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# Catechol-O-Methyltransferase Val158Met Polymorphism Is Associated with Anxiety, Depression, and Widespread Pressure Pain Sensitivity in Women with Chronic, but Not Episodic, Migraine

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### 41 Abstract

42 Objective: To analyse the association between the rs4680 Catechol-O-Methyltransferase 43 Val158Met polymorphism and to determine the association of this polymorphism with 44 clinical, psychological and pain sensitivity variables in women with episodic or chronic 45 migraine. Methods: Fifty women with episodic migraine, 50 with chronic migraine, and 46 50 matched healthy women participated. After amplifying Val158Met polymorphism by 47 polymerase chain reaction, we assessed genotype frequencies and allele distributions. 48 Participants were classified according to the Val158Met polymorphism genotype into 49 Val/Val, Val/Met, or Met/Met. A headache diary was used for collecting migraine pain 50 features. Disability was assessed with Migraine Disability Assessment Scale, trait/state 51 anxiety levels with the State-Trait Anxiety Inventory, and depression/anxiety with the 52 Hospital Anxiety and Depression Scale. Pressure pain thresholds (PPT) were bilaterally 53 assessed over the temporalis, the upper trapezius, the second metacarpal and the tibialis 54 anterior. Results: The distribution of rs4680 Val158Met genotype was not significantly 55 different between women with/without migraine (P=0.157). No differences in migraine 56 features were found depending on the Val158Met genotype. Women with the Met/Met 57 genotype showed higher migraine-related disability than those with Val/Val or Val/Met 58 genotype in both migraine groups (P < 0.01). Women with chronic, but not episodic, 59 migraine with the Met/Met genotype exhibited higher depressive and anxiety levels and 60 lower PPTs than those with Val/Val or Val/Met genotype. Conclusion: The Val158Met 61 rs4680 polymorphism does not appear to be involved in predisposition to suffer from 62 migraine; however, this genetic factor may be involved in the phenotypic expression of 63 chronic migraine, since anxiety, depression and widespread pressure pain sensitivity 64 was greater in those women with chronic, but not episodic, migraine with the Met/Met 65 genotype.

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# Catechol-O-Methyltransferase (COMT) Val158Met Polymorphism is Associated with Anxiety, Depression and Widespread Pressure Pain Sensitivity in Women with Chronic, but not Episodic, Migraine

# 83 Introduction

Migraine is a primary headache disorder with a worldwide prevalence of 11.6% (13.8% females/6.9% males) (1). In the last Global Burden of Disease Study, migraine and tension-type headache were found to be the second most prevalent pain conditions in the world (2). In fact, general costs of headaches in Europe (€13.8 billion) mainly account for migraine and tension type headache (3).

It is accepted that the pathophysiology of migraine is associated with abnormal neuronal excitability leading to cortical spreading depression and to central sensitization of trigemino-vascular pathways (4). There are several factors that could affect the pain processing. One of these factors is genetics. Different genetic epidemiological studies have investigated the familial aggregation in migraine and it seems that an hereditary component can be present in some migraine types, i.e. hemiplegic migraine (5,6).

95 The catechol-O-methyltransferase (COMT) gene is one of the potential genetic 96 determinants in chronic pain (7). The COMT is an enzyme involved in the metabolic 97 degradation of several neurotransmitters, e.g., dopamine, norepinephrine, or epinephrine 98 (8). The activity of the COMT gene can be affected by different polymorphisms such as 99 rs4680, rs6269, rs4633, or rs4818. It seems that the rs4680 genetic polymorphism due to 100 a  $G \rightarrow A$  substitution at codon 158 of this gene, leading to a value (Val) to methionine 101 (Met) substitution, will result in differences within COMT gene activity related to pain 102 sensitivity. In fact, a valine (Val) allele at codon 158 results in a high-activity variant 103 (Val/Val) whereas a methionine (Met) at this codon position (Val/Met, Met/Met) results 104 in low-activity variants (9). It has been found that subjects with the Met/Met genotype 105 exhibit higher pain sensitivity, that is, lower pain thresholds to different stimuli (10,11),

and different brain responses to painful stimuli (12) than those subjects with the Val/Val
genotype, supporting a role of this gene in nociceptive pain processing.

