COMT-Polymorphisms Modulated Functional Profile of the Fusiform Face Area Contributes to Face-Specific Recognition Ability

Chao Wu1, Zonglei Zhen2*, Lijie Huang3, Taicheng Huang3 & Jia Liu2*

Previous studies have shown that face-specific recognition ability (FRA) is heritable; however, the neural basis of this heritability is unclear. Candidate gene studies have suggested that the catechol-O-methyltransferase (COMT) rs4680 polymorphism is related to face perception. Here, using a partial least squares (PLS) method, we examined the multivariate association between 12 genotypes of 4 COMT polymorphisms (rs6269-rs4633-rs4818-rs4680) and multimodal MRI phenotypes in the human fusiform face area (FFA), which selectively responds to face stimuli, in 338 Han Chinese adults (mean age 20.45 years; 135 males). The MRI phenotypes included gray matter volume (GMV), resting-state fractional amplitude of low-frequency fluctuations (fALFF), and face-selective blood-oxygen-level-dependent (BOLD) responses (FS). We found that the first COMT-variant component (PLS1) was positively associated with the FS but negatively associated with the fALFF in the FFA. Moreover, participants with the COMT heterozygous-HEA-haplotype showed higher PLS1 FFA-MRI scores, which were positively associated with the FRA in an old/new face recognition task, than those with the COMT homozygous HEA haplotype and HEA non-carriers, suggesting that individuals with an appropriate (intermediate) level of dopamine activity in the FFA might have better FRA. In summary, our study provides empirical evidence for the genetic and neural basis for the heritability of face recognition and informs the formation of neural module functional specificity.
Candidate gene studies have demonstrated the association of the catechol-O-methyltransferase (COMT) gene, located on chromosome 22q11.2, with face recognition\(^\text{1}–\text{9}\). The COMT gene codes for the COMT enzyme, a protein that degrades catecholamine neurotransmitters and modulates dopamine metabolisms\(^\text{10}\). Among the COMT polymorphisms, Val158Met (rs4680) has been most reported to play a role in face processing\(^\text{11}–\text{24}\). For example, individuals with the Met/Met (rs4680) genotype are more sensitive to configurational changes in faces than those with other genotypes\(^\text{11}\). Healthy homozygous subjects with the Met allele of rs4680 are better at recognizing facial emotions\(^\text{16}\) and show a stronger bias to perceive neutral faces as expressions of anger\(^\text{15}\) than those with other genotypes. Previous studies also investigated the impact of the COMT gene on the brain activity during face emotional processing\(^\text{22}–\text{25}\) and demonstrated that the COMT Met allele seems to contribute to neural substrates of negativity biases\(^\text{22}–\text{24}\), and the COMT Val allele is associated with increased amygdala activity in response to fearful/angry facial expressions\(^\text{25}\). It should be noted that several meta-analyses suggest that the effects of COMT Val158Met on cognitive abilities are either very small or nil due to insufficient statistical power, unknown moderators including genetic variations at different loci, publication bias, or effect heterogeneity\(^\text{26}–\text{28}\). Therefore, individual studies should pay more attention to the methodological rigor and scientific nature. Accumulating studies have revealed that there are multi-to-multi relationships between genotypes and brain phenotypes linked to learning and cognitive ability\(^\text{29}–\text{33}\). Previous studies have indicated that four COMT polymorphisms (rs6269, rs4633, rs4818, and rs4680) have a strong linkage disequilibrium and always function as haplotype blocks to affect dopamine (DA) degrading enzyme activity\(^\text{29–33}\). However, little is known about whether certain patterns of the 4 COMT polymorphisms exert significant effects on the FFA and therefore affect the face recognition ability.

In this study, we employed a combined genetic and neuroimaging approach and incorporated three strategies to improve statistical power without dramatically increasing the number of participants. First, previous studies focused on a single measure of the FFA (i.e., either gray matter volume (GMV) or blood oxygen level-dependent [BOLD] responses); notably, multiple modals and joint modeling can improve sensitivity for identifying disease-associated biological pathways in common complex disorders\(^\text{34–37}\); accordingly, this study used multimodal MRI measures for the FFA (multimodal FFA-MRI data), including GMV, fractional amplitude of low-frequency fluctuations (fALFF) from resting-state fMRI, and BOLD responses specific to faces (face selectivity, FS) from task-state fMRI. Second, the partial least squares (PLS) approach was applied to examine relationships between sets of multivariate measures. This approach has demonstrated utility in studies with relative small samples and problems with multicollinearity\(^\text{37–40}\). Third, bilateral FFAs are involved in face processing simultaneously\(^\text{37,41}\), notwithstanding some hemispheric lateralization in their size and functional recruitments\(^\text{37,42}\). We conducted analyses for the right FFA (rFFA) and the left FFA (lFFA) separately to validate the genetic variants that modulate the FFA. Using these three strategies, we identified profiles of 4 COMT polymorphisms that would be robustly associated with profiles of structural or functional FFA, in a population of Han Chinese individuals.

