Amphetamines: Potent Recreational Drug of Abuse

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Abstract

Amphetamines are central nervous system (CNS) stimulants and belong to psychoactive drugs that affect chemicals in the brain, nerves and exert a constant contribution to hyperactivity and impulse control. The communal group of amphetamines comprises amphetamine, dextroamphetamine and methamphetamine. Amphetamines show its action on the mesolimbic dopaminergic reward system by inducing release of dopamine and to some extent norepinephrine, in the synaptic clefts of the nucleus accumbens and other terminal areas. They offer not only a sense of euphoric state, but also addiction. Amphetamines are frequently prescribed for the treatment of attention deficit hyperactivity disorder (ADHD) in children, adolescents and adults, narcolepsy and obesity. At therapeutic doses, amphetamine causes emotional and cognitive effects such as euphoria, change in desire for sex, increased wakefulness, improved cognitive control, etc. The balance of benefit/risk is the main challenge for its clinical use. When overused these drugs can be highly addictive, are often diverted from the user to be used as recreational drugs. Adverse effects include anxiety, aggression, headache, insomnia, hyperactivity, palpitations, increased breathing rate, increased blood pressure, tachycardia, arrhythmia, dilated pupils, paranoia, etc. However, at large doses, these drugs may impair cognitive function and induce rapid muscle breakdown. Even higher doses may give rise to hallucinations, paranoia, psychosis and potentially life-threatening conditions such as convulsions, stroke, kidney failure, etc. The foremost serious health implications of amphetamine resulting from chronic use are dependence, considered by compulsive drug-seeking and drug use and a phenomenon notorious as amphetamine psychosis. Therefore the objective of this study was to explore the functions of the amphetamines as recreational drug of abuse.

Keywords: Amphetamines; Psychoactive drug; Recreational drug; Abuse; Addiction

Abbreviations: CNS: Central nervous system; ADHD: Attention deficit hyperactivity disorder; US: United States; EU: European Union; 5-HT: Serotonin; MAO: Monoamine oxidase; VMAT: Vesicular monoamine transporter; DA: Dopamine; DAT: Dopamine transporter; CART: Cocaine and amphetamine regulated transcript; TAAR1: Trace amine-associated receptor 1; D: Dextrorotatory; L: Levorotary; PKA: Protein kinase A; PKC: Protein kinase C; ΔFosB: A member of the Fos family of transcription factors; FDA: Food and Drug Administration; DSM-IV-TR: Diagnostic and Statistical Manual of Mental Disorders, 4th edition, text revision; DSM-5: Diagnostic and Statistical Manual of Mental Disorders, 5th edition; APA: American Psychiatric Association; NF-kB: Nuclear factor kappa B; cAMP: Cyclic adenosine monophosphate; CREB: cAMP response element binding protein; GnRH: A histone methyltransferase enzyme; ΔUnD: A transcription factor; c-Fos: A proto-oncogene; DDS: Dopamine dysregulation syndrome; PMMA: p-Methoxyamphetamine; BDMPEA: 4-Bromo-2,5-dimethoxy-phenethylamine; MDMA: 3,4-Methylenedioxymethamphetamine; MDMA: 3,4-Methylenedioxymethamphetamine; MDEA: 3,4-Methyleneoxy-N-ethylamphetamine; DOB: Dimethyltryptamine; DOM: 2,5-Dimethoxy-4-methylamphetamine; TMA: 3,4,5-Trimethoxyamphetamine; Ca2+1.2: Calcium channel, voltage-dependent, L type, alpha 1C subunit; NMNA: N-methyl-D-aspartate; AMPA: a-amin-3-hydroxy-5-methyl-4-isoxazolepropionic acid; DRD: Dopamine receptor D1; CaM: Calmodulin; P2PB: Serine/threonine-protein phosphatase 2B; AC: Adenyl cyclase; DARPP32: Dopamine- and cAMP-regulated neuronal phosphoprotein 32; PP1: Protein phosphatase 1; CAMKII: Calcium/calmodulin-dependent protein kinase II; GS: A-G-protein; CREB: cAMP response element-binding protein; SIRT1: Siruin 1; HDAC1: Histone deacetylase 1; AADC: Aromatic L-amino acid decarboxylase.

