



## King's Research Portal

DOI:

[10.1016/j.biopsych.2020.06.003](https://doi.org/10.1016/j.biopsych.2020.06.003)

*Document Version*

Publisher's PDF, also known as Version of record

[Link to publication record in King's Research Portal](#)

*Citation for published version (APA):*

Zhang, Z., Robinson, L., Jia, T., Quinlan, E., Tay, N., Chu, C., Barker, E., Banaschewski, T., Barker, G., Bokde, A. L. W., Flor, H., Grigis, A., Garavan, H., Gowland, P., Heinz, A., Ittermann, B., Martinot, J-L., Stringaris, A., Penttilä, J., ... Desrivieres, S. (2020). Development of disordered eating behaviors and comorbid depressive symptoms in adolescence: neural and psychopathological predictors. *Biological psychiatry*, 0(0), 1-10.  
<https://doi.org/10.1016/j.biopsych.2020.06.003>

### **Citing this paper**

Please note that where the full-text provided on King's Research Portal is the Author Accepted Manuscript or Post-Print version this may differ from the final Published version. If citing, it is advised that you check and use the publisher's definitive version for pagination, volume/issue, and date of publication details. And where the final published version is provided on the Research Portal, if citing you are again advised to check the publisher's website for any subsequent corrections.

### **General rights**

Copyright and moral rights for the publications made accessible in the Research Portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognize and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the Research Portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the Research Portal

### **Take down policy**

If you believe that this document breaches copyright please contact [librarypure@kcl.ac.uk](mailto:librarypure@kcl.ac.uk) providing details, and we will remove access to the work immediately and investigate your claim.

1 **Development of disordered eating behaviors and comorbid depressive symptoms in**  
2 **adolescence: neural and psychopathological predictors**

3 Zuo Zhang, PhD<sup>1\*</sup>, Lauren Robinson, PhD<sup>2\*</sup>, Tianye Jia, PhD<sup>1,3,4</sup>, Erin Burke Quinlan, PhD<sup>1</sup>, Nicole  
4 Tay, PhD<sup>1</sup>, Congying Chu, PhD<sup>1</sup>, Edward D. Barker, PhD<sup>1,5</sup>, Tobias Banaschewski, MD, PhD<sup>6</sup>,  
5 Gareth J. Barker, PhD<sup>7</sup>, Arun L.W. Bokde, PhD<sup>8</sup>, Herta Flor, PhD<sup>9,10</sup>, Antoine Grigis, PhD<sup>11</sup>, Hugh  
6 Garavan, PhD<sup>12</sup>, Penny Gowland, PhD<sup>13</sup>, Andreas Heinz, MD, PhD<sup>14</sup>, Bernd Ittermann, PhD<sup>15</sup>,  
7 Jean-Luc Martinot, MD, PhD<sup>16</sup>, Argyris Stringaris, MD, PhD<sup>17</sup>, Jani Penttilä, MD, PhD<sup>18</sup>, Betteke  
8 van Noort, PhD<sup>19</sup>, Yvonne Grimmer, MD<sup>6</sup>, Marie-Laure Paillère Martinot, MD, PhD<sup>20</sup>, Corinna  
9 Isensee, PhD<sup>21</sup>, Andreas Becker, PhD<sup>21</sup>, Frauke Nees, PhD<sup>6,9,27</sup>, Dimitri Papadopoulos Orfanos,  
10 PhD<sup>11</sup>, Tomáš Paus, MD, PhD<sup>22</sup>, Luise Poustka, MD<sup>21</sup>, Sarah Hohmann, MD<sup>6</sup>, Juliane H. Fröhner,  
11 MSc<sup>23</sup>, Michael N. Smolka, MD<sup>23</sup>, Henrik Walter, MD, PhD<sup>14</sup>, Robert Whelan, PhD<sup>24</sup>, Gunter  
12 Schumann, MD<sup>1,25</sup>, Ulrike Schmidt, MD, PhD<sup>2,26</sup>, Sylvane Desrivières, PhD<sup>1</sup>

13  
14 1 Centre for Population Neuroscience and Precision Medicine (PONS), Institute of Psychiatry,  
15 Psychology & Neuroscience, Social, Genetic and Developmental Psychiatry Centre, King's  
16 College London, De Crespigny Park, London, SE5 8AF, United Kingdom; 2 Section of Eating  
17 Disorders, Department of Psychological Medicine, King's College London, London, United  
18 Kingdom; 3 Institute of Science and Technology for Brain-Inspired Intelligence, Fudan  
19 University, Shanghai, China; 4 Ministry of Education-Key Laboratory of Computational  
20 Neuroscience and Brain-Inspired Intelligence, Fudan University, Shanghai, China; 5

1 Developmental Psychopathology Lab, Department of Psychology, King's College London,  
2 London, United Kingdom; 6 Department of Child and Adolescent Psychiatry and  
3 Psychotherapy, Central Institute of Mental Health, Medical Faculty Mannheim, Heidelberg  
4 University, Square J5, 68159 Mannheim, Germany; 7 Department of Neuroimaging, Institute  
5 of Psychiatry, Psychology & Neuroscience, King's College London, United Kingdom; 8 Discipline  
6 of Psychiatry, School of Medicine and Trinity College Institute of Neuroscience, Trinity College  
7 Dublin, Dublin, Ireland; 9 Department of Cognitive and Clinical Neuroscience, Central Institute  
8 of Mental Health, Medical Faculty Mannheim, Heidelberg University, Square J5, Mannheim,  
9 Germany; 10 Department of Psychology, School of Social Sciences, University of Mannheim,  
10 68131 Mannheim, Germany; 11 NeuroSpin, CEA, Université Paris-Saclay, F-91191 Gif-sur-  
11 Yvette, France; 12 Departments of Psychiatry and Psychology, University of Vermont, 05405  
12 Burlington, Vermont, USA; 13 Sir Peter Mansfield Imaging Centre, School of Physics and  
13 Astronomy, University of Nottingham, University Park, Nottingham, United Kingdom; 14  
14 Charité - Universitätsmedizin Berlin, corporate member of Freie Universität Berlin, Humboldt-  
15 Universität zu Berlin, and Berlin Institute of Health, Department of Psychiatry and  
16 Psychotherapy, Campus Charite Mitte, Chariteplatz 1, Berlin, Germany; 15 Physikalisch-  
17 Technische Bundesanstalt (PTB), Braunschweig and Berlin, Germany; 16 INSERM U A10  
18 "Developmental Trajectories & Psychiatry"; Université Paris-Saclay, Ecole Normale Supérieure  
19 Paris-Saclay, CNRS, Centre Borelli; Gif-sur-Yvette, France; 17 National Institute of Mental  
20 Health / NIH, 15K North Drive, Bethesda MD, 20892, USA; 18 Department of Social and Health  
21 Care, Psychosocial Services Adolescent Outpatient Clinic Kauppakatu 14, Lahti, Finland; 19 MSB

1 Medical School Berlin, Calandrellistr. 1-9, 12247 Berlin, Germany; 20 Institut National de la  
2 Santé et de la Recherche Médicale, INSERM Unit 1000 “Neuroimaging & Psychiatry”, Université  
3 Paris Saclay, Université Paris Descartes; Sorbonne Université; and AP-HP, Department of Child  
4 and Adolescent Psychiatry, Pitié-Salpêtrière Hospital, Paris, France; 21 Department of Child and  
5 Adolescent Psychiatry and Psychotherapy, University Medical Centre Göttingen, von-Siebold-  
6 Str. 5, 37075, Göttingen, Germany; 22 Bloorview Research Institute, Holland Bloorview Kids  
7 Rehabilitation Hospital and Departments of Psychology and Psychiatry, University of Toronto,  
8 Toronto, Ontario, M6A 2E1, Canada; 23 Department of Psychiatry and Neuroimaging Center,  
9 Technische Universität Dresden, Dresden, Germany; 24 School of Psychology and Global Brain  
10 Health Institute, Trinity College Dublin, Ireland; 25 Department of Psychiatry and  
11 Psychotherapy, Campus Charite Mitte, Humboldt University, Berlin, Germany and Institute for  
12 Science and Technology of Brain-inspired Intelligence (ISTBI), Fudan University, Shanghai,  
13 China; 26 The Eating Disorders Service, Maudsley Hospital, South London and Maudsley NHS  
14 Foundation Trust, London, UK; 27 Institute of Medical Psychology and Medical Sociology,  
15 University Medical Center Schleswig Holstein, Kiel University, Kiel, Germany.

16

17 Address correspondence to Dr Sylvane Desrivières, Social, Genetic and Developmental  
18 Psychiatry Centre, Institute of Psychiatry, Psychology & Neuroscience, King’s College London,  
19 De Crespigny Park, London SE5 8AF, United Kingdom ([sylvane.desrivieres@kcl.ac.uk](mailto:sylvane.desrivieres@kcl.ac.uk)).

