

Multiscale neural gradients reflect transdiagnostic effects of major psychiatric conditions on cortical morphology

Bo-yong Park^{1,2,3}, Valeria Kebets¹, Sara Larivière¹, Meike D. Hettwer^{4,5,6,7}, Casey Paquola^{1,8}, Daan van Rooij^{9,10}, Jan Buitelaar¹¹, Barbara Franke^{11,12}, Martine Hoogman^{11,12}, Lianne Schmaal^{13,14}, Dick J. Veltman¹⁵, Odile A. van den Heuvel¹⁵, Dan J. Stein¹⁶, Ole A. Andreassen^{17,18}, Christopher R. K. Ching¹⁹, Jessica A. Turner^{20,21,22}, Theo G. M. van Erp^{23,24}, Alan C. Evans¹, Alain Dagher¹, Sophia I. Thomopoulos¹⁹, Paul M. Thompson¹⁹, Sofie L. Valk^{4,5}, Matthias Kirschner^{1,25,26} & Boris C. Bernhardt^{1,26}

It is increasingly recognized that multiple psychiatric conditions are underpinned by shared neural pathways, affecting similar brain systems. Here, we carried out a multiscale neural contextualization of shared alterations of cortical morphology across six major psychiatric conditions (autism spectrum disorder, attention deficit/hyperactivity disorder, major depression disorder, obsessive-compulsive disorder, bipolar disorder, and schizophrenia). Our framework cross-referenced shared morphological anomalies with respect to cortical myeloarchitecture and cytoarchitecture, as well as connectome and neurotransmitter organization. Pooling disease-related effects on MRI-based cortical thickness measures across six ENIGMA working groups, including a total of 28,546 participants (12,876 patients and 15,670 controls), we identified a cortex-wide dimension of morphological changes that described a sensory-fugal pattern, with paralimbic regions showing the most consistent alterations across conditions. The shared disease dimension was closely related to cortical gradients of microstructure as well as neurotransmitter axes, specifically cortex-wide variations in serotonin and dopamine. Multiple sensitivity analyses confirmed robustness with respect to slight variations in analytical choices. Our findings embed shared effects of common psychiatric conditions on brain structure in multiple scales of brain organization, and may provide insights into neural mechanisms of transdiagnostic vulnerability.

Mental illness refers to a wide range of psychiatric conditions affecting individuals, families, and health systems at large¹. While conventional psychiatric nosology classifies mental illness into distinct categories mainly based on descriptive symptoms and behaviors², high co-occurrence of symptoms across disorders as well as transdiagnostic risk factors have prompted reconceptualization of mental illnesses along symptom dimensions^{3–8}. Investigation of transdiagnostic effects may, thus, benefit detailed characterization of shared alterations across different psychiatric conditions and may identify direct brain-behavior associations that capture multiple symptom classes and mask clinical heterogeneity.

The shared components across major psychiatric diagnosis may be more clearly distinguishable at the neural level^{4,9}, as behavioral characterization likely involves complex interactions with society and the environment¹⁰. Structural magnetic resonance imaging (MRI), in particular, offers high spatial precision to help resolve the pattern of shared transdiagnostic effects across the cortical surface^{4,11–16}. Prior case-control studies have reported reproducible patterns of structural alterations in cohorts with psychiatric diagnoses relative to controls^{17–21}, often pointing to widespread changes in cortical morphology in these conditions. More recently, efforts have been expanded to a transdiagnostic perspective, aiming to identify structural compromise that is shared across different diagnoses^{22–24}. To ensure the sensitivity of such efforts and to strengthen reproducibility, it becomes increasingly relevant to pool these investigations across multiple sites. One such initiative, spearheaded by the Enhancing Neuroimaging Genetics through Meta-Analysis (ENIGMA) consortium, has aggregated MRI and phenotypic data in thousands of healthy individuals and those with a psychiatric diagnosis²⁵. Moreover, dedicated ENIGMA working groups have confirmed neuroanatomical disruptions in major psychiatric indications, including autism spectrum disorder (ASD)²⁶, attention deficit hyperactivity disorder (ADHD)²⁷, major depressive disorder (MDD)²⁸, obsessive-compulsive disorder (OCD)²⁹, bipolar disorder (BD)³⁰, and schizophrenia (SCZ)³¹, pointing to widespread changes in cortical morphology in each of these different conditions. Also fostered by the open dissemination of condition-related effect size maps through the primary ENIGMA papers and their aggregation within the recently developed ENIGMA toolbox³², it has now become possible to systematically study these effects.

In addition to providing robust evidence of neuroanatomical signatures associated with each of these conditions, an emerging body of studies has pooled data across different indications to identify shared anomalies of psychiatric conditions^{33,34}. In an effort to identify factors contributing to the topography of cross-disorder brain changes, a recent study has taken this approach one step further and examined associations to postmortem gene expression data, searching for spatially co-varying gene lists that may carry susceptibility to transdiagnostic disease effects. This study revealed that transdiagnostic effects may generally be more marked in regions with greater expression of CA1 pyramidal genes that were suggested to play a role in regulating cortical thickness. Beyond these molecular risk factors, a broad range of cellular, metabolic, and functional properties of brain regions may contribute to regional susceptibility, but such an association remains underexplored. An influential theory, also referred to as the “structural model”, posits that the internal microstructural and connective markup of different brain regions, in particular their laminar differentiation and cortico-cortical connectivity patterns, may represent mesoscale features associated with the potential of a region to show plasticity, and to be susceptible to pathological processes³⁵. According to this framework, paralimbic cortices with low laminar differentiation and associated

connectivity profiles may be more susceptible to effects of neurological as well as psychiatric disorders. Here, we tested this approach by aligning transdiagnostic effects with maps of microstructural variations derived from both in vivo imaging and 3D postmortem histology^{36–39}. In recent work, the application of nonlinear eigenvector decomposition techniques to imaging and histology datasets identified a sensory-fugal gradient that radiates from sensory and motor areas with strong laminar differentiation and higher myelination towards heteromodal association and paralimbic regions with less clear lamination and lower myelin content. Of note, similar yet not completely corresponding gradients have also been derived from the analysis of intrinsic functional connectivity patterns obtained from resting-state functional MRI^{38–40}. In line with foundational neuroanatomical conceptualization^{35,41,42}, an emerging literature has underscored a correspondence between such data-driven sensory-fugal gradients and region-to-region variations in cortical plasticity and genetic control^{40,43–47}, suggesting that these likely help understand susceptibility to common brain disorders as well^{40,43,48–52}.

Examining associations between shared morphological alterations and receptor architecture may provide additional opportunities for the contextualization of transdiagnostic effects. In the healthy brain, neurotransmitter systems are indeed broadly implicated in region-to-region variations of synaptic plasticity, neural dynamics, and inter-network communication. Moreover, recent initiatives have aggregated maps outlining the spatial distributions of different neurotransmitter systems in vivo, based on positron emission tomography (PET) and single photon computed emission tomography (SPECT) studies sensitive to different receptor and transporter types^{53–59}. Such mapping can complement microstructural and functional connectivity contextualization of transdiagnostic findings, promising insights into additional molecular factors contributing to regional susceptibility. Beyond the mapping of regional variations in the receptor architecture of the neurotypical brain, neurotransmitter imbalances have been described in several psychiatric conditions. Work in SCZ and depression, for example, implicated a role of dopamine and serotonin^{60–63}, and more recent work in BD and SCZ demonstrated associations between neurotransmitter and functional network imbalances⁶⁴.

Here, we studied the association between multiscale neural organization and transdiagnostic effects on cortical morphology across six major psychiatric conditions (ASD, ADHD, MDD, OCD, BD, and SCZ). Aggregating data from thousands of patients and healthy controls previously studied across several ENIGMA working groups^{26–31}, we defined the shared effect using principal component analysis. The robustness of the shared effect was further cross-validated based on openly aggregated effect size maps from the ENIGMA toolbox³². The shared dimension was contextualized across multiple neural scales. This involved systematic assessment of spatial associations to (i) in vivo myeloarchitecture and intrinsic functional connectivity, (ii) postmortem 3D cytoarchitecture, and (iii) in vivo maps of neurotransmitter distributions. Notably, in addition to assessing specific associations, we also adopted supervised machine learning to identify joint spatial associations between the above neural features and the common dimension of morphological alterations. Multiple sensitivity analyses verified robustness of our findings.

Results

Study overview and participants. We obtained case-control maps of cortical thickness differences in patients relative to controls, resulting from several ENIGMA working groups aggregated by a previous study that included a total of 28,546

participants across 145 independent cohorts (1821 ASD, 1815 ADHD, 2695 MDD, 2274 OCD, 1555 BD, 2716 SCZ; 15,670 site-matched controls Supplementary Table 1)³³. We then associated principal dimensions of morphological with (i) in vivo myeloarchitecture and functional connectivity gradients obtained from the Human Connectome Project (HCP)⁶⁵, (ii) postmortem cytoarchitecture, by cross-referencing data to a ultra-high resolution 3D histological human brain model⁶⁶, and (iii) in vivo neurotransmitter topographies provided by PET/SPECT studies^{53–59}. Approaches are openly available and replicable via the ENIGMA toolbox (<https://enigma-toolbox.readthedocs.io>)³². See Methods for more details.

Shared dimensions of structural alterations across psychiatric conditions. Following standardized ENIGMA protocols (<http://enigma.ini.usc.edu/protocols/imaging-protocols/>), gray matter thickness for 68 cortical regions of the Desikan–Killiany atlas⁶⁷ was calculated, and meta-analytic between-group differences in cortical thickness were assessed using inverse variance-weighted random-effects models (Fig. 1a)³³. Using principal component analysis adopted in a recent study³³, we estimated the shared disease dimensions explaining structural alterations across six conditions (Fig. 1b). The first dimension/component explained 55.7% of variance, and differentiated sensory/motor systems having positive scores from transmodal/paralimbic areas with negative scores (for details, and information on the other dimensions/components, see Supplementary Fig. 1a). Stratifying the first dimension according to intrinsic functional communities⁶⁸, it indeed differentiated somatomotor/visual from default/frontoparietal/limbic networks (Fig. 1b). Similar spatial patterns were observed across an atlas of the putative primate cortical hierarchy⁴¹, differentiating idiosyncratic/unimodal from heteromodal/paralimbic levels.

Several sensitivity analyses confirmed and further expanded these findings. Firstly, scores on the principal dimension translated into mean effect size of morphological alterations across case-control analyses, with paralimbic regions showing the strongest atrophy in patients relative to controls, while sensory/motor regions showed the least gray matter alterations (Supplementary Fig. 1b). Compared to the dimensional analysis that highlighted paralimbic as well as sensory/motor regions with opposite ends, this analysis confirmed that paralimbic regions are most vulnerable to the impact of neuropsychiatric conditions, and primary motor cortex is least vulnerable. Secondly, we directly ran principal component analysis on previously reported effect size maps (Cohen's *d*) concatenated across disorders, sourced from the ENIGMA toolbox³² (Supplementary Fig. 1c). The first principal dimension was highly similar to ours ($r = 0.552$, spin-test $p_{\text{spin}} < 0.001$), suggesting robustness. Thirdly, the shared disease effect resembled the effects of each condition, with the strongest spatial similarity to SCZ and BD, followed by MDD, ADHD, ASD, and OCD (spin-test followed by false discovery rate (FDR) correction, $p_{\text{spin-FDR}} < 0.001$; Supplementary Fig. 2), indicating that the shared effect captured structural alterations from each condition. Fourthly, we re-evaluated the shared dimension using leave-one-condition-out procedure (see Methods), and observed largely consistent results with the shared effect based on all conditions ($r > 0.9$, $p_{\text{spin-FDR}} < 0.001$; Supplementary Fig. 3), indicating that a single condition with strong meta-analytic profile did not determine the shared disease effect. Finally, we associated three alternative shared disease maps (including principal dimension (see Fig. 1b), mean effect (see Supplementary Fig. 1b) of meta-analytic profiles, and principal dimension of Cohen's *d* map (see Supplementary Fig. 1c)) with previously published maps of cortical expansion and functional

reconfiguration⁶⁹ to examine whether the shared disease dimension reflects evolutionary expansion. We found low-to-moderate correlations ($r = -0.256$, $p_{\text{spin}} = 0.036$ for shared disease dimension; $r = -0.193$, $p_{\text{spin}} = 0.087$ for mean effect size; $r = -0.480$, $p_{\text{spin}} < 0.001$ for principal component of Cohen's *d* maps).

