Nicotine degradation in smokers: will a new and potent enzymatic approach work where nicotine vaccines have failed?

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Abstract
Smoking and tobacco use continue to be the largest preventable causes of death. Although there are current pharmaceutical and behavioural therapies, the one-year sustained quit rate of these therapies is only 20–25% at best. Recently, an alternative biotherapeutic strategy has been proposed: enzymatic degradation of nicotine in the bloodstream preventing accumulation in the brain. The bacterial enzyme NicA2 oxidizes nicotine into pseudo-oxynicotine, a non-addictive compound already found in smokers. Proof-of-concept animal studies have shown that NicA2 can rapidly reduce the levels of nicotine accumulating in the brain after intravenous nicotine dosing, and NicA2 has shown to have efficacy in a continuous nicotine access self-administration rat model. Enzymatic elimination of nicotine upon smoke inhalation to combat tobacco addiction is an innovative therapeutic concept. However, it is in line with recent clinical studies demonstrating that reduction in nicotine content in cigarettes (to 2.5% of normal levels) lead to significant reduction in the number of cigarettes smoked and higher smoking cessation rates compared to a control group smoking normal nicotine level cigarettes. Enzymatic degradation of nicotine appears to be more potent than nicotine-specific antibodies or vaccines for reducing nicotine distribution to brain, and if this proves to be the case in humans, it could also be more effective for enhancing smoking cessation rates and succeed where nicotine vaccines have failed thus far. The work reviewed in this article constitutes promising initial steps towards an urgently needed new effective treatment strategy in smoking cessation therapy.

Introduction
Smoking and tobacco use continue to be the largest preventable causes of death [1]. In 2015, approximately 6.4 million deaths were attributed to smoking worldwide. Although most smokers are aware of the health risks, smoking cessation is usually difficult to maintain. Current pharmacological therapies for smoking cessation combined with counselling have significant clinical effects compared to counselling alone [2]. However, only 20–25% of smokers remain abstinent for at least 1 year after treatment [3]. This fact means that new, more efficacious drugs need to be developed.

Multiple meta-analyses have been conducted to investigate the pharmacological interventions for smoking cessation, and guidelines have been published by many organizations [2,4]. The first-line pharmacological therapy for smoking cessation are nicotine replacement products (patches, gums, inhalers, nasal sprays, tablets, and oral sprays). It evokes its effects by stimulating the nicotinic receptors in the ventral tegmental area of the brain releasing dopamine in the nucleus accumbens [5]. NRT can lead to a reduction in withdrawal symptoms in smokers who would like to quit. Varenicline works as a partial agonist of the nACh receptor also releasing dopamine [6]. Furthermore, Bupropion, a tricyclic antidepressant, can be used in smoking cessation therapy. It inhibits reuptake of dopamine, noradrenaline, and serotonin in the central nervous system, and is a non-competitive nicotine receptor antagonist. The inhibition of the levels of dopamine and noradrenaline are thought to be important for Bupropion to have its antismoking actions [7]. Varenicline and bupropion are usually prescribed and when used for 2-3 months achieve a doubling of the quit rate compared to placebo [8]. Furthermore, counselling should be given to help in smoking cessation. Brief advice alone given by a general practitioner result in a 2-3% increase in quit rates [9]. To stop smoking is to break a complex habit and addiction and, to achieve reasonable quit rates, it is necessary to provide psychological support combined with pharmacological drugs. However, even with optimal pharmacological therapies only 20–25% of smokers remain abstinent for at least 1 year after treatment. This means that new therapies need to be developed.

As an alternative to small-molecule-based therapies, immunotherapeutic approaches to smoking cessation and vaccination against nicotine were investigated in the last three decades [10]. Researchers showed that it is possible to link or conjugate psychoactive drugs (such as cocaine, heroin or nicotine) to carrier proteins, thus making these small molecules antigenic. This work led to the hypothesis that it may be possible to develop vaccines which can prevent or treat addiction to these drugs. The proposed mechanism of action is that vaccine-elicited antibodies target and capture the drug in the periphery, reducing the concentrations reaching the brain and reducing its reinforcing effects. Nicotine conjugate vaccines showed early promise preclinically but failed to demonstrate broad clinical promise.
nicotine degradation via an enzymatic approach, eliminating its exposure to the brain [14]. *Pseudomonas putida* S16 is an example of a nicotine-degrading bacterial strain that can use nicotine as its nitrogen and carbon source. It was originally isolated from a field underneath continuous tobacco cropping in China and is able to metabolise nicotine to fumaric acid [15]. The enzyme found in the first committed step of S16’s nicotine degradation is NicA2, an amine oxidase. NicA2 oxidises nicotine to N-methylmyosmine, which undergoes rapid, spontaneous hydrolysis to pseudooxynicotine, a non-addictive compound already found in smokers.