20

21 \*These authors contributed equally to this work.

1

2 **Running title:** Predictors of disordered eating and depression

3

4 **Keywords:** eating disorders, depression, attention deficit hyperactivity disorder, conduct  
5 disorder, biomarkers, grey matter volume

6

7

1 **Abstract**

2 **BACKGROUND:**

3 Eating disorders are common in adolescence, devastating and strongly comorbid with other  
4 psychiatric disorders. Yet, little is known about their etiology to develop effective preventive  
5 measures.

6 **METHODS:**

7 Longitudinal assessments of disordered eating behaviors (DEBs; binge-eating, purging and  
8 dieting) and comorbid psychopathology were measured in 1,386 adolescents from the  
9 IMAGEN study. Development of DEBs and associated mental health problems were  
10 investigated by comparing participants who reported symptoms at ages 16 or 19, but not at  
11 age 14 to asymptomatic controls. Voxel-based morphometry and psychopathological  
12 differences at age 14 were investigated to identify risk factors for the development of DEBs  
13 and associated mental health problems.

14 **RESULTS:**

15 DEBs and depressive symptoms developed together. Emotional and behavioral problems,  
16 including symptoms of attention deficit hyperactivity disorder (ADHD) and conduct disorder  
17 (CD), predated their development. Alterations in fronto-striatal brain areas also predated the  
18 development of DEBs and depressive symptoms. Specifically, development of binge-eating was  
19 predicted by higher grey matter volumes in the right putamen/globus pallidus at age 14.  
20 Conversely, development of purging and depressive symptoms was predicted by lower

1 volumes in the medial orbitofrontal, dorsomedial and dorsolateral prefrontal cortices. Lower  
2 grey matter volumes in the orbitofrontal and anterior cingulate cortices mediated the  
3 relationship between ADHD and CD symptoms and future purging and depressive symptoms.

4 **CONCLUSIONS:**

5 These findings suggest that alterations in frontal brain circuits are part of the shared etiology  
6 between eating disorders, ADHD, CD and depression and highlight the importance of a  
7 transdiagnostic approach to treating these conditions.

8

## 1 Introduction

2 Eating disorders (EDs), including anorexia nervosa (AN), bulimia nervosa (BN) and binge eating  
3 disorder (BED), are severe psychiatric disorders that affect up to 15% of young women and 3%  
4 of young men (1). The peak age of onset is from mid adolescence into emerging adulthood  
5 (age 15 to 19), i.e. at a developmentally sensitive time (2). EDs are characterized by disordered  
6 eating behaviors (DEBs), including dietary restriction, binge-eating and purging. Varying  
7 combinations of these DEBs occur in different EDs and across the weight spectrum from  
8 severely underweight to obese. Crucially, subclinical DEBs whose prevalence is particularly high  
9 (14%-22%) (3, 4) in children and adolescents predict development of full-syndrome EDs in later  
10 life (5, 6). Thus, identifying causal risk factors of DEBs and understanding their development is  
11 key to identifying high-risk groups and developing prevention strategies.

12  
13 Comorbid disorders are common in EDs. These include mood, anxiety and substance use  
14 disorders, which are common across all EDs (7) and impulse-control disorders like Attention  
15 Deficit Hyperactivity Disorder (ADHD), oppositional-defiant and conduct disorder (CD), which  
16 are prevalent in BN and BED (7, 8). Longitudinal studies have demonstrated that emotional and  
17 behavioral problems, including ADHD (9, 10) and CD symptoms (11) are risk factors of  
18 developing EDs. ADHD in childhood also confers higher risk of developing depression in  
19 adolescence/young adulthood (12, 13), suggesting that EDs and comorbid depression have  
20 shared psychopathological risk factors.



1  
2 EDs are widely accepted as brain-based disorders and neurobiological overlaps between EDs  
3 and addictions have been suggested (14). Neuroimaging studies have revealed structural and  
4 functional brain differences in ED patients compared with recovered patients and healthy  
5 controls. For example, meta-analyses combining adolescent and adult patients' data revealed  
6 reduced grey and white matter and increased cerebrospinal fluid (CSF) in AN patients  
7 compared with healthy individuals, such differences becoming less pronounced in recovered  
8 patients (15). However, most of these neuroimaging findings focus on AN or are based on small  
9 cross-sectional studies. The few studies comparing BN or BED patients to healthy individuals  
10 found regional volumetric or cortical thickness differences in the orbitofrontal cortex (OFC),  
11 insula, cingulate cortex and several other regions (16-20). Subcortical shape deformations (21)  
12 and white matter microstructure abnormality (22) were also found in BN patients and were  
13 associated with symptom severity.

14  
15 Critical questions remain as to whether any abnormalities displayed reflect predisposing risk  
16 factors or a consequence of prolonged eating disturbances. Should predisposing brain  
17 differences exist, it remains to be answered whether EDs and comorbid mental health  
18 problems have common neural underpinnings, and which neural mechanisms mediate the  
19 relationship between psychopathological risk factors and the development of EDs and  
20 comorbid mental health problems. We have recently demonstrated the value of longitudinal  
21 neuroimaging methods by showing that differences in neural responses to inhibitory control

1 can be detected 2 years before the onset of binge-eating or purging behaviors (23). Altogether,  
2 these findings suggest that advances in understanding and prevention of EDs are likely to  
3 benefit from an approach using dimensional and longitudinal assessments of DEBs, focusing  
4 on underlying neurobiological substrates (24).

5  
6 Building up on this, here we use the longitudinal design of IMAGEN, a large, prospective cohort  
7 of European adolescents, to investigate early psychopathological and neuroanatomical risk  
8 factors for the development of DEBs and comorbid mental health problems. Hypothesizing  
9 that structural brain alterations underlie shared risk for developing DEBs and comorbid mental  
10 health problems, we performed longitudinal and mediation analyses to lend insight into the  
11 etiology of EDs and evaluate underlying neural mechanisms through which psychopathological  
12 risk factors relate to the development of DEBs and comorbid mental health symptoms.

13

1 **Methods and Materials**

2 **Participants**

3 Questionnaire and neuroimaging data were obtained from IMAGEN – a longitudinal cohort  
4 acquired from 8 study sites in Europe. Each site received approval from their local research  
5 ethics committee. Written assent from the adolescents and written consent from the parents  
6 were obtained prior to participation. See (25) for details of the recruitment and assessment  
7 methods. In this study, we used questionnaire data acquired at ages 14, 16 and 19, and  
8 neuroimaging data acquired at age 14.

9

10 **Psychopathological assessments**

11 *Eating disorder symptoms:* Binge-eating, purging and dieting behaviors were assessed using  
12 the self-reports from the Development and Wellbeing Assessment (DAWBA) (26, 27). We used  
13 binary variables to indicate the presence or absence of binge-eating, purging and dieting  
14 symptoms at each age. A positive response to question 15, related to eating a large amount of  
15 food and losing control over eating was used to indicate the presence of binge-eating. A  
16 positive response to any of 3 questions (1c, 18f and 18g) related to self-induced vomiting or  
17 taking pills/medicines to lose weight was used to indicate the presence of purging behavior. A  
18 score of 3 (answer for ‘a lot’) for any of the 3 questions (question 18a, 18b and 18c) related to  
19 eating less at meals, skipping meals and fasting was used to indicate the presence of dieting

1 behavior. Dieting behaviors were also defined with more relaxed criteria (score  $\geq 2$ ), detailed  
2 in the Supplemental Information.

3

4 We categorized participants into groups who developed DEBs over time and those who  
5 remained asymptomatic based on each of the 3 DEBs. The “binge-eating developers” did not  
6 report binge-eating symptoms at age 14 but developed binge-eating symptoms at age 16 or  
7 19. The “non-bingers” did not report binge-eating at any of the 3 ages. In the same manner we  
8 defined “purging developers”, “non-purgers”, “dieting developers” and “non-dieters”. Besides  
9 these, we compared individuals who did not report any DEB at age 14 but reported at least  
10 one DEB at ages 16 or 19 (“any DEB developers”) to individuals without any DEB at any age  
11 (“non-DEB” group). We also defined groups based on Bartholdy et al. (2019) (23) who  
12 combined binge-eating and purging symptoms together. We report these as “binge-eating or  
13 purging (BoP)”, defining developers, maintainers and recoverers based on their longitudinal  
14 development (See Supplemental Methods).

15

16 As expected, these DEB developers and their asymptomatic controls showed significant sex  
17 differences (Supplemental Table S1). To obtain sex-balanced groups of cases and controls, each  
18 developer group and the corresponding asymptomatic group were matched for sex and  
19 acquisition site, by using the propensity score matching method implemented in the Matchit  
20 toolbox (28). Details are provided in the Supplemental Information.

