Cigarette Smoking: Neurobiology, Addiction and Treatment Implications

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ABSTRACT

Cigarette smoking is most common of all smoking devices both in terms of prevalence and health consequences. From the glowing tip of the lighted cigarette burning at a temperature of 800°C (1600-1800°F), the smoker with each puff draws into his mouth and lungs a hot potpourri of gases and many sized particles. Of the 7357 chemicals identified in cigarette smoke 70 are the known carcinogens. Nicotine is the main chemically active constituent in cigarette smoke responsible for causing addiction. Pulmonary absorption of nicotine, which is the most favored and perhaps commonest, occurs in a matter of seconds. Studies suggest that alpha-4 beta-2 nicotine acetylcholine receptor subtype is the main receptor that mediates nicotine addiction. Nicotine facilitates the release of dopamine and other neurotransmitters to induce pleasure and mood modulation. Smoking is also reinforced by conditioning. Pharmacotherapy for smoking cessation should reduce withdrawal symptoms and block the reinforcing effects of nicotine obtained from smoking without causing excessive adverse effects.

Keywords: Addiction, Cigarette Smoking, Neurobiology, Nicotine

Introduction

Tobacco smoking spread all over the world after Christopher Columbus, a Spanish navigator and his crewmen discovered the natives of America inhaling the smoke of burning leaves of a plant in 1492, which was later identified as tobacco.1 Introduction of tobacco from the Americas to the rest of the World some five hundred and twenty-five years ago generated the first drug controversy of a global dimension.2 Some of the early reports praised its medicinal virtue and called it herbal panacea while others dubbed it as an invention of the devil for its use in Natives’ pagan religions.3 However, the commercial forces managed to dictate its universal availability by way of engineering addiction among its users during these five centuries. Today, we know adequately that tobacco smoking causes not only addiction it harms every organ of the body, causes many diseases, reduces the health of the smokers in general, shortens their life-span and leads to premature deaths.1 Tobacco is a plant product obtained from the Nicotiana genus of the Solanaceae family of the plant kingdom. Unlike other members of this family, such as tomato and potato, which have an uncontroversial nutritional role, the tobacco plant carries into its leaves quantities of an alkaloid, nicotine, which gives it instead, power over man’s mind.2 Leaves of tobacco plant are harvested and cured to manufacture consumable tobacco products which are used for smoking or smokeless use. Smokeless tobacco products are used for chewing, snuffing or local application while...
smoking products are cigars, hookah, chillum, cheroot, cigarettes, beedis etc. which provide tobacco smoke to the smokers. Cigarette is a tobacco rod designed to deliver tobacco smoke in the body by puffing it from one end while the other end is lighted to smoulder. From the incandescent tip of the lighted cigarette burning at temperature of 800 °C (1600-1800 °F), the smoker with each puff draws along the tobacco rod and into his mouth a hot potpourri of gases and many sized particles. The smoke comprises of main-stream smoke inhaled from the cigarette and the side-stream smoke produced from the burning of cigarette paper.

**Cigarette Smoke**

Smoke constituents have organic matter, nicotine alkaloids, additives and pyrolyzed products obtained from the mainstream and the side-stream smokes. Estimated 7357 chemical compounds are detected in cigarette smoke 70 of which have confirmed carcinogenic activity. Amounts of these compounds vary in different brands of cigarettes. Nicotine is the main chemically active constituent in tobacco smoke responsible for a number of patho-physiological changes in the body. Carbon monoxide (CO), a highly toxic gas, combines with hemoglobin in the blood, reduces its oxygen carrying capacity and thus exposes the body to the risk of various respiratory disorders. Tar in tobacco smoke is a suspended particulate matter containing sticky, brown particles that stain teeth, fingers and lung tissue. Tar contains the carcinogen benzo(a)pyrene which triggers cancer development. Nitrogen oxide in tobacco smoke damages lung tissue and causes emphysema. Hydrogen cyanide stops the movement of cilia present in the lungs and thus impairs the foreign particle clearing capacity of the lungs and allows the poisonous tobacco smoke ingredients to accumulate in the lungs. Ammonia is a strong chemical that damages the lung tissue. Metals in tobacco smoke such as arsenic, cadmium and lead, are harmful to the human health and some of these are carcinogenic. Radioactive compounds found in tobacco smoke are proved carcinogens. In addition to nicotine, tobacco contains several other alkaloids that are structurally related to nicotine, giving rise to carcinogens N-nitrosornornicotine (NNN), 4-methyl-nitrosamino-1-butanone (NNK) and other toxic and carcinogenic nitrosamines. Tobacco-specific nitrosamines present in tobacco are formed during fermentation and curing of tobacco. Tobacco smoke is rich in naphthalene and polycyclic aromatic hydrocarbons (PAH), which are known carcinogens and produce serious adverse effects on the body.

