Genetic Variation in the Neuropeptide Y Gene Promoter Is Associated with Increased Risk of Tobacco Smoking

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Abstract
Background: Neuropeptide Y (NPY) is a strong candidate gene regarding the pathophysiology of tobacco dependence. It has been associated with various addictive and psychiatric disorders, and closely interacts with the brain reward system. The aim of the present study was to test for association between a functional genetic variant in the NP-Y promoter gene (SNP rs16147) and tobacco smoking. Methods: In a population-based case-control multicenter study designed for tobacco addiction research, a total of 550 Caucasian current smokers, and 544 never-smokers were genotyped for SNP rs16147 and behaviorally characterized with the State-Trait Anxiety Inventory (STAI). Results: Subjects with TT genotype of the SNP rs16147 were significantly more frequently smokers than never-smokers (p = 0.046). In addition, TT genotype exhibited increased state anxiety scores compared to carriers of the C allele (p = 0.037). Conclusions: Our results provide evidence for an involvement of the functionally relevant SNP rs16147 in the pathophysiology of tobacco dependence. Further studies are needed to confirm our findings.

Introduction
Tobacco smoking is a major health problem with extensive consequences for persons concerned, and the society in general [1, 2]. Biological, political and psycho-

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logical factors are involved in smoking behavior and tobacco dependence [3, 4]. Twin studies have consistently estimated heritability of around 50% for tobacco dependence [5, 6]. Thus, genetic factors have a substantial contribution to the tobacco dependence etiology. Several different genes are associated with tobacco smoking; recently, genome-wide association studies investigated several promising genetic determinants of tobacco dependence and found strongest genetic contribution to nicotine dependence from variation in the nicotinic receptor subunits and in nicotine metabolism [7–9]. However, variation within these genes accounts for only a small proportion of the risk for developing tobacco dependence, which implies that many genes seem to be involved in this complex behavior and disease [10]. Nicotine and other substances from tobacco affect brain functions through modifying different neurotransmitter systems, and nicotine is the major addictive substance in tobacco [11]. The rewarding effects of nicotine (e.g. relief of anxiety and arousal), withdrawal symptoms (e.g. craving, irritability, dysphoric or depressed mood, anxiety, restlessness, increased appetite or weight gain and insomnia), and environmental factors (e.g. stress) are the main factors which contribute to the consolidation of the tobacco dependence [11]. Nicotine mainly binds to nicotinic acetylcholine receptors containing the α4 and β2 subunits, often in combination with the α6 subunit, which are located with the highest concentrations in the mesolimbic dopaminergic reward system in the brain [12, 13]. Nicotine is the major addictive component of tobacco and a potent agonist of neuronal nicotinic receptors (AChRs) thereby stimulating dopamine (and norepinephrine) release [14]. Hence, the dopaminergic system is considered a major pathway for tobacco use and development of tobacco dependence [15].

Neuropeptide Y (NPY), a 36-amino-acid peptide, is expressed at high levels within the nervous system [16, 17]. The NPY gene (NP-Y) is a potential candidate gene involved in nicotine dependence because NPY has been shown to increase extracellular dopamine levels dose-dependently in the shell region of the nucleus accumbens in rats [18]. This indicates that NPY could play an important role in drug reinforcement by modulating dopamine levels in the brain reward system. On the other hand, the anxiolytic neuropeptide NP-Y is also known as endogenous regulator of appetite and anxiety [19–21], which is both affected through tobacco smoking, and nicotine withdrawal [11, 22, 23]. Hence, NPY gene variations have been associated with negative effects, and numerous psychiatric diseases in a number of studies [20, 24, 25]. Furthermore, several studies suggested a possible involvement of the NPY system in the neurobiological effects of different abused substances including alcohol [26], methamphetamine [27], and smoking behavior [28].

In summary, the NPY system has been shown to modulate drug intake behavior, anxiety, and positive rewarding properties. Taking these findings together, the NPY gene may be an interesting candidate gene to gain new insights into the pathophysiology of tobacco dependence. In particular, the (promotor) single nucleotide polymorphism (SNP) rs16147 has been reported to influence NPY gene expression on the level of mRNA and protein and accounts for a relevant part of differential expression of NPY in vitro [24, 25, 29]. We therefore analyzed whether this functional genetic variant in the NPY promoter gene is associated with cigarette smoking in an ethnically homogeneous sample of healthy Caucasian subjects from Germany.

