Evidence That Brain MAO A Activity Does Not Correspond to MAO A Genotype in Healthy Male Subjects

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Background: A functional polymorphism in the promoter region of the monoamine oxidase A (MAO A) gene has two common alleles that are referred to as the high and low MAO A genotypes. We report the first in vivo human study to determine whether there is an association between MAO A genotype and brain MAO A activity in healthy male subjects.

Methods: Brain MAO A activity was measured with positron emission tomography and [11C]clorgyline in 38 healthy adult male nonsmokers genotyped for MAO A polymorphism.

Results: There was no significant difference in brain MAO A activity between the high (n = 26) and low (n = 12) MAO A genotypes.

Conclusions: The lack of an association between the high and low MAO A genotype and brain MAO A activity suggests that this polymorphism by itself does not contribute to differences in brain MAO A activity in healthy adult male subjects.

Key Words: Brain MAO activity, MAO A genotype

here is a well-characterized variable number tandem repeat (VNTR) functional polymorphism in the promoter region of the monoamine oxidase A (MAO A) gene that has two common alleles (4-repeat and 3-repeat) that occur in a \sim 60:40 ratio in male humans (Sabol et al. 1998). These are referred to as high and low MAO A genotypes, defined by their significantly different transcriptional activities in human nonneuronal cell lines. Because of the importance of MAO in the regulation of monoamine levels and on mood and behavior, the MAO A genotype has been investigated as a variable in many clinical and behavioral phenotypes (Chen 2004; Shih et al. 1999). In particular, there appears to be a link between the low MAO A genotype and impulsive behavior (Manuck et al. 2000) through interaction with early environmental stressors (Caspi et al. 2002; Huang et al. 2004; Kim-Cohen et al. 2006). The relationship between MAO A genotype and brain MAO A activity has never been examined in vivo in healthy human subjects, however. The

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purpose of this study was to determine whether high and low MAO A genotypes correspond to high and low brain MAO A activity in normal healthy male subjects using the MAO A radiotracer, [11C]-clorgyline and positron emission tomography (PET; Fowler *et al.* 1996a, 2001).

Methods and Materials

Subjects

Thirty-eight healthy male nonsmokers were recruited specifically for this study. Nonsmoking status was ascertained by breath carbon monoxide measurement. Exclusion criteria included female gender due to a distribution of 9:1 of phenotypes (Caspi *et al.* 2002), current or past psychiatric or neurologic disease, history of drug or alcohol abuse, dependence on any substance other than caffeine, positive urine screen for drugs of abuse, history of head trauma with loss of consciousness, history of cardiovascular or endocrinologic disease, and current medical illness. Subjects were interviewed for age, education, and socioconomic status (SES) and assessed for intelligence (Table 1). Written informed consent was obtained as approved by the local institutional review board.

Procedures

Brain MAO A activity was measured in each subject with PET and [11 C]clorgyline (average dose 6.2 \pm .7 mCi; specific activity 250 mCi/ μ mol) using the scanning protocol reported previously (Fowler *et al.* 2001). Briefly, PET images were acquired using a whole body, high-resolution PET (Siemen's HR+) in 3-dimensional dynamic acquisition mode.

From each subject, DNA for MAO A genotyping was obtained from cheek swab samples (Freeman *et al.* 2003). Polymerase chain reactions (PCRs) were performed as described by Sabol *et al.* (1998), and PCR products were analyzed on an Applied Biosystems 3100 Genetic analyzer. Alleles were seen in expected ranges using Genescan version 3.7 and Genotyper version 3.6 software.

Data Analysis

Ten regions of interest (ROIs; medial frontal cortex, dorsolateral prefrontal cortex, anterior cingulate gyrus, primary visual

Table 1. Demographic Characteristics and Brain Monoamine Oxidase A (MAO A) Levels (λk_3) for 38 Study Participants