**Nicotine in Cigarette Smoke**

An alkaloid (1-methyl-2-[3-pyrdyl]) pyrrrolidine), having carbon, hydrogen and nitrogen in proportion C_{10}H_{14} and N_{2} forming a double ring like structure, nicotine is responsible for causing addiction to tobacco. It is a colorless, volatile, strongly alkaline liquid that turns pale yellow to dark brown on exposure to air giving it a characteristic tobacco smell. It is highly toxic and potentially lethal chemical and one drop of pure nicotine is sufficient to kill a dog (or a man) within minutes. Nicotine is dissolved in the moisture of tobacco leaf as a water-soluble salt and in a burning cigarette nicotine volatilizes and is present in the smoke as free nicotine suspended on minute droplets of tar.

From tobacco smoke the nicotine enters the blood stream through the lungs while nicotine in smokeless tobacco passes through the mucosal membrane of mouth and nose or the skin. Pulmonary absorption, which is the most favored and perhaps commonest, occurs in a matter of seconds. From the lungs, chemicals in the smoke are absorbed into body's systems and carried quickly to different parts of the body. Oral, sniffs and other smokeless tobacco products are absorbed more gradually. Amount of nicotine intake from one cigarette varies widely, in accordance with the smoker's latitude for adjusting the dose level. Nicotine intake ranges from 10 mg/day to 80 mg/day, or 0.4 mg to 1.6 mg/cigarette.

After absorption, nicotine travels rapidly and reaches the brain within seven seconds; it readily crosses the blood–brain barrier. There is elevation of blood pressure due sudden burst of nicotine in the brain because of stimulation of adrenal glands and discharge of epinephrine. There is also sudden release of glucose and increase in respiration, heart rate, constriction of arteries and increased alertness. Many of these effects are produced through its action on both the peripheral and central nervous system. Nicotine causes release of dopamine; therefore, the psycho-active rewards occur quickly and these rewards are highly reinforced. Nicotine is distributed throughout the body, mostly to skeletal muscles and brain and activates specific receptors known as cholinergic receptors.

In the brain, nicotine binds to the receptors and influences the cerebral metabolism. Nicotine is then distributed throughout the body. If nicotine were not absorbed quickly from the lungs, people would not take it in the form of smoke; if it were not taken up into the brain, it would not exert its psycho-pharmacological effects; if it were not rapidly metabolized and excreted, it would probably not be taken in such often-repeated doses.

Nicotine has structural similarity to acetylcholine (Ach), which conveys information from one neuron to another. When a nerve is stimulated, the excitement is initially propagated along the nerve fiber in the form of electrical impulse and at the nerve ending; acetylcholine is released from the synaptic vesicle into the synaptic cleft, which stimulates acetylcholine receptors in the next neuron, and the neurotransmitter is used as the messenger to pass on the information carried in the nerves. This way messages
are carried from the body to the brain, from the brain to the body and between different parts of the brain and spinal cord.\(^1\)

Acetylcholine is involved in systems concerned with mental and physical arousal, learning and memory and several aspects of emotion. There are also other receptors for acetylcholine in the body, apart from the ones at synapses. They are also found at junction of nerve and muscles and nerves and certain glands.\(^3\)

Acetylcholine receptors respond only to acetylcholine as they recognize acetylcholine molecule. Physiologically these receptors respond to the natural transmitter acetylcholine, however, nicotine also activates them but, unlike acetylcholine, nicotine remains for much longer and as a consequence a proportion of nAChRs are desensitized.\(^13\) Repeated exposure to nicotine leads to an increase in the number of nAChRs in the brains of laboratory animals and human smokers. The activation of nAChRs by nicotine can have a number of consequences in the recipient nerve cells, including the release of various other neurotransmitters.\(^14\)