Introduction

Amphetamines are potent CNS stimulant with sympathomimetic and adrenergic agonist activities [1]. This class of drugs was first synthesized in the late 19th century that includes amphetamine, dextroamphetamine and methamphetamine [2]. Although amphetamines had been available for research for many years, the first medical application of amphetamine was developed in the 1920s, when it's CNS and respiratory stimulant properties were discovered [2]. Amphetamines have been used as a treatment for cold and sinus symptoms (the original inhalers contained Benzedrine, an amphetamine), obesity, narcolepsy (i.e., a disease in which the patient uncontrollably falls asleep) and paradoxically, ADHD. Amphetamines also have a high potential for abuse [3]. Soldiers on both sides during World War II used these drugs for their stimulant properties. After the war, amphetamine abuse reached epidemic proportions in Japan, Sweden and other parts of Europe, yet the drugs were not recognized as dangerous in the United States (US) until the 1960s [2]. Complications of amphetamine were first monitored by US physicians when they prescribed methamphetamine as a treatment for heroin addiction [2]. In case of 15 to 34 year old adults, lifetime prevalence of

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Amphetamines use varies considerably between countries, from 0.1 to 12.4%, with a weighted European average of 5.5% [4]. The last year use of amphetamines in this age group ranges from 0 to 3.9%, with most countries reporting prevalence levels of 0.5 to 2.0%. It is estimated that approximately 1.7 million (i.e., 1.3%) young Europeans people have used amphetamine during the last year [5]. Study in the school going students aging between 15 to 16 years old, reported lifetime prevalence of amphetamine use ranged from 1 to 7% in the 24 European Union (EU) Member States and Norway. Countries like Belgium, Hungary also reported amphetamine prevalence levels of more than 4% [6].

It has been seen that single white men mainly addicted by amphetamine and its range high especially from 20 to 35 years [7]. Some research from rural populations showed that whites use amphetamine more than African Americans [8]. At present time women and other ethnic groups also addicted by amphetamine and this use become very common. Amphetamines stimulate the release of norepinephrine from central adrenergic receptors. It is reported that at high dose of amphetamine, dopamine can be secreted from the nigrostriatal and the mesocorticoclinic system [9]. Amphetamine may also act as a direct agonist on central 5-HT (i.e., serotonin) receptors and may inhibit monoamine oxidase (MAO) [10]. In the periphery, amphetamines are considered for the release of noradrenaline by acting on the adrenergic nerve terminals and α- and β-receptors [10]. Some pathways like serotonergic pathways may contribute to the calming effect. Amphetamine interacts with vesicular monoamine transporter (VMAT) enzymes to enhance release of non-exocytotic release of dopamine (DA) and 5-HT from vesicles [11,12].

Amphetamine is responsible for various complications like hypertension and tachycardia by its CNS stimulating action with feelings of increased confidence, sociability and energy [13]. It suppresses appetite and fatigue as well as leads to insomnia. Following oral use, the effects usually start within 30 min and last for many hours [13]. Amphetamine users may feel distressful conditions like irritable, restless, anxious and depression. This drug increases the activities of noradrenaline and dopamine neurotransmitter systems [14]. Amphetamine is less potent than methamphetamine, but in uncontrolled situations the effects are almost indistinguishable. After oral administration amphetamine is rapidly absorbed. Plasma levels of amphetamine may rise to maximum, 0.02 mg/L after a single oral dose of 10 mg [15]. The plasma half-life varies from 4 to 12 h and is dependent on the urinary pH (i.e., alkaline urine decreases the rate of elimination) [15].