Associations with cortical myeloarchitecture and functional connectivity gradients. To assess in vivo micro- and macroscopic properties of the shared disease dimension on cortical morphology, we first examined its spatial association with myeloarchitecture and intrinsic functional connectivity gradients^{38,40} (see Methods; Fig. 1c). The microstructural gradient was derived from inter-regional similarity matrices of intracortical profiles of myelin-sensitive MRI³⁸, and runs from sensory/motor regions with high laminar differentiation and high intracortical myelin content towards paralimbic cortices with reduced laminar differentiation and low myelin content³⁸. The intrinsic functional gradient was derived from resting-state functional MRI connectivity. While it also runs from sensory/motor to transmodal areas, it finds its apex in the heteromodal default mode and frontoparietal networks, and not in paralimbic cortices⁴⁰. Associating the patterns of shared dimension with these two in vivo gradients, we observed a negative association with the microstructural gradient ($r = -0.400$, $p_{\text{spin-FDR}} = 0.042$) and a negative trend with the functional connectivity gradient ($r = -0.247$, $p_{\text{spin-FDR}} = 0.090$; Fig. 1c). In other words, transdiagnostic morphological alterations follow sensory-fugal gradients of cortical organization, in particular, the microstructural gradient that differentiates sensory/motor areas with high myelination and distinct lamination from paralimbic areas with low myelin content and reduced laminar differentiation.

Cytoarchitectonic associations. We furthermore examined associations of the shared disease effect with inter-regional variations in cortical cytoarchitecture³⁷, using BigBrain, a 3D histological reconstruction of a postmortem human brain^{66,70}. We calculated cortex-wide variations in cytoarchitecture using two alternative approaches. First, we obtained intracortical intensity profiles and calculated their statistical moments, i.e., mean, SD, skewness, and kurtosis (Fig. 2a, b). In both classic cytoarchitecture analysis and more recent work, these features have been shown to relate to inter-areal microstructural differentiation^{39,71}. In particular, the skewness moment describes a robust spatial transition from areas with low laminar differentiation and negative skewness to those with high laminar differentiation and positive skewness^{71–73}. Moreover, we computed externopyramidization⁷⁴, describing a gradual shift of intensity profiles across cortical layers that has been suggested to also differentiate areas on the lower end of the cortical hierarchy from those that are higher up due to hierarchical shifts in laminar projection profiles⁷⁵ (Fig. 2a, b). Notably, while both skewness and externopyramidization describe overall sensory-fugal patterns, they do so in complementary ways ($r = 0.015$, $p_{\text{spin}} = 0.506$), with skewness differentiating mainly prefrontal and posterior cingulate regions from visual, auditory, and frontocentral regions while externopyramidization clearly differentiates postcentral and visual regions from the rest of the brain. Importantly, however, spatial correlations between these features and the principal disease dimension indicated relations to both features (skewness: $r = 0.400$, $p_{\text{spin-FDR}} = 0.015$; externopyramidization: $r = 0.472$, $p_{\text{spin-FDR}} = 0.015$; Fig. 2c). In other words, transdiagnostic cortical thickness decreases were more likely in paralimbic regions with low skewness and low externopyramidization, independently confirming that those areas with low laminar differentiation were more likely to show transdiagnostic cortical alterations. In addition to the associations with BigBrain cytoarchitectural features, we additionally examined associations of the shared disease effect with intracortical profile moments and

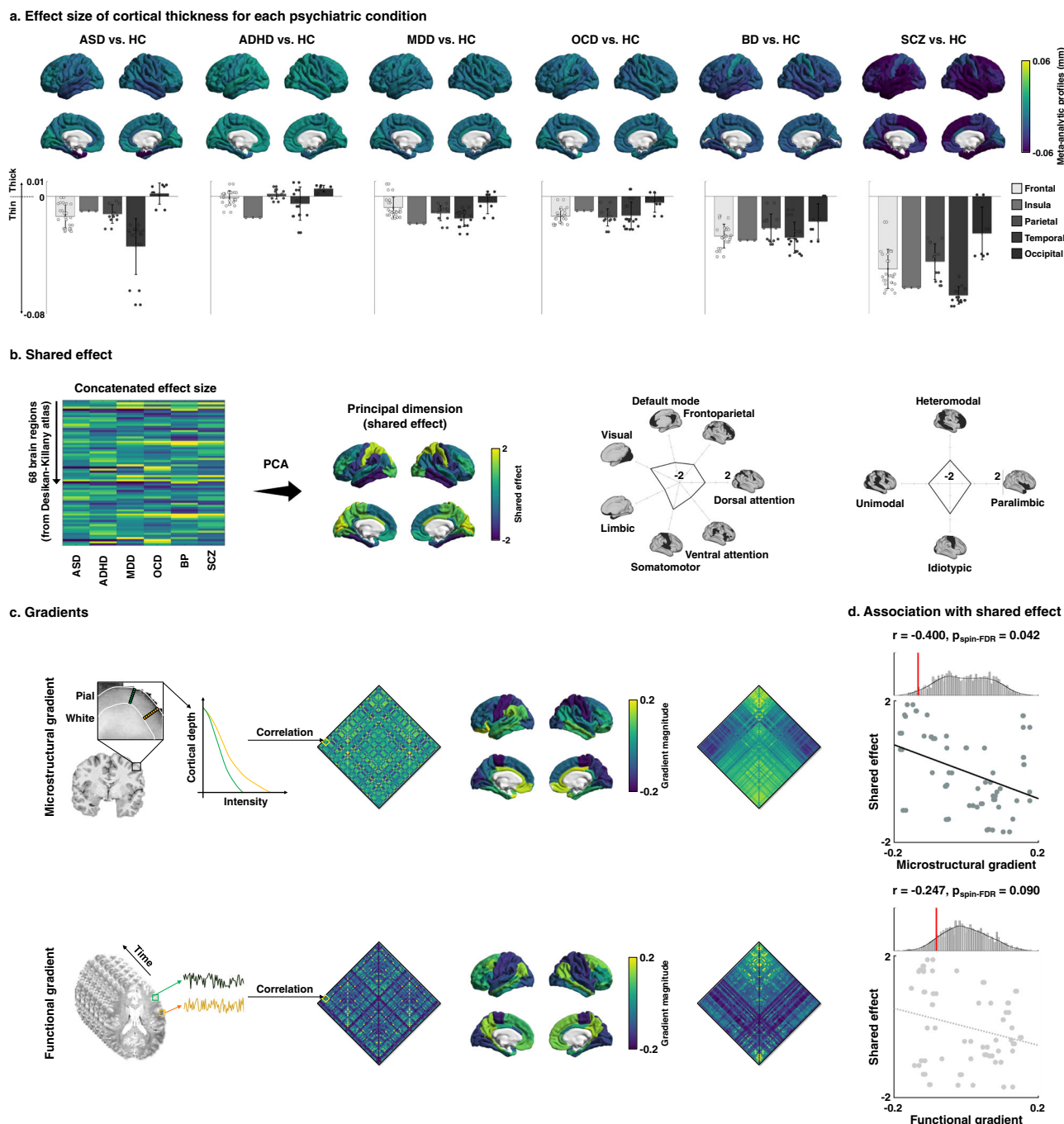


Fig. 1 Shared disease effect and associations to connectivity gradients. **a** Meta-analytic profiles of cortical thickness differences (unit in mm) in patients with each psychiatric condition relative to matched controls. Positive/negative values indicate increases/decreases in cortical thickness in patients relative to controls. Mean values of the regions involved in the same cortical lobes with SD are reported with bar plots. **b** The shared effect was identified through principal component analysis (PCA) applied to the concatenated effect size map. Spider/Radar plots stratify the effects according to functional communities⁶⁸ and cortical hierarchy levels⁴¹. **c** The microstructural and functional connectivity gradients were generated by applying nonlinear dimensionality reduction techniques to the group averaged connectivity matrix (middle left), and each connectivity matrix was reordered (right) according to the first gradients (middle right). **d** Spatial correlations of each gradient with the shared effect map are shown in the scatter plots. The distribution of correlation coefficients across 1,000 spin-tests are reported with histograms, and the actual r -values are represented with red bars. ASD autism spectrum disorder, ADHD attention deficit hyperactivity disorder, MDD major depressive disorder, OCD obsessive-compulsive disorder, BD bipolar disorder, SCZ schizophrenia, HC healthy controls, spin-FDR spin-test followed by false discovery rate.

externopyramidization calculated from in vivo myelin-sensitive MRI i.e., T1w/T2w measures obtained from the HCP database. We observed largely consistent results (Supplementary Fig. 4), suggesting robustness.

Associations with distributions of neurotransmitter systems. Neurotransmitter contextualization leveraged JuSpace⁵³, an open access toolbox that disseminates in vivo PET/SPECT data sensitive to ten different transmitters/transporters/receptors from independent studies in healthy human adults^{54–59} (Fig. 3a). Associating the shared

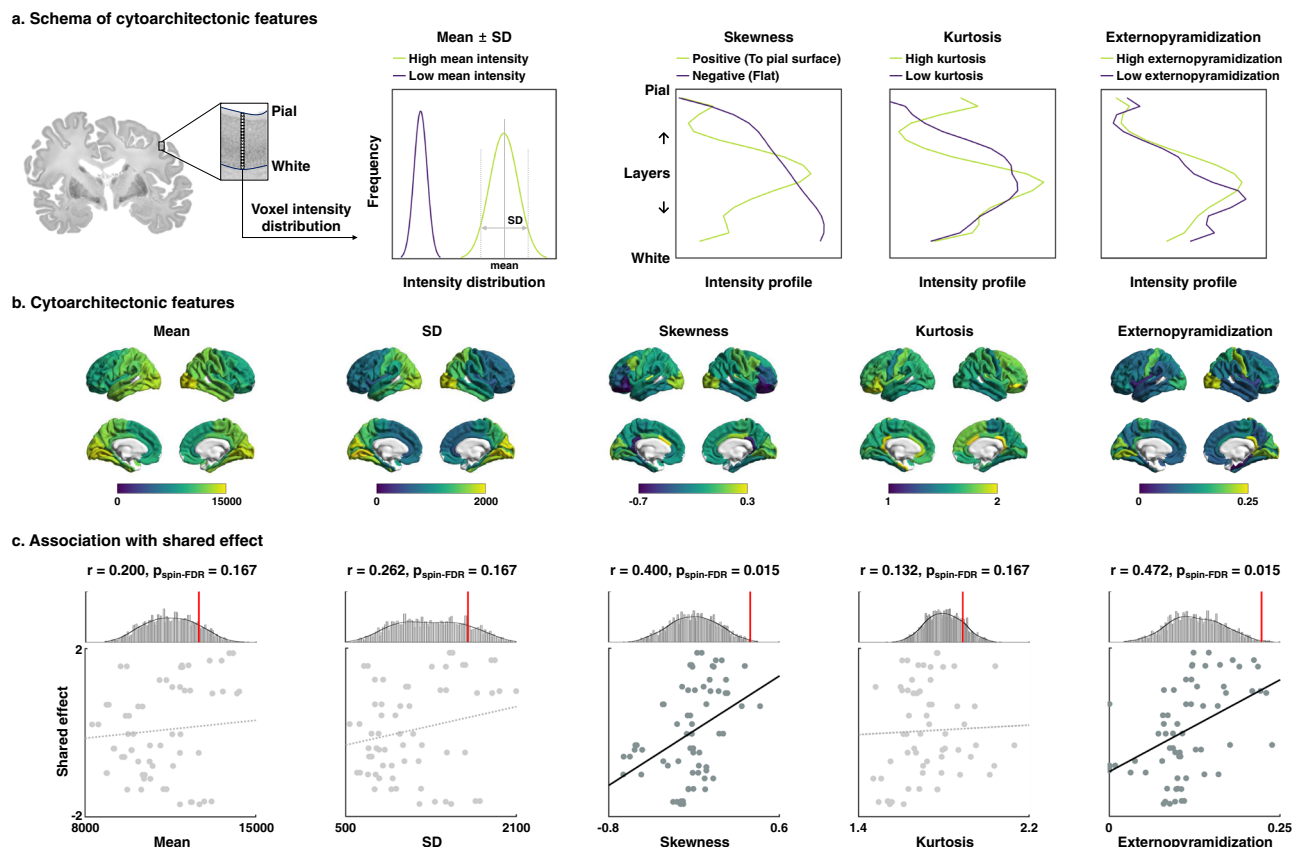


Fig. 2 Cytoarchitectonic associations with the shared disease effect. **a** Cytoarchitectonic moment features of mean, SD, skewness, and kurtosis, as well as externopyramidization of intracortical intensity profile were calculated from the postmortem human brain, and **b** plotted on brain surfaces. **c** Spatial correlations between the features and shared effects are shown on scatter plots. The distributions of correlation coefficients across 1000 spin-tests are reported with histograms, and the actual r -values are represented with red bars. SD standard deviation, spin-FDR spin-test followed by false discovery rate.

dimension with cortex-wide neurotransmitter maps, we observed positive associations with D2 and 5-HT1b receptor densities (D2: $r = 0.280$, $p_{\text{spin-FDR}} = 0.035$; 5-HT1b: $r = 0.349$, $p_{\text{spin-FDR}} = 0.025$), and negative correlations with dopamine transporter and 5-HT1a receptor density (DAT: $r = -0.240$, $p_{\text{spin-FDR}} = 0.041$; 5-HT1a: $r = -0.307$, $p_{\text{spin-FDR}} = 0.033$; Fig. 3b). The results indicate that common cortical alteration patterns across psychiatric and neurodevelopmental conditions may be reflected by serotonergic and dopaminergic systems. More specifically, higher transdiagnostic cortical atrophy was related to higher 5-HT1a and lower 5-HT1b, as well as higher DAT and lower D2 receptor density.