**Results**

Multivariate association between COMT polymorphisms and FFA phenotypes. A PLS correspondence analysis (PLSCA) was conducted to examine whether the 12 COMT genotypes contributed to the 3-modal right FFA-MRI phenotype in 338 participants. The PLSCA yielded 3 sets of paired latent variables (LVs) capturing the rFFA-SNP association, ordered by the size of the explained variance (91.51%, 7.0%, and 1.49%). Each component (LV pair) was a linear combination of the weighted COMT SNP scores that most strongly correlated with weighted MRI scores and was orthogonal to each other. The omnibus $p$ value of the model was 0.004 and only the top PLS component (PLS1) was significant (permutation test, $p = 0.001$). Thus, the PLS1 represented a significant association ($r = 0.258$, $p < 0.001$) between a specifically patterned COMT genotypic profile and a specifically patterned rFFA phenotype (middle panel of Fig. 1A). The power analysis\(^\text{43}\) for the PLS model (Supplementary Table S2) indicated a power of 0.713 for the path between the rFFA and the COMT polymorphisms, which means that we have the probability of 71.3% to detect a significant association between the rFFA and the COMT polymorphisms, given that the association is really there. The component reliability test showed that the rs4680 AA (Met) genotype (salience = 0.417, BSR = 3.33, FDR corrected $p = 0.007$, 95%CI [0.17, 0.72]), rs4818 AA genotype (salience = 0.110, BSR = 2.40, FDR corrected $p = 0.040$, 95%CI [0.02, 0.20]), rs4818 AG genotype (salience = −0.132, BSR = −2.94, FDR corrected $p = 0.013$, 95%CI [−0.22, −0.04]), rs4633 TT genotype (salience = 0.370, BSR = 3.23, FDR corrected $p = 0.007$, 95%CI [0.15, 0.59]), and rs6269 AG genotype (salience = −0.121, BSR = −2.66, FDR corrected $p = 0.024$, 95%CI [−0.21, −0.03]) reliably contributed to the PLS1 COMT SNP component (left panel of Fig. 1A); the FS (salience = 0.129, BSR = 3.21, FDR corrected $p = 0.007$, 95%CI [0.05, 0.21]) and fALFF (salience = −0.132, BSR = −2.84, FDR corrected $p = 0.004$, 95%CI [−0.22, −0.04]) reliably contributed to the PLS1 rFFA-MRI component (right panel of Fig. 1A). Moreover, the genetic regulation of the rFFA presented a pattern that rs4680 and rs4633 exhibited linear trend of modulation (from recessive homozygote to heterozygote and to dominant homozygote) on the PLS1 rFFA-MRI profile, whereas rs6269 and rs4818 showed a nonlinear relationship with the PLS1 rFFA-MRI profile.

To explain how this profile (linear combination) of the 12 COMT genotypes (i.e., PLS1 SNP profile) modulates the 3 raw MRI measures of the rFFA, we correlated the PLS1 COMT scores with FS, GMV, and fALFF of the rFFA. The results showed that the PLS1 COMT profile was positively correlated with FS ($r = 0.172, p = 0.002$, 95%CI [0.07, 0.27]), negatively correlated with fALFF ($r = -0.175, p = 0.001$, 95%CI [−0.28, −0.07]), and not correlated with GMV of the rFFA ($r = 0.075, p = 0.171$, 95%CI [−0.03, 0.18]) (Fig. 1B). Thus, the COMT polymorphism profile seems to be more likely to regulate the functional rather than structural profile of the right FFA.

To validate the results and improve reliability, we conducted the following control analyses. First, we randomly split the voxels within the rFFA into two halves and entered each half of the rFFA measures along with COMT polymorphisms into a PLSCA. We found that both of the rFFA-COMT associations were significant (Supplementary Fig. S1) and the difference between the two associations was not significant ($z = 0.227, p = 0.821$). Moreover, the intra-class correlations (ICC) for the two half GMV, fALFF, FS, PLS1-SNP scores, and PLS1-rFFA scores were...
0.985 (95%CI [0.981, 0.988]), 0.995 (95%CI [0.994, 0.996]), 0.989 (95%CI [0.986, 0.991]), 0.999 (95%CI [0.999, 1]), and 0.994 (95%CI [0.993, 0.996]), respectively. Thus, both the rFFA measures and the association pattern between the rFFA phenotypes and the COMT polymorphisms exhibit high reliability. Second, to test whether the threshold for identifying the rFFA affects the main results, we conducted control analyses with 4 different thresholds ($z = 2.71$, one-tailed uncorrected $p < 0.003$, FDR corrected $p < 0.05$; $z = 2.58$, one-tailed uncorrected $p < 0.005$; $z = 2.34$, one-tailed uncorrected $p < 0.01$; $z = 1.96$, one-tailed uncorrected $p < 0.05$) and found that the association between the 12 COMT genotypes and the rFFA-MRI phenotypes was significant in all of the 4 threshold conditions (Fig. S2 in Supplementary Material), suggesting that the threshold for defining the rFFA has little effect on the COMT-rFFA association. Third, to test whether total brain volume affects the current results, we conducted a control PLS analysis after regressing the total brain volume out of the rFFA and found the main results remained (Supplementary Fig. S3). Finally, to test whether the rFFA-COMT association can be reproduced for the left FFA (lFFA), we conducted a PLSCA to explore the association between the 12 COMT genotypes and the 3-modal lFFA-MRI phenotypes. The analysis yielded 3 sets of LVs capturing the lFFA-SNP association, ordered by the size of the explained variance (81.93%, 17.31%, and 0.07%). The covariation between the PLS1 COMT profile and the PLS1 lFFA-MRI profile was marginally significant (permutation test, $p = 0.053$; upper panel of Supplementary Fig. S4). The ICC for the PLS1 IFFA-SNP scores and PLS1 rFFA-SNP scores was 0.998 (95%CI [0.997, 0.998]), and the ICC for the PLS1 IFFA-MRI scores and rFFA-MRI scores was 0.787 (95%CI [0.736, 0.828]). Moreover, the difference between the IFFA-COMT association and the rFFA-COMT association was not significant ($z = 1.331$, $p = 0.183$). Thus, the IFFA exhibits a similar association pattern with the COMT polymorphisms as the rFFA does.