| | High MAO A Genotype ($n = 26$) | Low MAO A Genotype ($n = 12$) |
|---|----------------------------------|---------------------------------|
| Age | 31.15 ± 6.7 | 32.9 ± 6.4 |
| Education | 14.96 ± 2.3 | 15.4 ± 1.3 |
| Socioeconomic Status | | |
| (Hollingshead) | 44.04 ± 10.6 | 43.17 ± 14.04 |
| Intelligence | | |
| WRAT-3 Reading ^a | 102.76 ± 25 | 101.67 ± 13 |
| WASI MATRIX Reasoning ^b | 11.42 ± 2.02 | 11.4 ± 2.4 |
| Ethnicity | | |
| White | 18 | 4 |
| African American | 4 | 5 |
| Hispanic | 4 | 2 |
| Asian | 0 | 1 |
| MAO A (λk_3) mL g ⁻ min ⁻¹ | | |
| Medial Frontal Cortex | $.273 \pm .037$ | $.273 \pm .039$ |
| Dorsolateral Prefrontal | | |
| Cortex | $.279 \pm .046$ | $.264 \pm .049$ |
| Anterior Cingulate Gyrus | $.299 \pm .044$ | $.289 \pm .036$ |
| Primary Visual Cortex ^c | $.350 \pm .055$ | $.307 \pm .048$ |
| Temporal Cortex | $.293 \pm .039$ | $.289 \pm .034$ |
| Precuneus | .319 ± .047 | $.293 \pm .046$ |
| Caudate Nucleus | $.228 \pm .039$ | $.230 \pm .045$ |
| Putamen | $.266 \pm .045$ | $.262 \pm .042$ |
| Thalamus | $.389 \pm .053$ | $.376 \pm .062$ |
| Pons | .311 ± .057 | $.312 \pm .045$ |

 $^{^{}a}$ Wide Range Achievement Test III Reading subscale; estimate of verbal IO

cortex, temporal cortex, precuneus, caudate nucleus, putamen, thalamus, and pons) were selected. Right and left regions were averaged.

PET time-activity data for [11 C]clorgyline from different brain regions and time-activity data in arterial plasma were used to calculate the model term K_I , the plasma to brain transfer constant which is related to blood flow and λk_3 , which is a function of the concentration of catalytically active MAO A molecules (Fowler *et al.* 2001). The model term k_3 is related to the binding of [11 C]clorgyline by MAO A; λ is defined as K_I/k_2 and is independent

dent of blood flow and k_2 is related to the efflux of tracer from brain to blood.

In our statistical analysis, we first examined the normality assumption of the continuous variables using the Shapiro–Wilk test. Depending on the normality of the data, the parametric or nonparametric test was applied. We compared model terms K_I and λk_3 for the high and low MAO A genotypes using the Wilcoxon Rank Sum Test for the 10 brain regions (because most of these regions are not normally distributed) setting the significance level at p=.005 (Bonferroni corrected). The nonparametric Kruskal–Wallis test was applied to examine whether ethnicity would influence K_I and λk_3 among the high or low MAO genotype. All tests were two-tailed.

Results

The genotype distribution for the 38 male subjects recruited for this study was consistent with a previous study (Sabol *et al.* 1998) with 26 (68%) classified as the high and 12 (32%) classified as the low MAO A genotype. The two groups were well matched for age, socioeconomic status, education, and intelligence but not ethnicity (except for African Americans; Table 1).

For all subjects, [\$^{11}\$C]clorgyline binding was highest in the thalamus and visual cortex and lowest uptake was in the caudate nucleus (Figure 1). We found no significant difference in K_I or λk_3 (which is proportional to MAO A activity) between the high and low MAO A genotypes in any brain region (Table 1) although there was a trend for a small difference for λk_3 in the visual cortex (p = .026, high > low). There were no ethnic differences in absolute measures of λk_3 for all subjects and between the MAO A genotyped groups.

Discussion

The prevalence of the high and low MAO A genotypes in the male population has stimulated many studies on the association of MAO A genotype with impulsivity, inhibitory control, and aggression (Huang *et al.* 2004; Manuck *et al.* 2000; Passamonti *et al.* 2006). Of particular interest are a number of studies showing that MAO A genotype influences vulnerability to environmental stress both in humans (Caspi *et al.* 2002) and animals (Newman *et al.* 2005) and that this biological process can be initiated early in life (Kim-Cohen *et al.* 2006). In addition, the low MAO A genotype was recently reported to be associated with pro-

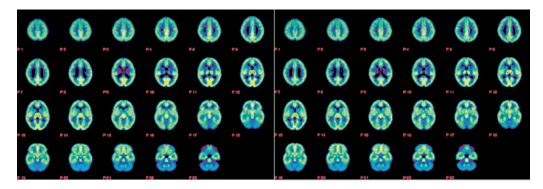


Figure 1. Averaged parametric images (λk_3) of the high (left panel) and the low (right panel) MAO A genotype groups showing 23 transaxial planes from the top of the head to the base of the skull and using a rainbow color scale where red indicates regions of highest monoamine oxidase A (MAO A) activity and blue indicates regions of lowest MAO A activity. Note the similarity in image intensity between the two groups, corroborating the findings of the region of interest analysis (Table 1).