Associations of multiscale features with other shared dimensions. We investigated associations of other principal components of cortical morphological alterations (Supplementary Fig. 1a), instead of first principal component, to multiscale neural features. We found that the first dimension showed significant associations to cytoarchitecture/microstructure features, while the second dimension showed higher (and inverted) associations to the functional gradient (Supplementary Table 2). The results indicate that the first principal dimension represents a more sensory-paralimbic axis, and the second dimension is a more sensory-heteromodal axis. In addition to cortical thickness, we investigated a shared disease dimension based on the surface area. The dimension showed a somatomotor-visual/frontoparietal pattern (Supplementary Fig. 5), which was different from the shared effect based on cortical thickness ($r = -0.030$, $p_{\text{spin-FDR}} = 0.446$). The surface area-based shared dimension was significantly associated with SERT ($r = 0.274$, $p_{\text{spin-FDR}} = 0.029$), 5-HT2a ($r = -0.297$,

$p_{\text{spin-FDR}} = 0.011$), and GABAA ($r = -0.325$, $p_{\text{spin-FDR}} = 0.005$; Supplementary Table 3), suggesting higher sensitivity to serotonergic and GABAergic systems.

Machine learning prediction of the shared disease effect. As a final analysis, we used supervised machine learning to predict the first shared dimension using the above multiscale features. Specifically, we leveraged least absolute shrinkage and selection operator (LASSO) regression⁷⁶ with five-fold nested cross-validation^{77–80} to predict the cross-condition effect using concatenated multiscale features (see Methods; Fig. 4a). Repeating the analysis for 100 times with different training and test dataset subsplits, we could reliably predict the spatial pattern of the shared disease dimension (mean $\pm$ SD, $r = 0.518 \pm 0.044$, mean absolute error (MAE) = 0.828 ± 0.039 , permutation-test $p_{\text{perm}} < 0.001$; Fig. 4b). Cytoarchitectural skewness and externopyramidization, followed by D2 and 5-HT1b receptors, as well as the microstructural gradient were frequently selected across cross-validations and repetitions (Fig. 4a). When considering each psychiatric condition separately, we could find significant prediction performances, but the features selected diverged across conditions (Supplementary Fig. 6).

Discussion

The current work determined cortex-wide variations in susceptibility to morphological alterations across six major psychiatric conditions (i.e., ASD, ADHD, MDD, OCD, BD, and SCZ), and cross-referenced these spatial patterns against multiscale cortical organization. Specifically, complementing earlier case-control MRI studies performed separately in common neuropsychiatric

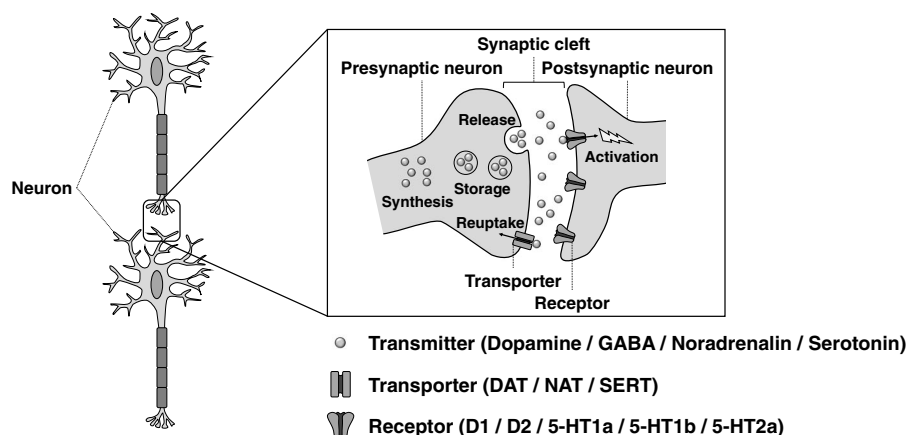
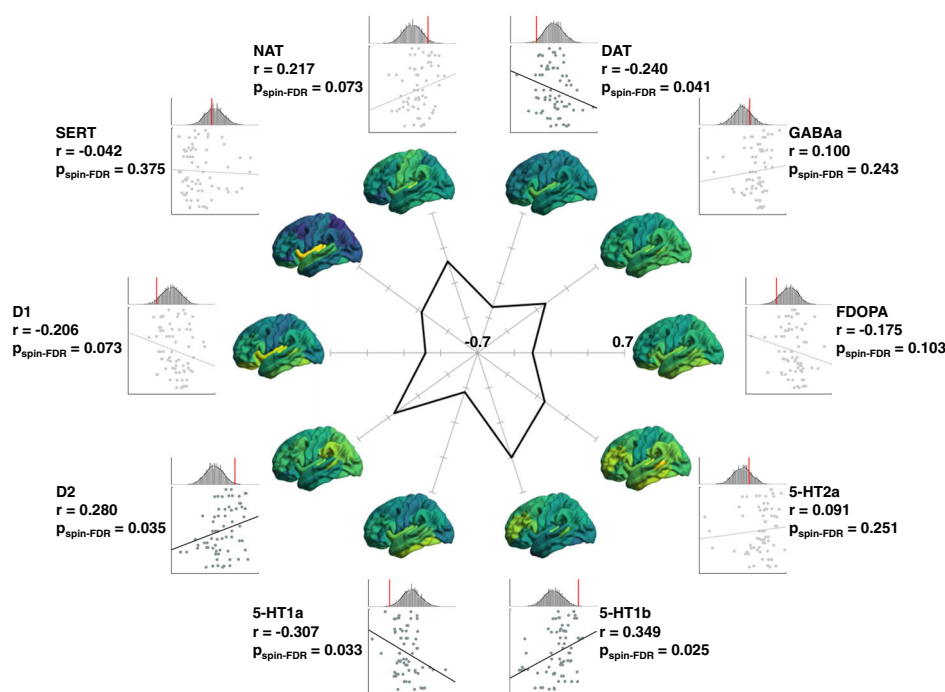
a. Schema of neurotransmitter systems**b. Association with shared effect**

Fig. 3 Associations of neurotransmitter systems with shared disease effect. a Schema of neurotransmitter systems of transmitters, transporters, and receptors. **b** Spatial correlations of each neurotransmitter map with shared effect are shown on scatter plots. The distributions of correlation coefficients across 1000 spin-tests are reported with histograms, and actual r -values are reported with red bars. The spider plot shows correlation coefficients. Cortex-wide spatial maps of the transmitter systems are reported on brain surfaces. FDOPA 18 F fluorodopa, DAT dopamine transporter, NAT noradrenaline transporter, SERT serotonin transporter, spin-FDR spin-test followed by false discovery rate.

conditions^{17,26–31,81}, we applied dimensional decomposition to cortical morphological data. We identified a shared dimension that followed a sensory-paralimbic pattern of increasing susceptibility to morphological alterations in paralimbic regions, which was robust across different data and approaches, and consistent with prior work on shared functional imbalances^{82,83}. Albeit prior functional connectivity-based findings highlighted heteromodal association and our structural morphological alterations highlighted paralimbic regions, the findings converge that higher order brain areas are vulnerable to multiple psychiatric conditions. Expanding from a prior study that reported transdiagnostic effects to be highest in brain regions expressing genes for pyramidal CA1 cells, pointing already to potentially increased susceptibility of limbic allocortices³³, here,

we characterized the transdiagnostic effects across multiple scales of neural organization. Specifically, we contextualized the shared disease effect on MRI-derived morphology with respect to (i) in vivo MRI measures sensitive to cortical myeloarchitecture and intrinsic functional connectivity^{38,40}, (ii) postmortem measures of cytoarchitecture, in particular laminar differentiation^{37,39,66,71}, and (iii) in vivo PET/SPECT maps of cortical neurotransmitter systems^{53–59}. Moreover, and in addition to studying specific associations between shared disease effects and individual features, we employed a supervised machine learning paradigm to synergistically assess the utility of multiscale neural features in explaining the shared disease effect. Our findings emphasize that microstructurally determined gradients, differentiating sensory/motor and paralimbic cortices^{38,40,84} can help to compactly

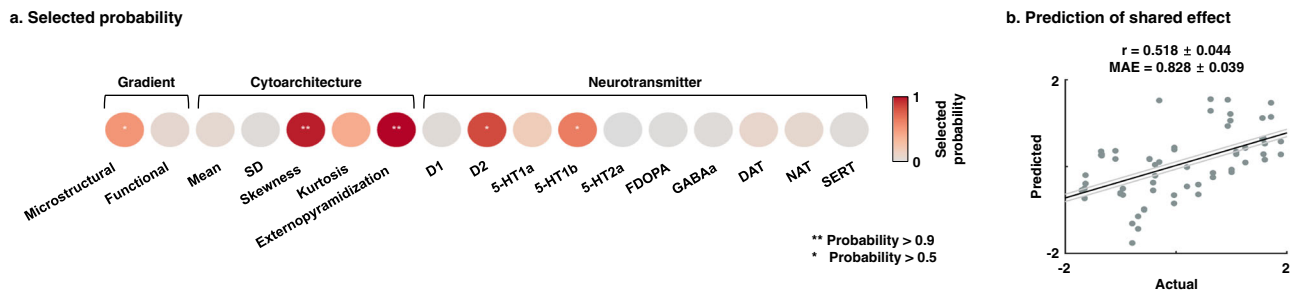


Fig. 4 Association between the shared disease effect and multiscale features using machine learning. **a** Probability of the selected features across five-fold nested cross-validations and 100 repetitions for predicting the shared disease effect. The frequently selected features are reported with asterisks. **b** Linear correlation between actual and predicted values of the effects is shown on a scatter plot. The black line indicates mean correlation and the gray lines represent the 95% confidence interval for 100 iterations with different training/test datasets. SD standard deviation, FDOPA 18 F fluorodopa, DAT dopamine transporter, NAT noradrenaline transporter, SERT serotonin transporter, MAE mean absolute error.

describe cortex-wide susceptibility to transdiagnostic effects of common mental health conditions. Overall heightened susceptibility was furthermore associated with two neurotransmitter markers, serotonin, and dopamine. Altogether, our work extended prior work in terms of (i) demonstrating robustness of a shared disease dimension across psychiatric conditions via multiple sensitivity analyses, (ii) providing a framework that integrates multiscale neural organization with the transdiagnostic disease effect on cortical morphology, and (iii) assessing the synergistic value of different cortical features to explain the shared effect, which may advance our understanding of neuropathology in psychiatry, and may inform the development of diagnostic and treatment strategies that cut across traditional disease boundaries.

