CYP2A6 Polymorphisms May Strengthen Individualized Treatment for Nicotine Dependence

Yawo Mawuli Akrodou

Department of Public Health, College of Health Sciences, Walden University, Minneapolis, MN 55401, USA

Correspondence should be addressed to Yawo Mawuli Akrodou; akym120@gmail.com

Received 20 November 2014; Revised 27 March 2015; Accepted 6 April 2015

Academic Editor: Nicolaas A. Franken

Copyright © 2015 Yawo Mawuli Akrodou. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Each CYP2A6 gene variant metabolizes nicotine differently depending on its enzymatic activities. The normal nicotine metabolizer CYP2A6*1A is associated with high scores of nicotine dependence (5–10) on the Fagerström Test for Nicotine Dependence (FTND) scale because it encodes for enzymes that catalyze nicotine 100%. Slow nicotine metabolizers (i.e., CYP2A6*1H, CYP2A6*4A, CYP2A6*9, and CYP2A6*12A) are associated with underrated nicotine metabolizing activity (50%–75%), linking them to low scores for nicotine dependence (0–4) on the FTND scale. In a clinical trial involving the use of bupropion, people who were carriers of slow nicotine metabolizers were found to have a tendency to maintain abstinence 1.7 times longer than people with normal nicotine metabolizers. An overview of CYP2A6 polymorphism enzymatic activities in nicotine dependence etiology and treatment revealed that slow nicotine metabolizers may strengthen the individualized treatment of nicotine dependence.

1. Introduction

CYP2A6 polymorphism consists of alleles with variable structures, ranging from deleted alleles (CYP2A6*2A/CYP2*4A) to fully functioning alleles (CYP2A6*1A), which determine the role that their level of enzymatic activity plays in nicotine etiology and the impact they have on nicotine dependence treatment [1, 2]. The nicotine candidate gene CYP2A6*1A has normally high enzymatic activity in nicotine digestion and is responsible, in part, for the high prevalence of smoking by predisposing people to nicotine addiction [3, 4], causing more than 5 million [5, 6] smoking-related deaths and disabilities each year. On the other hand, slow nicotine metabolizers may realize potential clinical benefits since they metabolize less nicotine [1]. This paper reviews the ability of CYP2A6 polymorphisms to digest nicotine differentially and investigates their impact on nicotine dependency treatment in the context of personalized treatments for nicotine dependence.

2. Study Background and Theoretical Foundation

The enzymatic activities of the normal nicotine metabolizer CYP2A6*1A can be considered to be responsible, in part, for the high prevalence of nicotine dependence due to CYP2A6*1A’s excessive metabolism of inhaled nicotine [1]. The slow nicotine metabolizer alleles of CYP2A6 catalyze less nicotine and positively influence nicotine-dependence treatment [7]. Nevertheless, current pharmacotherapy treatments are not tailored to the genetic makeup of nicotine-dependent people, limiting the odds of successful treatment outcomes in nicotine dependence. Unfortunately, more than 5 million deaths annually are linked to tobacco-related diseases. In the United States (US) alone, the smoking prevalence is still 20%; moreover, approximately 400,000 deaths each year are associated with tobacco-related diseases [5, 6], thereby compelling a search for a novel treatment for nicotine dependence.

In general, the implication of genes in addiction etiology and their possible use in treatment should be considered to
individualize addiction treatment. Behavioral Genetic Theory (BGT) underlines the existence of a correlation between genes and addresses the role that exposure to environmental agents [8–12] plays in an individual's predisposition to develop an addiction or in preventing an individual from becoming addicted to a substance. For example, the interaction between nicotine and the CYP2A6 polymorphism CYP2A6*1A (normal nicotine metabolizer) leads to nicotine addiction [13].

