



# Neurobehavioral correlates of obesity are largely heritable

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Recent molecular genetic studies have shown that the majority of genes associated with obesity are expressed in the central nervous system. Obesity has also been associated with neurobehavioral factors such as brain morphology, cognitive performance, and personality. Here, we tested whether these neurobehavioral factors were associated with the heritable variance in obesity measured by body mass index (BMI) in the Human Connectome Project ( $n = 895$  siblings). Phenotypically, cortical thickness findings supported the “right brain hypothesis” for obesity. Namely, increased BMI is associated with decreased cortical thickness in right frontal lobe and increased thickness in the left frontal lobe, notably in lateral prefrontal cortex. In addition, lower thickness and volume in entorhinal-parahippocampal structures and increased thickness in parietal-occipital structures in participants with higher BMI supported the role of visuospatial function in obesity. Brain morphometry results were supported by cognitive tests, which outlined a negative association between BMI and visuospatial function, verbal episodic memory, impulsivity, and cognitive flexibility. Personality–BMI correlations were inconsistent. We then aggregated the effects for each neurobehavioral factor for a behavioral genetics analysis and estimated each factor’s genetic overlap with BMI. Cognitive test scores and brain morphometry had 0.25–0.45 genetic correlations with BMI, and the phenotypic correlations with BMI were 77–89% explained by genetic factors. Neurobehavioral factors also had some genetic overlap with each other. In summary, obesity as measured by BMI has considerable genetic overlap with brain and cognitive measures. This supports the theory that obesity is inherited via brain function and may inform intervention strategies.

brain morphology | cortical thickness | cognition | twins | body mass index

Obesity is a widespread condition leading to increased mortality (1) and economic costs (2). Twin and family studies have shown that individual differences in obesity are largely explained by genetic variance (3). Gene enrichment patterns suggest that obesity-related genes are preferentially expressed in the brain (4). While it is unclear how these brain-expressed genes lead to obesity, several lines of research show that neural, cognitive, and personality differences have a role in vulnerability to obesity (5, 6). Here, we seek to test whether these neurobehavioral factors could explain the genetic variance in obesity.

In the personality literature, obesity is most often negatively associated with conscientiousness (self-discipline and orderliness) and positively with neuroticism (a tendency toward negative affect) (7). In the cognitive domain, tests capturing executive function, inhibition, and attentional control have a negative association with obesity (5–8). Neuroanatomically, obesity seems to have a negative association with the gray matter volume of prefrontal cortex and, to a lesser extent, the volume of parietal and temporal lobes, as measured by voxel-based morphometry (9). It has also been suggested that structural and functional asymmetry of the prefrontal cortex might underlie overeating and obesity (10). For genetic analysis, cortical thickness estimates of brain structure from magnetic resonance imaging (MRI) have been preferred over volumetric measures (11). However, to date,

reports of cortical thickness patterns associated with obesity have been inconsistent (12, 13). As a prerequisite to our goal of ascertaining the heritability of brain-based vulnerability to obesity, we sought to extend previous neurobehavioral findings in a large multifactor dataset from the Human Connectome Project (HCP). We also measured volumetric estimates of medial temporal lobe and subcortical structures, which have been implicated in appetitive control (e.g., ref. 14).

The main goal was to assess whether the aforementioned obesity–neurobehavioral associations are of genetic or environmental origin. Recent evidence from behavioral and molecular genetics suggests that there is considerable genetic overlap among obesity, cognitive test scores, and brain imaging findings (15–20). However, the evidence so far is not comprehensive across all neurobehavioral factors discussed. A recent paper assessed the heritability of obesity-associated regional brain volumes (21). However, the study did not analyze the heritability of the association between brain and obesity. The latter analysis is crucial for understanding whether brain anatomy and obesity could have a genetic overlap, which would suggest that the heritability of vulnerability to obesity is expressed in the brain.

In addition, we sought to estimate the genetic overlap between the different BMI-related neurobehavioral factors. Performance

## Significance

Obesity is a widespread heritable health condition. Evidence from psychology, cognitive neuroscience, and genetics has proposed links between obesity and the brain. The current study tested whether the heritable variance in body mass index (BMI) is explained by brain and behavioral factors in a large brain imaging cohort that included multiple related individuals. We found that the heritable variance in BMI had genetic correlations 0.25–0.45 with cognitive tests, cortical thickness, and regional brain volume. In particular, BMI was associated with frontal lobe asymmetry and differences in temporal-parietal perceptual systems. Further, we found genetic overlap between certain brain and behavioral factors. In summary, the genetic vulnerability to BMI is expressed in the brain. This may inform intervention strategies.

