in depression have been underpowered and yielded inconsistent and sometimes conflicting results (Glessner et al., 2010; Degenhardt et al., 2012; Perlis et al., 2012; Rucker et al., 2013, 2016; O’Dushlaine et al., 2014; Tansey et al., 2014; Yu et al., 2017).

Two recent large-scale efforts have assessed the contribution of rare CNVs to the genetic architecture of MDMDD. In a meta-analysis of four cohorts (5,780 cases, 6,626 controls), short deletions <100 kb were significantly enriched in individuals with MDMDD (Zhang et al., 2019). The enriched deletions mapped primarily to intergenic regions and, within those, to enhancers, suggesting that rare CNVs may increase risk of MDMDD by altering gene expression. A study of 407,074 individuals from the UKBB (23,979 cases) implicated 53 rare CNVs associated with neurodevelopmental conditions in risk of MD-major depression (OR=1.3–1.5) (Kendall et al., 2019). Three CNV loci—duplications at 1q21.1, the Prader-Willi Syndrome locus at 15q11-13, and at 16p11.2—were individually associated with self-reported depression. Outside of this group, there was no evidence for a residual burden of rare CNVs >100 kb in MDMDD depression. However, smaller CNVs were not tested in the study by Kendall and colleagues.
Rare CNVs therefore likely contribute to **MD-MDD** risk, but the exact nature of the association remains unclear.

### Genome-wide association studies

Many MDD GWAS have been conducted during the past decade, predominantly on subjects of European ancestry (*Table 3*). Therefore, European-ancestry studies are described first, followed by a review of GWAS in more diverse populations.

Initial **MD-MDD** GWAS of common variants (i.e., MAF ≥1%) were limited by small sample sizes and failed to identify genome-wide significant loci (*p*-value threshold <5×10⁻⁸) (*Table 3*). However, the first suggestive GWAS result implicated the gene encoding the presynaptic active zone component Piccolo (*PCLO*) (Sullivan *et al.*, 2009) and this has been confirmed as a risk gene in subsequent studies (Hek *et al.*, 2010; Wray *et al.*, 2018; Howard *et al.*, 2019). In 2011, the first genome-wide significant locus was reported - **SLC6A15** (a neuron-specific neutral amino acid transporter), although this was in a recessive model (Kohli *et al.*, 2011). This association did not reach genome-wide significance with additive models in larger follow-up GWAS (Wray *et al.*, 2018; Howard *et al.*, 2019). Recognising the importance of combining data across studies to increase statistical power, the PGC published its first **MD-MDD** GWAS meta-analysis in 2013, although no significant locus was identified (*Ripke et al.*, 2013).

Since then, large-scale MD-GWAS of major depression have been conducted with data from the direct-to-consumer company 23andMe and large-scale biobank efforts such as UKBB (*Table 3*). In 2016, a GWAS by 23andMe using self-reported depression diagnosis identified 15 genome-wide significant loci (Hyde *et al.*, 2016), confirming the estimate that >75,000 cases would be
required for genome-wide significant results (Levinson et al., 2014). A second large GWAS meta-analysis from the PGC described 44 MD-associated genetic loci, pooling cohorts with different phenotype definitions (Wray et al., 2018). The three datasets used in the latter studies were eventually combined in an analysis yielding 102 genome-wide significant variants, 87 of which replicated (Howard et al., 2019). The latest GWAS to date meta-analysed these datasets with data from the Million Veteran Program (MVP), yielding 178 associated loci (Levey et al., 2020).

Collectively, these GWAS confirmed that major depression is highly polygenic, with ORs of individual variants under 1.05 (Wray et al., 2018; Howard et al., 2019), as suggested previously (Nishino et al., 2018). The two loci with the overall most robust support for an association map to the genes NEGR1 and OLFM4, both with an OR of 1.04 and a MAF of ~40% (Wray et al., 2018; Howard et al., 2019). The neuronal growth regulator 1 (NEGR1) cell adhesion molecule modulates neurite outgrowth, synapse formation, and synaptic function (Wray et al., 2018; Noh et al., 2019). The secreted glycoprotein Olfactomedin 4 (OLFM4) has an established function in inflammation, innate immunity, and cancer (Liu and Rodgers, 2016; Wray et al., 2018). Several other MD-associated loci have functions in neurite outgrowth, synaptic function and plasticity, as well as in immunity and inflammation (Wray et al., 2018). To establish the functions of the proposed risk SNPs (single nucleotide polymorphisms) and their links to specific genes, analyses of cell-type-specific gene expression and epigenetic data are necessary (Zhong et al., 2019; Arloth et al., 2020; Chan et al., 2020; Li et al., 2020; Wang et al., 2020).

Given the small effect sizes of single SNPs, gene-level and pathway analyses that integrate the effects of many variants across genes and pathways may be more suitable for studying the
aetiology of MDD. Gene-level association analyses have implicated genes encoding pre- and post-synaptic proteins, especially receptor units, e.g. voltage-gated calcium channels (VGCC), the dopamine receptor DRD2, and glutamate receptors (Wray et al., 2018). Identified by both SNP- and gene-level analyses, the splicing regulator RBFOX1 may play an important role in the aetiology of major depression (Wray et al., 2018; Noh et al., 2019). Gene-set analyses implicated pathways with roles in neuronal differentiation and projection and, most prominently, synaptic formation and function, including gene sets linked to VGCC and synaptic RNA-binding proteins like FMRP and CELF4 (Wray et al., 2018; Noh et al., 2019; Levey et al., 2020).