**Methods**

**Data Collection**

For case-control association analysis, 550 current smokers (Fagerström test 5–10 points) and 544 never-smokers (max. 20 cigarettes/lifetime), of whom 464 subjects were male and 630 were female (entire sample n = 1,094), all were genotyped. Data were collected in seven recruitment centers throughout Germany (Departments of Psychiatry at the University of Aachen, Berlin, Bonn, Düsseldorf, Erlangen, Mainz, Mannheim) between the years 2007 and 2009. Inclusion criteria for subjects were: German origin and mother-tongue level German and grandparents of subjects were required to be born in Germany or in a country adjacent to Germany. Subjects with non-German ethnicity were not included. Further details of recruitment, testing procedures, inclusion/exclusion criteria, and characterization of this multicenter study on nicotine dependence and smoking-related behavior have been described previously [23, 30].

Inclusion/exclusion criteria were assessed through a medical examination, a standardized psychiatric interview (SCID-1), questionnaires, drug screen, alcohol testing and CO measurement. The present study was approved by the ethics committees of all participating centers; subjects were included in the study after they had given written informed consent.

**Psychometric Measures**

The absence of psychiatric axis-I comorbidity was verified with the Structured Clinical Interview [31] which is based on DSM-IV criteria (American Psychiatric Association, 2000). The Fagerström Questionnaire for Nicotine Dependence [32] was used as a measure for the level of nicotine dependence [33] and Heaviness of Smoking Index (HSI-G) [34]. Nicotine craving was measured via self-report with the Questionnaire of Smoking Urges (QSU) [35] (QSU German version [36]). Anxiety was as-
sessed with the trait version of the State-Trait Anxiety Inventory (STAI [37]; STAI German version [38]). Depressive symptoms were measured with the Beck Depression Inventory (BDI [39]; BDI German version [40]). Subjective experience of stress was measured by the Perceived Stress Scale (PSS [41]). To control for alcohol use the Alcohol Use Disorders Identification Test (AUDIT [42]; AUDIT German version [43]) was used.

**DNA Preparation and Genotyping**

Genotyping was done at the Cologne Center for Genomics (CCG), University of Cologne. DNA from fresh frozen EDTA blood was prepared with Qiagen FlexiGene DNA Kit according to the manufacturer protocol and normalized based on RNase P copy number measurement using the TaqMan RNase P assay from Applied Biosystems® (Foster City, Calif., USA). The SNP rs16147, a functional NP-Y promoter variant, was chosen to cover the functional expression of brain NP-Y. Genotyping was performed with SNPstream SNP genotyping assays. Genotyping call rates were 99%. All laboratory procedures were carried out blind to case-control status.

**Data Analysis**

The baseline demographics and clinical characteristics were compared between smokers and non-smokers. Descriptive statistics were calculated with t tests for independent samples and are given with mean and standard deviation. The Levene test was used to assess the homogeneity of variance. Associations between the genotype and smoking behavior were evaluated using the $\chi^2$ test, while the interaction effect of genotype and smoking behavior on the interval-scaled scores of the anxiety questionnaire (STAXI X1) was assessed with analysis of variance (univariate ANOVA). Where necessary, separate post-hoc t tests were performed for the three genotypes. The $p$ value for the post-hoc t tests was adjusted to 0.017 in order to control for the type I error. All other test statistics indicating a $p$ value of 0.05 or less were considered significant. Tests for deviation from Hardy-Weinberg equilibrium were performed using an exact test. Data analysis was performed using SPSS Statistics, version 19.

**Results**

A total of 1,094 subjects (550 current smokers and 544 never-smokers) were genotyped for SNP rs16147. Mean age was 34.7 ± 12.8 years. SNP rs16147 did not deviate from Hardy-Weinberg equilibrium (TT = 257, 23.6%, CT = 557, 51.1%, and CC = 277, 25.4%; for non-smokers p = 0.086, for smokers p = 0.444). Sociodemographic and clinical characteristics of the study population are presented in table 1.