Xue and colleagues studied the features of NicA2 in vitro to evaluate its potential as a starting point for the development of a nicotine-degrading drug for use in smoking cessation therapies [14]. They demonstrated that NicA2 has favourable characteristics such as high stability in buffer and mouse serum, as well as high catalytic activity at nicotine concentrations typically found in smokers’ blood [14].

NicA2 was subsequently evaluated in vivo through single-dose nicotine pharmacokinetic (PK) studies in rats pre-treated with a range of NicA2 doses [16,17]. Reduction in nicotine blood and brain levels was measured 1, 3 and 5 minutes after an intravenous bolus dose of 0.03 mg/kg nicotine. This nicotine dose is equivalent to 2 cigarettes with regard to milligrams of nicotine per kilogram of body weight. Short intervals were used, as the enzyme’s effectiveness is expected to be dependent on the rapid elimination of nicotine. While smokers achieve maximum levels of brain nicotine in 3 to 5 minutes, nicotine is initially detected in the brain 7 seconds after the first inhalation [18]. NicA2’s effects on nicotine distribution to the blood and brain were dependent on dose and time, as shown in the Figure 1 below [17,19]. When dosed at 5 mg/kg, blood levels of nicotine dropped below the limit of quantitation of the assay (2 ng/ml), virtually eliminating nicotine from the bloodstream within 1 minute as compared to the control group. The levels of nicotine in the brain were also assessed, with a 10-mg/kg NicA2 dose lowering brain nicotine levels by 95% at 3 and 5 minutes after nicotine dosing as compared to the control group, while a higher dose of 20 mg/kg was needed for reducing brain nicotine levels to the same extent within one minute. As one minute is a practical time limit to euthanise the rats and to collect blood and brain samples, the onset of enzyme activity was evaluated in blood samples in vitro, where typical maximum blood levels of nicotine were degraded to below the level of detection within 10 seconds [17].

In repeated nicotine dose experiments that simulated very heavy smoking, 5 doses of 0.03 mg/kg nicotine spaced 10 minutes apart (equivalent to 10 cigarettes over 40 minutes) were given intravenously to rats pre-treated with 10 mg/kg NicA2. Brain nicotine levels were lowered by the same degree after the 5th dose as after the 1st dose of nicotine, a potency never observed for immunotherapeutic approaches [17,19,20].

In order to enable longer-term in vivo testing, two different constructs fusing NicA2 to an albumin-binding domain (NicA2-ABD) [21] have been independently reported [17,22]. Circulating half-life was extended from a few hours to 2.5 days in rats, similar to that of endogenous serum albumin, without affecting its catalytic activity [16,17]. Consistent with the effects of NicA2 on reduced nicotine distribution in the brain, when such an enzyme fusion was administered to rats during a 7-day nicotine infusion, it reduced signs of withdrawal following termination of the nicotine infusion compared to the control group. A significant impact was observed on nicotine’s behavioral effects by preventing the development of irritability-like behavior, hyperalgesia and somatic signs of withdrawal in animals exposed to chronic nicotine, strongly supporting the theory that NicA2 may prevent the development of addiction-like behavior. Moreover, there was no nicotine detected in the blood or brains in the treated group, while the control group exhibited expected concentrations of nicotine in both blood and brains[22]. By contrast, nicotine vaccines were only partly able to reduce brain nicotine concentrations, probably leading to the observed lack of efficacy [23].

Importantly, a NicA2-ABD fusion was shown to decrease nicotine discrimination and reductions in nicotine reinforcement in a continuous nicotine access self-administration model which closely resembles human nicotine exposure. In this model [24], rats are trained to self-administer nicotine and develop a stable dependence on nicotine for several weeks before being tested. After a high dose of NicA2-ABD (70 mg/kg), nicotine-seeking behaviour was extinguished over 6 days of testing with rats having continuous access to nicotine as seen in Figure 1.

**Figure 1. Reduction of blood and brain nicotine concentrations by NicA2.** Rats were pre-treated with NicA2 IV and 5 minutes later received 0.03 mg/kg nicotine IV. Groups of eight rats had nicotine levels measured at 1, 3 or 5 minutes in blood (upper panel) and brain tissue (lower panel). Nicotine concentrations were rapidly reduced by NicA2 in a dose- and time-related manner. **p < 0.01, ***p < 0.001 compared to BSA using Bonferroni-corrected Welch’s t-tests. Reproduced with permission from [7].**
The dotted horizontal line represents baseline. The dashed horizontal line represents the mean (± SD) number of infusions during 23-hr access in vivo predicted on the basis of the reported treatment should reduce the risk of developing anti-NicA2 antibodies. is commonly observed with the 12-week treatment period of present enzyme treatments needed for smoking cessation is relatively brief, as authorities for human use [28]. In addition, the expected duration of origin or animal-human chimera have been approved by regulatory authorities for human use, this characteristic should be minimized. It is unfavorable for human use, this characteristic should be minimized. It should be noted that many engineered proteins of entirely non-human origin or animal-human chimera have been approved by regulatory authorities for human use [28]. In addition, the expected duration of enzyme treatments needed for smoking cessation is relatively brief, as is commonly observed with the 12-week treatment period of present smoking cessation drugs. This relatively short expected duration of treatment should reduce the risk of developing anti-NicA2 antibodies.