Despite a large number of identified genomic loci, the variance explained by major depression PRS is still very low (Nagelkerke’s pseudo-$R^2$ 1.5-3.2% on the liability scale) (Howard et al., 2019). This prediction accuracy was lower than achieved for, e.g. schizophrenia, where Nagelkerke’s pseudo-$R^2$ is 7.7% on the liability scale (Schizophrenia Working Group of the PGC et al., 2020). At much smaller sample sizes, the GWAS on schizophrenia also identified many more associated variants (Figure 1). The identification of risk variants has been more challenging than for schizophrenia because of its higher prevalence, lower heritability, and more heterogeneous samples (Wray et al., 2012; Levinson et al., 2014; Ormel et al., 2019). Schizophrenia GWAS saw a critical inflection point at approximately 15,000 cases and are currently discovering a new locus for, on average, every 230 new cases (Levinson et al., 2014). Such an inflection point was observed for major depression GWAS only at approximately 75,000 cases, as predicted (Wray et al., 2012; Levinson et al., 2014). MDD GWAS using liberal phenotype definitions of major depression now identify a new locus for every 1500 cases (Figure 1).
Importantly, phenotype definition matters. Given the strong genetic correlations across cohorts, recent GWAS meta-analyses combined cohorts with different ascertainment strategies to maximize the overall sample size. However, a comparison of genetic analyses using different phenotype definitions in UKBB revealed that GWAS of broad phenotype definitions might identify associated loci that are not specific to MDD, but more likely shared with other psychiatric disorders (Cai et al., 2020b). In order to identify robustly associated MDD variants, a balanced strategy of maximising sample size while focusing on more stringent phenotype definitions has been suggested (Davies et al., 2019; Schwabe et al., 2019; Byrne et al., 2020). Quantitative definitions of MDD have also been analysed in GWAS, and a continuum between depressive symptoms and MDD was suggested (Direk et al., 2017). Several variants were reported to be associated with depressive symptoms (Table 3), with limited replication success. Of these symptom-associated variants, rs62100776 (Okbay et al., 2016) maps to the gene DCC, coding for a neuronal cell adhesion molecule, which was a genome-wide significant locus in the 2018 PGC case-control GWAS (Wray et al., 2018). Further noteworthy GWAS were conducted on MDD stratified by the age at onset (Power et al., 2017), sex (Hall et al., 2018), or exposure to adversity (Peterson et al., 2018), or combined GWAS of different depression MDD phenotypes (Amare et al., 2020).

The studies discussed so far focused on patients of European ancestry. However, complex disorders show population-specific landscapes of genetic risk (Martin et al., 2017). Accordingly, GWAS in non-European populations have also been conducted (Table 3). The CONVERGE study on female Han Chinese recurrent in-patient MDD cases identified two significant loci near the SIRT1 and in the LHPP gene, and replicated them in the same population (CONVERGE consortium, 2015). Neither variant was significantly associated in GWAS on European samples.
(Wray et al., 2018; Howard et al., 2019), possibly due to large differences in allele frequency across ancestries – both variants exhibit markedly lower allele frequencies in Europeans. Follow-up studies estimated the genetic correlation between East Asian and European MDD cohorts at 0.33-0.41, while the average genetic correlations between European cohorts was 0.76 (Bigdeli et al., 2017; Wray et al., 2018), and identified trans-ancestral risk variants (Li et al., 2018). In a study of Hispanic/Latino individuals, two out of 25 European MDD risk GWAS loci were replicated (Dunn et al., 2018). The MVP study identified no genome-wide significant loci in over 20,000 African-ancestry MDD patients with major depression (Table 3), and only one of 206 analysed European risk SNPs formally replicated in the African sample (Levey et al., 2020). However, a trans-ancestry meta-analysis conducted within the same study increased the number of risk loci from 178 in Europeans alone to 183. These findings highlight the need for greater geographical, ethnic, and economic diversity in MDD-depression genetic studies (McIntosh et al., 2019).

**Genetic overlap between MDD and other traits**

Several lines of evidence, including cross-disorder analyses, support the notion that MDD shares genetic underpinnings with many other psychiatric phenotypes. A genetic overlap has also been demonstrated between MDD and non-psychiatric phenotypes. Common techniques for studying cross-disorder genetic overlap include genetic correlation \( r_g \) estimation and PRS analyses (Grotzinger et al., 2019).

Genetic correlation quantifies the genetic relationship between two traits, reflecting their underlying modes of pleiotropy – i.e. the same genetic variant influencing both traits (van Rheenen et al., 2019; Baselmans et al., 2020). Mathematically, it is derived as the genetic
covariance between the two traits scaled by the square roots of their genetic variances (van Rheenen \textit{et al.}, 2019). Two methods are commonly used to estimate $r_g$ – i) bivariate restricted maximum likelihood (REML) which uses individual-level genotype data as the input (Lee \textit{et al.}, 2012), and ii) linkage disequilibrium score regression (LDSC) (Bulik-Sullivan \textit{et al.}, 2015) and its recent extension, high-definition likelihood (HDL) method (Ning \textit{et al.}, 2020), which uses GWAS summary statistics. $r_g$ ranges between $-1$ and 1, with 1 meaning that shared liability is caused by the same risk SNPs and $-1$ meaning that identical SNPs increase the risk for one trait while decreasing risk for the other. It is important to note that $r_g$ estimates cannot infer causality.

Many studies have examined $r_g$ between \textit{MD-MDD} and both psychiatric and non-psychiatric disorders or traits. The strongest $r_g$ detected for \textit{MD-MDD} are with depressive symptoms measured as a continuous trait (0.63-1.0) (Direk \textit{et al.}, 2017; Turley \textit{et al.}, 2018), confirming the concept of a continuum between subthreshold depressive symptoms and clinical diagnoses of \textit{MDMDD}. Similarly, strong $r_g$ have been found for personality traits such as neuroticism ($r_g = 0.70-0.75$) and subjective well-being (-0.75) (Okbay \textit{et al.}, 2016; Howard \textit{et al.}, 2019). \textit{MD MDD}major depression was moderately correlated with several other psychiatric disorders, e.g. autism spectrum disorder (0.45), attention-deficit/hyperactivity disorder (0.44), bipolar disorder (BD) (0.36), and schizophrenia (0.34) (Cross-Disorder Group of the PGC, 2019). Further studies reported a weaker $r_g$ of \textit{MD-MDD major depression} with type-I (0.26-0.31) than with type-II BD (0.61-0.69) (Stahl \textit{et al.}, 2019; Coleman \textit{et al.}, 2020). Moreover, a positive $r_g$ was reported between the body-mass index (BMI) and atypical \textit{MD depression} with increased appetite/weight, which was not present for \textit{MD those} without changes in appetite (Milaneschi \textit{et al.}, 2017). An investigation of genetic correlations between neurological and psychiatric disorders found little evidence of shared genetic underpinnings, except for a significant but
small positive $r_g$ between MD-MDD and migraine (The Brainstorm Consortium et al., 2018). Negative genetic correlations have been detected for MD-major depression with educational attainment and cognitive performance (each $\sim$-0.15) (Howard et al., 2019).