21

1 **Emotional and behavioral problems:** The Strengths and Difficulties Questionnaire (SDQ) (29)  
2 was used to measure participants' emotional and behavioral symptoms. We used self-report  
3 scores at age 14 for the following 4 subscales: emotional symptoms, conduct problems (i.e.,  
4 CD symptoms), hyperactivity/inattention (i.e., ADHD symptoms) and peer relationship  
5 problems.

6  
7 **Mental health symptoms:** Computer-generated DAWBA diagnoses (DAWBA bands) (30)  
8 derived from the self-report questionnaire were used to measure the severity of  
9 psychopathology-related symptoms at the 3 ages. DAWBA bands comprised up to 6 levels  
10 (from 0 to 5) and indicated the probability of having a disorder (from <0.1% to >70% probability  
11 of DSM-IV based diagnoses). IMAGEN being a normative cohort, we focused on common  
12 mental health problems in our cohort, as defined by prevalence  $\geq 5\%$  at age 19 for participants  
13 scoring 3 and above (15%+ risk according to the DAWBA bands) for the disorder. DAWBA bands  
14 for depression and generalized anxiety disorder (GAD) passed this threshold and their  
15 associations with DEBs were investigated. We defined a group of "depression developers"  
16 (N=290) whose depression scores were below 3 at age 14, and above or equal to 3 at ages 16  
17 or 19. This was compared to controls (labeled as "non-depression", N=857) whose depression  
18 scores were below 3 across the 3 ages. Similarly, we defined "anxiety developers" (N=203) and  
19 "non-anxiety" (N=1107).

20

1 **Body mass index (BMI) and medication:** BMI (kg/m<sup>2</sup>) at age 14 was derived from height and  
2 weight measurements and transformed to age- and sex-adjusted z-scores based on the Centre  
3 for Disease Control and Prevention Growth Chart (31). A binary variable was created to indicate  
4 whether participants took any prescription medicine in the past 30 days based on the Timeline  
5 Followback Interview (32).

6

### 7 **Structural Magnetic resonance imaging (MRI) acquisition and processing**

8 MRI images were acquired with 3T MRI scanners (Siemens, Philips, General Electric) across all  
9 IMAGEN sites. A 3D magnetization scan based on the ADNI protocols  
10 (<http://adni.loni.usc.edu/methods/documents/mri-protocols/>) was used to acquire T1-  
11 weighted structural images. Quality control was performed through visual inspection to  
12 exclude images with movement artefacts, brace artefacts or field inhomogeneities. Voxel-  
13 based morphometry (VBM) analyses were conducted using the VBM8 toolbox  
14 (<http://www.neuro.uni-jena.de/vbm/>) in SPM8 (<https://www.fil.ion.ucl.ac.uk/spm/>) to obtain  
15 grey matter volumes (GMVs), as detailed in the Supplemental Information. Intracranial volume  
16 (ICV), estimated in VBM8 by summing up grey matter, white matter and CSF volumes, was used  
17 as a covariate in all the analyses involving GMVs (33).

18

### 19 **Statistical Analyses**

20 **DEB development and comorbid mental health symptoms.** To investigate if DEBs and mental  
21 health symptoms co-developed, we tested for associations between the development of DEBs

1 and the development of depression and anxiety symptoms while controlling for sex, acquisition  
2 site and depression or anxiety symptoms at age 14. These analyses were performed with the  
3 whole sample (i.e., without matching for sex and acquisition site).

4

5 ***Emotional and behavioral problems as predictors of DEBs.*** We investigated whether SDQ  
6 subscales at age 14 could predict DEB development using Firth logistic regression models (34)  
7 controlling for sex and acquisition site. Potentially confounding effects of BMI were tested by  
8 further controlling significant associations for BMI at age 14. The  $p$  values were corrected using  
9 the Holm-Bonferroni method. We also investigated whether emotional and behavioral  
10 problems that predicted DEBs also predicted the development of depression and anxiety  
11 symptoms, controlling for depression or anxiety symptoms at age 14, sex and acquisition site.

12

13 ***Voxel-wise GMV analyses.*** We investigated whether structural brain differences at age 14  
14 predated the development of DEBs. Generalized linear models (GLM) in SPM12 involved GMVs  
15 as the dependent variable, and the DEB group (developers vs. controls) as an independent  
16 variable. We tested whether shared anatomical differences underlay the development of DEBs  
17 and comorbid mental health problems as follows. First, we investigated voxel-wise GMV  
18 associations with development of depression and anxiety symptoms. We then tested if the  
19 associated brain regions overlapped with those associated with DEB development. We also  
20 investigated voxel-wise GMV associations with each of the 4 SDQ subscales at age 14. Control  
21 variables included in all analyses were sex, acquisition sites and total intracranial volumes (ICV).

1 Depression or anxiety symptoms at age 14 were also included as covariates in analyses  
2 involving development of depression and anxiety. The threshold for all the neuroimaging  
3 analyses was  $p < 0.001$  uncorrected at the voxel level, and  $p < 0.05$  corrected for family wise error  
4 (FWE) at the cluster level.

5  
6 **Mediation analyses.** We tested whether the brain regions associated with SDQ subscales  
7 mediated the associations between SDQ subscales at age 14 and development of DEBs and  
8 comorbid depression/anxiety symptoms. The brain regions associated with SDQ subscales  
9 were used as ROIs. Control variables included sex, acquisition site and ICV. The mental health  
10 symptom at age 14 was controlled for when investigating the development of  
11 depression/anxiety. The continuous variables were transformed to z-scores. Confidence  
12 intervals for the mediation effect were estimated from 5000 bootstrap samples by using the  
13 PROCESS macro (v3.2, <http://processmacro.org>) in SPSS (v25, IBM Corporation).

14



1 **Results**

2 A total of 1594 participants had non-missingness for the SDQ, DEB variables, BMI and  
3 neuroimaging data at age 14. Out of these, 1386 participants had DEB data at age 16 or 19 and  
4 were used to create DEB developer groups. Among the developers for binge-eating (n=115),  
5 purging (n=155), dieting (n=60), BoP (n=204) and any DEB (n = 138), between 60.9% and 79.1%  
6 were female. Taking this into account, case and control groups were matched based on sex  
7 and recruitment site (Supplemental Table S1 & Figure S1) for the analyses. Descriptive statistics  
8 for BMI and psychopathology scores are provided in Supplemental Table S2.

9

10 **DEBs co-develop with depression and anxiety symptoms**

11 We first established that DEBs co-developed with depression and anxiety by testing for  
12 associations between the development of DEBs and depression and anxiety symptoms.  
13 Development of DEBs was significantly associated with higher risk of developing depression  
14 and anxiety, after controlling for their levels at age 14 (Supplemental Table S3 and Figure S2).

15

16 **Emotional and behavioral problems predict the development of DEBs, depression and anxiety**  
17 **symptoms**

18 As emotional and behavioral problems are known predictors of eating disorders (9-11), we  
19 analyzed their associations with DEBs. Emotional problems at age 14 were significant  
20 predictors for the development of binge-eating (OR = 1.35,  $p = 4.4E-03$ ). ADHD symptoms and

1 CD symptoms at age 14 predicted the development of purging (OR = 1.35,  $p = 1.6E-03$  for ADHD  
2 symptoms; OR = 1.43,  $p = 8.5E-05$  for CD symptoms; Table 1) and BoP behaviors (OR = 1.28,  $p$   
3 = 4.7E-03 for ADHD symptoms; OR = 1.40,  $p = 1.0E-04$  for CD symptoms; Supplemental Table  
4 S4). CD symptoms at age 14 also predicted the maintenance of BoP (OR = 1.69,  $p = 5.8E-04$ ,  
5 Supplemental Table S4) and the development of “any DEB” (OR = 1.36,  $p = 3.9E-03$ ,  
6 Supplemental Table S5). These associations remained significant after controlling for BMI.  
7 Further analyses of the association with ADHD symptoms indicated that both the hyperactivity-  
8 impulsivity and inattention components contributed to the association with future purging  
9 behaviors (OR = 1.33,  $p = 2.5E-03$  and OR = 1.23,  $p = 0.028$  for hyperactivity-impulsivity and  
10 inattention, respectively). A significant predictor for dieting development was found only by  
11 using a more relaxed criterion for dieting (CD symptoms, OR = 1.27,  $p = 0.013$ , Supplemental  
12 Results).

13  
14 Emotional and behavioral problems at age 14 also predicted the development of depression  
15 and anxiety at later ages (Table 2). More specifically, emotional problems (OR = 1.56,  $p = 3.7E-$   
16 04) and ADHD symptoms (OR = 1.24,  $p = 0.045$ ) predicted the development of anxiety  
17 symptoms (controlling for depressive symptoms at ages 14, 16 and 19), while CD symptoms  
18 specifically predicted the development of depressive symptoms (OR = 1.22,  $p = 0.025$ ,  
19 controlling for anxiety symptoms at ages 14, 16 and 19).