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on cognitive tests and personality must originate from the brain (e.g., ref. 22) and, therefore, personality and cognition could be expected to explain brain–morphometry associations with BMI (6). However, brain–behavior associations are far from certain (23), and even different measurement traditions in both behavior (personality and cognitive tests) and brain morphometry (cortical thickness or brain volume) are often conceptualized as providing independent sources of information (7, 11). Documenting the degree of genetic overlap between behavioral and brain measures would shed light on whether similar underlying processes lead to obesity’s associations with different neurobehavioral factors.

Taken together, the goal of the current analysis was to use a large multifactor dataset to analyze the heritability of the associations between obesity and brain/behavior. We further tested genetic overlap between the different neurobehavioral factors themselves.

**Results**

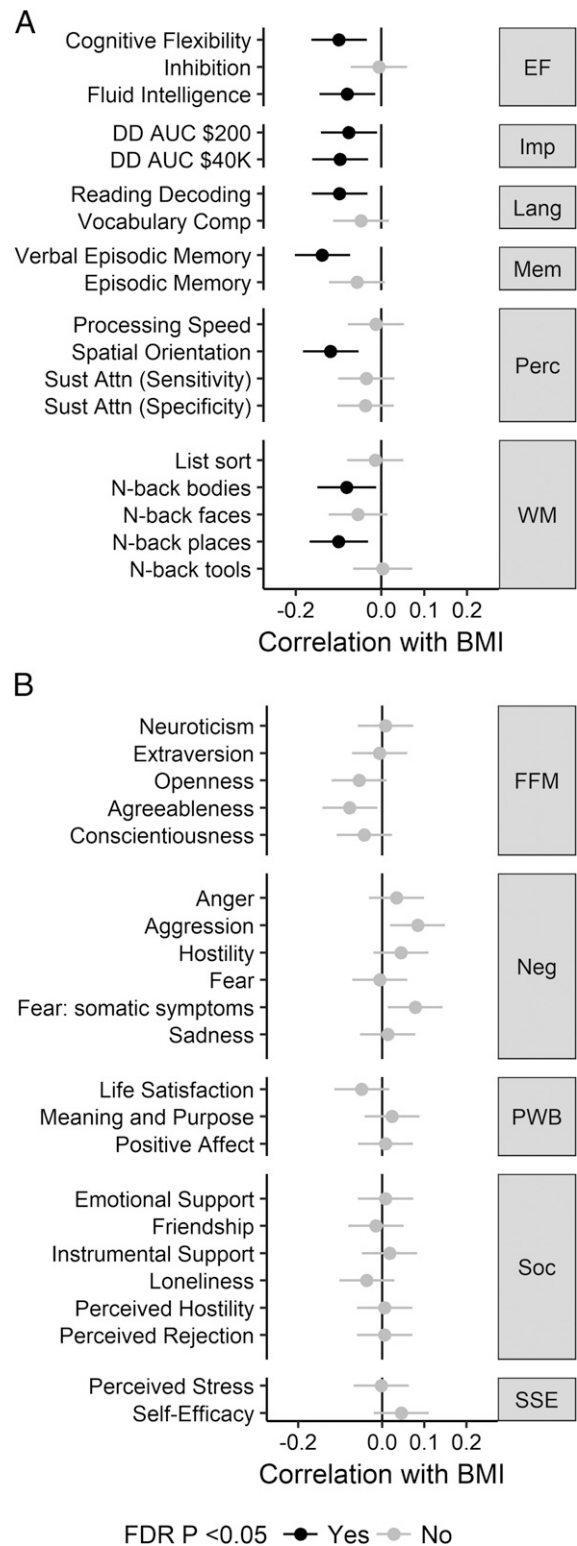
**Background.** We analyzed data from 895 participants from the Human Connectome Project S900 release (24), including 111 pairs of monozygotic twins and 188 pairs of dizygotic twins and siblings. Similar to many previous reports (3) we modeled BMI heritability with the AE model (A, additive genetics; E, unique environment), as opposed to the ACE model (C, common environment), as AE had the lowest Akaike Information Criterion ([Dataset S1, section 9](#)). BMI heritability was  $A = 71\%$  [95% CI: 61%; 78%], which is close to the published meta-analytic estimate ( $A = 75\%$ , ref. 3).