Haplotype analysis. Because the four COMT SNPs has been previously found to be in strong linkage disequilibrium and define three common haplotypes with functional consequences on dopamine (DA) degrading enzyme function, we performed a haplotype analysis on the 4 genotyped COMT SNPs to inform their functions as a haplotype block modulating the profile of the multimodal FFA phenotype. In our sample, the 4 SNPs were in strong linkage disequilibrium with all $D’ > 0.90$, and we reproduced the 3 haplotypes: (1) a high enzymatic activity (HEA) haplotype (rs6269/G-rs4633/C-rs4818/G-rs4680/G) with a frequency of 0.328, (2) an
intermediate enzymatic activity (MEA) haplotype (rs6269/A-rs4633/T-rs4818/C-rs4680/A) with a frequency of 0.239, and (3) a low enzymatic activity (LEA) haplotype (rs6269/A-rs4633/C-rs4818/C-rs4680/G) with the highest frequency of 0.389 (Table 1). Diplotype estimation revealed 34 homozygous HEA (low dopamine availability) haplotype carriers, 143 heterozygous HEA (intermediate dopamine availability) haplotype carriers, and 139 HEA haplotype non-carriers (high dopamine availability)33 in our sample (Table 1). A two-way ANOVA was carried out on PLS1-FFA scores by haplotype and hemisphere (Fig. 2). The interaction between the effects of haplotype and hemisphere on PLS1-FFA scores was not significant \[F(2,626) = 0.252, p = 0.777\]. The main effect of haplotype was significant \[F(2,626) = 9.436, p < 0.001, \eta^2_p = 0.031, \text{power} = 0.809\], but the main effect of hemisphere was not significant \[F(1,626) = 0.003, p = 0.958\]. Tukey’s HSD post hoc tests showed that

| SNPs                  | Genotype | GF  | N (Male) | Age years | Raven score | Face accuracy | Face sensitivity | Flower accuracy | Flower sensitivity |
|-----------------------|----------|-----|----------|-----------|-------------|---------------|-----------------|-----------------|-------------------|
| rs6269_G              | A/A      | 0.427 | 146 (55) | 20.3 (1.0) | 24.9 (6.8) | 76.3 (8.8) | 1.6 (1.0)          | 77.2 (9.7) | 2.1 (1.4)         |
| rs6269_G              | A/G      | 0.462 | 155 (65) | 20.5 (0.8) | 25.6 (6.0) | 75.8 (9.1) | 1.7 (1.1)          | 77.8 (8.3) | 2.1 (1.3)         |
| rs6269_G              | G/G      | 0.111 | 37 (14)  | 20.6 (1.2) | 25.7 (3.7) | 75.4 (10.7) | 1.7 (1.2)          | 77.5 (8.6) | 2.0 (1.2)         |
| rs4633_T              | C/C      | 0.557 | 187 (78) | 20.5 (1.0) | 28.0 (5.1) | 75.7 (10)  | 1.7 (1.2)          | 77.7 (8.9) | 2.1 (1.3)         |
| rs4633_T              | C/T      | 0.377 | 125 (45) | 20.3 (0.9) | 24.3 (6.3) | 76.7 (8.0) | 1.6 (0.9)          | 77.3 (9.0) | 2.0 (1.2)         |
| rs4633_T              | T/T      | 0.066 | 25 (11)  | 20.2 (1.0) | 24.8 (6.5) | 74.9 (10)  | 1.6 (1.1)          | 77.3 (10) | 2.1 (1.4)         |
| rs4818_G              | C/C      | 0.434 | 143 (53) | 20.3 (1.0) | 24.8 (6.9) | 76.2 (9.0) | 1.6 (1.0)          | 77.7 (9.8) | 2.1 (1.4)         |
| rs4818_G              | C/G      | 0.456 | 148 (63) | 20.5 (0.8) | 25.6 (5.0) | 75.9 (9.1) | 1.7 (1.1)          | 77.6 (8.4) | 2.1 (1.3)         |
| rs4818_G              | G/G      | 0.111 | 35 (14)  | 20.7 (1.3) | 25.9 (3.5) | 75.7 (9.1) | 1.7 (1.2)          | 77.8 (8.6) | 2.1 (1.2)         |
| rs4680_A              | G/G      | 0.582 | 193 (78) | 20.5 (1.0) | 26.0 (5.2) | 75.5 (9.7) | 1.7 (1.1)          | 77.9 (8.8) | 2.1 (1.3)         |
| rs4680_A              | A/G      | 0.358 | 121 (46) | 20.3 (0.9) | 24.4 (6.4) | 82.7 (8.2) | 1.6 (0.9)          | 76.9 (9.2) | 2.1 (1.3)         |
| rs4680_A              | A/A      | 0.060 | 21 (9)   | 20.5 (0.9) | 25.2 (5.7) | 76.8 (8.9) | 1.7 (1.1)          | 79.0 (8.7) | 2.2 (1.3)         |
| Haplotype combination | rs6269-rs4633-rs4818-rs4680 |     |          |           |             |             |                 |                 |                   |
| Homozygous-HEA (GCCG/GCGG) | 0.108 | 34 (13) | 20.6 (1.1) | 25.7 (3.4) | 75.7 (10.5) | 1.7 (1.2) | 77.7 (8.8) | 2.1 (1.2)         |
| Heterozygous-HEA      | 0.453 | 143 (57) | 20.5 (0.9) | 25.6 (5.1) | 76.0 (9.1) | 1.7 (1.1)          | 77.9 (8.3) | 2.1 (1.3)         |
| Non-carrier           | 0.439 | 139 (54) | 20.3 (1.0) | 24.8 (6.5) | 76.1 (8.9) | 1.6 (1.0)          | 77.7 (9.6) | 2.1 (1.4)         |