The capability of genes to metabolize drugs can strengthen or weaken nicotine dependence treatment by influencing nicotine pharmacodynamics and pharmacokinetics [5, 14]. Nicotine is an environmental agent; when it interacts with specific candidate nicotine genes, it can play an important role in the development of nicotine dependence [14], as shown in Figure 1. The normal nicotine metabolizer CYP2A6*1A is one of the first biological mediators of nicotine digestion [15]. The suppression or alteration of the CYP2A6*1A gene variant, which excessively metabolizes nicotine, should lead to the excretion of undigested nicotine in the urine by blocking cotinine from reaching the brain, thereby slowing down nicotine dependence [15, 16]. In fact, a slow nicotine metabolizer, such as CYP2A6*4A, has a deleted allele of CYP2A6, which does not metabolize nicotine. It disrupts the normal digestion process of nicotine, which is simply evacuated instead. Consequently, slow metabolizers have potential health benefits in nicotine dependence treatment for carriers [13, 16–20], acting as a type of nicotine-dependence treatment drug because the pharmacologic nicotine dependence treatment strategy consists of injecting small doses of nicotine into the blood, which limits the volume of nicotine needed to excite the brain [7, 21, 22].

3. Limitations of Current Nicotine-Dependence Therapies

Smoking cessation has major health benefits for men and women of all ages. However, a long-term cure for nicotine dependence remains difficult to achieve since more than 80% of moderate-to-heavy smokers who seek treatment relapse within 1 year [23]. Although many methods have been developed to aid smoking cessation (e.g., pharmacotherapy and behavioral counseling), currently available nicotine dependence treatments are still rather limited in efficacy [23]. While smoking behavior is sustained biologically by candidate nicotine genes, such as the normal nicotine metabolizer CYP2A6 as well as other candidate nicotine genes, including dopamine receptors and CYP2B6 gene variants [24, 25], information about these genes is not used in current nicotine dependence treatment.

There is a specific mass (1.0–1.5 mg) of nicotine that should be metabolized continuously while smoking in order to allow the tobacco user to enjoy a pleasurable feeling (triggered by the impact that nicotine has on the brain), which leads to his/her addiction [26, 27]. According to Hukkanen
et al. [22], 1.0–1.5 mg of nicotine is readily absorbed while smoking and easily passes through the lung alveoli membrane because nicotine is a weak base with a $pK_a = 0.8$. The absorption of nicotine through the lung membrane is facilitated by the pH level, and then nicotine is transported by the bloodstream to the brain within 10 seconds, where it induces pleasurable sensations. The mechanism of nicotine digestion occurs with the mediation of multiple candidate nicotine genes, but the most relevant nicotine gene metabolizers (and the most well-studied) are the polymorphic alleles of the \textit{CYP2A6} gene.

The role of \textit{CYP2A6} polymorphism is a determinant in the development of nicotine dependence and its treatment [4]. In treatment, a relapse of the nicotine addiction mechanism occurs because nicotine abstinence reduces dopamine release, triggering nicotine withdrawal symptoms, such as unpleasant mood states (irritability, anxiety, depression, and restlessness), poor concentration, hunger, sleep disturbance, and a craving for nicotine [28]. These withdrawal symptoms often defy the efficacy of current pharmacotherapies according to their structure (deleted/decreased or full function), thus confirming their propensity to predispose to or protect individuals from nicotine dependence [19]. Kubota et al. [2] found that the normal nicotine metabolizer \textit{CYP2A6*1A} gene variant, which has fully functional alleles, encodes the enzyme that metabolizes 100% the volume of inhaled nicotine. Consequently, the \textit{CYP2A6*1A} alleles are associated with a higher daily nicotine intake, indicating that they make the carrier highly susceptible to developing smoking behavior [5].