108 There are several studies investigating the role of Val158Met polymorphisms in 109 migraine; although the results are inconsistent. The most recent meta-analysis did not 110 observe a significant association between the Val158Met polymorphism and migraine 111 (13). Similarly, a recent study, not included in the abovementioned meta-analysis, did 112 not also reveal differences in Val158Met polymorphism distribution between subjects 113 with migraine and healthy controls (14). Based on current evidence, it would seem that 114 Val158Met polymorphism (rs4680) is not associated to a higher risk of suffering from 115 migraine. However, it should be noted that most studies did not differentiate between 116 episodic and chronic migraine. Similarly, another study including subjects with chronic 117 migraine did not also find an association of the rs4680 Val158Met polymorphism with 118 this subgroup (15).

119 Although no differences in Val158Met polymorphism distribution would exist 120 between individuals with and without migraine, there is evidence suggesting a genetic 121 influence of the COMT enzyme in several aspects of different chronic pain conditions, 122 e.g., related-fatigue and pressure pain sensitivity in breast cancer survivors (16) or mood 123 disorders (anxiety and depression) in women with fibromyalgia (17). Therefore, it is 124 possible that the Val158Met polymorphism can also influence some phenotypic aspects 125 in patients with migraine. In line with this hypothesis, Park et al found that individuals 126 with migraine carrying the Met allele experienced worse migraine-associated nausea 127 and vomiting and higher pain intensity of migraine attacks than those with the Val allele 128 (18). No previous study has investigated the role of the Val158Met polymorphism in 129 clinical, psychological and pain sensitivity outcomes in women with migraine.

Therefore, the aims of the current study were: 1) to investigate the association of the Val158Met polymorphism in women with episodic or chronic migraine; and 2) to determine the relevance of the Val158Met polymorphism with clinical, psychological, and pain sensitivity variables in women with migraine.

135

#### 136 Methods

# 137 **Participants**

138 One hundred and twenty consecutive women with migraine were recruited from 139 a Headache Unit located in a tertiary university-based hospital. They were diagnosed 140 following the third edition of International Headache Society (ICHD-III) criteria down 141 to third-digit level (code 1.1, 1.3) by an experienced neurologist (19). Migraine features 142 including location, quality of pain, years with disease, frequency and intensity of pain 143 attacks, family history, and medication intake were collected. To be included, subjects 144 had to describe typical pain features of migraine pain (unilateral location, pulsating 145 pain, high intensity, and aggravation during physical activity) and associated symptoms 146 including photophobia, phonophobia, mild nausea or vomiting (19).

147 Participants were excluded if they presented any of the following: 1) other primary 148 or secondary headache, including medication overuse headache; 2) history of cervical or 149 head trauma; 3) pregnancy; 4) history of cervical herniated disk or cervical osteoarthritis 150 on medical records; 5) any systemic medical disease, e.g., rheumatoid arthritis, lupus 151 erythematous; 6) comorbid fibromyalgia syndrome; 7) had received treatment including 152 anesthetic blocks, botulinum toxin or physical therapy within the previous 6 months; or, 153 8) male gender. All participants were carefully interviewed for assessing their medical history. Further a medical exam, including neuro-imaging examination (MRI or CT) of 154 155 the head, was performed in all patients in order to identify any exclusion criteria.

Age-matched healthy women without history of headache diagnosis and without reporting a headache pain attack during the previous year were also included. Exclusion criteria for the control group were the same as for the headache groups. All participants signed the informed consent form before their inclusion in the study. The local Ethics Committee of Hospital Rey Juan Carlos, Spain (HRJ 07/14) approved the study design.