Table 1. Demographic Information, Frequency of COMT Polymorphism, and Face Recognition Performance in COMT Genotype and Haplotype Groups. GF, frequency of genotype; N, number of subjects. Allele frequencies in this sample were very similar to those in the Han Chinese sample from the HapMap dataset (HapMap Data Release 27 Phase II + III). Standard deviation (SD) were put in the parentheses. *Participants with the rs4633 CC genotype showed higher Raven scores than those with the rs4633 CT genotype (p = 0.013, uncorrected); participants with the rs4680 GG genotype showed higher Raven scores than those with the rs4633 AG genotype (p = 0.016, uncorrected).

Figure 2. Comparisons of the group-mean PLS1 rFFA-MRI and PLS1 lFFA-MRI scores among three COMT haplotype groups. The difference in the PLS1 FFA-MRI scores was not significant between the hemispheres. Participants with the heterozygous-HEA COMT haplotype (intermediate dopamine availability) showed increased face selectivity (FS) and decreased fALFF (positive PLS1 FFA-MRI scores), whereas participants with the COMT homozygous HEA haplotype (high DA degrading enzymatic activity: a low level of DA) and HEA non-carriers (low DA degrading enzymatic activity: a high level of DA) showed decreased FS and increased rFFA-fALFF (negative FFA scores of PLS1). Error bars indicate the standard errors of the mean. *p < 0.05, **p < 0.01 (two-tailed).
COMT heterozygous-HEA carriers had significant larger PLS1 rFFA-MRI scores than COMT homozygous HEA carriers (mean difference = 0.011, 95%CI [0.002, 0.021], p = 0.005, pFDR = 0.017) and non-HEA carriers (mean difference = 0.010, 95%CI [0.004, 0.017], p < 0.001, pFDR < 0.001). Thus, participants with the COMT heterozygous-HEA-haplotype (intermediate DA degrading enzymatic activity therefore an intermediate level of DA) showed increased FS and decreased rFFA-fALFF (positive FFA scores of PLS1), whereas participants with the COMT homozygous HEA haplotype (high DA degrading enzymatic activity therefore a low level of DA) and HEA non-carriers (low DA degrading enzymatic activity therefore a high level of DA) showed decreased FS and increased rFFA-fALFF (negative FFA scores of PLS1).

Face-specific recognition ability is associated with a profile of FFA functional activity. To investigate whether the functional rFFA-SNP association contributes to the FRA, we correlated the face recognition accuracy score (hit rate + correct rejection rate) and flower recognition accuracy score with the PLS1 rFFA-MRI scores and the PLS1 COMT scores, respectively. The results showed that face recognition accuracy was positively correlated with the PLS1 rFFA-MRI scores (r = 0.119, uncorrected p = 0.029; left panel of Fig. 3A) when sex, age, and Raven scores were treated as nuisance covariates. No significant association was found between the flower recognition accuracy and the PLS1 rFFA-MRI scores (middle panel of Fig. 3A). Moreover, the association between the PLS1 rFFA-MRI profile and the face recognition accuracy was significantly greater than that between the PLS1 rFFA-MRI profile and the flower recognition accuracy (z = 2.94, two-tailed p = 0.003, Cohen’s d = 0.4, 95%CI = [0.06,0.28]), suggesting that the COMT-modulated rFFA-MRI profile was more relevant to the face recognition ability than to the flower recognition ability. Further, the association between the FRA (the normalized residual of the face recognition score after regressing out the flower recognition score) and the PLS1 rFFA-MRI scores was significant (r = 0.138, uncorrected p = 0.011, 95%CI [0.01,0.22], power = 0.721) when sex, age, and Raven scores were treated as nuisance covariates (right panel of Fig. 3A). The results suggested that a profile of increased FS and decreased fALFF in the right FFA might be associated with a better face-specific recognition ability. In the left FFA, although the association between the PLS1 lFFA-MRI scores and the FRA was not significant (r = 0.094, p = 0.084) (panel C of Supplementary Fig. S4), the associations of the FRA with the PLS1 lFFA-MRI scores and the PLS1 rFFA-MRI scores did not differ significantly (z = 0.966, p = 0.334), suggesting that the COMT-modulated bilateral FFA-MRI profile might have similar association patterns with the FRA. To make our results comparable to those in previous studies using the signal detection theory, we calculated the sensitivity index (d’) for face recognition and flower recognition. No significant correlations were found between face sensitivity (d’) and the PLS1 FFA-MRI score (Supplementary Table 3), suggesting that the COMT-profile modulated PLS1-FFA-MRI profile might be specific, but not sensitive enough, to face recognition.