Whether our findings also suggest a shared disease mechanism remains to be established. Prior histopathological assessments in people with ASD, MDD, OCD, BD, and SCZ have reported common cellular alterations, in particular, reduced neuronal and glial densities as well as neuronal size in different cortical areas^{85–92}. Moreover, a growing literature has shown shared genetic risk factors across major psychiatric conditions⁹³. Studies have implicated genes involved in several synaptic pathways⁹³, for example, common gene variants in cell adhesion and glutamate receptor pathways in ASD and SZ^{94–98}, as well as those in calcium signaling in BD and SZ⁹⁹. Together, the studies suggest that shared cellular and molecular risk factors may influence structural plasticity and lead increased disease susceptibility of psychiatric conditions⁹³, providing the rationale of investigating multiscale neural properties. As the first analysis, we defined the in vivo microstructural cortical gradient using a recently-introduced procedure³⁸, which identified axes of cortico-cortical differentiation based on the similarity of myelin-sensitive MRI profiles sampled across cortical depths. In healthy adults and adolescents^{38,71}, this approach has revealed a robust sensory-fugal cortical gradient running from sensory/motor areas with marked laminar differentiation and high myelin content towards paralimbic cortices with low overall myelination and rather agranular cortical profiles. By showing an association between the shared dimension and this microstructural gradient, we demonstrated a heightened susceptibility of paralimbic cortices to disease-related cortical thickness changes. Several features of the paralimbic cortex may underscore its increased susceptibility. On the one hand, paralimbic architecture may permit an increased potential for brain plasticity. This includes a lower neuronal density in paralimbic regions compared to eulaminate cortices, as well as increased dendritic arborization and synaptic density³⁵. Compared to other regions, paralimbic areas also continue to express developmental markers long into adulthood, such as growth-associated protein GAP-43¹⁰⁰. Furthermore, paralimbic cortices

have a protracted myelination and lower myelin content relative to sensory/motor areas. The role of intracortical myelination in plasticity is likely complex, but several streams of evidence point to the role of myelin acting as a buffer against plasticity. In addition to acting as an insulator for electrical transmission, myelin associated growth inhibitors limit activity and experience-induced axon sprouting, with downstream effects on synaptic plasticity¹⁰¹. Reduced myelin content, together with increased complexity of dendritic arborization in transmodal and paralimbic regions may render cortical microstructure in these regions more susceptible to pathological alterations, which would echo observations in other conditions. For example, the core pathological substrates of drug-resistant temporal epilepsy is thought to be localized in limbic/paralimbic regions^{102–104}, and prior work has suggested rather specific changes in myelin and microstructural proxies in these areas^{105,106}. Similar findings have been observed in neurodegenerative conditions such as Alzheimer's disease^{45,107,108}, where pathology spreads from disease epicenters in paralimbic allocortices to invade more widespread cortical/subcortical networks, but also depression¹⁰⁹ and autism^{110,111}. These findings collectively indicate that cellular and molecular features of paralimbic cortices and their cortico-cortical pathways promote brain plasticity as well as higher metabolic activity, and are, thus, likely more vulnerable to both developmental as well as acquired disruptions than other regions, supporting the hypothesis that their cortical type predisposes to a heightened vulnerability for an impact of neuropsychiatric conditions on alterations in brain morphology³⁵.

Studying a postmortem 3D model of the human brain, BigBrain⁶⁶, we obtained supporting confirmation for the above association between cortical microstructure and disease-related susceptibility. In particular, we discovered similarly marked associations between the shared disease dimension and laminar profile skewness as well as externopyramidization, two complementary features tapping into depth-dependent shifts in the distribution in cell densities^{39,74}. In prior work, profile skewness, in particular, was found to discriminate unimodal granular cortices from agranular/dysgranular paralimbic regions at a cortex-wide level⁷¹, and accurately delineated the iso-to-allocortical axis in the mesiotemporal lobe system⁷². Studying typical adolescent development, changes in profile skewness of myelin-sensitive MRI contrasts have furthermore been reported to spatially colocalize with expression patterns of genes enriched in oligodendrocytes⁷¹. As a complementary feature of the laminar organization, externopyramidization indexes the ratio of neuronal densities between supragranular and infragranular cortical layers. It increases when the cortex is cytoarchitecturally more differentiated, which happens in primary areas with a marked layer 4⁷⁴. Thus, the association of these cortical depth-dependent

cytoarchitectural features with the shared disease effect confirms the *in vivo* findings with ultra-high-resolution cytoarchitecture data suggesting that paralimbic areas, sensitive to transdiagnostic cortical alterations, are less laminarly differentiated. Furthermore, prior cellular and transcriptomic studies indicate regional susceptibility of synaptic elements as well as mutated genes in schizophrenia^{112,113} and bipolar disorder¹¹⁴. Indeed, major depression may be associated with atrophy of neurons in limbic regions¹¹⁵, pointing histopathological susceptibility of paralimbic areas in psychiatric conditions.

We also observed a marginal association between the transdiagnostic effect on brain structure and the principal functional connectivity gradient, but findings were overall weaker than for the above *in vivo* and postmortem derived microstructural gradients. Despite an overall convergence between structure and function in showing sensory-fugal gradients, prior work noted some divergence between sensory-heteromodal functional connectivity gradients⁴⁰ and sensory-paralimbic microstructure/cytoarchitecture gradients³⁸. This is in particular notable with respect to the heteromodal vs paralimbic anchor that these two gradients radiate towards. Mounting evidence suggests that heteromodal systems, such as the default mode network, decouple from microstructurally defined axes of brain organization that mainly describe differences in laminar differentiation^{38,116}. That work has also shown that regions with strong microstructure-function decoupling also host more flexible cognitive functions, and have marked cross-species differences between humans and nonhuman primates. Considering that difference, the current work shows an association between transdiagnostic disease effects and microstructural (i.e., sensory-paralimbic) gradients but not between transdiagnostic disease effects and functional (i.e., sensory-heteromodal) gradients. As such, the above divergence suggests increased specificity of the sensory-paralimbic gradient with respect to disease-related vulnerability, which is likely more determined by the microstructural context of cortical areas compared to their placement within cortical functional hierarchies. It is nevertheless important to underscore that vulnerability and susceptibility are likely affected by multiple factors and be reflected in different structural and functional gradient axes^{48,117–120}. These considerations collectively motivate caution in interpreting the observed associations, and also likely rule out a single mechanism underlying the observed association between the studied gradients and transdiagnostic effects.

In addition to our findings showing overall associations between the transdiagnostic effect and sensory-fugal microstructural gradients, we observed associations to the spatial distribution of different neurotransmitter systems derived from *in vivo* neuroimaging. Notably, associations were seen both to serotonin (5-HT1a and 5-HT1b) and dopamine receptors and transporters (DAT/D1 and D2), two previously reported markers of mental health and targets for pharmacological treatments^{64,121–128}. In both cases (i.e., 5-HT1a vs 5-HT1b, DAT/D1 vs D2), associations to the disease effect were of opposite polarity, confirming prior work in rodents^{129–133} and humans^{134–137}. Associations with *in vivo* neurotransmitter topographies provide a way of indirectly assessing the relationship between shared alterations of cortical morphology and neurotransmitter systems so that we can understand putative mechanisms of shared morphological alterations, extending prior work in rodents and humans. As different tracers may have variable sensitivity/specificity across cortical regions and, and as PET data may suffer from relatively coarse spatial resolution and partial volume effects, future studies are required to expand these findings based on a broader array of tracers and based on potentially more detailed techniques, such as 3D receptor autoradiography^{138,139}.

Our study has limitations. First, comorbidities and medication may contribute to the shared disease effect. Comorbidities are common in psychiatric conditions^{140–143}. Moreover, it has been shown that additional diagnosis beyond the primary diagnosis may affect the degree of cortical thickness changes, for example in ADHD with BD and MDD with anxiety^{144–146}. As data used in our study were collected from many independent research centers, comorbidities and medication effects could not be fully adjusted. Future studies need to consider controlling for such confounders. Second, we associated the shared disease dimension with features from multiple neural scales derived from independent cohorts, which precludes a direct interpretation of transdiagnostic effects with respect to histological as well as molecular mechanisms in the same subjects. Future studies, likely very challenging to accomplish, that measure multiscale features from the same individuals may set basis for more direct interpretations. Third, the cortical morphological data from the ENIGMA dataset were only available in the Desikan–Killiany parcellation⁶⁷, a macroscopic scheme following sulco-gyral patterns. In addition to not offering a high granularity on cortical arealization, the reliance on folding patterns alone may only provide rather limited sensitivity to contextualize our findings with respect to functional topographies. It would, thus, be relevant to re-evaluate functional gradient association based on functionally-defined parcellations^{147,148}, and/or to assess vertex-level feature data in future efforts. Lastly, through the investigation of the mean effect, we observed that the primary motor cortex is least vulnerable to the conditions we observed. However, the low absolute mean effect in sensory/motor regions does not per se indicate that these regions are less important in the understanding of psychiatric conditions. They may have potential effects that have not been detectable with our analyses. For example, prior functional connectivity studies observed that transdiagnostic effects may often be detectable in sensory and motor cortices^{149,150}, in addition to heteromodal and paralimbic areas^{82,83}.

As a final integrative analysis, we opted for a supervised statistical learning paradigm to predict the shared disease effect from combinations of neuroarchitectural features. This analysis indeed underscored that not a single feature, but rather combinations of microarchitectural and transmitter systems, have the highest utility in predicting the spatial pattern of the transdiagnostic morphological dimension. By, thus, highlighting microstructural and functional aspects of local cortical circuitry, our data-driven findings provide insights into potential determinants of transdiagnostic effects. Overall, our findings emphasize that an increasingly recognized principal gradient that differentiates sensory/motor networks from transmodal cortices in healthy brains^{38,40,84} also describes the main axis of cortex-wide susceptibility to transdiagnostic effects of common mental health conditions. Altogether, the findings may provide a potentially integrative framework for understanding neuropathology in psychiatry, and potentially inform the development of diagnostic and treatment strategies that cut across traditional disease boundaries.

Methods

Study dataset

ENIGMA data. We analyzed T1-weighted data from people with a diagnosis of ($n = 12,876$) ASD ($n = 1821$), ADHD ($n = 1815$), MDD ($n = 2695$), OCD ($n = 2274$), BD ($n = 1555$), and SCZ ($n = 2716$) and site-matched healthy controls ($n = 15,670$) from 145 independent cohorts participating in prior ENIGMA consortium studies^{26–31}. Demographic information is summarized in Supplementary Table 1 and available in a recent cross-condition study³³. Data from each center were processed using the standard ENIGMA workflow (<http://enigma.ini.usc.edu/protocols/imaging-protocols/>). Processing was conducted using FreeSurfer^{151–153} that involves magnetic field inhomogeneity correction, non-brain tissue removal, intensity normalization, and tissue segmentation. Estimated white and pial surfaces were inflated to spheres and registered to the *fsaverage* template. Based on the

Desikan–Killiany atlas⁶⁷, cortical thickness was measured for 68 gray matter brain regions. For each psychiatric condition, the ENIGMA groups performed multiple linear regression analyses to fit cortical thickness measures with age, age squared, sex, and site information. The meta-analytic profiles of between-group differences between patients and controls were estimated via an inverse variance-weighted random-effects model, which can be obtained from the previous study³³ (Fig. 1a). If the studies provided multiple effect sizes across children/adolescents/adults, only the effects from the adult sample were used, in order to match the age range across conditions. The positive/negative effects indicate increases/decreases in cortical thickness in patients relative to controls. Individual cohort investigators obtained approval from local institutional ethics boards, and informed consent was obtained from study participants or their guardians.

HCP data. To generate microstructural and functional connectivity gradients, we also studied 207 unrelated healthy young adults (60% females, mean age $\pm$ SD = 28.73 ± 3.73 years) from the HCP dataset⁶⁵. In the HCP, multimodal imaging data comprising T1- and T2-weighted as well as rs-fMRI were acquired on a Siemens Skyra 3 T at Washington University. The cohort selection is identical to our prior work^{32,154}. T1-weighted images were acquired using a magnetization-prepared rapid gradient-echo (MPRAGE) sequence (repetition time (TR) = 2400 ms; echo time (TE) = 2.14 ms; inversion time (TI) = 1000 ms; flip angle = 8°; field of view (FOV) = $224 \text{ mm}^2 \times 224 \text{ mm}^2$; voxel size = 0.7 mm isotropic; 256 slices). T2-weighted data were obtained using a T2-SPACE sequence, with the same acquisition parameters as for the T1-weighted data except for TR (3200 ms), TE (565 ms), and flip angle (variable). The rs-fMRI data were collected using a gradient-echo echo-planar imaging sequence (TR = 720 ms; TE = 33.1 ms; flip angle = 52°; FOV = $208 \text{ mm}^2 \times 180 \text{ mm}^2$; voxel size = 2 mm isotropic; the number of slices = 72; and 1200 volumes per time series), where participants were instructed to keep their eyes open looking at a fixation cross during the scan. Two sessions (left-to-right and right-to-left phase-encoded directions) of rs-fMRI data were acquired, providing up to four time series per participant. Participant recruitment procedures and informed consent forms, including consent to share de-identified data, were previously approved by the Washington University Institutional Review Board as part of the HCP.

Images underwent minimal preprocessing pipelines using FSL, FreeSurfer, and Workbench as follows^{155–157}:

T1- and T2-weighted data. Data were corrected for gradient nonlinearity and b0 distortions, and then T1- and T2-weighted data were co-registered using a rigid-body transformation. The bias field was adjusted based on the inverse intensities from the T1- and T2-weighting. The white and pial surfaces were generated^{151–153}, and the mid-thickness surface was generated by averaging them. The mid-thickness surface was inflated and the spherical surface was registered to the Conte69 template with 164k vertices¹⁵⁸ using MSMAll¹⁴⁸ and downsampled to a 32k vertex mesh.