Finally, improving catalytic activity to reduce the dose amounts predicted on the basis of the reported in vivo studies will be important.

![Figure 2. Effects of NicA2-ABD on self-administration of nicotine in rats having unlimited access to nicotine.](image)

**Figure 2.** Effects of NicA2-ABD on self-administration of nicotine in rats having unlimited access to nicotine. Mean (± SD) number of infusions during 23-hr access following pre-treatment with PBS vehicle (V) and NicA2-ABD over six consecutive test sessions, expressed as a percentage of baseline. Each point represents the mean of four rats. The dotted horizontal line represents baseline. The dashed horizontal line represents the 50% reduction criterion for extinction. Different from V by paired t-tests, **p < 0.01, ***p < 0.001. Reproduced with permission from [18].

Modern protein engineering techniques have been successful in optimising catalytic activities of enzymes [29,30]. Such improvements are needed for realistic dose levels, suitable routes and frequency of drug administration.

NicA2 promises a new treatment strategy in smoking cessation, since it rapidly degrades nicotine within a few minutes. In a clinical sense, it is envisioned to be administered intravenously by injection in a physician’s office during cessation counselling. As one of the most important predictors of success is self-motivation, it could be an important advantage over vaccines if patients can immediately initiate a quit attempt while the motivational level is at its peak without having to wait for months before an immune response builds up, as in the case of vaccines. Attempts to quit should become easier, since the nicotine does not reach the brain to evoke its rewarding effects and smoking will not result in a rewarding stimulus. Eventually, this process could lead to long-term abstinence, as smoking is no longer associated with a reward.

It is interesting to note the parallel between NicA2 treatment to reduce nicotine exposure and the effect of lowering the nicotine content in cigarettes themselves. In clinical studies, smokers who did not have an intention to quit were provided with a range of nicotine-containing cigarettes, including Very Low Nicotine Content (VLNC) cigarettes (0.3 mg/cig. 2% of Normal Nicotine Content (NNC) cigarettes). These smokers experienced reduced nicotine exposure and dependence, reduced cravings during abstinence from smoking and increased unprompted quit attempts in comparison with smokers who were assigned NNC cigarettes [31]. In treatment-seeking smokers, greater reductions in nicotine exposure while smoking VLNC cigarettes predicted smoking cessation [32]. Furthermore, withdrawal symptoms were mild to moderate, comparable to withdrawal symptoms when using a nicotine patch [33]. Moreover, minimal compensatory smoking was observed [34]. In a study where subjects had free access to VLNC cigarettes but not NNC cigarettes (sequestered in a hotel), a 92–94% reduction in nicotine exposure biomarkers was observed [35]. This approach has recently been embraced in the US by the FDA’s Center for Tobacco Products, proposing the regulation of tobacco products with the intention of lowering the nicotine content to non-addictive levels [36]. Whether it will ultimately be politically possible to eliminate the sale of tobacco containing addictive levels of nicotine is unknown. The timeline for the FDA plans is not yet defined and may take a decade or more to implement fully in the US, while it is uncertain how many other countries worldwide would or could follow a similar path. However, these clinical studies do emphasise that nicotine is a key addictive component and lend validity to the concept of enzymatically eliminating nicotine in the form of smoke inhalation to combat tobacco addiction.

Enzymatic degradation of nicotine appears to be more potent than nicotine-specific antibodies or vaccines for reducing nicotine distribution to the brain in rats. If this situation proves to be the case in humans as well, it could be more effective in order to enhance smoking cessation rates and succeed where nicotine vaccines have failed thus far. This development could be a major step forward in the race that we must win to reduce the number of smokers which die prematurely each year [37].

**Conflict of Interest**

Onno van Schayck has received consulting compensation for advice with respect to smoking cessation clinical study design and development planning as part of the NIH/NIDA Strategic Alliance grant to develop nicotine-specific monoclonal antibodies for smoking cessation (R01DA038877; Kalnik/PI, Antidote Therapeutics, Inc.).
Thomas Thisted and Matthew Kalnik are employees and shareholders of Antidote Therapeutics. TT and MK are inventors on nicotine-degrading enzyme patent applications assigned to Antidote Therapeutics. [16] Bo van Engelen interned at Antidote Therapeutics.

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