Finally, because the FFA has been demonstrated to be involved in the process of recognizing familiar and unfamiliar faces, we examined whether there is a difference in modulation of the COMT genotypic profile between old (familiar) face recognition and new (relatively unfamiliar) face recognition. To control the effect of response bias on the accuracy, we calculated the response bias |c = −Z(hit rate) + Z(false alarm rate)|.
ing state activity by synchronizing the low-frequency oscillation. Moreover, dopamine could stabilize the dopamine-function related generic variants imply that dopamine-related neurons may affect the baseline brain rest-activity in healthy people. Some studies focusing on the interaction effect between the possession of a functional characteristic profile of higher face-selective responses and lower intrinsic FFA activities, say, participants with an appropriate level of dopamine (i.e., the heterozygous HEA haplotype carries) in the FFA lobe, are likely modulated by changes in dopamine.

In our study, we explored the neural basis of the heritability of face recognition by investigating associations between 12 COMT genotypes and 3-modal FFA-MRI phenotypes. The results revealed that a profile of face-selective responses and resting-state neural fluctuations of the bilateral FFA were modulated by a profile of COMT polymorphisms. Participants with the COMT heterogeneous HEA haplotype showed higher PLS1 MRI scores, which represented an increased FS and a decreased fALFF and were associated with a better face-specific recognition ability, than those with the COMT homozygous HEA haplotype and HEA non-carrier, suggesting that participants with an appropriate (intermediate) level of dopamine in the bilateral FFA may have better face-specific recognition ability. Our study provides empirical evidence of genetic modulation of a functionally-specialized cortical region in humans.

Previous studies have found that participants with higher face selectivity in the FFA have higher face recognition ability than those with lower face selectivity, and this association is domain-specific. In this study, we revealed that a profile (linear combination) of higher face selectivity BOLD responses, as well as lower resting-state activities in the bilateral FFA, was associated with a higher face recognition (especially the new-face recognition) ability, and the association could not be accounted for by FFA responses to objects such as flowers, suggesting that the face-specific recognition ability might be modulated by a certain covariation pattern of resting-state and task-state activity in the FFA, which was modulated by a certain profile of the COMT polymorphisms.

The COMT Val/Met (rs4680) polymorphism has been suggested to play a role in face perception. Our study extends previous findings by identifying a neural site where COMT affects face recognition. Specifically, we investigated a broader modulation pattern of COMT genetic effects on the FFA by including multiple SNPs. We found that individuals with the heterozygous HEA haplotype exhibited higher face-selective responses but smaller fALFF. In contrast, individuals with the homozygous HEA haplotype or HEA non-carriers had lower face-selective responses and larger fALFF. Because COMT encodes catechol-O-methyltransferase, a protein that degrades catecholamine neurotransmitters and modulates dopamine metabolism, neural activity (including the face-recognition-task-state and resting-state activities) in the FFA is likely modulated by changes in dopamine. Moreover, we found that an intermediate level of dopamine in the FFA was associated with higher face-selective responses (in the task state) and smaller fALFF (in the resting state), and the profile of higher face-selective responses and lower fALFF in the FFA was associated with better face-specific recognition ability. Thus, an intermediate level of dopamine in the FFA is likely beneficial for behavioral performance in face recognition. That is to say, participants with an appropriate level of dopamine (i.e., the heterozygous HEA haplotype carriers) in the FFA possess a functional characteristic profile of higher face-selective responses and lower intrinsic FFA activities, therefore, they might have better face-specific recognition ability.

Previous studies have rarely reported the direct impact of COMT polymorphisms on resting-state FFA activity in healthy people. Some studies focusing on the interaction effect between the COMT rs4680 and other dopamine-function related generic variants imply that dopamine-related neurons may affect the baseline brain resting state activity by synchronizing the low-frequency oscillation. Moreover, dopamine could stabilize the balance between facilitating and suppressing neural signaling in brain networks responsible for behavior-stimuli oriented attention. Thus, an appropriate level of dopamine may improve the functional segregation-integration of a functional-specialized region by maintaining excitation-inhibition stability of neural signal transduction to ensure that individuals with the HEA heterozygous haplotype have a relatively higher rFFA activation in face recognition tasks (which has been found to be related to a better face-specific recognition ability), as well as a relatively lower rFFA activity during resting state (which was inferred from previous research that lower metabolism in the right temporal-parietal cortex is also associated with better cognitive reserve and memory ability in aging people). This is consistent with previous studies demonstrating associations between dopamine hyper- or hypo-function in cortical or subcortical regions and cognitive dysfunctions. Therefore, it would be interesting to examine the level of dopamine in the FFA in individuals with developmental prosopagnosia, who show specific deficits in face recognition with unclear genetic basis and those with mental disorders who might have disturbances in COMT modulation of face processing.