^bMatrix Reasoning from the Wechsler Abbreviated Scale of Intelligence; estimates of nonverbal/fluid IQ.

 $^{^{}c}p = .026 \text{ (high > low)}.$

nounced limbic volume reductions (ventral cingulate and amygdala; Meyer-Lindenberg et al. 2006).

A possible biochemical link underlying genotype-phenotype associations is that high and low MAO A genotypes may be associated with high and low levels of brain MAO A. Monoamine oxidase A is an enzyme that regulates the concentration of the MAO A-specific substrates-serotonin and norepinephrine, as well as dopamine. A number of direct studies in nonneuronal cell lines and in postmortem human brain provide conflicting results. For example, normalized MAO A specific activity from the high MAO A genotype has been reported to be significantly higher than that for the low MAO A genotype in human fibroblast cultures (Denney et al. 1999). In contrast, a direct assay of MAO A from cortical autopsy samples from 31 human subjects did not find a significant association of any single MAO A polymorphism with expression levels of MAO A (Balciuniene et al. 2002). Whether the high and low MAO A genotypes (which are defined in terms of their respective transcriptional activities) are associated with differences in MAO A (the gene product) has not been determined in vivo, however.

Here we report the first in vivo measurement of brain MAO A activity in 38 healthy adult male volunteers of high (n = 26) and low (n = 12) MAO A genotypes matched for age, SES, education, intelligence, and without smoking or drinking histories. We found no significant difference between MAO A genotype and brain MAO A activity in any brain region examined except for a trend for a small elevation in the visual cortex (high > low). The sample size in this study would guarantee a power of 76% for the Wilcoxon Rank Sum Test to detect the difference between the groups at the magnitude of effect size 1 (ratio between the mean difference and the pooled standard deviation) at the significance level of .05 (two-sided). In our study, the pooled standard deviation is about 13%-18% of the population mean. Thus, we have a power of 76% to detect a mean difference at the magnitude of 13%-18% of the individual population mean. This indicates that the effect of the MAO A genotype in MAO A activity in the healthy adult brain, if any, is smaller than the variability in MAO A brain concentration between adult subjects. This could reflect that either MAO A genotype has no effect on MAO A activity or that other factors (environmental, developmental, other genes) have a greater effect in modulating brain MAO activity in the adult brain. We note that other PET studies have also addressed the effect of genotype on brain levels of associated protein products such as the dopamine transporter, the dopamine D2 receptor, and the serotonin transporter. Overall, these studies, which used different tracers and different study populations, show that the effect of genotype on protein expression in the adult brain is inconsistent, and those that show differences report small differences (Martinez et al. 2001; Parsey et al. 2006; Shioe et al. 2003; van Dyck et al. 2004).

These findings suggest that variables other than baseline MAO A regional activity in the adult brain need to be considered in explaining gene behavior, gene-brain function, and gene-brain structure relationships in healthy individuals. One possibility is that the influence of the MAO A genotype may occur predominantly during brain development in the fetal and postnatal periods. The MAO A genotype may be the main variable regulating developmental catecholamine levels including those of serotonin, known to be crucial for brain development particularly because MAO B develops later than MAO A and would not be present to compensate for deficient MAO A (Shih et al. 1999). Compelling evidence suggests that the absence of MAO A during development results in an aggressive phenotype in both animals

(Cases et al. 1995; Mejia et al. 2002; Whitaker-Azmitia et al. 1994) and humans (Brunner et al. 1993). In contrast there is no evidence of aggression as a side effect in adults treated with MAO A-inhibiting drugs, and in rodents MAO A inhibition in adulthood reduces stress-induced aggression (Ossowska et al. 1999). Stress-induced monoamine surges could be particularly damaging during fetal and childhood development. Thus, the determination of brain MAO A levels corresponding to high and low MAO A genotype at different developmental stages as well as their interaction with environmental factors such as smoking that can further inhibit brain MAO (Fowler et al. 1996a, 1996b) during pregnancy merits further investigation. Similarly, future studies to evaluate the relationship between MAO A genotype and brain MAO A levels in diverse neuropsychiatric populations will allow determination of whether MAO A in the adult brain is differentially regulated in some neuropsychiatric diseases.

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