Microstructure data. Myelin-sensitive proxy was estimated based on the ratio of the T1- and T2-weighted contrast^{159,160}. We generated 14 equivolumetric surfaces within the cortex and sampled T1w/T2w intensity along these surfaces³⁸. A microstructural similarity matrix was constructed by calculating the linear correlation of cortical depth-dependent T1w/T2w intensity profiles between different cortical regions based on the Desikan–Killiany atlas⁶⁷, controlling for the average whole-cortex intensity profile³⁸. The matrix was thresholded at zero and log-transformed³⁸. A group matrix was constructed by averaging matrices across participants.

rs-fMRI data. Data were corrected for distortions and head motion, and registered to the T1-weighted data and subsequently to MNI152 standard space. Magnetic field bias correction, skull removal, and intensity normalization were performed. Noise components attributed to head movement, white matter, cardiac pulsation, arterial, and large vein related contributions were removed using FMRIB's ICA-based X-noiseifier (ICA-FIX)¹⁶¹. Preprocessed time series were mapped to the standard “grayordinate” space using a cortical ribbon-constrained volume-to-surface mapping algorithm. The total mean of the time series of each left-to-right/right-to-left phase-encoded data was subtracted to adjust the discontinuity between the two datasets and then concatenated to form a single time series. A functional connectivity matrix was constructed by calculating the linear time series correlations between Desikan–Killiany parcels⁶⁷, followed by Fisher's r-to-z transformation¹⁶². Individual connectivity matrices were averaged to construct a group level connectome.

Shared effects of cortical thickness differences across conditions. To assess transdiagnostic effects of cortical thickness differences in patients relative to controls, we applied principal component analysis to the concatenated effect size maps across six conditions¹⁶³ (Fig. 1b and Supplementary Fig. 1a). The first principal dimension was determined as the shared disease effect. We summarized the effects according to seven intrinsic functional communities⁶⁸, as well as four cortical hierarchical levels⁴¹. We additionally calculated the mean effect size across the conditions to intuitively interpret shared disease effect (Supplementary Fig. 1b) and also estimated the

principal dimension based on the data sourced from the ENIGMA toolbox (i.e., Cohen's d; Supplementary Fig. 1c). We compared the shared dimension and the effect size of each condition via linear correlations to assess the degree of contribution of each condition (Supplementary Fig. 2). The significance of the correlation was determined using 1000 nonparametric spin-tests, to account for spatial autocorrelation¹⁶⁴, and corrected for multiple comparisons using an FDR procedure¹⁶⁵. To assess robustness, we performed leave-one-condition-out cross-validation. Specifically, we estimated the shared dimension using five conditions by excepting for a single condition, and assessed similarity with the shared disease effect estimated based on the whole six conditions (Supplementary Fig. 3). We calculated the significance of the correlation using 1000 spin-tests and multiple comparisons were corrected using FDR^{164,165}. We furthermore obtained the map of cortical expansion and functional reconfiguration⁶⁹ and calculated correlations with three shared disease maps, where the significance was determined using a 1000 spin-test.

Associations to microstructural and functional connectivity gradients. We evaluated the underlying connectome organizations of the shared disease effects. Based on T1w/T2w and rs-fMRI data obtained from the HCP database⁶⁵, we estimated microstructural and functional gradients, the low dimensional representation of connectome organizations explaining spatial variation in the connectome data^{38,40}, using BrainSpace (<https://github.com/MICA-MNI/BrainSpace>)¹⁶⁶ (Fig. 1c). An affinity matrix was constructed with a normalized angle kernel from the group averaged connectivity matrix with the top 10% entries for each parcel. The connectome gradients were estimated using diffusion map embedding¹⁶⁷, which is robust to noise and computationally efficient compared to other nonlinear manifold learning techniques^{77,168}. It is controlled by two parameters α and t , where α controls the influence of the density of sampling points on the manifold ($\alpha = 0$, maximal influence; $\alpha = 1$, no influence) and t scales eigenvalues of the diffusion operator. The parameters were set as $\alpha = 0.5$ and $t = 0$ to retain the global relations between data points in the embedded space, following prior applications^{38,40,48,166,169}. We associated the shared effect with these gradients using linear correlation (Fig. 1d), where the significance was assessed using 1000 spin-tests followed by FDR^{164,165}.

Cytoarchitectonic associations with shared disease effects. We aimed to associate the shared dimensions with histology-driven cytoarchitectonic features derived from BigBrain surfaces with 62 cortical areas (<https://bigbrain.loris.ca/main.php>)⁶⁶. Specifically, BigBrain is a ultra-high resolution, 3D volumetric reconstruction of a postmortem Merker-stained and sliced human brain from a 65-year-old male, with specialized pial and white matter surface reconstructions⁶⁶. The postmortem brain was paraffin-embedded, coronally sliced into 7400 20- μm sections, silver-stained for cell bodies¹⁷⁰, and digitized. A 3D reconstruction was implemented with a successive coarse-to-fine hierarchical procedure, resulting in a full brain volume. Among 68 regions defined by the Desikan–Killiany atlas⁶⁷, three regions per hemisphere, including banks of the superior temporal sulcus, frontal pole, and temporal pole, were excluded as the BigBrain did not provide data for these regions. We generated 18 equivolumetric cortical surfaces within the cortex (<https://github.com/caseyapaola/BigBrainWarp>) and sampled the intensity values along these surfaces. Based on the intensity values, we calculated four moment features, including mean, SD, skewness, and kurtosis, as well as externopyramidization (Fig. 2a, b). The mean and SD represent the overall intensity distribution of cytoarchitecture across layers, skewness indicates shifts in intensity values towards supragranular layers (i.e., positive skewness) or flat distribution (i.e., negative skewness), and kurtosis identifies whether the tails of the intensity distribution contain extreme values. Externopyramidization reflects gradual shifts of intensity values from infragranular to supragranular layers defined as follows⁷⁵:

$$\text{Externopyramidization} = \frac{(\text{intensity})}{\text{mean}(\text{intensity})} \times \frac{1 - \text{thickness}_{\text{supra}}}{\text{thickness}_{\text{total}}} \quad (1)$$

To assess associations with shared disease effects, we calculated linear correlations between cytoarchitectonic features and shared effects (Fig. 2c). The significance of the correlations was assessed using 1,000 spin-tests followed by FDR across different cytoarchitectonic features^{164,165}. BigBrain is a 3D model of the human brain with microscopic resolution, enabling us to assess cellular organization in cortical layers⁶⁶. However, it is based a single and significantly older subject compared to our study participants. We thus additionally calculated intracortical profile moments as well as externopyramidization from in vivo myelin-sensitive MRI i.e., the ratio of T1- and T2-weighted contrast^{159,160}, obtained from the HCP database⁶⁵. We associated the features with shared disease effect, and the correlations were assessed using 1,000 spin-tests followed by FDR^{164,165}.

Associations between transmitter systems and shared effects. To provide underlying molecular properties of the shared effects in neuroanatomical disruptions across different psychiatric conditions, we associated the shared dimensions with ten different neurotransmitter maps of healthy controls provided by prior independent PET/SPECT studies^{54–59}, which contain neurotransmitters of FDOPA, GABAa, transporters of DAT, NAT, SERT, and receptors of D1, D2, 5-HT1a, 5-HT1b, and 5-HT2a (<https://github.com/juryxy/juSpace>)⁵³ (Fig. 3a). All PET maps were linearly rescaled to have intensity values between 0 and 100⁵³.

After mapping the neurotransmitter maps onto the Desikan–Killiany atlas⁶⁷, we calculated linear correlations between the shared effects and each neurotransmitter map (Fig. 3b), and assessed the significance using 1000 spin-tests followed by FDR to adjust for multiple comparisons across ten different maps^{164,165}.

Associations between multiscale features and other shared dimensions. We furthermore associated the second shared dimensions (Supplementary Fig. 1a) with microstructural and functional connectivity gradients, cytoarchitectural moments, and neurotransmitter system distributions to assess convergence or divergence across the shared dimensions (Supplementary Table 2). We also estimated a shared disease dimension based on surface area, sourcing Cohen's d maps from the ENIGMA toolbox³². Here, we applied principal component analysis to the concatenated effect size maps of surface area across five conditions, as ASD was not available (Supplementary Fig. 5 and Supplementary Table 3).

Prediction of shared effects using multiscale features. We associated multiscale features and shared effects using supervised machine learning to incorporate our findings (Fig. 4). Specifically, we aimed to predict the shared disease effects using concatenated multiscale features of microstructural and functional gradients, cytoarchitectonic (i.e., mean, SD, skewness, kurtosis, and extropy-pyramidalization), and transmitter maps (i.e., D1, D2, 5-HT1a, 5-HT1b, 5-HT2a, FDOPA, GABAa, DAT, NAT, and SERT). We used five-fold nested cross-validation^{78–80} with LASSO regression⁷⁶. Nested cross-validation split the dataset into training (4/5) and test (1/5) partitions, and each training partition was further split into inner training and testing folds using another five-fold cross-validation. The model with the best performance (lowest MAE) across the inner folds was applied to the test partition of the outer fold. Among the multiscale features, we selected performant features using LASSO regularization, and the effect size was predicted using linear regression with the selected features. The procedure was repeated 100 times with different training and test partitions. Prediction accuracy was evaluated with linear correlations between the actual and predicted effect size and the MAE, with their 95% confidence interval. Permutation-based correlations across 1000 tests were conducted by randomly shuffling cortical regions to verify whether the prediction performance exceeded chance levels. We also performed the prediction analysis using the effect size of each condition (Supplementary Fig. 6).

Statistics and reproducibility. The between-group differences in cortical thickness of each psychiatric condition were assessed using inverse variance-weighted random-effects models³³, and their shared disease effects were estimated via principal component analysis. We assessed associations between the shared disease dimension and microstructural and functional connectivity gradients, cytoarchitectonic features calculated from the BigBrain⁶⁶, and neurotransmitter maps obtained from independent PET/SPECT studies^{54–59} based on linear correlations with 1000 spin-tests followed by FDR^{164,165}. We opted for supervised machine learning to associate multiscale features and shared effects based on five-fold nested cross-validation^{78–80} with LASSO regression⁷⁶.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Disorder-related effect size measures analyzed in this project are openly available via <https://enigma-toolbox.readthedocs.io> and <https://doi.org/10.1001/jamapsychiatry.2020.2694>. Raw imaging data that support these findings are not publicly available in a repository as they contain information that could compromise the privacy of research participants. Although there are data sharing restrictions imposed by (i) ethical review boards of the participating sites, and consent documents; (ii) national and trans-national data sharing law, such as GDPR; and (iii) institutional processes, some of which require a signed MTA for limited and predefined data use, we welcome sharing data with researchers, requiring only that they submit an analysis plan for a secondary project to the leading team of the Working Group (<http://enigma.ini.usc.edu>). Once this analysis plan is approved, access to the relevant data will be provided contingent on data availability and local PI approval and compliance with all supervening regulations. If applicable, distribution of analysis protocols to sites will be facilitated. Source data (Supplementary Data 1) are provided with this paper.

Code availability

Codes for multiscale neural feature calculation and statistical analyses are provided at BrainSpace (<https://doi.org/10.1038/s42003-020-0794-7>; <https://github.com/MICA-MNI/BrainSpace>) and ENIGMA toolbox (<https://doi.org/10.1038/s41592-021-01186-4>; <https://github.com/MICA-MNI/ENIGMA>). Tutorials to carry out similar analyses are provided on <https://enigma-toolbox.readthedocs.io/en/latest/>.