Our study found that the 4 COMT-polymorphisms-modulated PLS1 rFFA-MRI profile mainly regulated the recognition ability of new faces. Although previous studies have revealed that it takes more time and is more likely to make mistakes to recognize faces of different identities than faces of same identities, there seems to be no difference in the increase of neural response activity of the FFA when recognizing familiar and unfamiliar faces. Compared with recognizing unfamiliar faces, recognizing familiar faces involves more activity in the anterior and mid-fusiform gyrus and extended face areas. In our study, the stimuli were not familiar to the subjects, and the social and emotional information involved in the face stimuli was relatively sparse; therefore, the recognition difference between the old face and the new face might be mainly reflected in the familiarity and unfamiliarity of the facial features. We found that the COMT-polymorphisms modulated PLS1 rFFA-MRI profile was more sensitive to the new-face recognition ability than to the old-face recognition ability.
fact that the novel signals can trigger the dopaminergic metabolic system\(^{35}\) in the task-related brain areas and enhance perception\(^{31,32}\).

The FFA is sensitive to the static or invariant properties of faces\(^{49}\). The posterior superior temporal sulcus (pSTS) is responsible for dynamic features\(^{63,64}\) and sensitive to emotional information\(^{65-46}\) from faces, and the prefrontal-amygdala circuit is involved in social and emotional regulation in face processing\(^{47}\). In fact, face perception as a whole is a network involving multiple brain regions\(^{49}\). Studies have found that the COMT Met allele is related to the modulation of neural substrates of negative face emotional bias\(^{22,23}\). Therefore, in future studies, it would be interesting to evaluate the role of multiple COMT polymorphisms in the core face network and its relation with face-specific recognition ability and face-motion recognition.

This study has some limitations. First, although we have adopted multiple strategies to improve the statistical power, the findings still need to be validated using data from other cohorts of participants. Second, measurements for the face recognition ability and the general mental ability are limited to the face memory test and the Raven test, respectively. More multivariate measurements could be introduced in the future to improve the reliability of the individual face recognition ability and general mental ability, and further elaborate the specificity of the effect of COMT on the face recognition ability. Third, all participants had the same ethnicity (i.e., Han Chinese); therefore, how the ethnicity of the individual participants affected the results needs to be examined using data from multiple ethnicities in the future. Fourth, the Caucasian children’ faces were used as stimuli to define the FFA in Han Chinese participants. As demonstrated in previous studies, there is an other-race effect in face recognition; therefore, how our findings are affected by stimulus attributes still needs to be investigated in further studies.

In conclusion, our study demonstrated that the COMT gene affected both resting- and task-state neural activity in the FFA through rs6269-rs4633-rs4818-rs6480 haplotypes, and the COMT-polymorphisms-modulated FFA was associated with face-specific recognition ability. The present findings provide a basis for the behavioral observation of face-specific recognition heritability and the approach used in this study might be a feasible method for exploring and validating the neural basis of heritable behavioral performance. In addition, this study may inform the genetic etiology of developmental prosopagnosia and other mental disorders showing abnormalities in face processing.

**Methods**

**Participants.** Three hundred and sixty-two Han Chinese college students (mean age, 20.47 ± 0.96 years; 144 males) were recruited from Beijing Normal University (BNU) in Beijing, China through campus advertisement. Ethnicity was determined by the ethnic information displayed on the ID card of the participants. All participants had normal or corrected-to-normal vision, and none of them reported a history of neurological or psychiatric disorders. All the experimental protocol and procedures were approved by the Committee for Protecting Human and Animal Subjects of the Faculty of Psychology at Beijing Normal University. All experiments of the study were conducted in accordance with the relevant guidelines and regulations of Beijing Normal University’s Institutional Review Board (Human Subjects Division). All participants gave written informed consent prior to their participation in this study.

**MRI Data acquisition.** Functional and structural MRI was performed at the BNU Imaging Center for Brain Research on a Siemens 3T scanner (MAGNETOM Trio, a Tim system) with a 12-channel phased-array head coil. Resting-state fMRI was acquired using a T2*-weighted gradient-echo-planar-imaging (GRE-EPI) sequence (repetition time [TR] = 2000 ms, echo time [TE] = 30 ms, flip angle = 90°, number of slices = 33, voxel size = 3.125 × 3.125 × 3.6 mm\(^3\)). Resting-state scanning lasted for 8 min and consisted of 240 contiguous volumes. Participants were instructed to relax and remain still with their eyes closed during resting state scanning\(^8\). A GRE-EPI sequence was also used to acquire task-state fMRI (TR = 2000 ms, TE = 30 ms, number of slices = 30, voxel size = 3.125 × 3.125 × 4.8 mm\(^3\)). A high-resolution structural T1-weighted image was acquired with a 3D magnetization-prepared rapid acquisition gradient echo (MP-RAGE) sequence (TR = 2530 ms, TE = 3.39 ms, inversion time = 1100 ms, flip angle = 7°, matrix = 256 × 256, number of slices = 128, voxel size = 1 × 1 × 1.33 mm\(^3\)). Earplugs were used to attenuate scanner noise, and a foam pillow and extendable padded head clamps were used to restrain head motion\(^1\). None of the participants were excluded due to excessive head motion (>2 mm in translation or 2° in rotation) during MRI scanning.