In all analyses below, we controlled for age, gender, race, ethnicity, handedness, and evidence of drug consumption on day of testing, which mostly associated with BMI ([SI Appendix, SI Results and Fig. S2](#)). When presenting and interpreting phenotypic associations, we controlled for family structure to avoid inflated effect sizes and SEs (e.g., ref. 25). The behavioral genetics analysis did not control for family structure, since this information is needed for modeling heritability. As socio-economic status (SES) is intertwined with cognitive test scores (26), personality (27), and brain morphometry (28), we also present phenotypic associations controlling for SES (education and income) in [SI Appendix, Supplementary Material](#). All in-text *P* values are provided without correcting for multiple comparisons. False discovery rate (FDR) correction was applied when screening for features within cognitive, personality, and brain factors (Figs. 1 and 2).

**Cognitive and Personality Factors.** BMI was negatively correlated with the following tests of executive function: cognitive flexibility, fluid intelligence, inability to delay gratification, reading abilities, and working memory. Intriguingly, the strongest effects were present for nonexecutive tasks measuring visuospatial ability and verbal memory (Fig. 1A). These tasks remained associated with BMI after controlling for SES; controlling for SES reduced the number of executive function tests involved with BMI to cognitive flexibility and inability to delay gratification ([SI Appendix, Fig. S3A, Left](#)). No personality test score correlated with BMI when FDR correction was applied (Fig. 1B).

**Brain Morphology.** Cortical thickness was estimated from each T1-weighted MRI using CIVET 2.0 software (29). Parcel-based analysis identified negative associations with BMI in right inferior lateral frontal cortex and bilateral entorhinal-parahippocampal cortex (Figs. 2A and 3A). Positive associations with BMI were found with the left superior frontal cortex, left inferior lateral frontal cortex, and bilateral parietal cortex parcels. Controlling for SES did not change these results ([SI Appendix, Fig. S4A, Left](#)). The frontal lobe asymmetry in the BMI association (thinner on the right, thicker on the left) mostly involved the inferior lateral prefrontal areas, such as inferior frontal gyrus.

Regional brain volumes were measured for estimation of brain morphology–obesity associations in brain structures not covered by the CIVET cortical thickness algorithm. Medial temporal



**Fig. 1.** Associations between BMI and cognitive test scores (A) and personality traits (B). Error bars represent 95% confidence intervals. See [Dataset S1, section 1](#) for explanation of cognitive tests. Numerical values are reported in [Dataset S1, section 2](#). EF, executive function; FDR, false discovery rate; FFM, Five-Factor Model; Imp, (lack of) impulsivity; Lang, language; Mem, memory; Neg, negative affect; Perc, perception; PWB, psychological well-being; Soc, social relationships; SSE, stress and self efficacy; WM, working memory.

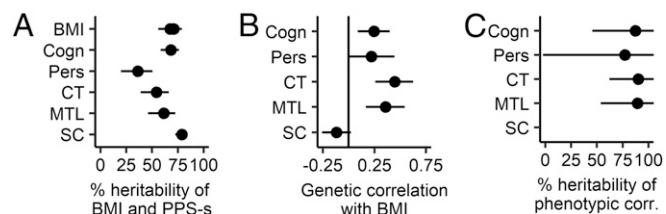




for cortical thickness ( $r = 0.26$ ,  $P < 0.001$ ,  $n = 591$ ) and medial temporal brain volume ( $r = 0.23$ ,  $P < 0.001$ ,  $n = 594$ ). There was no association between BMI and subcortical brain volume ( $r = -0.05$ ,  $P = 0.169$ ,  $n = 828$ ). To test the generalizability of the PPS approach, we used weights obtained from the full S900 release (SI Appendix, Fig. S3, Right and SI Appendix, Fig. S4, Right) to test PPS–BMI correlation among the unseen additional participants in the S1200 release (referred to as S1200n,  $n = 236$ ). Cortical thickness PPS had essentially unchanged effect size when correlated with BMI in S1200n (SI Appendix, SI Results and Fig. S7). At the same time, cognitive and personality PPS-s were less stable (SI Appendix, SI Results and Fig. S7), likely because the smaller effect sizes of individual features need larger training datasets to reduce inaccuracies, or that the true PPS–BMI effect size was too small to be found just within the S1200n sample.