Analyses of PRS is another method used to assess the polygenic overlap between MD-MDD and other traits. PRS use SNP effect sizes from a training GWAS to compute the cumulative genetic risk burden of each individual in a target sample. PRS are calculated as the weighted sum of the allele count multiplied by its effect size across all selected SNPs (Andlauer and Nöthen, 2020; Lewis and Vassos, 2020). Therefore, they capture the combined effects of many SNPs and are used to examine the polygenic liability for a disorder or trait or to examine the genetic overlap across traits (Wray et al., 2014). Although the generation of PRS requires large training GWAS, it can be applied to much smaller target sets. For examples, a MD-MDD PRS was predictive of population-based obsessive-compulsive symptoms (Zilhão et al., 2018). Clinically, PRS may be used for diagnosis, prognosis, and treatment prediction (Schijven et al., 2020). However, PRS for schizophrenia have, so far, shown greater potential for such applications, given their higher variance explained (Schizophrenia Working Group of the PGC et al., 2020). Importantly, PRS were used to demonstrate that MD-MDD subtypes may diverge in their genetic overlaps with other disorders. For example, early-onset MD-MDD cases show a stronger association with PRS for schizophrenia and BD than late-onset MD-MDD patients do (Power et al., 2017). In the future, PRS might also be used to identify aetiologically more homogeneous MD-MDD subtypes. For example, subgrouping on the basis of treatments such as electroconvulsive therapy might be informative (Foo et al., 2019) and provide further insights into the underlying genetic differences of the severity of the disorder. Finally, MD-MDD PRS may also be used to study PRS-
environment (PRS×E) interactions, to support treatment choice, and to refine the penetrance of high-risk variants (Lewis and Vassos, 2020).

Gene environment interaction

In addition to the heritable component of MDMDD, environmental factors such as stressful life events (e.g. childhood trauma) play a major role in the aetiology of MDMDD, both independently and in combination with genetic factors. One active area of research has been the study of gene-environment interactions (G×E). Early G×E efforts examined specific candidate genes with putative biological significance for MDMDD, e.g. the 5-HTTLPR polymorphism of the serotonin transporter gene, and found evidence of interaction with stressful life events (Caspi et al., 2003). However, these initial reports showed inconsistent replicability (Duncan and Keller, 2011), and recent meta-analytic evidence indicated no robust evidence of candidate G×E effects on MDMDD (Van der Auwera et al., 2018; Border et al., 2019a). Studies using genome-wide data have reported varied G×E results. First, stratified GWAS results from the CONVERGE study have suggested that MDMDD may have a stronger genetic basis when it occurs in the absence of environmental adversity, with genome-wide significant hits for MDMDD identified only among women not exposed to adversity (Peterson et al., 2018). Data from the UKBB, however, have suggested the opposite, with a higher SNP-based heritability for major depression-MD observed among individuals reporting lifetime trauma exposure (Coleman et al., 2020). Second, PRS have been used to aggregate genome-wide effects for MDMDD and examine their interactions with stressful life events. Again, this revealed discrepant findings where MDMDD PRS were more strongly associated with MDMDD among individuals both exposed (Peyrot et al., 2014) and unexposed (Mullins et al., 2016) to childhood trauma. A later but still modestly powered meta-analysis (n=5,765) identified limited
evidence for the interaction of PRS and childhood trauma on MD-MDD (Peyrot et al., 2018), which was echoed in epidemiological and clinical cohorts (n=184-1,450) (Halldorsdottir et al., 2019). However, large-scale studies like the UKBB have provided the opportunity to re-examine PRSxE effects with greater power. One study showed stronger PRS effects on MD-major depression among individuals exposed to childhood trauma and socioeconomic adversity (Shen et al., 2020), though not other types of exposures such as adulthood stressful life events. These findings support the notion that genetic influences could vary by specific environments. Third, genome-wide GxE analyses have identified few variants with genome-wide significant GxE effects (Dunn et al., 2016; Arnau-Soler et al., 2019), although these studies have been largely underpowered and results have not been further replicated. Together, current evidence suggests that genome-wide influences on MD-MDD may vary by stressful life exposures but that the detection of GxE may be sensitive to the research design (e.g. how depression and adversity are defined).

The role of stressful life events, specifically childhood trauma, was the predominant focus of GxE in MD-MDD, but other environmental exposures have also received attention. For example, studies examining social support in combination with MD-MDD PRS in vulnerable populations have consistently shown that social support appears to reduce risk of depression. This effect remained even after accounting for genetic vulnerability for depression, but did not vary by or modify the genetic effects (Choi et al., 2020a; Stringa et al., 2020). Similar observations of independent effects have been made for physical activity and MD-major depression, suggesting that even among individuals at high genetic vulnerability for MD-depression, physical activity levels may still be protective (Choi et al., 2020c). Another genetically informed approach to estimating environmental effects on MD-MDD is Mendelian randomisation (MR), which uses
genetic instruments to identify potential causal relationships between environmental exposures and MD risk. MR methods have emerged as an alternative strategy to examine causal inference, obviating typical sources of confounding (Smith and Ebrahim, 2003). Here, strongly associated genetic variants are selected as statistical instruments — or proxies — to estimate the effect of an exposure of interest (e.g., substance use) on an outcome (e.g., MDD) under specific assumptions. MR studies have supported the role of educational attainment in reducing the depression risk and of BMI in increasing the risk (Wray et al., 2018). These studies have also suggested a potential role of lifestyle exposures in shaping the risk of major depression, such as social connections and sedentary media use (Choi et al., 2020b). Establishing stronger causal evidence for different environmental factors with updated methods may help to clarify inconsistencies across G×E findings (Ni et al., 2019). An additional source of potential bias and inconsistency in G×E research are underlying gene-environment correlations, as the presence of gene-environment correlations may lead to false GxE (Purcell and Sham, 2002; Ni et al., 2019).

Discussion

In summary, MDD is a common, debilitating, phenotypically heterogeneous disorder, with a heritability of ranges between 30-50%. MDD with early-onset, recurrence, comorbid anxiety, greater severity, and postpartum depression may be more heritable subtypes. Compared to other psychiatric disorders, its high prevalence, relatively low moderate heritability, and strong polygenicity have posed major challenges for gene-mapping. Recently, rare CNVs have been implicated in risk of MDD. Studies of common variation in MDD, driven forward by large international collaborations such as the PGC, confirmed the highly polygenic nature of the disorder and implicated over 100 loci in disorder risk. These studies identified genes involved in neuronal growth, synaptic function, and inflammation in the
pathophysiology of the disorder. Finally, gene-environment interaction and Mendelian randomisation studies have implicated childhood trauma, socioeconomic adversity, social support, physical activity, and their interactions with genetic factors, in risk of MDD.