We found a significant difference in the number of allele carriers between the group of smokers and non-smokers: $\chi^2$ analysis revealed a significantly higher probability for TT genotype of being a smoker compared to being a non-smoker ($\chi^2 = 4.74; p = 0.029$). However, comparisons of CC and CT alleles between the two groups did not show significant differences (CC: d.f. = 1, p = 1.0; CT: d.f. = 1, p = 0.271) (fig. 1; table 2). The ratio between smok-
ers and non-smokers in CC/CT allele carriers is 0.94, while the ratio between smokers and non-smokers in TT allele carriers is 1.28. The OR and CI for CC/CT compared to TT allele carriers is OR = 0.732, CI = 0.553–0.970.

In an omnibus interaction analysis, smokers compared to non-smokers showed a significantly higher state anxiety score (STAI X1) (F = 11.790; d.f. = 1; p = 0.001) but not trait anxiety score (STAI X2) (F = 2.833; d.f. = 1; p = 0.093). This effect was specifically accentuated within the TT allele carriers: the interaction effect (smoking status × genotype) indicates significantly higher scores of the STAI X1 questionnaire in TT allele carriers who smoked compared to the TT allele carriers who never smoked (p < 0.05). Comparisons of CC and CT alleles between the two groups did not show significant differences (p > 0.05).

We did not observe significant differences between the different alleles of the SNP rs16147 concerning the age of onset of tobacco dependence, FTND scores, pack-years, amount cigarettes smoked per day, and tobacco craving (QSU) (data not shown).
Discussion

The main result of our study is the finding that the risk variant of the SNP rs16147 (TT allele), located in the NP-Y gene promoter, is associated with increased probability of tobacco smoking behavior. TT allele (rs16147) carriers have a significantly higher probability of being a smoker, compared to CC and CT allele carriers. However, we did not find differences concerning the age of onset of tobacco dependence, amount of cigarettes smoked per day, and tobacco craving (QSU) between the different genotypes. Numerous studies on the effects of rs16147 on NP-Y gene expression have been reported in the literature, reporting rather mixed and conflict results [24, 44–46]. However, the C allele was identified as the high NP-Y expressing allele in most reports. Our results are consistent with reports suggesting that TT allele carriers (rs16147) show reduced gene expression of the anxiolytic protein NPY. We found significantly higher scores of the STA11 questionnaire in TT allele carriers who smoked compared to the TT allele carriers who never smoked. Recently published data support our results, reporting the T allele of rs16147 is associated with anxiety and depressive symptoms in young adults [24, 29]. This might be a consequence of the reduced gene expression of the anxiolytic protein NPY in T(T) allele carriers (rs16147). It is well known that patients with psychiatric disorders (e.g. anxiety disorders) have a higher risk to develop tobacco dependence. In addition to genetic risk factors of psychiatric disorders and tobacco dependence, the anxiety and stress attenuation effects of nicotine are known to play an important role in the development of tobacco addiction. Especially in pathophysiological mechanisms of anxiety/depression and substance abuse, NP-Y has been implicated repeatedly [20, 26–28, 47]. Since the biological effects of NP-Y are substantially controlled by the NP-Y gene, the increased state anxiety and associated risk of tobacco smoking might be a compensation of the impaired NP-Y expression of TT allele carriers (rs16147). This hypothesis is based on the observation that NPY has an important function in counter-regulating states of anxiety [20].

Taken together, our results suggest an involvement of the NPY system in tobacco smoking. We found individuals carrying the risk genotype TT of rs16147, associated with reduced NP-Y levels in former studies, have significantly higher state anxiety and have a higher individual vulnerability for tobacco smoking. However, the TT allele of rs16147 was not associated with higher tobacco craving, earlier age of onset of tobacco dependence, and higher amount of cigarettes smoked per day, implying additional risk factors which contribute to the complex disease of tobacco dependence. Our study has some limitations: Whether rs16147 directly or indirectly influences NP-Y expression, or whether it acts through other pathways has not been investigated so far and should be targeted in further studies. However, in our large case-control study, we found that a mutation in the NP-Y gene promoter may constitute a genetic risk factor for tobacco dependence. Further studies are needed to replicate our findings and to further elucidate other genetic risk variants of tobacco dependence.

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Disclosure Statement

The authors have no conflicts of interest to disclose.

References

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