# 161 DNA Collection and COMT Genotyping

Non-stimulated whole saliva samples were collected into collection tubes (passive drooling technique) according to standardized procedures. Saliva collections were made when participants were headache-free, or with a migraine intensity of less than 3 points (in those patients with high frequency of attacks). Immediately after collection, samples were centrifuged at 3000 rpm for 15min to obtain the cell sediment and they were stored at -20° C until the analysis. We prefer to use saliva instead of blood sampling because salivary collection is a non-invasive, stress-free and ethic suitable assessment method.

169 Laboratory technicians were blinded to the subject's condition. Genomic DNA 170 was hence extracted from saliva cell sediments using the "Genomic DNA extraction and 171 purification Kit" (Real Molecular Biology) following the manufacturer instructions. The 172 single Val158Met (rs4680) nucleotide polymorphism was genotyped using a TaqMan® 173 Drug Metabolism Genotyping Assays on a Real Time PCR ABI Prism 7000 Sequence 174 Detection System (APPLIED BIOSYSTEM, USA) in the Genomic Unit at the Centro 175 de Apoyo Tecnológico Universidad Rey Juan Carlos, Madrid (Spain). The 3 possible 176 allelic variants were associated with different fluorescent dyes to proper identification 177 of the different genotype forms: Val/Val (H/H), Val/Met (H/L), or Met/Met (L/L). The 178 results are derived from a  $G \rightarrow A$  substitution at the following sequence:

179 CCAGCGGATGGTGGATTTCGCTGGC [A/G] TGAAGGACAAGGTGTGCATGCCTGA

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## 182 Migraine Features

A 4-weeks headache diary was used to register clinical features of migraine (20). The diary was used to calculate the following variables: 1, migraine intensity, calculated from the mean intensity of the days with s migraine attack as assessed with a 11-point numerical pain rate scale (21) (NPRS; 0: no pain, 10: maximum pain); 2, migraine frequency (days/month); and 3, migraine duration, calculated by dividing the total hours of the attack by the number of days with migraine (hours/attack).

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# Psychological and Disability Variables

The Hospital Anxiety and Depression Scale (HADS) was used to evaluate the levels of anxiety and depression. This questionnaire includes 7-items scored at a 4points scale for assessing anxiety (HADS-A) and other 7-items for assessing depressive (HADS-D) symptoms (22). Both scales are considered reliable and valid for assessing anxiety (Cronbach's  $\alpha$ : 0.83) and depression (Cronbach's  $\alpha$ : 0.82) (23). In subjects with headache, the HADS has shown good internal consistency (Cronbach's  $\alpha$ : 0.84) (24).

The State-Trait Anxiety Inventory (STAI) is a 40-items scale assessing separate dimensions of state anxiety (items 1-20, STAI-S) and trait anxiety (items 21-40, STAI-T) (25,26). The STAI-S assesses relatively enduring symptoms of anxiety at a moment, and the STAI-T scale measures a stable propensity to experience anxiety and tendencies to perceive stressful situations as threatening. Both scales have exhibited good internal consistency ( $\alpha$ : 0.89) and high reliability (ICC: 0.88) (27). Higher scores in both scales indicate greater levels of state or trait anxiety.

To assess the degree of related-disability in daily activities (work or school, family and social) caused by migraine, we used the Migraine Disability Assessment Scale (MIDAS) questionnaire. It consists of 5 questions related to days of partial or total loss within the last 3 months regarding 3 main activities: 1, paid work or school; 207 2, household chores; 3, family, social, or leisure activities (28). The final score comes 208 from the sum of the missed days regarding the 3 activities.

209

# Widespread Pressure Pain Sensitivity

210 The evaluation was held when patients were headache-free or, in those with high 211 frequency of migraine, when the intensity of headache was less than 3 points on NPRS. 212 Participants were asked to avoid any analgesic or muscle relaxant 24 hours prior to the 213 examination. No change was made on the prophylactic treatment of the patients. All the 214 participants attended a session for familiarization with the pressure test procedure over 215 the wrist extensor muscles.