Evidence on the genetic basis of MDD has multiple potential applications including the development of pharmacological treatments, informing discussions regarding disorder risk, and treatment choice. A greater understanding of the biological processes underlying MDD may facilitate the development of new, potentially more effective medications. MDD GWAS have implicated biological processes underlying the disorder including neurotransmission and synaptic structure as well as genetically informed pharmacological modes of action (Howard et al., 2019). MDD is a very heterogeneous disorder and genetic research will likely benefit from the study of more homogeneous disorder subtypes. For example, research on patients with atypical features of increased appetite or weight suggested that this patient subgroup might benefit from treatments specifically targeting immunometabolic pathways (Milaneschi et al., 2017).

An individual’s genetic make-up likely impacts the disease course – from prevention to diagnosis, treatment, and prognosis (Lewis and Vassos, 2020). Whilst the application of MDD PRS to predict individual risk is currently limited by its low discriminative accuracy, in the future MDD-PRS might be used for risk stratification at the population level (Andlauer and Nöthen, 2020). This is supported by recent evidence that MDD PRS – well-established to predict prevalent depression – also predict incident cases in the general population (Musliner et al., 2019). For other complex disorders like coronary artery disease and type-2 diabetes, PRS-based risk stratification is already powerful (Khera et al., 2018). Further development of PRS to allow
the identification of individuals at heightened genetic risk, coupled with targeted prevention and intervention measures, holds potential to mitigate depression risk (McIntosh et al., 2019). Knowledge of the genetic architecture of \textbf{MDD} and its links with other psychiatric and non-psychiatric disorders might inform genetic counselling and clinical care. For example, identified genetic correlations with physical health phenotypes might facilitate the development of clinical guidelines for physical health monitoring in individuals with \textbf{MDMDD}.

\textbf{MDD} is a multifactorial disorder. Other levels of genetic variation, such as DNA methylation (Barbu et al., 2020; Van Assche et al., 2017b) and subsequent gene-expression differences (Arloth et al., 2015; Schiweck et al., 2020), and also the telomere length (Lin et al., 2016), influence depression risk and severity. Importantly, epigenetic mechanisms serve as an interface between the individual and its environment (Provencal and Binder, 2015). A better understanding of the role of epigenetic variation and an improved integration of multi-omics data may enhance our understanding of the aetiology and treatment of MDD. In addition, this review focuses on genetic factors influencing the risk or severity of depression. We did not discuss how genetic variants affect response to treatment, especially the metabolism of antidepressant drugs via cytochrome P450 genes. This important topic was recently reviewed by Bousman and colleagues (Bousman et al. 2021).

Here, we highlighted the state of the art, the promises, and the pitfalls of genetic research in \textbf{MDMDD}. The field would benefit from further exploration of the stratification properties of genetics – both for the understanding of its aetiology, including the role of environmental factors, and for the development of targeted treatment opportunities. Rich datasets with high-quality and deep phenotyping are crucial for studying heterogeneous disorders like MDD. Combining deep, harmonized phenotypes with molecular data will greatly aid the genetic
stratification of MDD. Improving our insights into the impact of genetic factors on MDD holds great promise for improving clinical care, and this intense and fast-developing field has a bright future.

Acknowledgments

We thank Andrew McIntosh, Cathryn Lewis, and Naomi Wray for critical comments on our manuscript.
### Tables and Figures

Table 1. MDD heritability estimates from selected publications

<table>
<thead>
<tr>
<th>Key paper</th>
<th>Type</th>
<th>$h^2$ (95% CI)</th>
<th>$c^2$ (95% CI)</th>
<th>$e^2$ (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Twin data</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Sullivan et al., 2000)</td>
<td>Meta-analysis</td>
<td>0.37 (0.31-0.42)</td>
<td>0 (0-0.05)</td>
<td>0.63 (0.58-0.67)</td>
</tr>
<tr>
<td>(Kendler et al., 2006)</td>
<td>Largest twin study (15.5K pairs)</td>
<td>male: 0.29 (0.19-0.38)</td>
<td>NA$^a$</td>
<td>0.71 (0.62-0.81)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.42 (0.36-0.47)</td>
<td>NA$^a$</td>
<td>0.58 (0.53-0.64)</td>
</tr>
<tr>
<td>(Polderman et al., 2015)</td>
<td>Meta-analysis of 50 years of twin studies</td>
<td>depressive episode: 0.34 (0.31-0.38)</td>
<td>0.11 (0.08-0.13)</td>
<td>0.55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recurrent MDD: 0.45 (0.36-0.55)</td>
<td>0.03 (0-0.08)</td>
<td>0.52</td>
</tr>
<tr>
<td>(Kendler et al., 2018a)</td>
<td>Twins identified from SWE register; Treated cases</td>
<td>male: 0.41 (0.19-0.49)</td>
<td>0 (0-0.13)</td>
<td>0.59 (0.51-0.66)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.49 (0.31-0.56)</td>
<td>0.02 (0-0.17)</td>
<td>0.49 (0.44-0.55)</td>
</tr>
<tr>
<td><strong>EHR or register-based data</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Wray and Gottesman, 2012)</td>
<td>DEN register, parent-offspring</td>
<td>0.32 (0.30-0.34)</td>
<td>NA$^b$</td>
<td>NA$^b$</td>
</tr>
<tr>
<td>(Kendler et al., 2018a)</td>
<td>SWE register, siblings (full vs half, reared together vs apart)</td>
<td>male: 0.36 (0.31-0.38)</td>
<td>0.05 (0.05-0.05)</td>
<td>0.59 (0.58-0.61)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>female: 0.51 (0.51-0.53)</td>
<td>0.02 (0.02-0.02)</td>
<td>0.47 (0.46-0.49)</td>
</tr>
<tr>
<td>(Polubriaginof et al., 2018)</td>
<td>US EHR, inferred extended pedigree</td>
<td>depressive episode: 0.25 (0.17-0.30)</td>
<td>NA$^a$</td>
<td>0.75 (0.70-0.83)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recurrent MDD: 0.36 (0.21-0.51)</td>
<td>NA$^a$</td>
<td>0.64 (0.49-0.79)</td>
</tr>
</tbody>
</table>

$h^2$ narrow-sense heritability; $c^2$ shared environment; $e^2$ unique environment. SWE: Sweden; DEN: Denmark. EHR: Electronic Health Records.