**Heritability.** Bivariate heritability was similarly conducted with the AE model, since the main goal was to explain variance in BMI, for which AE was the best model. All PPS-s were found to be highly heritable, with the A component explaining 36–79% of the variance (Fig. 4A and Dataset S1, section 10). Significant genetic correlations ( $r_g$ ) were found between BMI and cognitive test scores [ $r_g = 0.25$  ( $P = 0.002$ ), cortical thickness ( $r_g = 0.45$ ,  $P < 0.001$ ), and medial temporal brain volume ( $r_g = 0.36$ ,  $P < 0.001$ ) (Fig. 4B and Dataset S1, section 11). The personality PPS genetic correlation with BMI was not significant ( $r_g = 0.22$ ,  $P = 0.052$ ). Molecular evidence relying on linkage disequilibrium score regression has reported effects of similar magnitude between higher cognitive test scores and BMI ( $r_g = -0.22$ , ref. 15;  $r_g = -0.18$ , ref. 18). Environmental correlations (i.e., correlations between environmental variances) were small and not significant (Dataset S1, section 11). As expected from high heritability of the traits and high genetic correlations, the phenotypic BMI–PPS correlations described in the previous sections were 77–89% explained by genetic factors (Fig. 4C and Dataset S1, section 10).

The results broadly replicated when repeating the analysis with just the top features within a PPS, suggesting that PPS-based findings summarize the effects of the underlying individual features (SI Appendix, Fig. S8). We further replicated the heritability patterns in a separate analysis focused only on the additional participants from the S1200 HCP release (SI Appendix, Fig. S9). Additionally, controlling for SES (education and income) did not change the results for brain-based PPS-s. However, the estimates for cognitive test scores and personality became lower and not significant in the S900 release (SI Appendix, Fig. S10). However, the same estimates were significant in the combined sample S900+S1200, suggesting that the effects of cognition and personality were reduced but not eliminated when controlling for SES.



**Fig. 4.** Heritability analysis of the association between PPS and BMI. (A) Heritability of each trait. BMI has multiple estimates, since it was entered into a bivariate analysis with each PPS separately. (B) Genetic correlations between BMI and each PPS. The genetic correlations are positive, because the PPS-s are designed to positively predict BMI. (C) Heritability of the significant phenotypic correlation between BMI and PPS. Horizontal lines depict 95% confidence intervals. Cogn, PPS of cognitive tests; corr, correlation; CT, PPS of cortical thickness; MTL, PPS of medial temporal lobe volume; Pers, PPS of personality tests; SC, PPS of subcortical structure volumes.

**Genetic Overlap Between Neurobehavioral Factors.** Phenotypically, certain PPS-s had small but significant intercorrelations (SI Appendix, Fig. S11, upper triangle). After FDR correction, we were able to find two genetic correlations between PPS-s of cognition and cortical thickness ( $r_g = 0.35$ ), as well as cognition and personality ( $r_g = 0.33$ , SI Appendix, Fig. S11, lower triangle). Taken together, while the neurobehavioral factors have mostly independent effects on BMI, cognitive test scores may have a small genetic overlap with brain structure and personality.

## Discussion

Cortical thickness, medial temporal lobe volume, and cognitive measures all had covariation with BMI, and their effect on BMI was almost entirely heritable. Similarly, we found genetic correlations between obesity risk scores of cognition, cortical thickness, and personality. Together, our results from a large sample support the role of brain and psychological constructs in explaining genetic variance in BMI.

BMI correlated with increased cortical thickness in the left prefrontal cortex and decreased thickness in the right prefrontal cortex, supporting the “right brain” hypothesis for obesity (10). The effect was most prominent in the inferior frontal gyrus (Figs. 24 and 34). Only preliminary support for the right brain hypothesis has been previously available (13). Right prefrontal cortex has been implicated in inhibitory control (22) and possibly bodily awareness (10). Many neuromodulation interventions (e.g., transcranial magnetic stimulation) aimed at increasing self-regulation capacity often target right prefrontal cortex. However, effects have also been demonstrated in studies targeting left prefrontal cortex (31).

Cortical thickness results also highlighted the role of temporo-parietal perceptual structures in obesity. Namely, BMI was associated with bilaterally decreased thickness of the parahippocampal and entorhinal cortices, and with mostly right-lateralized increased thickness of parietal and occipital lobes. Volumetric results within the medial temporal lobe supported the role of entorhinal cortex and also suggested that obesity is positively associated with the volume of left amygdala. Emergence of the effects of the right parietal structures together with right prefrontal structures hint at the role of the ventral frontoparietal network, thought to be especially important for detection of behaviorally relevant visual stimuli (32). The parahippocampal and entorhinal cortex are associated with episodic memory and context mediation (33). Similarly, the hippocampus has been associated with the modulation of food cue reactivity by homeostatic and contextual information, and hippocampal dysfunction is postulated to promote weight gain in the western diet environment (34). The amygdala is implicated in emotional and appetitive responses to sensory stimuli, including food cues (35).