216 Pressure pain thresholds (PPTs), i.e. the minimal amount of pressure where a 217 sensation of pressure changes to pain (29), were assessed with an electronic algometer 218 (Somedic AB, Farsta, Sweden). The pressure was applied perpendicularly to the point at 219 a rate of approximately 30 kPa/s. Participants were instructed to press the "stop button" 220 when the sensation first changed from pressure to pain. The mean of 3 trials on each 221 point was calculated and used for the main analysis. A 30sec resting period was allowed 222 between trials for avoiding temporal summation (30). The reliability of pressure 223 algometry has been found to be high (31,32).

224 To determine widespread pressure pain sensitivity, PPTs were bilaterally assessed 225 over a trigeminal point (i.e. temporalis muscle), an extra-trigeminal point (i.e. C5/C6 226 joint), and two distant pain-free points (i.e. the second metacarpal and tibialis anterior 227 muscle) by an assessor blinded to the individual's condition. The order of assessment 228 was randomized between participants. Since no side-to-side differences were observed, 229 mean of both sides were used in the analysis.

230

#### 232 Sample Size Calculation

Sample size determination and calculations were based on detecting a moderate-large effect size of 0.7 on PPTs between migraine and healthy controls related to Val158Met genotype distribution, a 2-tailed test, with an alpha level ( $\alpha$ ) of 0.05, and a desired power ( $\beta$ ) of 90%. This generated a sample size of, at least, 42 participants per group.

237 Statistical Analysis

238 Data were analyzed with the SPSS statistical package (22.0 Version). Results are 239 expressed as mean and 95% confidence interval (95% CI). The Kolmogorov-Smirnov 240 test showed that all quantitative variables showed a normal distribution of the data (P >241 0.05). Comparisons of genotype distribution and allele frequency among groups were performed on raw frequencies using an extended chi-squared test ( $\chi^2$ ). A  $\chi^2$  analysis of 242 243 the Hardy-Weinberg equilibrium for the genotypes was conducted to determine whether 244 the allele frequencies were stable within all groups. A 2x2 analysis of variance ANOVA 245 was used to compare clinical and psychological variables according to the Val158Met 246 polymorphism genotype (Val/Val, Val/Met, Met/Met) in women with migraine (episodic, 247 chronic). A 3x3 mixed-model ANOVA was used to investigate differences in PPTs over 248 each point (temporalis, C5-C6 joint, second metacarpal, tibialis anterior) according to 249 the Val158Met genotype (Val/Val, Val/Met, Met/Met) and group (episodic migraine, 250 chronic migraine, healthy control). Post-hoc analyses comparisons were conducted with 251 the Bonferroni test. The statistical analysis was conducted at a 95% confidence level. A 252 P value less than 0.05 was considered statistically significant.

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254

# 256 **Results**

257 One hundred and twenty (n=120) consecutive women presenting with headache 258 were screened for eligibility criteria. Twenty (17%) were excluded for the following 259 reasons: co-morbid headaches (n=7); previous head or neck trauma (n=6); receiving 260 anaesthetic block in the past 3 months (n=5) or pregnancy (n=2). Finally, 50 women 261 with chronic migraine (age:  $43\pm12$  years), 50 with episodic migraine (age:  $42\pm13$  years) 262 satisfied all criteria, signed the informed consent, and agreed to participate. Further, 50 263 age-matched women without headache (age: 43±11 years) were also included. Table 1 264 summarizes clinical, psychological and pain sensitivity data of the sample. Women with 265 chronic migraine exhibited significant higher headache frequency (P<0.001) and higher 266 migraine-related disability (P=0.04) than those with episodic migraine. Further, women 267 with episodic or chronic migraine exhibited higher widespread pressure pain sensitivity 268 (P < 0.001) than healthy women, without differences between them (P > 0.9).

# 269 Distribution of Val158Met Polymorphism in migraine

The genotype distributions in women with and without migraine did not deviate from those expected based on the Hardy-Weinberg equilibrium. The distribution of the Val158Met genotypes ( $\chi^2$ =6.63; P=0.157) was not significantly different among women with episodic or chronic migraine and healthy women (**table 2**).