### Neurobiology of Nicotine Addiction

#### Nicotine Acetylcholine Receptors

Traditionally classified as nicotine receptors (those that respond to nicotine) and muscarine receptors (those that respond to muscarine), acetylcholine receptors are present throughout the body.\(^8\) The ability of nicotine to combine with acetylcholine receptors means that it can exert actions like acetylcholine at all synapses where nicotine acetylcholine receptors (nAChRs) are present. They trigger impulses down postsynaptic nerve fibers resulting in effects that otherwise occur only when acetylcholine is released following stimulation of the pre-synaptic nerve. Synapses involving acetylcholine are very widespread in the body, affecting systems ranging from the cardiovascular to the psychological and also interacting with other transmitter systems showing that nicotine has multifarious actions. Electrical excitation of a nerve produces not just one impulse but a whole train of impulses. This multiplicity imposes the requirement at the synapse that the combination of transmitter with the receptor must be quickly reversed between each impulse, leaving the receptor free to combine with the next pack of acetylcholine released.\(^9\)

Nicotine binds to the acetylcholine receptors (nAChRs) in the brain and influences the cerebral metabolism by stimulating these receptors. The stimulation of presynaptic nAChRs on the neurons increases the transmitter release as well as the metabolism. Chronic administration of nicotine results in desensitization and inactivation of nAChRs with subsequent up-regulation of nAChRs sites.\(^13,14\)

Cholinergic receptors are concentrated in the midbrain areas, such as mid-brain tegmentum, the striatum, nucleus accumbens (NAc) and the ventral tegmentum as well as in muscles, adrenal glands, the heart and other organs.\(^15\) These receptors are normally activated by acetylcholine. Besides binding to acetylcholine receptors (nAChRs) nicotine also binds to the cholinergic receptors in the autonomic ganglia, adrenal medulla, chemoreceptors of the carotid bodies and aortic body and neuromuscular junction. The specific sites of binding in the brain are the hypothalamus, thalamus, midbrain, brainstem and cerebral cortex. Nicotine also binds to the receptors in nigrostriatal and mesolimbic dopaminergic neurons. On stimulation of dopaminergic receptors acetylcholine, norepinephrine, dopamine serotonin, vasopressin, growth hormone and ACTH are released. Nicotine is one of the most potent stimulants of the midbrain dopamine reward pathway.\(^16,17\) Nicotine acts on locus coeruleus regulating vigilance, arousal, concentration and stress reactions making the tobacco users more alert. Owing to the interaction between nicotine and neuronal high-affinity nicotine acetylcholine receptors (nAChRs), nicotine affects learning, memory and other functions.\(^18\)

Nicotine also alters the function of some of the neurotransmitters implicated in the pathogenesis of some of the major psychiatric disorders. These include dopamine, norepinephrine, serotonin (5-HT), glutamate, gamma-aminobutyric acid (GABA) and endogenous opioid peptides.\(^10,20\) These effects could be presynaptic, preterminal, or cell body nicotine receptors, rather than mediated through neurotransmission wherein presynaptically released acetylcholine acts on postsynaptic, junctional nAChRs to cause neuronal firing.\(^21\)

The cholinergic receptors are relatively large structures consisting of several components known as subunits. Nicotine receptors are composed of 12 subunits in mammalian brain, 9 alpha subunits (alpha\(_2\) to alpha\(_9\)) and 3 beta subunits (beta\(_2\) to beta\(_4\)), which play the central role in autonomic transmission.\(^22\) The AChRs complex is composed of 5 subunits and is found both in peripheral and central nervous system.\(^23\) Different combinations make different receptors, which vary in terms of affinity and localization within the brain.\(^7\) The most abundant receptor subtypes in the brain of humans are alpha\(_2\)-beta\(_2\), alpha\(_7\)-beta\(_2\) and alpha\(_9\) (hommeric).\(^4\) The beta subunit has role in nicotine addiction. The alpha\(_4\)-beta\(_4\) subunit combination has greatest sensitivity to nicotine.\(^6\) In mice, knocking out beta\(_2\) subunit gene eliminates the behavioral effects of nicotine, including self-administration.\(^24\) Reinserting the beta\(_2\) subunit gene into VTA of a beta\(_2\) knocked out mouse restores behavioral responses to nicotine.\(^25\) The alpha\(_4\)-beta\(_4\) and alpha\(_9\) (hommeric) receptor subtypes mediate the cardiovascular effects of nicotine.\(^26\) The alpha\(_2\) subunit is also thought to be involved in rapid synaptic transmission and may play a role in learning and sensory gating.\(^27,28\)