Integrating these findings, one could envision a model where obesity is associated with a certain cognitive profile (36). The model starts with a hyperactive visual attention system attributing heightened salience to food stimuli, implicating the ventral visual stream and amygdala. These signals are then less optimally tied into relevant context by the parahippocampal and entorhinal structures, and less well moderated (or filtered) by the prefrontal executive system. This could result in consummatory behavior driven by the presence of appetitive food signals, which are ubiquitous in our obesogenic environment. An impaired response inhibition and salience attribution model of obesity has been suggested based on the functional neuroimaging literature. Namely, functional MRI studies have consistently identified obesity to associate with heightened salience response to food cues, coupled with reduced activation in prefrontal and executive systems involved in self-regulation and top-down attentional control (e.g., ref. 35). A similar conclusion emerged from a recent resting state network analysis of the HCP data (37), in which obesity was associated with alterations in perceptual networks and decreased activity of default mode and central executive networks.



This brain morphology-derived model has some support from cognitive tests. The role of prefrontal executive control is outlined by our finding of a negative association between BMI and scores on several executive control tasks. Surprisingly, there was no effect of motor inhibition as measured by the Flanker inhibitory task. A relation between obesity and reduced motor inhibition, while often mentioned, has been inconsistent even across meta-analyses (7, 8). However, we found a relationship between decisional impulsivity, measured by delay discounting, and BMI, replicating previous literature (6, 7, 18). While controlling for education reduced the number of executive tasks associated with BMI, the overall pattern remained the same, suggesting that education level is a proxy for certain executive function abilities.

Intriguingly, BMI was found to be negatively associated with spatial orientation and verbal episodic memory. These tasks tap into the key functions associated with entorhinal and parahippocampal regions implicated in our study (33). Therefore, both cognitive and brain morphology features propose that the increased salience of food stimuli could be facilitated by dysregulated context representation in obesity.

Regarding personality, we were unable to find any questionnaire-specific effects, notably with respect to neuroticism and conscientiousness, both often thought to be associated with obesity (5–7). There are potential explanations for this negative finding. First, the meta-analytical association between various personality tests and BMI is small ( $r = 0.05$ , ref. 7), for which we might have been underpowered after  $P$  value correction. Second, controlling for family structure likely further reduced the effect sizes (25). Third, the personality–obesity associations tend to pertain to more specific facets and nuances than broad personality traits (38), therefore, further analysis with more detailed and eating-specific personality measures is needed in larger samples.

All of the associations discussed here were largely due to shared genetic variance between neurobehavioral factors and BMI. This is in accordance with recent molecular genetics evidence that 75% of obesity-related genes express preferentially in the brain (4). Similarly, the genetic correlation between cognition and BMI uncovered in our sample is at the same magnitude as molecular estimates of associations between more specific cognitive measures and BMI (15, 18). The current evidence further supports the brain–gene association with obesity vulnerability.

A possible explanation of the genetic correlations is pleiotropy—the existence of a common set of genes that influence variance in both obesity and brain function. It is possible that people with a higher genetic risk for obesity also have genetic propensity for the brain and cognitive patterns outlined here. It is also likely that interventions could influence both obesity and brain function. For instance, regular exercise can support weight management (39), reduce the heritability of obesity (40), and improve cognitive health (41).

However, our results could also support a causal relationship—that the genetic correlation is due to a persistent effect of heritable brain factors on overeating and, hence, BMI. For instance, we could hypothesize that the heritable obesity-related cognitive profile promotes overeating when high-calorie food is available. As high-calorie food is abundant and inexpensive, the cognitive risk profile could lead to repeated overeating, providing an opportunity for genetic obesity proneness to express. Such longitudinal environmental effects of a trait need not to be large, they just have to be consistent (ref. 42, see discussion in ref. 43). Of course, a reverse scenario is also possible—obesity leads to alterations in cortical morphology due to the consequences of cardiometabolic complications, including low-grade chronic inflammation, hypertension, and vascular disease (reviewed in refs. 9 and 44). However, we find this hypothesis less plausible as global brain atrophy due to metabolic syndrome is mostly seen in older participants, whereas the current sample had a mean age of

29. Young adults often experience “healthy or transitional obesity,” where clinical inflammation levels (45) and other cardiometabolic comorbidities have not yet developed (46).