The study showed that at normal therapeutic doses, the physical side effects of amphetamine vary by age and from people to people [16]. Some cardiovascular side effects include hypertension, Raynaud's phenomenon (i.e., reduced blood flow to extremities) and tachycardia (i.e., increased heart rate) [17]. Men may affected by sexual side effects including erectile dysfunction, frequent erections, or prolonged erections. Abdominal side effects may include abdominal pain, loss of appetite, weight loss, nausea, etc. [18]. Blurred vision, dry mouth, excessive grinding of the teeth, rhinitis medicamentosa (i.e., drug-induced nasal congestion), reduced seizure threshold, etc. are some other potential side effects [19]. Dangerous physical side effects are rare at typical pharmaceutical doses [20]. The therapeutic doses of amphetamine can include following psychological effects: increased alertness, apprehension, concentration, decreased sense of fatigue, mood swings (i.e., elated mood followed by mildly depressed mood), increased initiative, insomnia or wakefulness, self-confidence and sociability [20]. Other than these patients also suffer some other less common effects such as, anxiety, change in libido, grandiosity, irritability, repetitive or obsessive behaviors and restlessness [21]. Amphetamine psychosis can occur if patients use heavy doses of amphetamine [22]. Even though very rare, this psychosis may occur in case of long-term use of therapeutic doses [23,24].

Amphetamine overdose can lead to a number of symptoms [25]. Overdose symptoms increase with dosage of amphetamine and decreases with tolerance to amphetamine [25,26]. Patient who is tolerant can consume approximately 5 g of amphetamine per day, which may lead to state like coma [24]. A research data showed that due to overdose of amphetamine, [18] more than 3,788 patients died worldwide in 2003 (i.e., 3,425–4,145 deaths, 95% confidence) [27]. Therefore, the purpose of this study was to show the potentiality of the amphetamines as drug of abuse.

History of Amphetamine

Amphetamine was synthesized at first in Germany in 1887 by Romanian chemist Lazăr Edeleanu who named it phenylisopropylamine; the excitatory effect of its remained unknown until 1927, when it was independently resynthesized by Gordon Alles and claimed it's sympathomimetic properties [28]. In the early 1930s, when amphetamine's CNS and respiratory stimulant effects were discovered, it was commercialized as an inhaler for nasal congestion (i.e., Benzedrine) [29]. Meanwhile, it was also suggested by the health practitioners as a cure for a range of ailments like alcohol hangover, narcolepsy, depression, weight reduction, hyperactivity in children and vomiting associated with pregnancy. Furthermore, the use of amphetamine grew rapidly because of its easy abundance, cheap price, long lasting effects and good appreciation from the physicians as an addiction risk medicament [29]. Whilst, amphetamine and its methamphetamine derivatives were also introduced and became available as oral and intravenous formulations for therapeutic purposes. At the time of World War II to boost up alertness and endurance and to improve mood amphetamine was hugely utilized by the military of the US Great Britain, Germany and Japan [10,30]. But during the 1960s and 1970s abuse began with the discovery of its euphoric effects upon intravenous injection of methamphetamine with a more rapid onset than oral administration. Being structurally similar, methamphetamine has more prominence action on the CNS compared to amphetamine [31].

In the early 1970s according to the Controlled Substances Act of the US, amphetamine was claimed as a schedule II controlled substance owing to aforementioned health hazard effects [32]. But despite strict government controls, amphetamine has been frequently used legally or illicitly by people, mainly musicians, mathematicians, authors and athletes [33-35].

Chemistry of Amphetamine

The 2D and 3D structure of D-amphetamine and L-amphetamine are presented in Figure 1. The chemical formula of amphetamine is C_{10}H_{15}N. It is a methyl homolog of the mammalian neurotransmitter phenethylamine. The carbon atom adjacent to the primary amine has a stereogenic center and actually it is composed of a racemic mixture (1:1) of two enantiomeric mirror images. This racemic mixture can be separated into its optical isomers: D-amphetamine and L-amphetamine [10]. The pure free base of amphetamine is volatile, colorless with a characteristically strong amine odor and acrid burning taste at room temperature [36]. The salts form of amphetamine comprises amphetamine aspartate, hydrochloride, phosphate, saccharate, sulfate, etc. [15]. Amphetamine is also the parent compound of its own structural class, which contains various psychoactive derivatives as well as admirable chiral ligand for the stereo-selective synthesis [37,38].