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Repeated inhalation of tobacco generates bolus of nicotine delivered into the brain, superimposed on a relatively stable level of plasma nicotine maintained by the smoker throughout the smoking day. This basal level of nicotine keeps a proportion of nAChRs in a desensitized state, while the remaining are available for activation by nicotine bolus, if appropriate concentrations are achieved. This way smokers manipulate their plasma nicotine profile to achieve balance desensitization versus activation. When a smoker is asleep, plasma level of nicotine decreases and the nicotine receptors gradually recover their active function. In the morning, a smoker has a greater number of active nAChR sites (up-regulation) contributing to withdrawal symptoms and craving. It is, therefore, the first cigarette of the day is, most satisfying, as overnight abstinence allows a substantial recovery from nAChRs desensitization. Post-mortem findings in smokers brain show increased number of nAChR binding sites. With repeated exposure to nicotine, there is neuroadaptation to some of the effects of nicotine, leading to an increase of nAChRs to up-regulate the nicotine-mediated desensitization. This desensitization plays role in nicotine tolerance and dependence. Craving and withdrawal symptoms begin in chronic smokers when previously desensitized alpha-β nAChRs become unoccupied and recover to a responsive state during the period of abstinence i.e. night sleep. Electrophysiological studies demonstrate that nicotine agonists stimulate the release of GABA from rodent brain and this release are Ca²⁺-dependent. The actions of nicotine on ventral tegmental GABAergic innervation, which modulates the mesolimbic dopamine excitability, have been studied. Nicotine was found to increase firing rate of dopamine and non-dopamine neurons, while the former was more vigorous. These findings suggest that nicotine stimulates the firing rate of dopaminergic neurons of VTA and also the GABAergic neurons, which may be an important target for the effects of nicotine on the central nervous system. Acute nicotine administration stimulates the release of noradrenalin (NA) in the different parts of the brain, primarily at locus coeruleus level. Chronic nicotine administration decreases the concentration of 5-HT in the hippocampus because it is associated with selective increase in the density of 5-HT1A receptors in this area. Hippocampus receives serotonergic innervation from the median raphe nucleus. Suppression of 5-HT release brings about anxiolytic response to nicotine microinjection into the dorsal hippocampus. The effects of nicotine on 5-HT are difficult to dissociate from those on dopamine neurons. Increased exposure to stressful stimuli is likely to increase the desire to smoke as reported by smokers. The effects of nicotine withdrawal on dopamine release in the brain may be exacerbated by exposure to stressful stimuli and may underlie the role of stress as a factor in tobacco smoking, as well as the role of nicotine on reducing the effects by acting on 5-HT neurons within the hippocampus.

Animal studies suggest commonalities between nicotine withdrawal and opiate abstinence syndrome. Nicotine stimulation induces the release of endogenous opioid peptides in various brain regions resulting in over-activation of opiate receptors resembling opiate dependence. Abrupt termination of nicotine stimulation may precipitate an opiate abstinence like syndrome. A recent study suggests that cotinine, a metabolite of nicotine stimulates nicotine receptors to evoke the release of dopamine (DA) in a calcium-dependent manner from super-fused rat striatal slices.

Neuro-imaging technique shows dramatic effect of tobacco smoking on the brain of awake human beings. On positron emission tomography (PET), it was found that cigarette smoking decreases the levels of monoamine oxidase (MAO), which is responsible for breakdown of dopamine. Decrease in the levels of MAO-A and MAO-B results in increase of dopamine levels.