Received: 31 May 2022; Accepted: 7 September 2022;

Published online: 27 September 2022

References

- Whiteford, H. A. et al. Global burden of disease attributable to mental and substance use disorders: findings from the Global Burden of Disease Study 2010. *Lancet* **382**, 1575–1586 (2013).
- American Psychiatric Association. *Diagnostic and Statistical Manual of Mental Disorders* (American Psychiatric Press, 2013).
- Marshall, M. The hidden links between mental disorders. *Nature* **581**, 19–21 (2020).
- Xia, C. H. et al. Linked dimensions of psychopathology and connectivity in functional brain networks. *Nat. Commun.* **9**, 3003 (2018).
- Insel, T. et al. Research domain criteria (RDoC): toward a new classification framework for research on mental disorders. *Am. J. Psychiatry* **167**, 748–751 (2010).
- Cuthbert, B. N. The RDoC framework: facilitating transition from ICD/DSM to dimensional approaches that integrate neuroscience and psychopathology. *World Psychiatry* **13**, 28–35 (2014).
- Kotov, R., Krueger, R. F. & Watson, D. A paradigm shift in psychiatric classification: the Hierarchical Taxonomy of Psychopathology (HiTOP). *World Psychiatry* **17**, 24–25 (2018).
- Kotov, R. et al. The Hierarchical Taxonomy of Psychopathology (HiTOP): a dimensional alternative to traditional nosologies. *J. Abnorm. Psychol.* **126**, 454–477 (2017).
- Parkes, L. et al. Transdiagnostic dimensions of psychopathology explain individuals' unique deviations from normative neurodevelopment in brain structure. *Transl. Psychiatry* **11**, 232 (2021).
- Borsboom, D. A network theory of mental disorders. *World Psychiatry* **16**, 5–13 (2017).
- Kaczurkin, A. N. et al. Evidence for dissociable linkage of dimensions of psychopathology to brain structure in youths. *Am. J. Psychiatry* **176**, 1000–1009 (2019).
- Kaczurkin, A. N. et al. Neurostructural heterogeneity in youths with internalizing symptoms. *Biol. Psychiatry* **87**, 473–482 (2020).
- Linden, D. E. J. The challenges and promise of neuroimaging in psychiatry. *Neuron* **73**, 8–22 (2012).
- Hong, S. J. et al. Toward neurosubtypes in autism. *Biol. Psychiatry* **88**, 111–128 (2020).
- Castellanos, F. X., Di Martino, A., Craddock, R. C., Mehta, A. D. & Milham, M. P. Clinical applications of the functional connectome. *Neuroimage* **80**, 527–540 (2013).
- Milham, M. P., Vogelstein, J. & Xu, T. Removing the reliability bottleneck in functional magnetic resonance imaging research to achieve clinical utility. *JAMA Psychiatry* **39**, 4213–4227 (2021).
- Valk, S. L., Di Martino, A., Milham, M. P. & Bernhardt, B. C. Multicenter mapping of structural network alterations in autism. *Hum. Brain Mapp.* **36**, 2364–2373 (2015).
- Hong, S. J., Valk, S. L., Di Martino, A., Milham, M. P. & Bernhardt, B. C. Multidimensional neuroanatomical subtyping of autism spectrum disorder. *Cereb. Cortex* **28**, 3578–3588 (2018).
- Hajek, T., Carrey, N. & Alda, M. Neuroanatomical abnormalities as risk factors for bipolar disorder. *Bipolar Disord.* **7**, 393–403 (2005).
- Khundrakpam, B. S., Lewis, J. D., Kostopoulos, P., Carbonell, F. & Evans, A. C. Cortical thickness abnormalities in autism spectrum disorders through late childhood, adolescence, and adulthood: a large-scale MRI study. *Cereb. Cortex* **27**, 1721–1731 (2017).
- Lorenzetti, V., Allen, N. B., Fornito, A. & Yücel, M. Structural brain abnormalities in major depressive disorder: a selective review of recent MRI studies. *J. Affect. Disord.* **117**, 1–17 (2009).
- Wise, T. et al. Common and distinct patterns of grey-matter volume alteration in major depression and bipolar disorder: evidence from voxel-based meta-analysis. *Mol. Psychiatry* **22**, 1455–1463 (2017).
- Goodkind, M. et al. Identification of a common neurobiological substrate for mental illness. *JAMA Psychiatry* **72**, 305–315 (2015).
- Gong, Q. et al. A transdiagnostic neuroanatomical signature of psychiatric illness. *Neuropsychopharmacology* **44**, 869–875 (2019).
- Thompson, P. M. et al. ENIGMA and global neuroscience: a decade of large-scale studies of the brain in health and disease across more than 40 countries. *Transl. Psychiatry* **10**, 100 (2020).
- van Rooij, D. et al. Cortical and subcortical brain morphometry differences between patients with autism spectrum disorder and healthy individuals across the lifespan: results from the ENIGMA ASD working group. *Am. J. Psychiatry* **175**, 359–369 (2018).

27. Hoogman, M. et al. Brain imaging of the cortex in ADHD: a coordinated analysis of large-scale clinical and population-based samples. *Am. J. Psychiatry* **176**, 531–542 (2019).
28. Schmaal, L. et al. Cortical abnormalities in adults and adolescents with major depression based on brain scans from 20 cohorts worldwide in the ENIGMA Major Depressive Disorder Working Group. *Mol. Psychiatry* **22**, 900–909 (2017).
29. Boedhoe, P. S. W. et al. Cortical abnormalities associated with pediatric and adult obsessive-compulsive disorder: findings from the ENIGMA obsessive-compulsive disorder working group. *Am. J. Psychiatry* **175**, 453–462 (2018).
30. Hibar, D. P. et al. Cortical abnormalities in bipolar disorder: an MRI analysis of 6503 individuals from the ENIGMA Bipolar Disorder Working Group. *Mol. Psychiatry* **23**, 932–942 (2018).
31. van Erp, T. G. M. et al. Cortical brain abnormalities in 4474 individuals with schizophrenia and 5098 control subjects via the enhancing neuro imaging genetics through meta analysis (ENIGMA) consortium. *Biol. Psychiatry* **84**, 644–654 (2018).
32. Larivière, S. et al. The ENIGMA Toolbox: multiscale neural contextualization of multisite neuroimaging datasets. *Nat. Methods* <https://doi.org/10.1038/s41592-021-01186-4> (2021).
33. Patel, Y. et al. Virtual histology of cortical thickness and shared neurobiology in 6 psychiatric disorders. *JAMA Psychiatry* **78**, 47 (2021).
34. Opel, N. et al. Cross-disorder analysis of brain structural abnormalities in six major psychiatric disorders: a secondary analysis of mega- and meta-analytical findings from the ENIGMA consortium. *Biol. Psychiatry* **88**, 678–686 (2020).
35. García-Cabezas, M. Á., Zikopoulos, B. & Barbas, H. The structural model: a theory linking connections, plasticity, pathology, development and evolution of the cerebral cortex. *Brain Struct. Funct.* **224**, 985–1008 (2019).
36. Amunts, K., Mohlberg, H., Bludau, S. & Zilles, K. Julich-Brain: A 3D probabilistic atlas of the human brain's cytoarchitecture. *Science* **369**, 988–992 (2020).
37. Wagstyl, K. et al. Mapping cortical laminar structure in the 3D bigbrain. *Cereb. Cortex* **28**, 2551–2562 (2018).
38. Paquola, C. et al. Microstructural and functional gradients are increasingly dissociated in transmodal cortices. *PLoS Biol.* **17**, e3000284 (2019).
39. Schleicher, A., Morosan, P., Amunts, K. & Zilles, K. Quantitative architectural analysis: a new approach to cortical mapping. *J. Autism Dev. Disord.* **39**, 1568–1581 (2009).
40. Margulies, D. S. et al. Situating the default-mode network along a principal gradient of macroscale cortical organization. *Proc. Natl Acad. Sci. USA* **113**, 12574–12579 (2016).
41. Mesulam, M. M. From sensation to cognition. *Brain* **121**, 1013–1052 (1998).
42. Sanides, F. *Die Architektonik des Menschlichen Stirnhirns* Vol. 98 (Springer, 1962).
43. Huntenburg, J. M., Bazin, P. L. & Margulies, D. S. Large-scale gradients in human cortical organization. *Trends Cogn. Sci.* **22**, 21–31 (2018).
44. Goulas, A. et al. The natural axis of transmitter receptor distribution in the human cerebral cortex. *Proc. Natl Acad. Sci. USA* **118**, e2020574118 (2021).
45. Vogel, J. W. et al. A molecular gradient along the longitudinal axis of the human hippocampus informs large-scale behavioral systems. *Nat. Commun.* **11**, 960 (2020).
46. Valk, S. L. et al. Shaping brain structure: genetic and phylogenetic axes of macroscale organization of cortical thickness. *Sci. Adv.* **6**, 1–15 (2020).
47. Vainik, U. et al. Heritability of cortical morphology reflects a sensory-fugal plasticity gradient. Preprint at *bioRxiv* (2020).
48. Hong, S.-J. et al. Atypical functional connectome hierarchy in autism. *Nat. Commun.* **10**, 1022 (2019).
49. Lowe, A. J. et al. Targeting age-related differences in brain and cognition with multimodal imaging and connectome topography profiling. *Hum. Brain Mapp.* **40**, 5213–5230 (2019).
50. Bijsterbosch, J. et al. Challenges and future directions for representations of functional brain organization. *Nat. Neurosci.* **23**, 1484–1495 (2020).
51. Haak, K. V., Marquand, A. F. & Beckmann, C. F. Connectopic mapping with resting-state fMRI. *Neuroimage* **170**, 83–94 (2018).
52. Mars, R. B. et al. Whole brain comparative anatomy using connectivity blueprints. *Elife* **7**, 1–15 (2018).
53. Dukart, J. et al. JuSpace: a tool for spatial correlation analyses of magnetic resonance imaging data with nuclear imaging derived neurotransmitter maps. *Hum. Brain Mapp.* **42**, 555–566 (2021).
54. Alakurtti, K. et al. Long-term test-retest reliability of striatal and extrastriatal dopamine D2/3 receptor binding: study with [¹¹C]raclopride and high-resolution PET. *J. Cereb. Blood Flow. Metab.* **35**, 1199–1205 (2015).
55. Hesse, S. et al. Central noradrenaline transporter availability in highly obese, non-depressed individuals. *Eur. J. Nucl. Med. Mol. Imaging* **44**, 1056–1064 (2017).
56. Kaller, S. et al. Test-retest measurements of dopamine D1-type receptors using simultaneous PET/MRI imaging. *Eur. J. Nucl. Med. Mol. Imaging* **44**, 1025–1032 (2017).
57. Savli, M. et al. Normative database of the serotonergic system in healthy subjects using multi-tracer PET. *Neuroimage* **63**, 447–459 (2012).
58. García Gómez, F., Huertas, I., Lojo Ramírez, J. & García Solís, D. Elaboración de una plantilla de SPM para la normalización de imágenes de PET con 18F-DOPA. *Imagen Diagn.óstica* **9**, 23–25 (2018).
59. Dukart, J. et al. Cerebral blood flow predicts differential neurotransmitter activity. *Sci. Rep.* **8**, 4074 (2018).
60. Tost, H., Alam, T. & Meyer-Lindenberg, A. Dopamine and psychosis: theory, pathomechanisms and intermediate phenotypes. *Neurosci. Biobehav. Rev.* **34**, 689–700 (2010).
61. Howes, O. D. & Kapur, S. The dopamine hypothesis of schizophrenia: Version III - The final common pathway. *Schizophr. Bull.* **35**, 549–562 (2009).
62. Wainwright, S. R. & Galea, L. A. M. The neural plasticity theory of depression: assessing the roles of adult neurogenesis and psc-nam within the hippocampus. *Neural Plast.* **2013**, 805497 (2013).
63. Mulinari, S. Monoamine theories of depression: historical impact on biomedical research. *J. Hist. Neurosci.* **21**, 366–392 (2012).
64. Conio, B. et al. Opposite effects of dopamine and serotonin on resting-state networks: review and implications for psychiatric disorders. *Mol. Psychiatry* **25**, 82–93 (2020).
65. Van Essen, D. C. et al. The WU-Minn Human Connectome Project: an overview. *Neuroimage* **80**, 62–79 (2013).
66. Amunts, K. et al. BigBrain: an ultrahigh-resolution 3D human brain model. *Science* **340**, 1472–1475 (2013).
67. Desikan, R. S. et al. An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. *Neuroimage* **31**, 968–980 (2006).
68. Buckner, R. L., Krienen, F. M., Castellanos, A., Diaz, J. C. & Yeo, B. T. T. The organization of the human cerebellum estimated by intrinsic functional connectivity. *J. Neurophysiol.* **106**, 2322–45 (2011).
69. Xu, T. et al. Cross-species functional alignment reveals evolutionary hierarchy within the connectome. *Neuroimage* **223**, 117346 (2020).
70. Paquola, C. et al. The BigBrainWarp toolbox for integration of BigBrain 3D histology with multimodal neuroimaging. *Elife* **10**, e70119 (2021).
71. Paquola, C. et al. Shifts in myeloarchitecture characterise adolescent development of cortical gradients. *Elife* **8**, e50482 (2019).
72. Paquola, C. et al. Convergence of cortical types and functional motifs in the human mesiotemporal lobe. *Elife* **9**, e60673 (2020).
73. Royer, J. et al. Myeloarchitecture gradients in the human insula: histological underpinnings and association to intrinsic functional connectivity. *Neuroimage* **216**, 116859 (2020).
74. Goulas, A., Zilles, K. & Hilgetag, C. C. Cortical gradients and laminar projections in mammals. *Trends Neurosci.* **41**, 775–788 (2018).
75. Paquola, C. et al. A multi-scale cortical wiring space links cellular architecture and functional dynamics in the human brain. *PLoS Biol.* **18**, e3000979 (2020).
76. Tibshirani, R. Regression shrinkage and selection via the Lasso. *J. R. Stat. Soc. Ser. B* **58**, 267–288 (1996).
77. Tenenbaum, J. B., Silva, V. de & Langford, J. C. A global geometric framework for nonlinear dimensionality reduction. *Science* **290**, 2319–2323 (2000).
78. Varma, S. & Simon, R. Bias in error estimation when using cross-validation for model selection. *BMC Bioinforma.* **7**, 91 (2006).
79. Cawley, G. C. & Talbot, N. L. C. On over-fitting in model selection and subsequent selection bias in performance evaluation. *J. Mach. Learn. Res.* **11**, 2079–2107 (2010).
80. Parvande, S., Yeh, H. W., Paulus, M. P. & McKinney, B. A. Consensus features nested cross-validation. *Bioinformatics* **36**, 3093–3098 (2020).
81. Bedford, S. A. et al. Large-scale analyses of the relationship between sex, age and intelligence quotient heterogeneity and cortical morphometry in autism spectrum disorder. *Mol. Psychiatry* **25**, 614–628 (2020).
82. Sha, Z., Wager, T. D., Mechelli, A. & He, Y. Common dysfunction of large-scale neurocognitive networks across psychiatric disorders. *Biol. Psychiatry* **85**, 379–388 (2019).
83. Schwarz, K. et al. Transdiagnostic prediction of affective, cognitive, and social function through brain reward anticipation in schizophrenia, bipolar disorder, major depression, and autism spectrum diagnoses. *Schizophr. Bull.* **46**, 592–602 (2020).
84. Park, B. et al. An expanding manifold in transmodal regions characterizes adolescent reconfiguration of structural connectome organization. *Elife* **10**, e64694 (2021).
85. Edmonson, C., Ziats, M. N. & Rennert, O. M. Altered glial marker expression in autistic post-mortem prefrontal cortex and cerebellum. *Mol. Autism* **5**, 1–9 (2014).
86. Cotter, D., Mackay, D., Landau, S., Kerwin, R. & Everall, I. Reduced glial cell density and neuronal size in the anterior cingulate cortex in major depressive disorder. *Arch. Gen. Psychiatry* **58**, 545–553 (2001).