**MRI Data processing.** Structural MRI. Voxel-based morphometry (VBM) was performed using the SPM8 (Statistical Parametric Mapping, Wellcome Department of Imaging Neuroscience, London, UK). First, unified segmentation in SPM8 was used to segment T1-weighted anatomical images into gray matter (GM), white matter (WM), and cerebral spinal fluid (CSF). Second, the Diffeomorphic Anatomical Registration through Exponential Lie algebra (DARTEL) registration method\(^{68}\) was used to build a study-specific GM template and normalize individual GM images into the template. Third, GM voxel values were modulated by multiplying the Jacobian determinants derived from the normalization procedure in order to preserve the tissue volume of each structure. Fourth, modulated GM images were smoothed using an 8-mm full width at half maximum (FWHM) isotropic Gaussian kernel. Finally, the modulated images were masked using absolute masking with a threshold of 0.2 to exclude noisy voxels. Masked-modulated GM images were used for further statistical analyses\(^{69}\).

**Task-state fMRI.** A dynamic face localizer was used to define the face-selective area and derive F5\(^{83}\). Movie clips of close-up faces of 7 Caucasian children (filmed when children were dancing or playing), objects (moving toys), scenes (moving view of a suburb, canyons, or tunnels), and scrambled objects (constructed by scrambling each frame of the object movie clips) were included to examine category selectivity in the ventral visual cortex\(^9\). Participants were instructed to passively view movie clips during scanning. Because Adults’ FFA have been effectively identified by children’ faces in previous studies\(^{45,43,45,67}\) and the FFA is sensitive to the invariant properties
of faces, but not to the dynamic or emotional information of faces\textsuperscript{45–47,70}, we expected that the dynamic children’s faces would accurately and unbiasedly localize the FFA in adults.

In total, participants completed three 198-s runs. Each run consisted of 2 block sets intermixed with 3 18-s rest blocks at the beginning, middle, and end of the run. Each block set contained four 18-s (including six 3-s movie clips) blocks corresponding to 4 stimulus categories. The order of category blocks in each run was palindromic and randomized across runs (Fig. 4A).

Functional images were analyzed with the FMRI Expert Analysis Tool version 5.98, part of FMRIB’s Software Library (www.fmrib.ox.ac.uk/fsl). Preprocessing included head-motion correction, brain extraction, spatial smoothing (Gaussian kernel; FWHM = 6 mm), grand-mean intensity normalization, and high-pass temporal filtering (120-s cutoff). A first-level analysis was performed separately on each run for each participant using FMRIB’s Improved Linear Model with a local autocorrelation correction. A boxcar kernel was convolved with a gamma hemodynamic response function and its temporal derivative was used to model BOLD signal. Six parameters from motion-correction were also included in the model as regressors of no interest to account for the effect of head movement. A second-level fixed-effect analysis was conducted to combine all runs. Specifically, the parameter (i.e., beta) image from the first-level analysis was first aligned to the individual’s structural images using FMRIB’s linear image registration tool with 6 degrees of freedom and then warped to the MNI152 template using FMRIB’s nonlinear image registration tool with default parameters. The spatially normalized parameter images (resampled to 2-mm isotropic voxels) were then summarized across all runs using a fixed-effect model\textsuperscript{3,9,69}. In this study, statistical images from the contrast between faces and objects were used to define the right fusiform face area and derive its face selectivity (FS), which was calculated as the average z-score (faces vs. objects) across all voxels in the defined face-selective region.

**Resting-state fMRI.** Resting-state functional images were preprocessed with a procedure similar to task-state fMRI. Motion correction parameters, mean signals of the CSF and the WM, and the first derivatives of these signals were regressed out to remove the signal fluctuations caused by head motion, cardiac cycle and respiration. Then, the fractional amplitude of low-frequency fluctuations (fALFF), which is defined as the fractional sum of the amplitudes within the low frequency range divided by the sum of amplitude across the entire frequency range (0–0.25 Hz)\textsuperscript{71}, was calculated for each GM voxel. Participant-level voxel-wise fALFF maps were further standardized by subtracting the mean whole-brain voxel-wise fALFF from the participant-level voxel-wise fALFF and dividing it by the standard deviation. Finally, the standardized fALFF was normalized to the MNI152 space with the same normalization procedure as that used in the task-state fMRI analysis.

**Identification of fusiform face area.** A threshold of $Z > 2.58$ (one-tailed $p < 0.005$, uncorrected) was used to define the bilateral FFA based on the random-effect group-level Z-statistic image for faces vs. objects\textsuperscript{3}. Specifically, all contiguous voxels that surpassed the threshold within the posterior fusiform area (defined by the anatomical automatic labeling [AAL] template; http://www.gin.cnrs.fr/AAL) were defined as the FFA (right panel of Figs. 1A and S4).