- a. Missing estimate of $c^2$ as a result of AE models (i.e., quantitative genetic models with only additive genetics and unique environment components) in the original study.
- b. Only the heritability was derived from the summary level data.
Table 2. Whole-exome sequencing and exome chip studies assessing the role of rare variants in subphenotypes of depression.

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Method</th>
<th>Sample size</th>
<th>Ancestry</th>
<th>Implicated gene or variant</th>
<th>Implicated pathway</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population isolate</td>
<td>WES, exome chip</td>
<td>1999 (discovery) 1604 (replication)</td>
<td>European</td>
<td>NKPD1</td>
<td>de novo synthesis of sphingolipids</td>
<td>(Amin et al., 2017a)</td>
</tr>
<tr>
<td>Depressive symptoms in families &amp; Familial MD &amp; population isolate</td>
<td>Linkage, haplotype analysis, WES, exome chip</td>
<td>1336 WES 1527 exome chip</td>
<td>European</td>
<td>RCLI p.Leu186Phe (rs115482041)</td>
<td>interlaminar astrocytes</td>
<td>(Amin et al., 2018)</td>
</tr>
<tr>
<td>Familial &amp; early-onset MDD</td>
<td>targeted resequencing of 1742 synaptic genes</td>
<td>259 cases 334 controls</td>
<td>European</td>
<td>Cav2-adapter gene set</td>
<td>calcium signaling, actin polymerisation, spine formation</td>
<td>(Pirooznia et al., 2016)</td>
</tr>
<tr>
<td>Familial &amp; early-onset MDD</td>
<td>WES</td>
<td>12</td>
<td>European</td>
<td>ZNF34</td>
<td>brain TFs</td>
<td>(Subaran et al., 2016)</td>
</tr>
<tr>
<td>Sex differences in MDD</td>
<td>WES and WGS</td>
<td>1000 cases (70.7% female) 72 controls</td>
<td>Asian</td>
<td>PDE4A (rs201432982), FDX1L (rs62640397, rs79442975), MYO15B (rs820182, rs820148) more frequent in female cases</td>
<td>none</td>
<td>(Kang et al., 2020)</td>
</tr>
<tr>
<td>Suicide in MDD</td>
<td>WES specifically with 5'/3'-UTRs and promoters</td>
<td>23 cases 21 controls (post-mortem brain samples)</td>
<td>European</td>
<td>C0L6A6 variants in 17% of suicide completers; 3 calcium channel genes highlighted (CACNA1B, -1C, -2D4) 86 non-synonymous variants in 42 genes found in at least two suicide victims but not controls</td>
<td>TGF-β pathway</td>
<td>(Tomáč et al., 2017)</td>
</tr>
<tr>
<td>MRI grey matter volume changes in MDD</td>
<td>WES</td>
<td>77 cases 245 controls</td>
<td>Asian</td>
<td>CSMD1, CNTNAP5</td>
<td>neuroactive ligand receptor interacting pathway</td>
<td>(Zhang et al., 2020)</td>
</tr>
<tr>
<td>Study Description</td>
<td>Method</td>
<td>Sample Size</td>
<td>Region</td>
<td>Gene/Variant</td>
<td>Pathways/Function</td>
<td>Reference</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------------</td>
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<td>-----------------------------------------------</td>
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<td>-----------------------------</td>
<td>--------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>Depressive symptoms, general population</td>
<td>WES</td>
<td>3612</td>
<td>European</td>
<td>LIPG p.Asn396Ser (rs77960347)</td>
<td>steroid and cholesterol biosynthesis</td>
<td>(Amin et al., 2017b)</td>
</tr>
<tr>
<td>Depressive symptoms in context of environmental stress</td>
<td>exome chip, WGS</td>
<td>203 cases, 196 controls (discovery) 473 cases, 497 controls (replication)</td>
<td>Hispanic</td>
<td>PHF21B</td>
<td>9 pathways including innate immune response, glutamate signalling, sensory perception</td>
<td>(Wong et al., 2017)</td>
</tr>
<tr>
<td>Treatment resistance</td>
<td>GWAS, WES, exome chip</td>
<td>1209 cases (discovery) 1128 cases (replication)</td>
<td>European</td>
<td>response predictor in resistance vs. response to treatment with serotonergic antidepressants using rare variants</td>
<td>cell survival and proliferation, neurodegeneration, immune response</td>
<td>(Fabbri et al., 2020)</td>
</tr>
</tbody>
</table>

MDD=major depressive disorder; WES=whole-exome sequencing; WGS=whole-genome sequencing; GWAS=genome-wide association study; TFs=transcription factors; UTR=untranslated region; TGF-ß=Transforming growth factor beta; MRI=magnetic resonance imaging
Table 3. Published genome-wide association studies on depression.

<table>
<thead>
<tr>
<th>Year</th>
<th>Discovery: Nca/Nco or Ntot</th>
<th>Discovery: Loci</th>
<th>Replication: Cases/Controls</th>
<th>Replicated loci</th>
<th>MD-d</th>
<th>Definition of depression</th>
<th>Ancestry</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>2008</td>
<td>1738/1802</td>
<td>0</td>
<td>6079/5893</td>
<td>0*1</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Sullivan et al., 2009)‡</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>1359/1782</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Muglia et al., 2010)‡</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>3957/3428</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Shyn et al., 2011)‡</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>1636/1594</td>
<td>0</td>
<td>1418/1918</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Lewis et al., 2010)‡</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>604/1364</td>
<td>0</td>
<td>409/541</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Rietschel et al., 2010)‡</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>1020/1636</td>
<td>0</td>
<td>NA</td>
<td>NA</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Shi et al., 2011)‡</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>2431/3673</td>
<td>0</td>
<td>3332/3228</td>
<td>0</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Wray et al., 2012)‡</td>
<td></td>
</tr>
<tr>
<td>2011</td>
<td>353/366</td>
<td>0</td>
<td>3735/10635</td>
<td>1*2</td>
<td>Structured interview: MDD</td>
<td>European</td>
<td>(Kohli et al., 2011)‡</td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>MA/NA</td>
<td>N</td>
<td>MA/NA</td>
<td>N</td>
<td>Study Description</td>
<td>Location</td>
<td>Reference</td>
<td></td>
</tr>
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<td>-------------</td>
<td>-----------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>9240/9519</td>
<td>0</td>
<td>6783/50695</td>
<td>0</td>
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<td>Hispanic/ Latino</td>
<td>(Dunn et al., 2018)</td>
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<td>(Howard et al., 2018)</td>
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<td>Sample Size</td>
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<td>NA</td>
<td>Mixed</td>
<td>European and African American</td>
<td>(Levey et al., 2020)</td>
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_Year_ indicates the year in which the study was first published (including online ahead of print and, were no published version as available, preprint), which may deviate from the year the article was published.
in print. Nca/Nco for number of cases and controls in case-control design; Ntot as total sample size for quantitative trait.