87. Rajkowska, G. et al. Morphometric evidence for neuronal and glial prefrontal cell pathology in major depression. *Biol. Psychiatry* **45**, 1085–1098 (1999).
88. de Oliveira, K. C. et al. Layer-specific reduced neuronal density in the orbitofrontal cortex of older adults with obsessive-compulsive disorder. *Brain Struct. Funct.* **224**, 191–203 (2019).
89. Rajkowska, G., Halaris, A. & Selemon, L. D. Reductions in neuronal and glial density characterize the dorsolateral prefrontal cortex in bipolar disorder. *Biol. Psychiatry* **49**, 741–752 (2001).
90. Benes, F. M., Davidson, J. & Bird, E. D. Quantitative cytoarchitectural studies of the cerebral cortex of schizophrenics. *Arch. Gen. Psychiatry* **43**, 31 (1986).
91. Iritani, S. What happens in the brain of schizophrenia patients?: an investigation from the viewpoint of neuropathology. *Nagoya J. Med. Sci.* **76**, 11–28 (2014).
92. Selemon, L. D., Rajkowska, G. & Goldman-Rakic, P. S. Abnormally high neuronal density in the schizophrenic cortex. *Arch. Gen. Psychiatry* **52**, 805 (1995).
93. Forrest, M. P., Parnell, E. & Penzes, P. Dendritic structural plasticity and neuropsychiatric disease. *Nat. Rev. Neurosci.* **19**, 215–234 (2018).
94. Fromer, M. et al. De novo mutations in schizophrenia implicate synaptic networks. *Nature* **506**, 179–184 (2014).
95. Marshall, C. R. et al. Contribution of copy number variants to schizophrenia from a genome-wide study of 41,321 subjects. *Nat. Genet.* **49**, 27–35 (2017).
96. Ripke, S. et al. Biological insights from 108 schizophrenia-associated genetic loci. *Nature* **511**, 421–427 (2014).
97. Sanders, S. J. et al. De novo mutations revealed by whole-exome sequencing are strongly associated with autism. *Nature* **485**, 237–241 (2012).
98. Yuen, R. K. C. et al. Whole genome sequencing resource identifies 18 new candidate genes for autism spectrum disorder. *Nat. Neurosci.* **20**, 602–611 (2017).
99. Sklar, P. et al. Large-scale genome-wide association analysis of bipolar disorder identifies a new susceptibility locus near ODZ4. *Nat. Genet.* **43**, 977–985 (2011).
100. Benowitz, L. I. & Routtenberg, A. GAP-43: an intrinsic determinant of neuronal development and plasticity. *Trends Neurosci.* **20**, 84–91 (1997).
101. Dinse, J. et al. A cytoarchitecture-driven myelin model reveals area-specific signatures in human primary and secondary areas using ultra-high resolution in-vivo brain MRI. *Neuroimage* **114**, 71–87 (2015).
102. Thom, M. Review: hippocampal sclerosis in epilepsy: a neuropathology review. *Neuropathol. Appl. Neurobiol.* **40**, 520–543 (2014).
103. Bernhardt, B. C. et al. The spectrum of structural and functional imaging abnormalities in temporal lobe epilepsy. *Ann. Neurol.* **80**, 142–153 (2016).
104. Blümcke, I. et al. International consensus classification of hippocampal sclerosis in temporal lobe epilepsy: a Task Force report from the ILAE Commission on Diagnostic Methods. *Epilepsia* **54**, 1315–1329 (2013).
105. Adler, S. et al. Topographic principles of cortical fluid-attenuated inversion recovery signal in temporal lobe epilepsy. *Epilepsia* **59**, 627–635 (2018).
106. Bernhardt, B. C. et al. Preferential susceptibility of limbic cortices to microstructural damage in temporal lobe epilepsy: a quantitative T1 mapping study. *Neuroimage* **182**, 294–303 (2018).
107. La Joie, R. et al. Intrinsic connectivity identifies the hippocampus as a main crossroad between alzheimer's and semantic dementia-targeted networks. *Neuron* **81**, 1417–1428 (2014).
108. Shafiei, G. et al. Spatial patterning of tissue volume loss in schizophrenia reflects brain network architecture. *Biol. Psychiatry* **87**, 727–735 (2020).
109. Mayberg, H. S. et al. Deep brain stimulation for treatment-resistant depression. *Neuron* **45**, 651–660 (2005).
110. Zikopoulos, B., García-Cabezas, M. Á. & Barbas, H. Parallel trends in cortical gray and white matter architecture and connections in primates allow fine study of pathways in humans and reveal network disruptions in autism. *PLoS Biol.* **16**, e2004559 (2018).
111. Zikopoulos, B. & Barbas, H. Changes in prefrontal axons may disrupt the network in autism. *J. Neurosci.* **30**, 14595–14609 (2010).
112. van Berlekom, A. B. et al. Synapse pathology in schizophrenia: a meta-analysis of postsynaptic elements in postmortem brain studies. *Schizophr. Bull.* **46**, 374–386 (2020).
113. Mizutani, R. et al. Three-dimensional alteration of neurites in schizophrenia. *Transl. Psychiatry* **9**, 85 (2019).
114. Toker, L., Mancarci, B. O., Tripathy, S. & Pavlidis, P. Transcriptomic evidence for alterations in astrocytes and parvalbumin interneurons in subjects with bipolar disorder and schizophrenia. *Biol. Psychiatry* **84**, 787–796 (2018).
115. Duman, R. S., Sanacora, G. & Krystal, J. H. Altered connectivity in depression: GABA and glutamate neurotransmitter deficits and reversal by novel treatments. *Neuron* **102**, 75–90 (2019).
116. Valk, S. L. et al. Genetic and phylogenetic uncoupling of structure and function in human transmodal cortex. *Nat. Commun.* **13**, 2341 (2022).
117. Dong, D. et al. Compression of cerebellar functional gradients in schizophrenia. *Schizophr. Bull.* **46**, 1282–1295 (2020).
118. Meng, Y. et al. Systematically disrupted functional gradient of the cortical connectome in generalized epilepsy: initial discovery and independent sample replication. *Neuroimage* **230**, 117831 (2021).
119. Tian, Y., Zalesky, A., Bousman, C., Everall, I. & Pantelis, C. Insula functional connectivity in schizophrenia: subregions, gradients, and symptoms. *Biol. Psychiatry Cogn. Neurosci. Neuroimaging* **4**, 399–408 (2019).
120. Park, B. et al. Differences in subcortico-cortical interactions identified from connectome and microcircuit models in autism. *Nat. Commun.* **12**, 2225 (2021).
121. Marazziti, D. Understanding the role of serotonin in psychiatric diseases. *F1000Res.* **6**, 180 (2017).
122. Lin, S. H., Lee, L. T. & Yang, Y. K. Serotonin and mental disorders: a concise review on molecular neuroimaging evidence. *Clin. Psychopharmacol. Neurosci.* **12**, 196–202 (2014).
123. Lopez-Ibor, J. J. The involvement of serotonin in psychiatric disorders and behaviour. *Br. J. Psychiatry* **153**, 26–39 (1988).
124. Volkow, N. D. et al. Evaluating dopamine reward pathway in ADHD: Clinical Implications. *JAMA* **302**, 1084–1091 (2009).
125. Pine, A., Shiner, T., Seymour, B. & Dolan, R. J. Dopamine, time, and impulsivity in humans. *J. Neurosci.* **30**, 8888–8896 (2010).
126. Cropley, V. L., Fujita, M., Innis, R. B. & Nathan, P. J. Molecular imaging of the dopaminergic system and its association with human cognitive function. *Biol. Psychiatry* **59**, 898–907 (2006).
127. Ashok, A. H. et al. The dopamine hypothesis of bipolar affective disorder: the state of the art and implications for treatment. *Mol. Psychiatry* **22**, 666–679 (2017).
128. Dunlop, B. W. & Nemeroff, C. B. The role of dopamine in the pathophysiology of depression. *Arch. Gen. Psychiatry* **64**, 327–337 (2007).
129. Zhuang, X. et al. Altered emotional states in knockout mice lacking 5-HT1A or 5-HT1B receptors. *Neuropsychopharmacology* **21**, 52S–60S (1999).
130. Yohn, C. N., Gergues, M. M. & Samuels, B. A. The role of 5-HT receptors in depression. *Mol. Brain* **10**, 28 (2017).
131. Seamans, J. K., Gorelova, N., Durstewitz, D. & Yang, C. R. Bidirectional dopamine modulation of GABAergic inhibition in prefrontal cortical pyramidal neurons. *J. Neurosci.* **21**, 3628–3638 (2001).
132. Gorelova, N., Seamans, J. K. & Yang, C. R. Mechanisms of dopamine activation of fast-spiking interneurons that exert inhibition in rat prefrontal cortex. *J. Neurophysiol.* **88**, 3150–3166 (2002).
133. Trantham-Davidson, H., Neely, L. C., Lavin, A. & Seamans, J. K. Mechanisms underlying differential D1 versus D2 dopamine receptor regulation of inhibition in prefrontal cortex. *J. Neurosci.* **24**, 10652–10659 (2004).
134. Parsey, R. V. et al. Altered serotonin 1A binding in major depression: a [carbonyl-C-11] WAY100635 positron emission tomography study. *Biol. Psychiatry* **59**, 106–113 (2006).
135. Lemond, S. et al. Impaired repression at a 5-hydroxytryptamine 1A receptor gene polymorphism associated with major depression and suicide. *J. Neurosci.* **23**, 8788–8799 (2003).
136. Salatino-Oliveira, A., Rohde, L. A. & Hutz, M. H. The dopamine transporter role in psychiatric phenotypes. *Am. J. Med. Genet. B Neuropsychiatr. Genet.* **177**, 211–231 (2018).
137. Fernández-Jaén, A. et al. Cortical thickness differences in the prefrontal cortex in children and adolescents with ADHD in relation to dopamine transporter (DAT1) genotype. *Psychiatry Res. Neuroimaging* **233**, 409–417 (2015).
138. Zilles, K. & Palomero-Gallagher, N. Multiple transmitter receptors in regions and layers of the human cerebral cortex. *Front. Neuroanat.* **11**, 78 (2017).
139. Khan, A. F. et al. Personalized brain models identify neurotransmitter receptor changes in Alzheimer's disease. *Brain* <https://doi.org/10.1093/brain/awab375> (2021).
140. Ross, S. & Peselow, E. Co-occurring psychotic and addictive disorders: neurobiology and diagnosis. *Clin. Psychopharmacol.* **35**, 235–243 (2012).
141. Al-Asadi, A. M., Klein, B. & Meyer, D. Multiple comorbidities of 21 psychological disorders and relationships with psychosocial variables: a study of the online assessment and diagnostic system within a web-based population. *J. Med. Internet Res.* **17**, e55 (2015).
142. Plana-Ripoll, O. et al. Exploring comorbidity within mental disorders among a Danish national population. *JAMA Psychiatry* **76**, 259–270 (2019).
143. Simonoff, E. et al. Psychiatric disorders in children with autism spectrum disorders: prevalence, comorbidity, and associated factors in a population-derived sample. *J. Am. Acad. Child Adolesc. Psychiatry* **47**, 921–929 (2008).
144. Hegarty, C. E. et al. ADHD comorbidity can matter when assessing cortical thickness abnormalities in patients with bipolar disorder. *Bipolar Disord.* **14**, 843–855 (2012).