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Mechanism of action of amphetamine

Amphetamine mainly works by increasing the action of the neurotransmitters dopamine and norepinephrine in the brain [39,40]. It also initiates the release of various other hormones (e.g. epinephrine) and neurotransmitters (e.g. serotonin and histamine) as well as the biosynthesis of particular neuropeptides (e.g. cocaine and amphetamine regulated transcript [CART] peptides) [36]. In case of Adderall both active ingredients such as D-amphetamine and L-amphetamine, bind to the same receptor, their binding affinities might vary. D-amphetamine and L-amphetamine are both potent agonists of trace amine-associated receptor 1 (TAAR1) and react with VMAT2, with D-amphetamine being the more effective agonist of TAAR1 [20,41]. Whereas, D-amphetamine generates more CNS stimulation than L-amphetamine; however, L-amphetamine has merely greater cardiovascular and peripheral action [42]. It has been noticed that certain children have a better clinical response to L-amphetamine [43].

Usually VMAT2 do the task of transporting the monoamine (e.g. dopamine, histamine, serotonin, norepinephrine, etc.) from the intracellular fluid to the synaptic vesicles (i.e., chemical storage units inside a neuron) of neuron in the absence of amphetamine [44]. Upon entry of amphetamine into neuron and interaction with VMAT2, the transporter reverses its approach of transport, which intern release stored monoamines inside synaptic vesicles back into the neuron’s intracellular fluid (Figure 2) [44]. At that time activation of TAAR1 by amphetamine, the receptor induces the neuron’s cell membrane-bound monoamine transporters (like the dopamine transporter, norepinephrine transporter, or serotonin transporter) to either stop transporting monoamines altogether via transporter internalization or transport monoamines out of the neuron [45]. In other words, the reversed membrane transporter will expel dopamine, norepinephrine and serotonin out of the neuron's intracellular fluid and into the synaptic cleft [45]. At the end, upon interaction with VMAT2 and TAAR1, amphetamine initiates neurotransmitter release from the synaptic vesicles initiated by VMAT2 into the intracellular fluid where they subsequently eliminate the neuron with the help of membrane-bound, reversed monoamine transporters initiated by TAAR1 [45].

Side effects of amphetamine

The side effects of amphetamine are diverse and the amount of amphetamine used is the primary factor in determining the likelihood and severity of side effects [20]. Amphetamine products such as Adderall, Dexedrine, and their generic equivalents are currently approved by the Food and Drug Administration (FDA) for long-term therapeutic use [24,46]. Recreational use of amphetamine generally involves much larger doses, which have a greater risk of serious side effects than the dosages used for therapeutic reasons [20]. The physical and psychological side effects of amphetamine are:

Physical side effects: Amphetamine stimulates the medullary respiratory centers, producing faster and deeper breaths (Table 1) [20]. In a normal person at therapeutic doses, this effect is usually not noticeable, but when respiration is already compromised, it may be evident [20]. Amphetamine also induces contraction in the urinary bladder sphincter, the muscle which controls urination, which can result in difficulty urinating [20]. This effect can be useful in treating bed wetting and loss of bladder control [20]. The effects of amphetamine on the gastrointestinal tract are unpredictable [20]. If intestinal activity is high, amphetamine may reduce gastrointestinal motility (i.e., the rate at which content moves through the digestive system); [20] however, amphetamine may increase motility when the smooth muscle of the tract is relaxed [20]. Amphetamine also has a slight analgesic effect and can enhance the pain relieving effects of opioids [20].

FDA-commissioned studies from 2011 indicate that in children, young adults and adults, there is no association between serious adverse cardiovascular events (i.e., sudden death, heart attack and stroke) and the medical use of amphetamine or other ADHD stimulants [47,48]. However, amphetamine pharmaceuticals are contraindicated in individuals with cardiovascular disease [49].