Studies used for Figure 1.

+ Significant in the meta-analysis of discovery and replication cohorts in a recessive model. This association did not replicate in subsequent GWAS in additive models. Replication has, to our knowledge, not been attempted using a recessive model.
+ Significant in the meta-analysis of discovery and replication cohorts.
+ This association did not replicate in the follow-up Hispanic ancestry GWAS meta-analysis Dunn et al. 2018.
+ Meta-analysis of 23andMe and PGC MDD cohorts.
+ Replication of top results from the meta-analysis of 23andMe and PGC MDD cohorts.
+ Only SNPs not reaching genome-wide significance in the discovery-stage were analysed in the replication phase. Six loci (of which two were not significant in the discovery GWAS) covered by these selected SNPs reached significance in the meta-analysis of discovery and replication cohorts.
+ From a meta-analysis of overlapping subjects (PGC and UKBB), case and control numbers may thus be inflated.
+ The study identified 102 independent variants at 101 loci.

Figure 1: Identified GWAS loci in relation to the number of cases.
The relationship of identified loci to the number of cases based on published studies on schizophrenia, bipolar disorder (BD), and major depression (MDD). To simplify the presentation, studies on both MDD and MD with different depression definitions have been combined into the MD category. The plot shows the number of cases and independent significant loci in the discovery-stage analyses (without replication). Schizophrenia GWAS saw a critical inflection
point at approximately 15,000 cases and are currently discovering a new locus for, on average, every 230 new cases (Levinson et al., 2014). BD currently discovers a new locus for every 400 new cases. Such an inflection point was observed for MD GWAS only at approximately 75,000 cases, as predicted (Wray et al., 2012; Levinson et al., 2014), and MD GWAS using liberal phenotype definitions now identify a new locus every 1500 cases. Slopes starting at the inflection point were estimated using linear regression. The MD studies used for the plot are provided in Table 3.
References


Thank you for the careful review of our manuscript and the helpful feedback. In this response, we have addressed all issues raised, point-by-point. We have also carefully revised the manuscript and submitted a track-changed copy (MDDreview_R1_trackchanged) and a cleaned copy (MDDreview_R1_clean). We believe the manuscript is substantially improved with the constructive suggestions from the editor and the reviewers. We confirm this revised manuscript has met all the journal's requirements.

Reviewers' and editor's comments:
Both reviewers were broadly positive but had some concerns. PM does not take a strong position on MD vs. MDD initials. But what is important is to define what you mean - especially to divide studies that apply full (typically DSM) criteria or not and to distinguish whether that is done in a clinical interview or be questionnaire and whether you are assessing current mood state or a lifetime history of episodes of MD. Both reviewers felt more attention to measurement issues was warranted. Other concerns were also raised. Your MS is already a bit long for us - typical upper limit 4500. So take a few more hundred words if needed, but not more than that in your revision.

We agree with the editor and the reviewers that a clear definition is of vital importance, given that the construct definition is considered as one of the key sources of heterogeneity in MDD (Cai N, Choi KW, Fried EI. Human Molecular Genetics, 2020; PMID: 32568380). Therefore, we clarified the depression definitions in the Introduction (Page 5):

“In line with previous studies (Wray et al., 2018), we will use 'MDD' where the study subjects met the clinical criteria for major depressive disorder, and 'major depression' where the subjects were ascertained using alternative strategies, but nevertheless likely to have met the diagnostic criteria for MDD.”

Throughout the review, we have adjusted the use of the terms MDD and major depression to reflect the specific ascertainment strategies used in each cited study.

For more details, please also refer to our responses to Reviewer 1 comment 1 (i, iv, v), and Reviewer 2 comment 1.

Reviewer #1: Kendall et al., The Genetic Basis of Major Depression
This is a well-written and comprehensive review article on genetics of depression, with the major focus on molecular genetic methods and findings. The authors have read the literature carefully and they generally review it with good perspective on the differences among methods and the strength of evidence for various findings. This would be a good place for a non-specialist to find an overview and a perspective of research to date, and for researchers to find references on the topics that are covered here.

Thank you for this positive assessment.

The following are mostly relatively minor comments that might help to improve the review. The first item is, in my view, a slightly more major point that I think should be corrected - recognizing that it is not easy to judge when it is necessary to use the term "MDD" vs. "MD," so perhaps the editor will weigh in on the journal's preference here.

We have now carefully revised the definitions as suggested. Please refer to our response to the editor's comment for further details.
1. The introductory paragraphs on defining depression are important but need improvement: (i) It should be stated that the definition here is for the major depressive disorder (MDD) category (DSM-V, ICD-10). The alternative of studying continuous depression questionnaire scores is discussed later but should be mentioned here.

We agree and thank the reviewer for this suggestion. In response, we have added the following text to the Definition section (Page 4-5):

“The term ‘depression’ most often refers to the clinical diagnosis of MDD, as defined by the International Classification of Diseases (ICD) and the Diagnostic and Statistical Manual of Mental Disorders (DSM) diagnostic criteria. [...] Apart from such a dichotomous, diagnosis-based definition, the severity of depressive symptoms is often studied as a quantitative trait, so the full variation of depressive symptoms can be taken into account.”

(ii) Consider mentioning that the MDD category is used by most studies because it predicts familial risk (relatives or twins), antidepressant treatment response and future course (which does not prove it is the best way to measure depression, but most genetic researchers have concluded that it is the best available way).

We added a brief comment in the Definition section (Page 5):

“Despite the shortcomings of dichotomizing a heterogeneous disorder like MDD, a categorical definition of depression has been widely used in research.”

The specific examples mentioned by the reviewer were brought up in later sections.

(iii) The Cai et al. and PGC perspectives are not presented clearly. Correctly or not, both prefer the MDD category. PGC papers assert that because GWAS using "light phenotyping" methods (self-report, records) have high genetic (SNP) correlation with more rigorous interview-based studies, thus it is valid and more powerful to meta-analyze them. Cai asserts that examination of data for different strategies suggest that less rigorous methods risk missing genetic risk factors for the most severe forms of depressive illness.