In an \textit{in vivo} study, Benowitz et al. [33] administered nicotine to clinical trial participants and found that those who carried fully functioning alleles of \textit{CYP2A6*1A} secreted a small quantity of excreted, unchanged nicotine and cotinine N-glucuronide. This quantity represented 25%–30% of the excreted urinary metabolites. They also found a high quantity of cotinine-derived metabolites, consisting of 58%–67% of the excreted urinary metabolites. That experiment provides clues that subjects carrying \textit{CYP2A6*1A} normal nicotine metabolizers are highly susceptible to nicotine dependence. On the other hand, the \textit{CYP2A6*2A} variant is a deleted allele of \textit{CYP2A6} gene, and it does not produce enzymes [5]. Therefore, it is inactive and cannot metabolize nicotine [34,35]. As a result, carriers of \textit{CYP2A6*2A} are poor nicotine metabolizers, smoke less, and recover quickly following nicotine dependence treatment.

In \textit{in vitro} studies, participant carriers of the homozygous \textit{CYP2A6} deleted alleles (\textit{CYP2A6*4}) received nicotine [36]. After two hours, they were found to have very low plasma-cotinine levels and high levels of unchanged nicotine. These different enzymatic activities of the \textit{CYP2A6} polymorphisms result from structural change at the nucleotide level in each of the alleles of \textit{CYP2A6}, as depicted in Table 1. For example, \textit{CYP2A6*1A} is a wild-type allele of \textit{CYP2A6}, which did not evolve; on the other hand, \textit{CYP2A6*1H} to \textit{CYP2A6*12} underwent an evolution or mutation expressed by a change in their nucleotides, thereby affecting their level of enzymatic activities.

<table>
<thead>
<tr>
<th>CYP2A6 gene variants</th>
<th>Nucleotide change</th>
<th>Protein alteration</th>
<th>Enzymatic activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>CYP2A6*1A (wild type)</td>
<td>None</td>
<td>None</td>
<td>Normal</td>
</tr>
<tr>
<td>CYP2A6*1H</td>
<td>-745A&gt;G</td>
<td>Disruption of CCAAT box</td>
<td>Decrease</td>
</tr>
<tr>
<td>CYP2A6*4A</td>
<td>CYP2A6 deleted</td>
<td>No produced protein</td>
<td>None</td>
</tr>
<tr>
<td>CYP2A6*9A</td>
<td>-1013A&gt;G, -48T&gt;G; 51G&gt;A</td>
<td>Change in TATA box</td>
<td>Decrease</td>
</tr>
<tr>
<td>CYP2A6*12A</td>
<td>Change Exon 3–8 of CYP2A6 origin</td>
<td>10 amino acid substitution</td>
<td>Decrease</td>
</tr>
</tbody>
</table>

Note: change in the nucleotide of these gene variants alters or inhibits their enzymatic activities decreasing their nicotine metabolism activities in comparison with \textit{CYP2A6*1A} (wild type) gene variants [22].

4. \textit{CYP2A6} Polymorphism’s Impact on Nicotine Dependence

Among the currently identified nicotine gene candidates, the \textit{CYP2A6} gene is the most studied and has been found to be principally responsible for nicotine metabolism, uptake, distribution, and clearance [19]. The \textit{CYP2A6} gene is located on the long arm of chromosome 19 [31]. \textit{CYP2A6} belongs to the family of genes known as cytochrome P450, which has a mixed-function oxidase system and is involved in the metabolism of xenobiotics in the body. \textit{CYP2A6} variants have properties that induce or inhibit nicotine according to their structural function [32]; thus, they are hypothesized to enhance or alter nicotine dependence treatment [3].

The evaluation of the enzymatic activities of \textit{CYP2A6} in \textit{in vitro} and \textit{in vivo} experiments has shown that \textit{CYP2A6} polymorphisms can alter or enhance the pharmacokinetics of nicotine according to their structure (deleted/decreased or full function), thus confirming their propensity to predispose to or protect individuals from nicotine dependence [19].
5. CYP2A6 Polymorphism Enzymatic Activities and Smoking Behavior