Psychological side effects: At normal therapeutic doses, the most common and less common psychological side effects of amphetamine are stated earlier (Table 1) [20]. Amphetamine psychosis (e.g. delusions and paranoia) can occur in heavy users [22]. Although very rare, this psychosis can also occur at therapeutic doses during long-term therapy [23]. According to the FDA, “there is no systematic evidence” that stimulants produce aggressive behavior or hostility [24]. Amphetamine has also been shown to produce a conditioned place preference in humans taking therapeutic doses, [50] meaning that individuals acquire a preference for spending time in places where they have previously used amphetamine [39].
Amphetamine overdose

As the doses of amphetamine increases, there is risk of developing impaired cognitive function and induce rhabdomyolysis (Table 2). Although there is a perceived risk of drug addiction along with amphetamine use, however, it is unlikely to happen for typical long-term medical uses at therapeutic doses [51]. Recreational amphetamine use often involves much higher doses than therapeutic doses, which involves risk of adverse drug reactions, including psychosis (e.g. hallucinations and neurosis) [52,53].

In amphetamine addiction, an excessive pathological activation of the mesolimbic pathway (a dopamine pathway that connects the ventral tegmental area to the nucleus accumbens) is found to play critical role [54,55]. Accumbal ΔFosB is recognized as “molecular switch” and considered as the “master control protein” for addiction. Regular recreational users often overdose themselves with amphetamines, as a result accumbal ΔFosB level slowly increases [56,57]. Consequently, induction of nucleus accumbens ΔFosB takes place which in turn trigger addictive behaviors like compulsive desire to take the drug. Still there is no effective therapeutic agent available which can treat addiction of amphetamine [58,59]. However, regular involvement with sustained aerobic exercises are found to not only lessen the development of amphetamine addiction, but also effective in the treatment of amphetamine addiction [60]. It has also been found that clinical treatment outcomes can be enhanced by exercise therapy and can also be incorporated with cognitive behavioral therapy, currently this combination therapy is considered as the best clinical treatment [39,61].

Amphetamine addiction and dependence

The acute reinforcing effects of amphetamine lead to patterns of drug use that, in epigenetically vulnerable individuals, result eventually in addiction, a state hypothesized to be the result of plastic changes in multiple neural circuits [64]. The types of plasticity that underlie addiction can be divided conceptually into three groups: compensatory adaptations in neural systems that regulate autonomic and other somatic functions leading to physical dependence and withdrawal; adaptations in the mesocorticolimbic reward system that mediate the subjective reinforcing effects of the drug, and adaptations in the mesolimbic dopamine system that underlie the subjective reinforcing effects of the drug.

Figure 2: The mechanism of action of amphetamine in a dopamine neuron. Amphetamines pass in the presynaptic neuron by means of the neuronal membrane or through dopamine transporter (DAT). After its entry, it binds to TAAR1 or arrives synaptic vesicles using VMAT2. When amphetamine reaches into the synaptic vesicles by the VMAT2, dopamine is started to release into the cytosol (yellow-orange area). When amphetamine fixes to TAAR1, it decreases the firing rate of postsynaptic neuron through potassium channels and finally triggers protein kinase A (PKA) also protein kinase C (PKC) signaling, afterward phosphorylation of DAT. PKA-phosphorylation of DAT is accountable for the withdraw of DAT into the presynaptic neuron (internalize) and eventually stop transport. PKC-phosphorylated DAT may work in reverse or similar PKA-phosphorylated DAT, internalize and terminate transport. Amphetamine is also recognized to raise the level of intracellular calcium, which is related to DAT phosphorylation by a CAMKIIα-dependent pathway, successively causes efflux of dopamine [Adapted from 20,21].
brain reward circuitry itself resulting in the emotional and motivational aspects of dependence and withdrawal; plasticity involving both the mesoaccumbens circuitry and other limbic circuits yielding sensitization and cue-dependent, positively biased emotional memories of drug use that may predispose to relapse. Unlike the opiates and ethanol, amphetamine does not produce physical dependence; they are nonetheless among the most reinforcing and addictive drugs known, underscoring the importance of plasticity in emotional circuits [64].