We thank the reviewer for raising this point, and agree that the relevant text can be improved so that the two perspectives are more clearly presented.

However, we prefer to discuss these perspectives fully in a later section (e.g., after we introduced the GWAS, Page 15), and limit the Introduction section to broadly presenting the different strategies.

We now added the PGC perspective on Page 14 where we discussed this topic:

“[Importantly, phenotype definition matters]. Given the strong genetic correlations across cohorts, recent GWAS meta-analyses combined cohorts with different ascertainment strategies to maximize the overall sample size.”

(iv) PGC's "major depression" terminology is only partially explained. It does refer to the increased diagnostic uncertainty when combining diverse studies; but it also rests on the fact that all of the included strategies are targeting individuals likely to meet MDD criteria, as opposed to SSGAC which specifically includes studies based on symptom scores.

We added the following clarification on Page 5:
“[…] and 'major depression' where the subjects were ascertained using alternative strategies, but nevertheless likely to have met the diagnostic criteria for MDD.”

(v) The authors err in using "MD" when citing research on rigorously diagnosed MDD. "MDD" should be used when it was explicitly studied (i.e., for most of the literature cited here), "MD" or "proxy for MDD" when self-reported diagnoses or the like were used or were part of the range of definitions that were used; and "symptom score" or something comparable where appropriate; with a comment if the phenotyping strategy falls somewhere in between. The PGC's use of "MD" is an informal, descriptive term; it is not an historical, validated concept. An example of the negatives effects of this imprecision is a sentence on p 14, "A comparison of genetic analyses using different phenotype definitions in UKBB revealed that GWAS of broad phenotype definitions might identify associated loci that are not specific to MD (Cai et al., 2020b)." Cai's point is rather, that MD≠MDD, and that analyses of liberally-defined MD might miss critical loci associated with severe MDD. (I am agnostic personally about that conclusion, but it should be made more clear here.) On p 17, where it is stated that the high rg between MD and symptom score studies confirms "the concept of a continuum between subthreshold depressive symptoms and clinical diagnoses of MD." There is no such thing as a "clinical diagnosis of MD," only of MDD.

We thank the reviewer for raising this important point. As the reviewer correctly pointed out, the majority of the cited studies used strict criteria to define MDD. It was inappropriate to combine all studies into a single category under the liberally-defined term MD. Now, we explicitly defined the two terms, MDD and major depression (see our response to the editor's comment) and used them correspondingly throughout the review.

In addition, we clarified the text on Page 14:
“[...] However, a comparison of genetic analyses using different phenotype definitions in UKBB revealed that GWAS of broad phenotype definitions might identify associated loci that are not specific to MDD, but, more likely, shared with other psychiatric disorders (Cai et al., 2020b).”

And on Page 16:
“[...] confirming the concept of a continuum between subthreshold depressive symptoms and clinical diagnoses of MDD.”

(vi) The Fried and Hesse create a "straw man" regarding "unique combinations" of MDD symptoms. No combination is empirically unique or is a valid individual characteristic (i.e., endorsed symptoms show some variation across "test/re-test" interviews during an episode, or during different episodes, or during one interview that explicitly inquires about different episodes). But meeting MDD criteria has validated (if imperfect) significance.

This reference is not closely related to the key topics of our review. Therefore, we decided to remove the respective sentence and the reference in order to shorten the manuscript (Page 4 in the Introduction).

2. In the paragraph on CNVs, the Zhang et al. and Kendall et al. studies are described, but it is not made clear whether Kendall et al. actually tested the finding reported in Zhang et al. (increased burden for deletion < 100 kb) (I think they did not, but the authors of this review also wrote the excellent Kendall et al. CNV paper and could clarify.)

Essentially, the analyses of Zhang et al. and Kendall et al. overlapped only partially. CNVs <100kb, which were significant in Zhang et al., were not tested in Kendall et al. Conversely,
the neurodevelopmental CNVs assessed in Kendall et al. were not specifically screened in Zhang et al. The relevant sentences were changed, which now read as follows (Page 11):

“Outside of this group, there was no evidence for a residual burden of rare CNVs >100 kb in major depression. However, smaller CNVs were not tested in the study by Kendall and colleagues.”

3. It would be helpful to state at the beginning of the GWAS review that European-ancestry studies will be reviewed first. Otherwise it is a bit glaring that the CONVERGE GWAS paper is missing from the initial review of the history of significant MD/MDD GWAS findings.

We agree with the reviewer and added the following sentence to the start of the GWAS section (Page 11):

“Many MDD GWAS have been conducted during the past decade, predominantly on subjects of European ancestry (Table 3). Therefore, European-ancestry studies are described first, followed by a review of GWAS in more diverse populations.”

4. Mendelian Randomization is first introduced on p 20 in the context of GxE analysis. It would be helpful to introduce more broadly what the method does, and then discuss it in the specific GxE context. The MR analyses in the Wray et al. (2018) paper, specifically of obesity, are not mentioned, but should be.

We thank the reviewer for this suggestion. We have now added a brief introduction to Mendelian randomization methods and discuss them in the context of studying environmental influences on depression on Page 20:

“[Another genetically informed approach to estimating environmental effects on MDD is Mendelian randomisation (MR).] MR methods have emerged as an alternative strategy to examine causal inference, obviating typical sources of confounding (Smith and Ebrahim, 2003). Here, strongly associated genetic variants are selected as statistical instruments — or proxies — to estimate the effect of an exposure of interest (e.g., substance use) on an outcome (e.g., MDD) under specific assumptions. MR studies have supported the role of educational attainment in reducing the depression risk and of BMI in increasing the risk (Wray et al., 2018). Studies have also supported the potential role of various lifestyle exposures in shaping the risk of MDD, such as social connections and sedentary media use (Choi et al., 2020b).”

5. Parts of the Discussion read a bit too much like an advertisement for genetic studies - it is stated that the current state of this research does not adequately predict individual risk, and then all sorts of future benefits are described that would depend on much better individual predictions. It would be better to briefly note some of the uses that have been proposed (especially if supported by any empirical data), but not to oversell whether they will become reality. For example, I do not think that people with or without severe MDD need to know their PRS score to know that they might do well to eat better and exercise more. The problem isn't knowing that it might help, it is figuring out how to get oneself to actually do it, which is not a genetics problem.