20

1 **Structural brain differences at age 14 predate the development of DEBs and depressive**  
2 **symptoms**

3 Whole brain VBM analyses demonstrated that binge-eating developers had higher GMVs in a  
4 subcortical cluster comprising the right putamen and globus pallidus at age 14 (Figure 1A &  
5 Supplemental Table S6). Conversely, purging developers had lower GMVs in a cluster  
6 encompassing the medial OFC (mOFC), the gyrus rectus, the anterior and middle cingulate  
7 cortex (ACC and MCC), the left dorsomedial and dorsolateral prefrontal cortex (Figure 1B).  
8 Similarly, BoP developers had smaller GMVs at age 14 in the mOFC, gyrus rectus, ACC and MCC  
9 (Figure 1C). No significant GMV differences were associated with dieting developers, “any DEB”  
10 developers, BoP maintainers or BoP recoverers. Repeating analyses by controlling for BMI at  
11 age 14, or removing participants who took any medication in the past 30 days, or removing  
12 individuals reporting other DEBs at age 14 did not substantially change main associations (see  
13 Supplemental Results, Tables S6-S7 & Figures S3-S4). However, repeating the VBM analyses in  
14 girls only did not yield significant results on the whole brain level.

15  
16 We tested if brain regions associated with DEB development were also associated with  
17 developing symptoms of depression or generalized anxiety. Whole brain VBM analyses  
18 demonstrated that depression developers had lower GMVs in two clusters comprising the  
19 medial and left lateral OFC, ACC, MCC, the supplementary motor area, the dorsomedial PFC  
20 and left dorsolateral PFC (Figure 1D & Supplemental Table S8). These clusters overlapped with  
21 those associated with purging development in the ACC, MCC, mOFC, dorsomedial and left

1 dorsolateral PFC (Figure 1E). Similar overlaps were found for brain regions associated with  
2 development of BoP and depressive symptoms (Figure 1F). No significant results were found  
3 for the development of anxiety on the whole-brain level.

4

5 **Brain structure underlying CD and ADHD symptoms mediate the development of purging and**  
6 **comorbid depressive symptoms**

7 The analyses presented above suggested that neural correlates of SDQ traits – should they be  
8 detected – may serve as potential brain-based mediators for DEBs. To test this, we first  
9 analyzed associations between GMVs and DEB-associated SDQ measures at age 14, using the  
10 whole sample of 1594 participants. Higher CD symptoms were associated with lower grey  
11 matter volumes in the ACC, mOFC and the superior and middle frontal gyrus (Figure 2A &  
12 Supplemental Table S6). The ACC/mOFC region partly overlapped with those associated with  
13 purging development (262 voxels), BoP development (59 voxels) and depression development  
14 (74 voxels, Figure 2A). In contrast, higher ADHD symptoms were significantly associated with  
15 lower GMVs in a cluster encompassing the mOFC, gyrus rectus and anterior orbital gyrus  
16 (Figure 2B & Supplemental Table S6). These regions overlapped with those associated with  
17 purging development (461 voxels), BoP development (853 voxels) and depression  
18 development (148 voxels, Figure 2B). No significant GMV associations were observed for  
19 emotional problems.

20

1 Next, we investigated whether the GMV differences identified above mediated the relationship  
2 between CD or ADHD symptoms at age 14 and the development of purging, BoP and  
3 depression. Brain regions associated with CD symptoms (ACC/mOFC) and ADHD symptoms  
4 (labelled as OFC) were used as ROIs. Lower GMVs in the ACC/mOFC mediated the relationship  
5 between CD symptoms and the development of purging (indirect effect = 0.021, bootstrap 95%  
6 CI = 7.3E-04-0.056) and depression (indirect effect = 0.021, bootstrap 95% CI = 0.0023-0.050,  
7 Figure 3A). Likewise, lower GMVs in the OFC mediated the association between ADHD  
8 symptoms and the development of purging (indirect effect = 0.024, bootstrap 95% CI = 0.0016-  
9 0.058) and depression (indirect effect = 0.025, bootstrap 95% CI = 0.0039-0.053, Figure 3B).  
10 Similarly, these ROIs significantly mediated the effects of CD and ADHD symptoms on the  
11 development of BoP (Supplemental Figure S5). No significant mediation effects were found for  
12 the development of anxiety symptoms.

1 **Discussion**

2 Our longitudinal analyses, aimed at identifying early predictors of the development of DEBs  
3 and comorbid depression/anxiety symptoms in adolescence, identified shared neural  
4 substrates underlying the psychopathological risk for the development of DEBs and depression.  
5 We show that higher GMVs in the putamen and globus pallidus and emotional difficulties  
6 predate the development of binge-eating. We similarly demonstrated that lower GMVs in  
7 frontal and cingulate cortices, and ADHD and CD symptoms were associated with development  
8 of purging, BoP and depression. Importantly, the lower GMVs associated with ADHD and CD  
9 symptoms mediated the relationships between these symptoms and future purging, BoP and  
10 depression. These results support previous research showing high prevalence of ADHD, CD,  
11 depression and GAD in EDs, particularly in bulimia nervosa (7, 8), and extend our knowledge of  
12 the neural underpinning of these disorders.

13  
14 We have identified several neural substrates as early markers for the development of DEBs.  
15 Lower GMVs across the prefrontal cortex, including the mOFC, ACC, MCC, the dorsomedial  
16 and dorsolateral PFC are early indicators for the development of purging and BoP behaviors.  
17 Our mediation analyses also implicate the OFC and ACC/mOFC in the neural mechanisms  
18 underlying the associations between ADHD and CD symptoms and future purging and BoP  
19 behaviors. These results are consistent with previous suggestions of overlapping neural circuits  
20 involved in cognitive control and reward systems between ADHD and EDs (35). For example,

1 reduced GMVs in the OFC are consistently found in ADHD patients (36), which correlate with  
2 their functional impairments in emotion regulation, reward-related decision making and  
3 control of motivation (37). Shared neurobiological mechanisms (35) are further supported by  
4 observations of overlapping treatment responses to psychostimulant medications in ADHD and  
5 BN/BED patients (38, 39).

6

7 Youths with ADHD are at higher risk of developing depression (40), and their common  
8 neurobiological mechanisms have been suggested (41). For example, resting-state functional  
9 connectivity between the left OFC and left hippocampus is reduced in children with ADHD;  
10 furthermore, this connectivity is also negatively associated with depressive symptom severity in  
11 children with ADHD (41). However, research on neurobiological mechanisms of ADHD and  
12 depression has been largely separate, and the neural mediators between the two have been  
13 unclear until now. Our results highlight structural differences in the OFC to be a neural substrate  
14 that confers higher risk for depression in youths with greater ADHD symptoms. As the OFC plays  
15 important roles in emotion regulation (42), our result concurs with the finding that emotion  
16 regulation deficiencies mediate the association between ADHD and depressive symptoms (43, 44).  
17 By demonstrating that reduced GMVs in the OFC mediate the effect of ADHD symptoms on both  
18 future purging and depressive symptoms, our results further suggest that ADHD, depression and  
19 EDs may have a common neurobiological basis.

20

1 The observed reduced GMVs in the ACC/mOFC was consistent with recent neuroimaging  
2 findings in CDs (45). Both the ACC and mOFC are involved in reward-based decision making by  
3 encoding action-reward associations (46, 47) and also in top-down emotion regulation (48).  
4 Our results suggest that impairments in reward processing and/or emotion regulation may  
5 underlie the link between conduct problems and future purging behaviors and depression.

6  
7 Other brain regions associated with purging/BoP symptoms, i.e., the dorso-lateral, dorso-  
8 medial PFC and ACC, are part of the inhibition control system responsible for regulatory control  
9 and response inhibition (49). Hypoactivation of these regions has been associated with  
10 substance use and behavioral addictions (49). In line with our findings, a cluster in the ACC was  
11 also found by Bartholdy *et al.* (2019) to be hypo-activated in BoP developers during successful  
12 (vs. failed) response inhibition (23). Conversely, activating the dorsolateral PFC with high-  
13 frequency repetitive transcranial magnetic stimulation (rTMS) can reduce food craving and  
14 binge-eating frequencies in bulimic disorders (50). Based on these findings, lower GMVs in the  
15 dorsolateral and dorsomedial PFC in purging developers suggest weakened functions in  
16 inhibition control, which may underlie their vulnerability to impulsive behaviors like purging.