**Biology of Nicotine Reinforcement: Dopamine and Reward Pathways**

Brain imaging studies show that nicotine acutely increases activity in prefrontal cortex, thalamus and visual system consistent with activation of cortico-basal ganglia-thalamic brain circuits. Nicotine is a powerful reinforcing agent in both animals and humans. The mesolimbocortical dopamine system consists of neurons with cell bodies localized in ventral tegmental area (VTA) and axon projections to nucleus accumbens (NAc) and medial prefrontal cortex. Nicotinic receptors concentrated in VTA and NAc activate the mesolimbic dopamine system, which is responsible for reinforcing behavior like other dependence producing drugs. VTA and its projections to NAc are involved in reward and mediate the reinforcing actions of drug abuse. Nicotine stimulates the release of dopamine in the pleasure circuit and increases extracellular level of dopamine in NAc. Lesions of mesolimbic dopamine neurons attenuate nicotine self-administration in rats. It also attenuates locomotor stimulant effect of systemically administered nicotine. Stimulation of central nAChRs by nicotine results in the release of a variety of neurotransmitters in the brain, most importantly dopamine in the mesolimbic area, the corpus striatum and the frontal cortex. Of importance are the dopaminergic neurons in the VTA of the midbrain and release of dopamine in the shell of NAc, which is critical in drug-induced reward. Other neurotransmitters, including norepinephrine, acetylcholine, serotonin, gamma-aminobutyric acid (GABA), glutamate and endorphins, are released as well, mediating various behaviors of nicotine release of neurotransmitters occurs via modulation by presynaptic nAChRs along with the direct release.

**Dopamine**
release is facilitated by nicotine-mediated augmentation of glutamate release and long-term treatment by inhibition of GABA release.

**Genetics of Nicotine Addiction**

CYP2A6 is an enzyme responsible for the majority of inactivation of nicotine in humans; it is also responsible for activating tobacco-related precarcinogens such as the nitrosamines. A common genetic defect in nicotine metabolism decreases smoking. Genetic variation in CYP2A6 gene may protect individuals from becoming nicotine-dependent smokers. Mimicking this gene defect by inhibiting CYP2A6 decreases nicotine metabolism. In cigarette smoking like other behaviors shows evidence of heterogeneity. Dopamine transporter (DAT) gene (SLC6A3) encodes protein that regulates the synthetic levels of dopamine in the brain and leads to addictive behavior. It is expected that recent advances in molecular biology, including the completion of draft sequence of the human genome may help in identifying gene markers that predict a heightened risk of using tobacco to increase our understanding of nicotine dependence.

**Learning and Memory**

Nicotine plays a potential role in cognitive enhancement, while the earlier studies reported improvement in learning and performance with nicotine; in the recent studies nicotine has been shown to produce a sort of place preference in rats and mice. The self-administration of nicotine has also been demonstrated. In rodents, nicotine has anxiolytic-like action on various behavioral tests namely the mirror chambered, the elevated plus maze, the two compartment light dark transition test and fear-potentiated startle. Nicotine and nicotine agonists improve performance in a variety of cognitive tasks by animals with basal forebrain lesion.

**Tolerance and Dependence**

Tolerance to nicotine and onset and persistence of withdrawal on cessation of nicotine treatment has been extensively studied in experimental animals. Chronic exposure to nicotine increases high affinity binding of nicotine agonists to brain tissue and induces chronic tolerance to many of the drug’s behavioral and physiological effects. The increase in receptor number (upregulation) has been interpreted as compensation for agonist-induced desensitization of nAChRs, and this prolonged desensitization has been viewed as a mechanism for chronic tolerance to nicotine. Acute and chronic nicotine administration in experimentally naïve rats depresses locomotor activity.