In fact, CYP2A6 polymorphism consists of alleles (i.e., CYP2A6*2A and CYP2A6*4A) that have a complete loss-of-function or have an enzymatic activity less than 50%, alleles (i.e., CYP2A6*9A and CYP2A6*12A) with a decrease of function having enzymatic activity less than 75%, and alleles (i.e., CYP2A6*1A wild type) that are fully functional with 100% enzymatic activity [26, 33]. The correlation of CYP2A6 alleles’ enzymatic activities with the rate of their markers of 3HC/COT production has led to a method to determine smokers’ phenotype [37]. Thus, different CYP2A6 allele enzymatic activities have permitted researchers to classify individual smokers into three CYP2A6 phenotype groups: slow nicotine metabolizers (people who smoke less), intermediary nicotine metabolizers (moderate smokers), and normal metabolizers (heavy smokers) [37], showing an extreme interindividual variability in smoking behavior. Table 2 summarizes the classification of CYP2A6 polymorphism according to the patients’ enzymatic activities in the two groups.

6. CYP2A6 Polymorphism Impact on Nicotine Dependence Treatment

CYP2A6 polymorphism enzymatic activities have been correlated with nicotine dependence, but their activities also influence the nicotine dependence treatment outcome [33]. In clinical trials, individuals who carried CYP2A6 variants with diminished enzymatic function activity rates and a low 3HC/COT production ratio while taking bupropion/nicotine replacement therapy (NRT) had the highest cessation rates: 32% compared with those with normal CYP2A6 allele activities and a high 3HC/COT production ratio, who had only a 10% cessation rate at the end of a 6-month treatment [36]. Using data generated by three clinical trials among Caucasian populations, Patterson et al. [38] demonstrated that individuals using transdermal nicotine patch treatment with a low 3HC/COT ratio and slow CYP2A6 variant nicotine-metabolism activity were significantly more likely to quit smoking than those with high 3HC/COT ratios (indicative of normal CYP2A6 nicotine-metabolism activities) at week eight end-of-treatment (46% as opposed to 28%) and 6 month follow-up visits (30% as opposed to 11%). Therefore, it may be beneficial to translate relevant functions of CYP2A6 polymorphisms into clinical practice, as they variably convert nicotine to cotinine at 78%, which is further transformed into trans-3-hydroxycotinine (3HC) [39]. CYP2A6 gene variant functions may be harnessed to tailor nicotine dependence treatment for individual treatment needs and should not be neglected in the case of nicotine dependence.

7. Nicotine Dependence Treatment One-Size-Fits-All Paradigm

Currently, many new advanced technologies are allowing researchers to understand more complex interactions among genes and genetic interactions with environmental agents [40, 41] and to translate their findings into clinical treatment. In addition, the emerging sciences of pharmacogenetic and pharmacogenomic research have revealed gene functions and the mechanism of their potential clinical utility and validity, which are being incorporated into drug making and disease treatment management [42].

Unfortunately, nicotine dependence treatment programs (cognitive behavioral therapy and pharmacologic therapy) have been rooted in a ”one-size-fits-all” paradigm and cannot be readily tailored to the unique needs of individual patients based on their unique constellation of behavioral, biological, and clinical characteristics [43]. Munafò et al. [44] proposed that the one possibility to increase and enhance treatment of nicotine dependence is to offer nicotine dependence treatments tailored to an individual’s genotype. Since 70%–80% of inhaled nicotine is biologically converted into cotinine and 90% of this reaction occurs under the influence of the enzyme cytochrome P450 2A6 (CYP2A6), the cotinine is then completely transformed by CYP2A6 into trans-3-hydroxycotinin [22].