17  
18 The neural substrates associated with development of binge-eating encompassed the right  
19 putamen and globus pallidus, with greater GMVs implicating higher risk. Previous studies of ED  
20 patients found reduced GMVs in the bilateral striatum in BN (18) and BED (51, 52) and  
21 increased GMVs in the OFC for BN (18), which is at odds with our findings. The differences may



1 be due to sample characteristics and the study design, previous neuroimaging studies being  
2 cross-sectional and involving adult patients with EDs. In comparison, the present study  
3 investigated onset of DEBs symptoms in a longitudinal cohort of healthy adolescents. It is  
4 possible that higher putamen volumes confer risk for future binge-eating, while reduced  
5 volumes ensue from suffering these conditions over the years. Clearly, more longitudinal  
6 developmental research is needed to study abnormal developmental patterns associated with  
7 DEBs and depression.

8

### 9 **Strengths and Limitations**

10 Our study has several strengths. First, it is one of the only two longitudinal studies (23) to  
11 examine the neurobiological predictors of DEBs in a well-characterized population-based  
12 adolescent cohort. The sample size and multi-modality of the data (questionnaires and  
13 neuroimaging) were other strengths of our study. Matching sex and acquisition sites between  
14 case and control groups removed confounding effects from these variables.

15

16 There are also several weaknesses to consider. First, the very limited number of males in the  
17 case groups did not enable us to investigate sex-specific effects. Second, the effects of other  
18 confounding factors (e.g., parental and social) were not ruled out. Thirdly, DEBs were assessed  
19 through self-reports only. Fourthly, it cannot be concluded that the risk factors identified here  
20 are associated exclusively with a single DEB as our analyses did not exclude coexisting DEBs.  
21 Fifth, the mediation analysis included two variables (SDQ symptoms and brain structure)

1 assessed at one time point and a third variable (development of DEB or depression) assessed  
2 at another, rather than all variables assessed at separate time points. Lastly, as we we did not  
3 find significant structural brain risk factors for dieting, the neurobiological risk factors for  
4 restrictive eating behaviors remain unclear (53).

5

6 Overall, our study highlights psychopathological traits that may be causal risk factors of DEBs,  
7 including emotional problems, ADHD and CD symptoms, which can help identify high-risk  
8 groups for targeted prevention. The identification of neurobiological substrates of DEBs and  
9 comorbid depression, specifically in the OFC and ACC, provides promising therapeutic  
10 strategies for EDs and comorbid conditions (54, 55).

11

## 1 **Acknowledgements**

2 This work received support from the following sources: the Medical Research Council and  
3 Medical Research Foundation (grants MR/R00465X/1 and MRF-058-0004-RG-DESRI: 'ESTRA:  
4 Neurobiological underpinning of eating disorders: integrative biopsychosocial longitudinal  
5 analyses in adolescents'; MR/S020306/1 and MRF-058-0009-RG-DESR-C0759: 'Establishing  
6 causal relationships between biopsychosocial predictors and correlates of eating disorders and  
7 their mediation by neural pathways'), the European Union-funded FP6 Integrated Project  
8 IMAGEN (Reinforcement-related behavior in normal brain function and psychopathology)  
9 (LSHM-CT- 2007-037286), the Horizon 2020 funded ERC Advanced Grant 'STRATIFY' (Brain  
10 network based stratification of reinforcement-related disorders) (695313), ERANID  
11 (Understanding the Interplay between Cultural, Biological and Subjective Factors in Drug Use  
12 Pathways) (PR-ST-0416-10004), BRIDGET (JPND: Brain Imaging, cognition Dementia and next  
13 generation Genomics) (MR/N027558/1), Human Brain Project (HBP SGA 2, 785907), the FP7  
14 project MATRICS (603016), the Medical Research Council Grant 'c-VEDA' (Consortium on  
15 Vulnerability to Externalizing Disorders and Addictions) (MR/N000390/1), the National  
16 Institute for Health Research (NIHR) Biomedical Research Centre at South London and  
17 Maudsley NHS Foundation Trust and King's College London, the Bundesministerium für Bildung  
18 und Forschung (BMBF grants 01GS08152; 01EV0711; Forschungsnetz AERIAL 01EE1406A,  
19 01EE1406B), the Deutsche Forschungsgemeinschaft (DFG grants SM 80/7-2, SFB 940/2, NE  
20 1383/14-1), the National Institutes of Health (NIH) funded ENIGMA (grants 5U54EB020403-05

1 and 1R56AG058854-01). Further support was provided by grants from: the ANR (ANR-12-  
2 SAMA-0004, AAPG2019–GeBra), the Eranet Neuron (AF12-NEUR0008-01–WM2NA; and ANR-  
3 18-NEUR00002-01–ADORe), the Fondation de France (00081242), the Fondation pour la  
4 Recherche Médicale (DPA20140629802), the Mission Interministérielle de Lutte-contre-les-  
5 Drogues-et-les-Conduites-Addictives (MILDECA), the Assistance-Publique-Hôpitaux-de-Paris  
6 and INSERM (interface grant), Paris Sud University IDEX 2012, the Fondation de l’Avenir (grant  
7 AP-RM-17-013), the Fédération pour la Recherche sur le Cerveau; the National Institutes of  
8 Health, Science Foundation Ireland (16/ERCD/3797), U.S.A. (Axon, Testosterone and Mental  
9 Health during Adolescence; RO1 MH085772-01A1), and by NIH Consortium grant U54  
10 EB020403, supported by a cross-NIH alliance that funds Big Data to Knowledge Centers of  
11 Excellence. Ulrike Schmidt is supported by a Senior Investigator award from the National  
12 Institute for Health Research (NIHR). This research was reviewed by a team with experience of  
13 mental health problems and their carers who have been specially trained to advise on research  
14 proposals and documentation through the Young Person’s Mental Health Advisory Group: a  
15 free, confidential service in England provided by the National Institute for Health Research  
16 Maudsley Biomedical Research Centre via King’s College London.

17

18 IMAGEN Consortium authors: Tobias Banaschewski, M.D., Ph.D., Gareth J. Barker, Ph.D., Arun  
19 L.W. Bokde, Ph.D., Erin Burke Quinlan, Ph.D., Sylvane Desrivières, Ph.D., Herta Flor, Ph.D.,  
20 Antoine Grigis, Ph.D., Hugh Garavan, Ph.D., Penny Gowland, Ph.D., Andreas Heinz, M.D., Ph.D.,

1 Bernd Ittermann, Ph.D., Jean-Luc Martinot, M.D., Ph.D., Marie-Laure Paillère Martinot, M.D.,  
2 Ph.D., Frauke Nees, Ph.D., Dimitri Papadopoulos Orfanos, Ph.D., Tomáš Paus, M.D., Ph.D., Luise  
3 Poustka, M.D., Sarah Hohmann, M.D., Juliane H. Fröhner, MSc, Michael N. Smolka, M.D., Henrik  
4 Walter, M.D., Ph.D., Robert Whelan, Ph.D., Gunter Schumann M.D.

5  
6 Other IMAGEN consortium members: Uli Bromberg, Ph.D., University Medical Centre  
7 Hamburg-Eppendorf, House W34, 3.OG, Martinistr. 52, 20246, Hamburg, Germany. Vincent  
8 Frouin, Ph.D., NeuroSpin, CEA, Université Paris-Saclay, F-91191 Gif-sur-Yvette, France. Eric  
9 Artiges, M.D., Ph.D., Institut National de la Santé et de la Recherche Médicale, INSERM Unit  
10 1000 “Neuroimaging & Psychiatry”, University Paris Sud, University Paris Descartes - Sorbonne  
11 Paris Cité; and Psychiatry Department 91G16, Orsay Hospital, France. Herve Lemaitre, Ph.D,  
12 NeuroSpin, CEA, Université Paris-Saclay, F-91191 Gif-sur-Yvette, France; Institut National de la  
13 Santé et de la Recherche Médicale, UMR 992 INSERM, CEA, Faculté de médecine, Université  
14 Paris-Sud, Université Paris-Saclay, NeuroSpin, F-91191 Gif-sur-Yvette, France. Sabina Millenet,  
15 Dipl.-Psych., Department of Child and Adolescent Psychiatry and Psychotherapy, Central  
16 Institute of Mental Health, Medical Faculty Mannheim, Heidelberg University, Square J5, 68159  
17 Mannheim, Germany.

18

1 **Disclosures**

2 Dr. Banaschewski served in an advisory or consultancy role for Lundbeck, Medice, Neurim  
3 Pharmaceuticals, Oberberg GmbH, Shire. He received conference support or speaker's fee by  
4 Lilly, Medice, Novartis and Shire. He has been involved in clinical trials conducted by Shire &  
5 Viforpharma. He received royalties from Hogrefe, Kohlhammer, CIP Medien, Oxford University  
6 Press. The present work is unrelated to the above grants and relationships. Dr. Barker has  
7 received honoraria from General Electric Healthcare for teaching on scanner programming  
8 courses. The other authors report no biomedical financial interests or potential conflicts of  
9 interest.