The reviewer’s point is well-taken. Indeed, genetic risk prediction is in its infancy, with only 1.5-3% of the MDD liability currently explained by PRS. Thus, at this current stage, the outlook regarding future benefits is not clearly convincing. As the reviewer suggested, we have highlighted a few use cases in the Discussion, Pages 21-22:

“MDD is a very heterogeneous disorder and genetic research will likely benefit from the study of more homogeneous disorder subtypes. For example, research on patients with
atypical features of increased appetite or weight suggested that this patient subgroup might benefit from treatments specifically targeting immunometabolic pathways (PMID: 29049554).

 [...] Whilst the application of MDD PRS to predict individual risk is currently limited by its low discriminative accuracy, in the future PRS might be used for risk stratification at the population level (Andlauer and Nöthen, 2020). This is supported by recent evidence that MDD PRS of MDD – well-established to predict prevalent depression – also predict incident cases in the general population (PMID: 30698613). For other complex disorders like coronary artery disease and type-2 diabetes, PRS-based risk stratification is already powerful (PMID: 30104762)."

Reviewer #2:
Kendall and colleagues have submitted a very broad overview of the genetics of depression including sections on twin studies, candidate gene studies, rare variants, GWAS, PRS, and GxE. If the goal is to provide a cursory review, then they have done it by providing text on a wide range of topics. MDD/MD genetics is quite nuanced and more attention could be made on effects of phenotyping (depression symptoms, versus clinical interviews), subtypes (PPD, atypical depression), and population genetics (severe under-representation of diverse pops, transferability & precision medicine), and pharmacogenetics (cytochrome P450 enzyme genes & antidepressant drug metabolism). Could the authors also outline how this review significantly expands on previous reviews on the genetics of depression (Flint & Kendler 2014, Smoller 2016, McIntosh 2019, Ormel 2019, Cai Choi et al 2020)?

We thank the reviewer for this suggestion.

First, we have extensively revised the definitions in light of the comments and suggestions by the editor and Reviewer 1. As outlined above, we have now specifically defined and adjusted the use of the terms ‘MDD’ and ‘major depression’.

We agree that phenotyping, subtypes, population genetics, and pharmacogenetics are important topics. In fact, we have reviewed some of these topics wherever relevant. For example, we have summarized the literature on MDD subtypes related to the topics of Genetic Epidemiology, Rare Genetic Variants, GWAS, and Genetic Overlap; studies on population genetics are presented in the GWAS section.

We further added the following sentence about phenotyping on Page 5 (depressive symptoms vs clinical interview):

“Apart from such a diagnosis-based definition of depression, the severity of depressive symptoms is often studied as a quantitative trait, so the full variation of depressive symptoms can be taken into account.”

And about pharmacogenetics on Page 22:

“In addition, this review focuses on genetic factors influencing the risk or severity of depression. We did not discuss how genetic variants affect response to treatment, especially the metabolism of antidepressant drugs via cytochrome P450 genes. This important topic was recently reviewed by Bousman and colleagues (PMID: 33147643).”

We also thank the reviewer for the question regarding how our review expands on previous reviews on the genetics of MDD. The intention of our review was not to provide an entirely novel perspective compared to previous literature. Instead, we aimed to provide a comprehensive, up-to-date overview of the current knowledge in the field, from the perspective of the PGC. To this end, we focused on, in our opinion, the key topics related to MDD genetics. Our review could be considered as a primer, aiming to answer the question “What is MDD and
what is known about its genetics?” As Reviewer 1 commented, this review “… would be a good place for a non-specialist to find an overview and a perspective of research to date, and for researchers to find references on the topics that are covered here.”

A few minor comments:

*Abstract - consider editing from (a heritability) to 'heritability ranges from… and is moderate’ (not low)

The sentence was modified in the Abstract (Page 3):
“[…] with heritability ranges from 30% to 50%. […], moderate heritability, …”

and in the Discussion (Page 20):
“The heritability ranges between 30-50%. […] Compared to other psychiatric disorders, its high prevalence, moderate heritability, and strong polygenicity have posed major challenges for gene-mapping in MDD.”

*Definition - consider editing the name of 'somatic domain' - are sleep features considered somatic?

On page 4, ‘somatic domain’ has been replaced by ‘vegetative symptoms’, a terminology often used in this context.

*Consider adding the range of prevalences & factors that impact it (ex: country) in the Epidemiology section

We have added the range of prevalence across countries to the Epidemiology section (Page 5):
“The 12-month prevalence of MDD is estimated to be ~6% …, although these figures vary considerably across countries (e.g., the 12-month prevalence ranged from 2.2% in Japan to 10.4% in Brazil).”

A range of factors including age and gender are likely to contribute to the cross-country differences (Bracke, et al. 2020; PMID: 32980172). These factors are also presented in the Epidemiology section on Pages 5-6.

*Error in CONVERGE citation - ultra rare variant burden scores WERE associated with MD (PMID: 28002544)

We thank the reviewer for bringing to our attention that this point was not entirely clear. We changed the text as follows on Page 10:

“However, the largest WGS study of MDD to date, by the CONVERGE consortium (5,303 cases and 5,337 controls), failed to find evidence for the association of individual rare or low-frequency variants with MDD (CONVERGE consortium, 2015). Using the same data set, it was subsequently shown that individuals with MDD had a significantly higher overall burden of singleton exonic variants predicted to be deleterious (Peterson et al., 2017).”

*Consider adding a few sentences on other genetic approaches like methylation, telomere, mitochondrial genome

We thank the reviewer for this suggestion. We added a brief paragraph on Page 22:
“MDD is a multifactorial disorder. Other levels of genetic variation, such as DNA methylation (Barbu et al., 2020; Van Assche et al., 2017b) and subsequent gene-expression differences (Arloth et al., 2015; Schiweck et al., 2020), and also the telomere length (Lin et al., 2016), influence depression risk and severity. Importantly, epigenetic mechanisms serve as an interface between the individual and its environment (Provencal and Binder, 2015). A better understanding of the role of epigenetic variation and an improved integration of multi-omics data may enhance our understanding of the aetiology and treatment of MDD.”

*Consider expanding discussion on what is needed for the field to move forward - areas in need of further research & implications

We also thank the reviewer for the suggestion. We extended the relevant text on Page 23:

“[The field would benefit from further exploration of the stratification properties of genetics – both for the understanding of its aetiology, including the role of environmental factors, and for the development of targeted treatment opportunities.] Rich datasets with high-quality and deep phenotyping are crucial for studying heterogeneous disorders like MDD. Combining deep, harmonized phenotypes with molecular data will greatly aid the genetic stratification of MDD.”