8. Context of Individualized Nicotine Dependence Treatment

Twin studies have demonstrated that a person’s genetic makeup contributing to his or her liability to nicotine, alcohol, and other drug addictions can also shield the individual from the development of addiction [38, 41]. Therefore, treating nicotine patients on an individual basis according
Individualized medicine practices
(1) Individual prognostics and diagnostics
(2) Gene testing and screening interpretation and counseling
(3) Gene therapies

Difficulties achieving individualized medicine “one-size-fits-all.”
(1) Inconsistent findings
(2) Inadequate studies designs
(3) Unsuccessful replications of genes functions
(4) Epistasis phenomena

Global biologic system
Phenotype development
Disease/not diseased

Gene-gene interaction factors influences
Gene-environmental factors influences

Figure 2: Schema illustrating personalized medicine concept and its difficulties. Note. Dashed lines indicate difficulties achieving personalized medicine. The figure is the illustration of classic “one-size-fits-all” pharmacological therapies to individualized treatment which takes to account the individual genotype makeup.

to their genotype may help increase a successful nicotine dependence treatment outcome [41]. For example CYP2A6 polymorphism provides information that governs the basics if biological and physiological activities in nicotine dependence that can be correlated to individual genetic makeups [38]. In addition, the human genome shows that the DNA sequences of any two individuals are 99.9% identical and the 0.1% difference in DNA constitutes the origin of profound genetic material diversity. Therefore, it is very important that nicotine addiction treatment be individualized according to the patient’s genetic makeup [38] as summarized in Figure 2 illustrating personalized medicine concept.

A large interindividual variability of nicotine metabolism underlines that the treatment of nicotine dependence should consider information regarding candidate nicotine genes. In a quantitative study, Nakajima et al. [35] compared different alleles of CYP2A6 genes’ nicotine-metabolism activities and found large interindividual variability in the cotinine/nicotine ratios in races with the following means: Caucasian $7.1 \pm 4.7$; Black $7.2 \pm 5.0$; Chinese, $8.7 \pm 11.9$; and Japanese, $3.8 \pm 3.1$. These different nicotine metabolism activities of CYP2A6 alleles in races confirmed the varying prevalence of nicotine dependence across a race spectrum [18, 22, 45].

9. Finding the Impact of CYP2A6 Polymorphisms

To date, few emerging smoking-cessation clinical trials have reported that CYP2A6 activity is associated with nicotine-quitting success [30]. Unraveling the role of gene variants and determining how to use them to design effective therapy are difficult [46] but not unachievable [40].

This goal can be realized through studies that may provide accurate assessment through multivariable analyses of CYP2A6 genes variants in correlation with nicotine dependence and withdrawal syndrome, other genes, and different forms of therapies [42]. A correlational study could also help examine to what extent CYP2A6 major variants are correlated with other nicotine dependence risk factors and how they can influence nicotine dependence treatment.

10. Summary

CYP2A6 polymorphisms mediate the metabolism of nicotine according to their ability to encode for the enzyme catalyzer of nicotine to cotinine [13]. They serve as a biological gateway for nicotine to reach the brain by the bloodstream [13]. The fully functional allele CYP2A6*1A metabolizes enough nicotine each time a tobacco user inhales the smoke of cigarette to lead to nicotine dependence. In contrast, slow nicotine metabolizers catalyze less nicotine, which prevents addiction [22]. In nicotine dependence treatment, carriers of slow nicotine metabolizer genes are 1.7 times more likely to quit smoking or maintain abstinence than those who have a normal nicotine metabolizer underlining their potential health benefit; these metabolizers can therefore be used in nicotine dependence treatment [30].

The role of CYP2A6 polymorphism capability to catalyze nicotine provides clues that nicotine treatment should be individualized. Twin, adoption, clinical trial, and case-control studies suggested that there is strong evidence of
the influence of genetic variants in nicotine-dependence etiology and treatment, where genetic variants contributions were evaluated at more than 50% [1, 32], conveying the possibility that CYP2A6 polymorphism information analysis may strengthen nicotine dependence treatment.

Conflict of Interests

The author declares that there is no conflict of interests regarding the publication of this paper.

Acknowledgments

The author thanks Drs. Peter Anderson and Sandra Rasmussen for their advice in the writing of this paper.

References


[40] N. Cohen, Pharmacogenomics and Personalized Medicine, Humana Press, Louisville, Ky, USA, 2008.