10

11

1       **References**

- 2       1.   Allen KL, Byrne SM, Crosby RD, Stice E (2016): Testing for interactive and non-linear effects  
3       of risk factors for binge eating and purging eating disorders. *Behav Res Ther* 87:40-47.
- 4       2.   Micali N, Hagberg KW, Petersen I, Treasure JL (2013): The incidence of eating disorders in  
5       the UK in 2000-2009: findings from the General Practice Research Database. *BMJ Open*  
6       3:e002646.
- 7       3.   Jones JM, Bennett S, Olmsted MP, Lawson ML, Rodin G (2001): Disordered eating attitudes  
8       and behaviours in teenaged girls: a school-based study. *CMAJ* 165:547-552.
- 9       4.   Holling H, Schlack R (2007): Eating disorders in children and adolescents. First results of  
10       the German Health Interview and Examination Survey for Children and Adolescents (KiGGS).  
11       *Bundesgesundheitsblatt Gesundheitsforschung Gesundheitsschutz* 50:794-799.
- 12       5.   Dakanalis A, Clerici M, Bartoli F, Caslini M, Crocamo C, Riva G, et al. (2017): Risk and  
13       maintenance factors for young women's DSM-5 eating disorders. *Arch Womens Ment Health*  
14       20:721-731.
- 15       6.   Stice E, Gau JM, Rohde P, Shaw H (2017): Risk factors that predict future onset of each  
16       DSM-5 eating disorder: Predictive specificity in high-risk adolescent females. *J Abnorm Psychol*  
17       126:38-51.
- 18       7.   Hudson JI, Hiripi E, Pope HG, Jr., Kessler RC (2007): The prevalence and correlates of eating  
19       disorders in the National Comorbidity Survey Replication. *Biol Psychiatry* 61:348-358.
- 20       8.   Yao S, Kuja-Halkola R, Martin J, Lu Y, Lichtenstein P, Noring C, et al. (2019): Associations  
21       Between Attention-Deficit/Hyperactivity Disorder and Various Eating Disorders: A Swedish

- 1 Nationwide Population Study Using Multiple Genetically Informative Approaches. *Biol*  
2 *Psychiatry* 86:577-586.
- 3 9. Yilmaz Z, Javaras KN, Baker JH, Thornton LM, Lichtenstein P, Bulik CM, et al. (2017):  
4 Association Between Childhood to Adolescent Attention Deficit/Hyperactivity Disorder  
5 Symptom Trajectories and Late Adolescent Disordered Eating. *J Adolesc Health* 61:140-146.
- 6 10. Yoshimasu K, Barbaresi WJ, Colligan RC, Voigt RG, Killian JM, Weaver AL, et al. (2012):  
7 Childhood ADHD is strongly associated with a broad range of psychiatric disorders during  
8 adolescence: a population-based birth cohort study. *J Child Psychol Psychiatry* 53:1036-1043.
- 9 11. Hilbert A, Pike KM, Goldschmidt AB, Wilfley DE, Fairburn CG, Dohm FA, et al. (2014): Risk  
10 factors across the eating disorders. *Psychiatry Res* 220:500-506.
- 11 12. Chronis-Tuscano A, Molina BS, Pelham WE, Applegate B, Dahlke A, Overmyer M, et al.  
12 (2010): Very early predictors of adolescent depression and suicide attempts in children with  
13 attention-deficit/hyperactivity disorder. *Arch Gen Psychiatry* 67:1044-1051.
- 14 13. Biederman J, Ball SW, Monuteaux MC, Mick E, Spencer TJ, Mc CM, et al. (2008): New  
15 insights into the comorbidity between ADHD and major depression in adolescent and young  
16 adult females. *J Am Acad Child Adolesc Psychiatry* 47:426-434.
- 17 14. Smith DG, Robbins TW (2013): The neurobiological underpinnings of obesity and binge  
18 eating: a rationale for adopting the food addiction model. *Biol Psychiatry* 73:804-810.
- 19 15. Seitz J, Herpertz-Dahlmann B, Konrad K (2016): Brain morphological changes in adolescent  
20 and adult patients with anorexia nervosa. *J Neural Transm* 123:949-959.



- 1 16. Donnelly B, Touyz S, Hay P, Burton A, Russell J, Caterson I (2018): Neuroimaging in bulimia  
2 nervosa and binge eating disorder: a systematic review. *J Eat Disord* 6:3.
- 3 17. Berner LA, Stefan M, Lee S, Wang Z, Terranova K, Attia E, et al. (2018): Altered cortical  
4 thickness and attentional deficits in adolescent girls and women with bulimia nervosa. *J*  
5 *Psychiatry Neurosci* 43:151-160.
- 6 18. Frank GK, Shott ME, Hagman JO, Mittal VA (2013): Alterations in brain structures related  
7 to taste reward circuitry in ill and recovered anorexia nervosa and in bulimia nervosa. *Am J*  
8 *Psychiatry* 170:1152-1160.
- 9 19. Schafer A, Vaitl D, Schienle A (2010): Regional grey matter volume abnormalities in bulimia  
10 nervosa and binge-eating disorder. *Neuroimage* 50:639-643.
- 11 20. Marsh R, Stefan M, Bansal R, Hao X, Walsh BT, Peterson BS (2015): Anatomical  
12 characteristics of the cerebral surface in bulimia nervosa. *Biol Psychiatry* 77:616-623.
- 13 21. Berner LA, Wang Z, Stefan M, Lee S, Huo Z, Cyr M, et al. (2019): Subcortical Shape  
14 Abnormalities in Bulimia Nervosa. *Biol Psychiatry Cogn Neurosci Neuroimaging* 4:1070-1079.
- 15 22. He X, Stefan M, Terranova K, Steinglass J, Marsh R (2016): Altered White Matter  
16 Microstructure in Adolescents and Adults with Bulimia Nervosa. *Neuropsychopharmacology*  
17 41:1841-1848.
- 18 23. Bartholdy S, O'Daly OG, Campbell IC, Banaschewski T, Barker G, Bokde ALW, et al. (2019):  
19 Neural Correlates of Failed Inhibitory Control as an Early Marker of Disordered Eating in  
20 Adolescents. *Biol Psychiatry* 85:956-965.

- 1 24. Dunlop KA, Woodside B, Downar J (2016): Targeting Neural Endophenotypes of Eating  
2 Disorders with Non-invasive Brain Stimulation. *Front Neurosci* 10:30.
- 3 25. Schumann G, Loth E, Banaschewski T, Barbot A, Barker G, Buchel C, et al. (2010): The  
4 IMAGEN study: reinforcement-related behaviour in normal brain function and  
5 psychopathology. *Mol Psychiatry* 15:1128-1139.
- 6 26. Goodman R, Ford T, Richards H, Gatward R, Meltzer H (2000): The Development and Well-  
7 Being Assessment: description and initial validation of an integrated assessment of child and  
8 adolescent psychopathology. *J Child Psychol Psychiatry* 41:645-655.
- 9 27. Bartholdy S, Allen K, Hodsoll J, O'Daly OG, Campbell IC, Banaschewski T, et al. (2017):  
10 Identifying disordered eating behaviours in adolescents: how do parent and adolescent reports  
11 differ by sex and age? *Eur Child Adolesc Psychiatry* 26:691-701.
- 12 28. Ho DE, Imai K, King G, Stuart EA (2011): MatchIt: Nonparametric Preprocessing for  
13 Parametric Causal Inference. *J Stat Softw* 42:1-28.
- 14 29. Goodman R (1997): The Strengths and Difficulties Questionnaire: a research note. *J Child*  
15 *Psychol Psychiatry* 38:581-586.
- 16 30. Goodman A, Heiervang E, Collishaw S, Goodman R (2011): The 'DAWBA bands' as an  
17 ordered-categorical measure of child mental health: description and validation in British and  
18 Norwegian samples. *Soc Psychiatry Psychiatr Epidemiol* 46:521-532.
- 19 31. Flegal KM, Cole TJ (2013): Construction of LMS parameters for the Centers for Disease  
20 Control and Prevention 2000 growth charts. *Natl Health Stat Report*:1-3.