**Psychoactive Effects of Nicotine and Nicotine Withdrawal**

Nicotine in humans induces stimulation and pleasure and reduces stress and anxiety. Nicotine use modulates the level of arousal and controls mood in daily life. Smoking may improve concentration, reaction time and performance of certain tasks. On stoppage of smoking, nicotine withdrawal symptoms emerge, which include irritability, depressed mood, hedonic dysregulation, restlessness, anxiety, difficulty concentrating, increased hunger, insomnia and craving for tobacco. Relative deficiency of dopamine release following long-standing exposure accounts for mood disturbances, anhedonia and craving for tobacco that may persist for long time after quitting. The pharmacological basis of nicotine addiction can be viewed as the combination of positive reinforcement, such as enhancement of mood or functioning, as well as avoidance of negative consequences of prior drug use - the relief of withdrawal symptoms.

**Conditioning Behavior in Nicotine Addiction**

Drug taking is a learned behavior occurring due to conditioning, reinforced by consequences of pharmacological actions, and association with specific moods, situations, or environmental factors with the rewarding effects of the drug. Respiratory sensory cues of tobacco smoking represent a conditioned reinforcement that play an important role in smoke intake, craving and rewarding effects.

**Mechanism of Pharmacotherapy for Smoking-Cessation**

Pharmacological effects of nicotine play a crucial role in tobacco addiction, and pharmacotherapy has to address this component of tobacco dependence. A pharmacological treatment for smoking cessation should both block the positive reinforcing effects of nicotine and prevent or reduce the development of withdrawal symptoms. Pharmacotherapy should also target the receptor subtypes involved in nicotine addiction without affecting the receptors that, it activated, would produce unwanted adverse effects.

FDA (Food and Drug Administration) of USA approved the medications used for smoking-cessation, which include Nicotine Replacement Therapy (NRT) (transdermal patch, gum, nasal spray, inhaler and lozenges); bupropion and varenicline. Nortriptyline and clonidine, though not approved by the FDA, are clinically effective in smoking-cessation.

**Nicotine Replacement Therapy**

Nicotine Replacement Therapy (NRT) acts in several ways - it relieves craving and withdrawal symptoms, which are relieved with relatively low blood nicotine levels, and causes positive reinforcement for arousal and stress relief. Nicotine desensitizes α₄β₂ nAChRs resulting in reduced effects of cigarette smoking.

**Non-nicotine Agents**

**Antidepressant Agents**

Bupropion blocks dopamine reuptake and, to a lesser
extent, norepinephrine reuptake and it has some nicotine receptor-blocking activity. Bupropion increases dopamine and norepinephrine blood levels, similar to the effect of nicotine on these neurotransmitters. In rats, bupropion in low doses blocks the rewarding effects of nicotine. The blockade of nicotine receptors could contribute to reduced reinforcement from a cigarette in the case of a lapse. Nortriptyline is a norepinephrine reuptake blocker and as such simulates noradrenergic actions of nicotine in the brain.

Clonidine
It is an alpha₂-adrenergic receptor agonist and reduces sympathetic neural outflow resulting into sedation, anxiolysis, potential hypotension, bradykinesia and dry mouth. Calming and anxiolytic effect helps in smoking cessation, particularly in those who are very anxious while quitting smoking.72

Varenicline
It is an analog of plant alkaloid, cytisine, reported to have some benefit in smoking cessation. Varenicline has high and selective activity at alpha₂-beta₂ receptor. It is a partial agonist at this receptor in vivo producing lesser response than that of nicotine (30-60%) but also blocks the effect of any nicotine added to the system. Thus, varenicline maintains a moderate level of dopamine release, which reduces craving and withdrawal symptoms during abstinence. It also blocks the reinforcing effects of nicotine obtained from cigarette smoke in the case of relapse.73

NicVAX
It works by stimulating the immune system to make antibodies that bind to nicotine molecules, making them too big to cross the blood–brain barrier and preventing them from reaching nicotine receptors and triggering the pleasure sensation that smokers and users of nicotine experience and become addicted to. Data from pre-clinical trials suggest that the injectable vaccine would be effective not only in preventing relapse but also in relapse prevention because the nicotine antibodies last a long time.74,75

Conclusion
Nicotine in cigarette smoke causes addiction like other drugs causing addiction. Nicotine addiction is a complex process involving the interplay of pharmacology, conditioned factors, personality and social setting. Therefore, the ideal treatment for tobacco-cessation involves a comprehensive approach that addresses all major issues of tobacco addiction both pharmacological and non-pharmacological.

Conflict of Interest: None

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