**Genotyping.** Genomic DNA was extracted from peripheral blood samples of each subject using QuickGene-Mini80 equipment and the QuickGene DNA whole blood kit S (Fujifilm). Sixty-four cognition-related candidate SNPs (including the four COMT polymorphisms of rs6269, rs4633, rs4818, and
rs4680 examined in this study; please refer to Supplementary Table S1 for details of other SNPs) were automatically genotyped using a customized 64 TaqMan® OpenArray® GT Kit (Applied Biosystems; Foster City, CA, USA). Quality control was performed using PLINK software version 1.07. All the 4 COMT SNPs met the following criteria: genotyping call rate >0.95, minor allele frequency (MAF) >0.05, and Hardy–Weinberg equilibrium (HWE) P > 0.05. Participants were excluded from analyses if they had a missing genotype rate of >25% (i.e., the participants who had one of the four COMT polymorphisms missing). Twenty-four participants were excluded, and the remaining 338 participants (mean age, 20.45 ± 0.96 years; 135 males) were included in the data analyses. Allele frequencies in this sample were very similar to those in the Han Chinese sample from the HapMap dataset (HapMap Data Release 27 Phase II + III).

**Face recognition task.** We used an old/new cognitive memory task to measure the face-specific recognition ability (FRA). There was a face block (containing grayscale face pictures of adult Chinese faces with the external contours removed) and a flower block (containing grayscale pictures of common flowers) in this task (face and flower stimuli were shown in Fig. 4B) that were counterbalanced across participants. Each block consisted of one study segment and one test segment. In the study segment, each stimulus (picture) was presented twice (1 second each time), with an inter-stimulus interval of 0.5 seconds. Then, in the test segment, half of the studied images were randomly intermixed with new images from the same category. For each stimulus picture, participants were instructed to indicate whether the image had been shown in the study segment. For a participant, a face recognition accuracy score was calculated as the average proportion of hits and correct rejections for faces, and an object (flower) recognition accuracy score was calculated as the average proportion of hits and correct rejections for flowers. The FRA was calculated as the normalized residual of the face-recognition score after regressing out the flower recognition score. D’ (sensitivity index) was also calculated for face recognition and flower recognition.

**Statistical analyses.** The PLS correspondence analysis (PLSCA) was conducted using the R Package ExPosition (https://cran.r-project.org/web/packages/ExPosition/). PLS is particularly suitable for identifying associations between 2 sets of variables, especially when variables of each set are highly interdependent or multi-collinear. The PLS approach maximizes the covariance between a set of variables (a linear combination of X measures) and another set of variables (a linear combination of Y measures). More specifically, the cross-correlation matrix between measures of SNPs and measures of brain-MRI was first computed. Then, a set of paired latent variables (LVs), which were uncorrelated (orthogonal) with each other and ordered by the explained covariance of brain and SNP measures, were generated via singular value decomposition. LVs are linearly weighted by the raw variables, and these weights or saliences measure the contribution of each raw variable to the LV. Finally, the significance of the variance of singular values was tested by comparison with the distribution of variance arising from random permutation tests (1000 repetitions), and a bootstrap procedure (500 times) was used to test the reliability (i.e., bootstrap ratio, BSR) of each variable in the significant PLS component.

**SNPs-FFA association analysis.** The SNPs-FFA multivariate association analysis was used to examine whether specific patterns of COMT polymorphisms contributed to the multimodal FFA-MRI phenotypes. First, GM, fALFF, and FS within the right FFA and the left FFA ROI were extracted for each participant. Then, each SNP was coded into 3 genotypes (e.g., AA, AG, or GG) as categorical variables, and each variable was weighted according to the information it provided. The weight of a variable was defined as the inverse of its relative frequency because a rare variable provides more information than does a frequent variable. This fully categorical coding scheme allows us to detect linear and nonlinear relationships within (e.g., genotypic) and between datasets. Thus, treating each genotype as a level of a categorical variable is suitable for genetic association studies, especially when the inheritance pattern, direction, and/or size of the effect are unknown. Accordingly, a matrix of measures of 12 COMT genotypes estimated in the 338 participants and a matrix of measures of 3-modal FFA-MRI phenotype estimated in the 338 participants were entered into the PLS analysis.

**Haplotype analysis.** A haplotype block analysis was conducted to investigate how SNPs in strong linkage disequilibrium work as blocks to modulate FFA phenotypes. Haplotype blocks of SNPs affecting the FFA phenotype were assessed and illustrated using Haploview software version 4.2 (https://sourceforge.net/projects/haploview/) with the solid spine of the linkage disequilibrium method and parameter 0.80. Associations of haplotypes with the FFA were examined using R software version 3.5.2 with generalized linear models.

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Author contributions

C.W., Z.Z. and J.L. designed the experiments; C.W., L.H. and T.H. conducted the experiments; C.W. and Z.Z. analyzed the data; C.W., Z.Z. and J.L. wrote the manuscript; and J.L. supervised the project.

Competing interests

The authors declare no competing interests.
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Correspondence and requests for materials should be addressed to Z.Z. or J.L.

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