- 1 32. Sobell LC, Cunningham JA, Sobell MB (1996): Recovery from alcohol problems with and  
2 without treatment: prevalence in two population surveys. *Am J Public Health* 86:966-972.
- 3 33. Sargolzaei S, Sargolzaei A, Cabrerizo M, Chen G, Goryawala M, Noei S, et al. (2015): A  
4 practical guideline for intracranial volume estimation in patients with Alzheimer's disease. *BMC*  
5 *Bioinformatics* 16:S8.
- 6 34. Firth D (1993): Bias reduction of maximum likelihood estimates. *Biometrika* 80:27-38.
- 7 35. Seymour KE, Reinblatt SP, Benson L, Carnell S (2015): Overlapping neurobehavioral circuits  
8 in ADHD, obesity, and binge eating: evidence from neuroimaging research. *CNS Spectrums*  
9 20:401-411.
- 10 36. Norman LJ, Carlisi C, Lukito S, Hart H, Mataix-Cols D, Radua J, et al. (2016): Structural and  
11 Functional Brain Abnormalities in Attention-Deficit/Hyperactivity Disorder and Obsessive-  
12 Compulsive Disorder: A Comparative Meta-analysis. *JAMA Psychiatry* 73:815-825.
- 13 37. Wilbertz G, van Elst LT, Delgado MR, Maier S, Feige B, Philipsen A, et al. (2012):  
14 Orbitofrontal reward sensitivity and impulsivity in adult attention deficit hyperactivity disorder.  
15 *Neuroimage* 60:353-361.
- 16 38. Hudson JI, McElroy SL, Ferreira-Cornwell MC, Radewonuk J, Gasior M (2017): Efficacy of  
17 Lisdexamfetamine in Adults With Moderate to Severe Binge-Eating Disorder: A Randomized  
18 Clinical Trial. *JAMA Psychiatry* 74:903-910.
- 19 39. McElroy SL, Hudson J, Ferreira-Cornwell MC, Radewonuk J, Whitaker T, Gasior M (2016):  
20 Lisdexamfetamine Dimesylate for Adults with Moderate to Severe Binge Eating Disorder:

- 1 Results of Two Pivotal Phase 3 Randomized Controlled Trials. *Neuropsychopharmacology*  
2 41:1251-1260.
- 3 40. Daviss WB (2008): A review of co-morbid depression in pediatric ADHD: etiology,  
4 phenomenology, and treatment. *J Child Adolesc Psychopharmacol* 18:565-571.
- 5 41. Posner J, Siciliano F, Wang Z, Liu J, Sonuga-Barke E, Greenhill L (2014): A multimodal MRI  
6 study of the hippocampus in medication-naïve children with ADHD: what connects ADHD and  
7 depression? *Psychiatry Res* 224:112-118.
- 8 42. Ochsner KN, Gross JJ (2005): The cognitive control of emotion. *Trends Cogn Sci* 9:242-249.
- 9 43. Seymour KE, Chronis-Tuscano A, Iwamoto DK, Kurdziel G, Macpherson L (2014): Emotion  
10 regulation mediates the association between ADHD and depressive symptoms in a community  
11 sample of youth. *J Abnorm Child Psychol* 42:611-621.
- 12 44. Anastopoulos AD, Smith TF, Garrett ME, Morrissey-Kane E, Schatz NK, Sommer JL, et al.  
13 (2011): Self-Regulation of Emotion, Functional Impairment, and Comorbidity Among  
14 Children With AD/HD. *J Atten Disord* 15:583-592.
- 15 45. Rogers JC, De Brito SA (2016): Cortical and Subcortical Gray Matter Volume in Youths With  
16 Conduct Problems: A Meta-analysis. *JAMA Psychiatry* 73:64-72.
- 17 46. O'Doherty J, Kringelbach ML, Rolls ET, Hornak J, Andrews C (2001): Abstract reward and  
18 punishment representations in the human orbitofrontal cortex. *Nat Neurosci* 4:95-102.
- 19 47. Chudasama Y, Daniels TE, Gorrin DP, Rhodes SE, Rudebeck PH, Murray EA (2013): The role  
20 of the anterior cingulate cortex in choices based on reward value and reward contingency.  
21 *Cereb Cortex* 23:2884-2898.

- 1 48. Stevens FL, Hurley RA, Taber KH (2011): Anterior cingulate cortex: unique role in cognition  
2 and emotion. *J Neuropsychiatry Clin Neurosci* 23:121-125.
- 3 49. Luijten M, Machielsen MW, Veltman DJ, Hester R, de Haan L, Franken IH (2014): Systematic  
4 review of ERP and fMRI studies investigating inhibitory control and error processing in people  
5 with substance dependence and behavioural addictions. *J Psychiatry Neurosci* 39:149-169.
- 6 50. Van den Eynde F, Claudino AM, Mogg A, Horrell L, Stahl D, Ribeiro W, et al. (2010):  
7 Repetitive transcranial magnetic stimulation reduces cue-induced food craving in bulimic  
8 disorders. *Biol Psychiatry* 67:793-795.
- 9 51. Woolley JD, Gorno-Tempini ML, Seeley WW, Rankin K, Lee SS, Matthews BR, et al. (2007):  
10 Binge eating is associated with right orbitofrontal-insular-striatal atrophy in frontotemporal  
11 dementia. *Neurology* 69:1424-1433.
- 12 52. Voon V, Derbyshire K, Ruck C, Irvine MA, Worbe Y, Enander J, et al. (2015): Disorders of  
13 compulsivity: a common bias towards learning habits. *Mol Psychiatry* 20:345-352.
- 14 53. Allen KL, Byrne SM, Forbes D, Oddy WH (2009): Risk factors for full- and partial-syndrome  
15 early adolescent eating disorders: a population-based pregnancy cohort study. *J Am Acad Child*  
16 *Adolesc Psychiatry* 48:800-809.
- 17 54. Fettes P, Schulze L, Downar J (2017): Cortico-Striatal-Thalamic Loop Circuits of the  
18 Orbitofrontal Cortex: Promising Therapeutic Targets in Psychiatric Illness. *Front Syst Neurosci*  
19 11:25.
- 20 55. Dalton B, Bartholdy S, McClelland J, Kekic M, Rennalls SJ, Werthmann J, et al. (2018):  
21 Randomised controlled feasibility trial of real versus sham repetitive transcranial magnetic

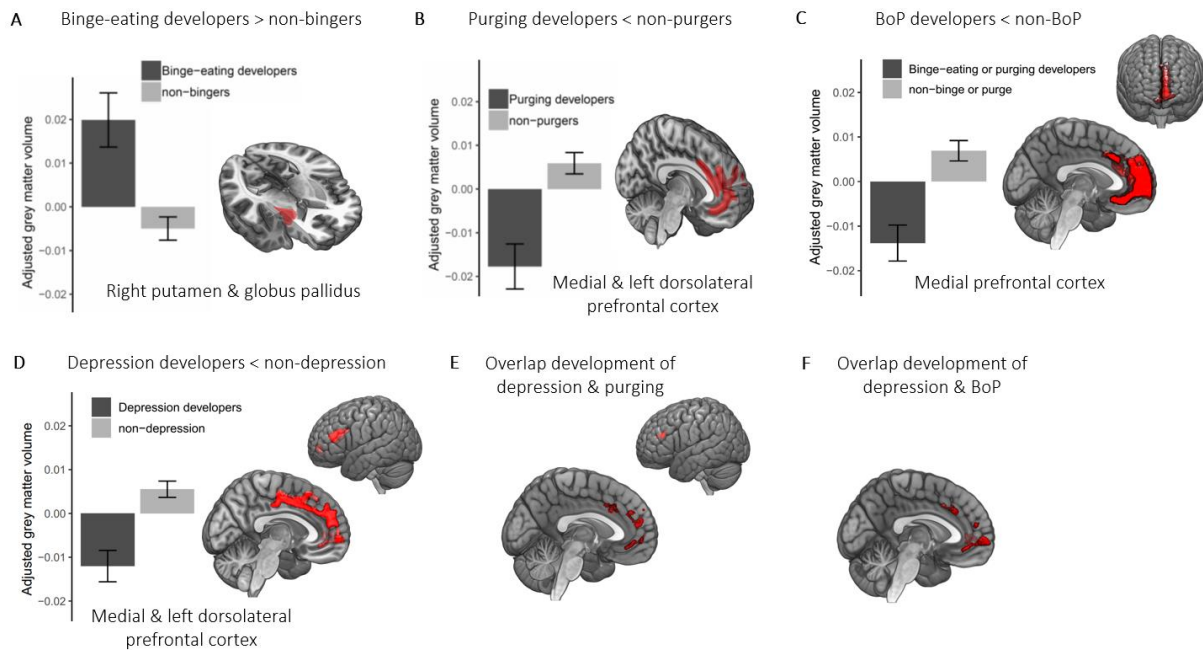
1 stimulation treatment in adults with severe and enduring anorexia nervosa: the TIARA study.