# 274 Clinical and psychological measures and Val158Met Polymorphism

The mixed-model ANOVA did not reveal significant differences depending on the Val158Met polymorphism genotype (**table 3**) in both groups of migraine women for years with pain (F=0.874; P=0.420), migraine intensity (F=0.172; P=0.842), migraine frequency (F=1.986; P=0.143), and migraine duration (F=0.308; P=0.736).

280 Similarly, no significant differences depending on the Val158Met polymorphism 281 genotype were either found (table 4) in both women with episodic or chronic migraine 282 for STAI-T (F=0.340; P=0.712), and HADS-A (F=1.494; P=0.188). A significant group 283 \* Val158Met genotype interaction was observed for HADS-D (F=4.369; P=0.015) and 284 STAT-S (F=3.219; P=0.045): women with chronic migraine, but not those with episodic 285 migraine, carrying the Met/Met genotype showed higher depressive and anxiety state 286 levels than those carrying the Val/Val (P=0.01) or Val/Met (P=0.04) genotype. Finally, 287 significant differences based on the Val158Met polymorphism genotype for the MIDAS 288 (F=7.078 P<0.001) were found in both migraine groups: women carrying the Met/Met 289 genotype exhibited higher levels of related-disability than those with the Val/Val or the 290 Val/Met genotype (P<0.01) in both episodic and chronic migraine groups (table 4).

# 291 Pressure pain sensitivity and Val158Met polymorphism

292 All patients with episodic migraine and 45 (90%) patients with chronic migraine 293 were headache-free during PPT examination. The 3x3 mixed-model ANOVA revealed 294 significant group\*Val158Met polymorphism genotype interactions for PPTs over the 295 temporalis muscle (F=3.714; P=0.025), the second metacarpal (F=3.641; P=0.024), and 296 tibialis anterior (F=3.431; P=0.03), but not for the C5-C6 zygapophyseal joint (F=1.479; 297 P=0.212). Women with chronic migraine with the Met/Met genotype showed lower PPT 298 than women with chronic migraine with the Val/Met or Val/Val genotype (P<0.001). 299 No significant differences existed in PPTs between women with chronic migraine with 300 the Val/Val or Val/Met genotypes (P>0.5). Table 5 shows PPT according to Val158Met 301 polymorphism in women with episodic and chronic migraine and healthy women. 302

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# 305 **Discussion**

The current study found no differences in the genotype distribution and allele frequency of the Val158Met polymorphism between those women with migraine, either episodic or chronic, and healthy women. Further, the presence of the Met/Met genotype was associated to higher levels of anxiety, depression, disability and greater widespread pressure hyperalgesia, in women with the chronic, but not episodic, form of the disease.

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# Val158Met polymorphism and migraine

312 We did not observe significant differences in the distribution of the Val158Met 313 polymorphism between women with episodic or chronic migraine and healthy women, 314 supporting the assumption that this polymorphism is not involved in a predisposition to 315 suffer from migraine. Our results agree with a recent systematic review concluding that 316 the Val158Met polymorphism was not associated with migraine risk (13). Additionally, 317 Takigawa et al did not also observe differences in the presence of other haplotypes of 318 the COMT gene, e.g., rs4633, rs6267, rs6270 between individuals with migraine and 319 healthy people (14). Nevertheless, since the rs4680 Val158Met polymorphism has been 320 associated, in some studies, to different conditions, e.g., fibromyalgia syndrome (33) or 321 temporomandibular pain (34), it is possible that it could be associated to some particular 322 pain conditions rather than to chronic pain in general. Furthermore, since migraine is 323 comorbid with other chronic pain syndromes, i.e. fibromyalgia (35), we do not know if 324 different subgroups of patients with migraine and co-morbid conditions would lead to 325 different associations. Obviously, the fact that the rs4680 Val158Met polymorphism is 326 not associated with migraine does not exclude the role of genetics in this headache 327 form. Therefore, future studies investigating the role of other genetic components in 328 migraine are guaranteed.