145. Makris, N. et al. Further understanding of the comorbidity between attention-deficit/hyperactivity disorder and bipolar disorder in adults: an MRI study of cortical thickness. *Psychiatry Res. Neuroimaging* **202**, 1–11 (2012).
146. Canu, E. et al. Brain structural abnormalities in patients with major depression with or without generalized anxiety disorder comorbidity. *J. Neurol.* **262**, 1255–1265 (2015).
147. Schaefer, A. et al. Local-global parcellation of the human cerebral cortex from intrinsic functional connectivity MRI. *Cereb. Cortex* **28**, 3095–3114 (2018).
148. Glasser, M. F. et al. A multi-modal parcellation of human cerebral cortex. *Nature* **536**, 171–178 (2016).
149. Keller, S. S. & Roberts, N. Voxel-based morphometry of temporal lobe epilepsy: an introduction and review of the literature. *Epilepsia* **49**, 741–757 (2008).
150. Kebets, V. et al. Somatosensory-motor dysconnectivity spans multiple transdiagnostic dimensions of psychopathology. *Biol. Psychiatry* **86**, 779–791 (2019).
151. Fischl, B., Sereno, M. I., Tootell, R. B. H. & Dale, A. M. High-resolution inter-subject averaging and a surface-based coordinate system. *Hum. Brain Mapp.* **8**, 272–284 (1999).
152. Dale, A. M., Fischl, B. & Sereno, M. I. Cortical surface-based analysis: I. Segmentation and surface reconstruction. *Neuroimage* **9**, 179–194 (1999).
153. Fischl, B., Sereno, M. I. & Dale, A. M. Cortical surface-based analysis: II. Inflation, flattening, and a surface-based coordinate system. *Neuroimage* **9**, 195–207 (1999).
154. Larivière, S. et al. Network-based atrophy modelling in the common epilepsies: a worldwide ENIGMA study. *Sci. Adv.* **6**, eabc6457 (2020).
155. Glasser, M. F. et al. The minimal preprocessing pipelines for the Human Connectome Project. *Neuroimage* **80**, 105–124 (2013).
156. Jenkinson, M., Beckmann, C. F., Behrens, T. E. J., Woolrich, M. W. & Smith, S. M. Fsl. *Neuroimage* **62**, 782–790 (2012).
157. Fischl, B. FreeSurfer. *Neuroimage* **62**, 774–781 (2012).
158. Van Essen, D. C., Glasser, M. F., Dierker, D. L., Harwell, J. & Coalson, T. Parcellations and hemispheric asymmetries of human cerebral cortex analyzed on surface-based atlases. *Cereb. cortex* **22**, 2241–2262 (2012).
159. Glasser, M. F., Goyal, M. S., Preuss, T. M., Raichle, M. E. & Van Essen, D. C. Trends and properties of human cerebral cortex: correlations with cortical myelin content. *Neuroimage* **93**, 165–175 (2014).
160. Glasser, M. F. & Essen, Van. D. C. Mapping human cortical areas in vivo based on myelin content as revealed by T1- and T2-weighted MRI. *J. Neurosci.* **31**, 11597–11616 (2011).
161. Salimi-Khorshidi, G. et al. Automatic denoising of functional MRI data: combining independent component analysis and hierarchical fusion of classifiers. *Neuroimage* **90**, 449–468 (2014).
162. Thompson, W. H. & Fransson, P. On stabilizing the variance of dynamic functional brain connectivity time series. *Brain Connect.* **6**, 735–746 (2016).
163. Wold, S., Esbensen, K. & Geladi, P. Principal component analysis. *Chemom. Intell. Lab. Syst.* **2**, 37–52 (1987).
164. Alexander-Bloch, A. F. et al. On testing for spatial correspondence between maps of human brain structure and function. *Neuroimage* **178**, 540–551 (2018).
165. Benjamini, Y. & Hochberg, Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J. R. Stat. Soc.* **57**, 289–300 (1995).
166. Vos de Wael, R. et al. BrainSpace: a toolbox for the analysis of macroscopic gradients in neuroimaging and connectomics datasets. *Commun. Biol.* **3**, 103 (2020).
167. Coifman, R. R. & Lafon, S. Diffusion maps. *Appl. Comput. Harmon. Anal.* **21**, 5–30 (2006).
168. von Luxburg, U. A tutorial on spectral clustering. *Stat. Comput.* **17**, 395–416 (2007).
169. Park, B. et al. Signal diffusion along connectome gradients and inter-hub routing differentially contribute to dynamic human brain function. *Neuroimage* **224**, 117429 (2021).
170. Tsamardinos, I., Rakhshani, A. & Lagani, V. Performance-estimation properties of cross-validation-based protocols with simultaneous hyper-parameter optimization. *Int. J. Artif. Intell. Tools* **24**, 1540023 (2015).

Acknowledgements

The authors would like to express their gratitude to the open science initiatives that made this work possible: (i) The ENIGMA-Epilepsy consortium and, in particular, the ASD, ADHD, MDD, OCD, BD, and SCZ working groups (<http://enigma.ini.usc.edu/ongoing/>) and (ii) The Human Connectome Project (Principal Investigators: David Van Essen and Kamil Ugurbil; 1U54MH091657) funded by the 16 NIH Institutes and Centers that support the NIH Blueprint for Neuroscience Research; and by the McDonnell Center for Systems Neuroscience at Washington University. Bo-yong Park was funded by the National Research Foundation of Korea (NRF-2021R1F1A1052303 and NRF-2022R1A5A7033499), Institute for Information and Communications Technology Planning and Evaluation (IITP) funded by the Korean Government (MSIT) (No. 2022-0-00448, Deep Total Recall: Continual Learning for Human-Like Recall of Artificial Neural Networks; No. 2020-0-01389, Artificial Intelligence Convergence Research Center (Inha University); No. RS-2022-00155915, Artificial Intelligence Convergence Innovation Human Resources Development (Inha University); No. 2021-0-02068, Artificial Intelligence Innovation Hub), and Institute for Basic Science (IBS-R015-D1). Valeria Kebets was funded by the Quebec Autism Research Training Fellowship of the Transforming Autism Care Consortium (TACC). Sara Larivière was funded by the Canadian Institutes of Health Research (CIHR). Meike D. Hettwer was funded by the Max Planck Society and the German Federal Ministry of Education and Research (BMBF). Martine Hoogman is supported by a personal Veni grant from the Netherlands Organization for Scientific Research (NWO, grant number 91619115). Lianne Schmaal was funded by NHMRC Career Development Fellowship (1140764), National Institute of Mental Health of the National Institutes of Health R01MH117601. Ole A. Andreassen was funded by Research Council of Norway (#223273, #283798), KG Jebsen Stiftelsen. Dan J Stein is funded by the SA MRC. Dr. Matthias Kirschner acknowledges support from the National Bank Fellowship (McGill University) and the Swiss National Foundation (P2SKP3_178175). Jessica Turner was funded by the National Institutes of Health (NIH 5R01MH121246). Paul Thompson and Sophia Thomopoulos are funded in part by the US National Institutes of Health, under grants R01MH116147, R01MH111671, R01AG058854, R01MH123163, and U54 EB020403. Boris C. Bernhardt acknowledges research support from the National Science and Engineering Research Council of Canada (NSERC Discovery-1304413), the CIHR (FDN-154298, PJT), SickKids Foundation (NI17-039), Azrieli Center for Autism Research (ACAR-TACC), BrainCanada, Fonds de la Recherche en Santé (FRQ-S), and the Tier-2 Canada Research Chairs program. Alan Evans, Boris C. Bernhardt, and Casey Paquola were funded in part by Helmholtz Association's Initiative and Networking Fund under the Helmholtz International Lab grant agreement InterLabs-0015, and the Canada First Research Excellence Fund (CFREF Competition 2, 2015–2016) awarded to the Healthy Brains, Healthy Lives initiative at McGill University, through the Helmholtz International BigBrain Analytics and Learning Laboratory (HIBALL).

Author contributions

B.-y.P., M.K., and B.C.B. designed the study, conducted the experiments, researched the data, and wrote the manuscript. V.K., S.L., M.D.H., and C.P. aided the experiments. D.v.R., J.B., B.F., M.H., L.S., D.J.V., O.A.v.d.H., D.J.S., O.A.A., C.R.K.C., J.A.T., T.G.M.v.E., A.C.E., A.D., S.I.T., P.M.T., and S.L.V. reviewed the manuscript. B.-y.P. and B.C.B. assume all the responsibilities for the accuracy of all content.

Competing interests

Ole A. Andreassen received speaker's honorarium from Lundbeck and Sunovion, Consultant to HealthLytix. Paul M. Thompson received grant support from Biogen, Inc., and consulting payments from Kairos Venture Capital for work unrelated to the current manuscript. The remaining authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s42003-022-03963-z>.

Correspondence and requests for materials should be addressed to Bo-yong Park or Boris C. Bernhardt.

Peer review information *Communications Biology* thanks the anonymous reviewers for their contribution to the peer review of this work. Primary Handling Editor: George Inglis.

Reprints and permission information is available at <http://www.nature.com/reprints>

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2022

¹McConnell Brain Imaging Centre, Montreal Neurological Institute and Hospital, McGill University, Montreal, QC, Canada. ²Department of Data Science, Inha University, Incheon, Republic of Korea. ³Center for Neuroscience Imaging Research, Institute for Basic Science, Suwon, Republic of Korea. ⁴Forschungszentrum Jülich, Jülich, Germany. ⁵Max Planck Institute for Cognitive and Brain Sciences, Leipzig, Germany. ⁶Max Planck School of Cognition, Leipzig, Germany. ⁷Institute of Systems Neuroscience, Heinrich Heine University Düsseldorf, Düsseldorf, Germany. ⁸Institute of Neuroscience and Medicine (INM-1), Forschungszentrum Jülich, Jülich, Germany. ⁹Center for Cognitive Neuroimaging, Donders Institute for Brain, Cognition and Behavior, Radboud University, Nijmegen, The Netherlands. ¹⁰Department of Psychiatry, University of Groningen, University Medical Center Groningen, Groningen, The Netherlands. ¹¹Donders Institute for Brain, Cognition and Behaviour, Radboud University, Nijmegen, The Netherlands. ¹²Department of Human Genetics and Department of Psychiatry, Radboud university medical center, Nijmegen, The Netherlands. ¹³Centre for Youth Mental Health, The University of Melbourne, Melbourne, VIC, Australia. ¹⁴Orygen, Parkville, VIC, Australia. ¹⁵Departments of Psychiatry and Anatomy and Neuroscience, Amsterdam University Medical Centers, Vrije Universiteit Amsterdam, Amsterdam, The Netherlands. ¹⁶SA Medical Research Council Unit on Risk & Resilience in Mental Disorders, Dept of Psychiatry and Mental Health, University of Cape Town, Cape Town, South Africa. ¹⁷NORMENT Centre, Division of Mental Health and Addiction, Oslo University Hospital & Institute of Clinical Medicine, University of Oslo, Oslo, Norway. ¹⁸KG Jebsen Centre for Neurodevelopment, Institute of Clinical Medicine, University of Oslo, Oslo, Norway. ¹⁹Imaging Genetics Center, Mark and Mary Stevens Neuroimaging and Informatics Institute, Keck School of Medicine, University of Southern California, Los Angeles, CA, USA. ²⁰Department of Psychology and Behavioral Health, The Ohio State University, Columbus, OH, USA. ²¹Tri-institutional Center for Translational Research in Neuroimaging and Data Science (TReNDS), Georgia State University, Georgia Institute of Technology and Emory University, Atlanta, GA, USA. ²²Neuroscience Institute, Georgia State University, Atlanta, GA, USA. ²³Clinical Translational Neuroscience Laboratory, Department of Psychiatry and Human Behavior, University of California Irvine, Irvine, CA, USA. ²⁴Center for the Neurobiology of Learning and Memory, University of California Irvine, Irvine, CA, USA. ²⁵Division of Adult Psychiatry, Department of Psychiatry, Geneva University Hospitals, Geneva, Switzerland. ²⁶These authors contributed equally: Matthias Kirschner, Boris C. Bernhardt. ✉email: boyong.park@inha.ac.kr; boris.bernhardt@mcgill.ca