2 BMJ Open 8:e021531.

3

4

1 **Figure Legends**



2

3 **FIGURE 1.** Structural brain associations with the development of binge-eating (A), purging (B),

4 binge-eating or purging (BoP, C) and depressive symptoms (D). The bar plots show the regional

5 means of GMVs, adjusted by sex, acquisition site and total intracranial volumes. Error bars

6 represent standard errors. Statistical parametric maps were thresholded at voxel-level  $p < 0.001$

7 (uncorrected) and cluster-level  $p < 0.05$  (FWE corrected). Overlapping brain regions associated

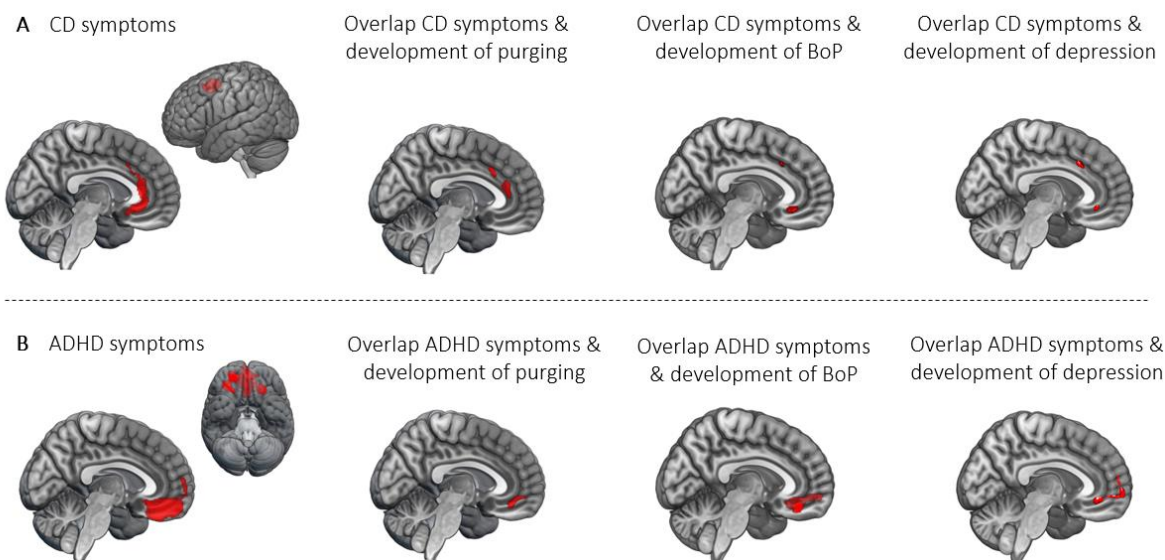
8 with the development of purging and depressive symptoms are presented in E. Overlapping

9 brain regions for the development of BoP and depressive symptoms are presented in F. The

10 3D rendered views are generated by MRICroGL

11 (<https://www.mccauslandcenter.sc.edu/mricrogl/>).

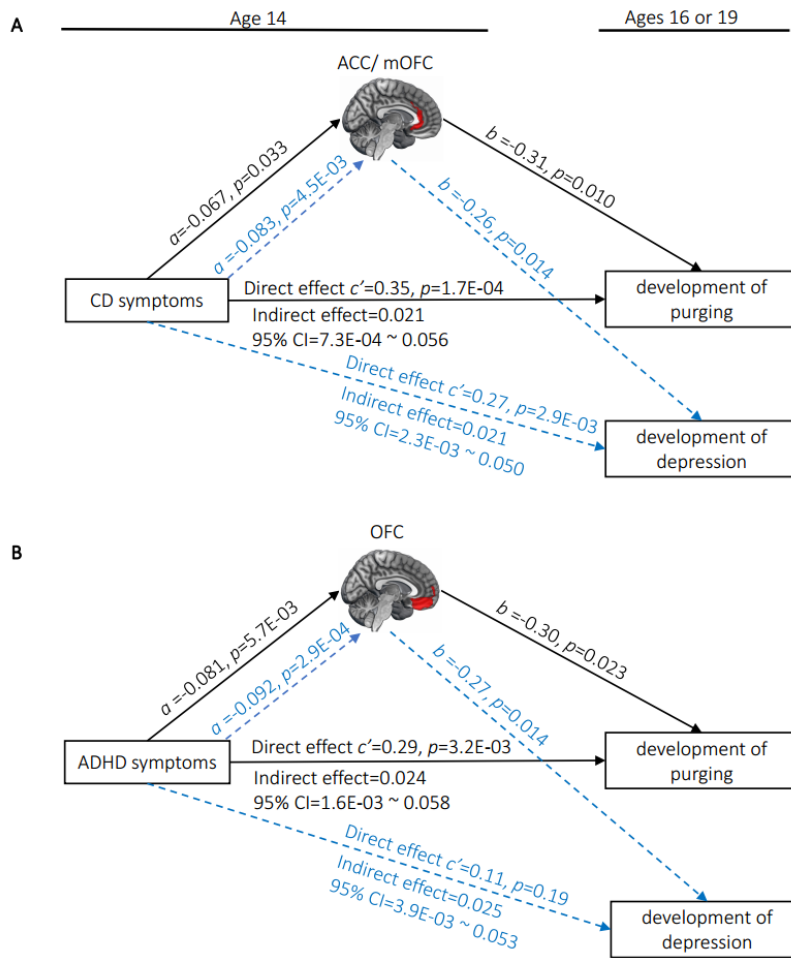
12



1  
2 **FIGURE 2.** Grey matter volume associations with CD symptoms (A, negative association) and  
3 ADHD symptoms (B, negative association) at age 14. Statistical parametric maps were  
4 thresholded at voxel-level  $p < 0.001$  (uncorrected) and cluster-level  $p < 0.05$  (FWE corrected).  
5 The columns to the right show overlapping brain regions associated with the development of  
6 purging, BoP and depressive symptoms.

7





1

2 **FIGURE 3.** Results for the mediation analysis, using the ACC/mOFC ROI linked to CD symptoms  
 3 (A) and the OFC ROI linked to ADHD symptoms (B) as mediators. For analyses on the  
 4 development of purging, control variables included sex, acquisition sites and the total  
 5 intracranial volume. For analyses on the development of depression (blue dashed lines), the  
 6 depression symptom at age 14 was involved as an additional control variable. ROI: region of  
 7 interest; CI: confidence interval; ACC: anterior cingulate cortex; OFC: orbitofrontal cortex.

1 **Tables**

2 Table 1. Psychopathological predictors of DEB symptoms

developers of DEBs	SDQ symptoms at age 14	Without controlling for BMI			After controlling for BMI		
		OR	95% CI	P	OR	95% CI	P
binge-eating developers vs. non-bingers	conduct problems	1.29	1.06-1.57	1.1E-02			
	emotional symptoms	<b>1.35</b>	<b>1.10-1.66</b>	<b>4.4E-03</b>	1.34	1.09-1.65	5.7-03
	hyperactivity/ inattention	1.27	1.03-1.57	2.4E-02			
	peer problems	1.24	1.01-1.51	3.7E-02			
purging developers vs. non-purgers	conduct problems	<b>1.43</b>	<b>1.20-1.71</b>	<b>8.5E-05</b>	1.41	1.18-1.70	1.5E-04
	emotional symptoms	1.24	1.03-1.49	2.5E-02			
	hyperactivity/inattention	<b>1.35</b>	<b>1.12-1.64</b>	<b>1.6E-03</b>	1.37	1.13-1.66	1.2E-03
	peer problems	0.98	0.81-1.18	8.6E-01			
dieting developers vs. non-dieters	conduct problems	1.22	0.92-1.59	1.6E-01			
	emotional symptoms	1.22	0.91-1.63	1.8E-01			
	hyperactivity/inattention	1.17	0.88-1.57	2.7E-01			
	peer problems	1.23	0.94-1.59	1.2E-01			

3 The p values shown in bold survived the Holm-Bonferroni correction, correcting for 12 tests (3 DEBs ×  
 4 4 SDQ subscales). Significant associations were further controlled for BMI. DEB: disordered eating  
 5 behavior.

6

7

1 **Table 2.** Psychopathological predictors for the development of depression and generalized  
 2 anxiety symptoms

Development of mental health symptoms	SDQ symptoms at age 14	Controlling for the corresponding mental health symptom at age 14			Additionally controlling for the other mental health symptom at all ages		
		OR	95% CI	P	OR	95% CI	P
Depression	hyperactivity/inattention	1.25	1.08-1.44	<b>2.5E-03</b>	1.10	0.93-1.29	0.27
	conduct problems	1.32	1.13-1.54	<b>4.6E-04</b>	1.22	1.03-1.46	0.025
	emotional symptoms	1.39	1.19-1.62	<b>2.8E-05</b>	0.98	0.79-1.20	0.82
Generalized anxiety	hyperactivity/inattention	1.33	1.13-1.57	<b>5.4E-04</b>	1.24	1.00-1.53	0.045
	conduct problems	1.26	1.06-1.49	<b>9.3E-03</b>	0.97	0.77-1.22	0.79
	emotional symptoms	1.79	1.46-2.19	<b>1.0E-08</b>	1.56	1.22-2.00	3.7E-04

3 The p values shown in bold survived the Holm-Bonferroni correction, correcting for 6 tests (2 mental  
 4 health symptoms × 3 SDQ subscales). Significant associations were further controlled for the other  
 5 mental health symptom at ages 14, 16 and 19 (i.e., controlling depression development for anxiety  
 6 symptoms at all ages, and vice-versa).

7  
 8  
 9