329 It has been previously that the Val158Met can be associated with worse clinical 330 presentation of migraine. For instance, individuals with migraine carrying the Met allele 331 experienced higher pain intensity and worse migraine-associated symptoms than those 332 with Val allele (18). We observed that women with migraine, either episodic or chronic, 333 with the Met/Met genotype exhibited higher migraine-related disability as assessed with 334 the MIDAS than those with Val/Met or Val/Val genotype. Further, a Met/Met genotype 335 was also associated with higher depressive and anxiety state levels, but only within the 336 chronic migraine group, suggesting that the Val158Met polymorphism can play a role in 337 different psychological aspects. In fact, our results agree with previous studies showing 338 that the Met allele is associated with anxiety-related behaviors in healthy women (36), 339 with higher stress responses after a whiplash injury (37), or with higher psychological 340 distress in fibromyalgia syndrome (38). A potential explanation for these findings could 341 be related to the fact that individuals carrying the Met/Met genotype had greater brain 342 activation of the limbic region as response to emotionally challenging situations (39,40). 343 Additionally, Met/Met carriers exhibited lower activation of the dorso-lateral pre-frontal 344 cortex and cingulate cortex than Val/Val carriers (41). Therefore, it is also possible that 345 individuals with the Met/Met genotype exhibit different cortical activation patterns than 346 those carrying the Val/Val genotype.

# 347 Val158Met polymorphism and pain hyperexcitability

Another relevant finding of the current study is that women with chronic, but not episodic, migraine carrying the Met/Met genotype exhibited higher widespread pressure pain sensitivity than those with the Val/Val or Val/Met genotype. These findings would suggest that the Val158Met polymorphism could play a role within the nociceptive pain processing in the chronic form of the disease. A potential association of the Val158Met polymorphism with higher sensitivity to pressure pain has been previously observed in 354 children with chronic tension type headache (42) and women with fibromyalgia (43). 355 Our study is the first reporting an association between the Val158Met polymorphism 356 and widespread pressure pain sensitivity in chronic migraine. Several mechanisms could 357 explain this association. For instance, a reduction within COMT gene activity associated 358 with the Met allele at codon 158 of the Val158Met leads to a reduction in the content of 359 enkephalins in some regions of the central nervous system associated with pain (9). This 360 hypothesis would correlate with the presence of hyper-excitability of the central nervous 361 system and of endogenous inhibitory pain pathways previously observed in adults with 362 chronic migraine (44). In fact, migraine has been associated with a non-physiological 363 production of some neuromodulators (45). Therefore, another potential mechanism may 364 be an increase of catecholamine levels, which will promote stimulation of β2-adrenergic 365 receptors in the central nervous system, associated with a reduced COMT gene activity 366 (46). Since individuals with migraine exhibit hyper-excitability of the central nervous 367 system, it is possible that the presence of the Met/Met genotype, in some predisposed 368 subjects, could contribute to this process. In fact, this hypothesis is also suggested in 369 subjects with fibromyalgia (47).

# 370 Limitations

371 Although the results of this study are informative, potential limitations should be 372 considered. First, we included women with migraine and derived from a specialized 373 tertiary hospital center. Therefore, our results should be not extrapolated to men and to 374 other primary headaches such as tension-type headache. Second, it is possible that the 375 study was underpowered for other outcomes different than PPTs. Therefore, a greater 376 sample size including patients from the general population would be needed to further 377 confirm these results. Third, we only investigated the rs4680 nucleotide of Val158Met 378 polymorphism. Future studies should include a greater number of polymorphisms and

other genes to further clarify their potential role in the phenotypic expression of chronicmigraine.

# 382 Conclusions

No differences were found in the genotype distribution and allele frequency of the Val158Met polymorphism between women with migraine, either episodic or chronic and healthy women. The presence of the Met/Met genotype was associated with higher related-disability in both episodic and chronic migraine, and with higher depressive and anxiety levels, and higher pressure pain hyperalgesia but only in the chronic migraine group. Our results suggest that the rs4680 Val158Met polymorphism may contribute to the altered nociceptive pain processing in women with chronic migraine and may contribute to the chronification process.

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