We found neurobehavioral PPS-s to have occasional phenotypic and genetic correlations with each other. Here, it is hard to argue against pleiotropy playing a role. While one could reasonably expect that at least part of the variation in cognitive performance would be shaped by brain morphometry (22), it is also the case that engaging in education leads to improvement in cognitive test scores (26) and might also lead to changes in cortical thickness (47). The small genetic overlap between cognition, cortical thickness, and personality can probably be explained by common pleiotropic roots. At the same time, integrating morphometry and cognitive findings is difficult with this dataset.

From a practical point of view, our work suggests that evidence from psychology and neuroscience can be used to design intervention strategies for people with higher genetic risk for obesity. One way would be modifying neurobehavioral factors, e.g., with cognitive training, to improve people’s ability to resist the obesogenic environment (31, 36). Another path could be changing the immediate environment to be less obesogenic (e.g., ref. 48) so that individual differences in neurobehavioral factors would be less likely to manifest. In any case, obesity interventions should not focus solely on energy content, but also acknowledge the certain neurobehavioral profile that obesity is genetically intertwined with.

The current analysis has limitations. Due to the cross-sectional nature of the dataset, causality between neurobehavioral factors and BMI is only suggestive—longitudinal designs would enable better insight into the causal associations between brain morphology, psychological measures, and BMI or weight gain. BMI is a crude proxy for actual eating behaviors or health status. In addition, there were more normal-weight than obese participants. However, the 25% obesity rate in this sample is close to the published obesity rate of the state of Missouri (31.7%) and the United States (36.5%, ref. 49). Also, we expect that BMI itself and the neurobehavioral mechanisms behind it are continuum processes, therefore all variation in the range from normal weight to obesity is likely helping to uncover underlying associations. While the measurement of cognition and personality was exhaustive, it lacked some common behavioral tasks like the stop-signal task, or common questionnaires measuring self-control, impulsivity, and eating-specific behaviors, that have been previously associated with body weight (5, 6). Particularly, the common eating-specific behaviors such as uncontrolled eating (50) are likely better candidates for explaining brain morphology–BMI associations as they are more directly related to the hypothesized underlying behavior.

One has to be careful in translating individual differences in cortical thickness in normal populations to underlying neural mechanisms. Diverse biological processes have been suggested to influence MRI-based cortical thickness measures, ranging from synaptic density to apparent thinning due to synaptic pruning and myelination (summarized in refs. 51 and 52). A definitive model of the underlying mechanism that links normal variations in cortical thickness to differences in brain function cannot be given, as cortical thickness has not been mapped with both MRI and histology in humans (52). Still, the associations between cortical thickness and BMI in one sample were able to predict BMI in a new separate sample, suggesting that the pattern is robust. Our conceptual interpretation of the significance of cortical thickness patterns has support from measures of both brain structure and cognitive function.

Relying on PPS-s prevented us from analyzing detailed interactions between cortical thickness and cognitive function and their genetic overlap with each other. However, given the relatively small associations between PPS-s and the number of candidate measures that could be expected to interact with one another, we believe it would have been hard to find an association that would have survived multiple testing corrections.

Future, focused, hypothesis-driven studies have to further elucidate the neurobehavioral mechanisms behind obesity proneness.

In summary, the current analysis provides comprehensive evidence that the obesity-related differences in brain structure and cognitive tests are largely due to shared genetic factors. Genetic factors also explain occasional overlap between neurobehavioral factors. We hope that increasingly larger longitudinal datasets and dedicated studies will help to outline more specific neurobehavioral mechanisms that confer vulnerability to obesity and provide a basis for designing informed interventions.

## Methods

Data were provided by the Human Connectome Project (24). Certain people were excluded due to missing data or not fulfilling typical criteria. Exclusion details,

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demographics, and family structure are summarized in *SI Appendix, SI Methods and Table S1*. Software pipelines for obtaining features of cortical thickness and brain volume are described in *SI Appendix, SI Methods*. Analysis scripts to reproduce results presented are available at: <https://osf.io/hxt7u>.

*SI Appendix, Fig. S1* provides a schematic pipeline for data analysis. Details of each data analysis step are outlined in *SI Appendix, SI Methods*. We describe how PPS weights are obtained through cross-validation and how the weights generalize to a separate dataset (S1200n). We further describe the main principles of twin and sibling-based heritability analysis and replication of these findings using individual features instead of PPS-s and replication in a separate dataset (S1200n). Finally, the software and packages used are listed.

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