It may not possible to recognize the signs of amphetamine addiction at first, but as the addiction progresses and the effects of amphetamine use set in, the signs of addiction become more and more evident [65]. Some of the early signs of addiction are tolerance and physical dependence or an urge to use amphetamines (Table 3) [65].

As mentioned previously that amphetamines are used to medically treat such disorders as narcolepsy and ADHD [66]. Amphetamines may be illegally obtained and abused for the euphoric effects. When abused there are a number of side effects that can occur. The side effects experienced depend upon the individual and the amount of time the substance has been used. Some of the symptoms of amphetamine abuse are presented in Tables 4 and 5 [67,68].

| Body System | Minor or Moderate Overdose | Severe Overdose |
|-------------|---------------------------|----------------|
| Central nervous | Confusion, Abnormally fast reflexes, Severe agitation | Acute amphetamine psychosis (e.g., delusions and paranoia), Compulsive and repetitive movement |
| Cardiovascular | Abnormal heartbeat, High or low blood pressure | Cardiogenic shock (i.e., heart not pumping enough blood), Cerebral hemorrhage (i.e., bleeding in the brain) |
| Respiratory | Rapid breathing | Pulmonary edema (i.e., fluid accumulation in the lungs), Pulmonary hypertension (i.e., high blood pressure in the arteries of the lung) |
| Musculoskeletal | Muscle pain | Rhabdomyolysis (i.e., rapid muscle breakdown) |
| Urinary | Painful urination, Urinary retention | No urine production, Kidney failure |
| Other | Elevated body temperature, Mydriasis (i.e., dilated pupils) | Elevated or low blood potassium, Hyperpyrexia (i.e., extremely elevated core body temperature) |

Table 2: Clinical manifestations of amphetamine overdose [20,24,26,82,63].

| Signs of Amphetamine Addiction |
|-------------------------------|
| Compulsive use of amphetamines |
| Using amphetamines to feel good |
| Lack of personal appearance or ill desire to take care of one’s self |
| Taking amphetamines to cope with everyday activities |
| Using amphetamines to socially interact |
| Lack of pleasure when not taking amphetamines |
| Taking amphetamines despite problems in a relationship or with loved ones |
| Taking amphetamines despite the known consequences of taking the drugs |
| Withdrawal symptoms when not taking amphetamines |

Table 3: The signs of amphetamine addiction [65].
According to the Diagnostic and Statistical Manual of Mental Disorders, 4th edition, text revision (DSM-IV-TR) and Diagnostic and Statistical Manual of Mental Disorders, 5th edition (DSM-5); manual of the American Psychiatric Association (APA) diagnostic features of amphetamine use disorder is listed in Table 6 [69,70]. Consistent with DSM-5, it is necessary to specify the current severity: mild (presence of 2-3 symptoms), moderate (4-5 symptoms) or severe (6 or more symptoms) [70].

Although there is a substantial risk of developing addiction with larger doses of amphetamine, however this is unlikely to happen with typical long-term therapeutic uses [71]. On the other hand, drug tolerance is likely to take place with regular abuse of amphetamine, henceforth eventually larger doses of amphetamine will be needed to get the same effect [72] (Table 7).

Nuclear accumbens is the particular portion of the brain where altered gene expression takes place due to long-term use of amphetamine [73]. Nuclear factor kappa B (NF-κB), ΔFosB and cAMP response element binding protein (CREB) are the noteworthy transcription factors which are found to induce these alterations (Figure 3) [57]. Among these transcription factors, ΔFosB plays critical role in the development of drug addiction. The reason behind this pivotal role is due to the neural and behavioral adaptations that takes place due to the involvement of adequate ΔFosB overexpression in D1-type medium spiny neurons in the nucleus accumbens. An increment in the ΔFosB expression is found to be associated with the ΔFosB induced addictive state which is primarily due to the adequate ΔFosB overexpression. Similar conditions are seen in addictions like methylphenidate, substituted amphetamines, alcohol, cocaine, propofol, nicotine, cannabinoids, phencyclidine, etc. [74].

A transcription factor, ΔJunD and a histone methyltransferase enzyme, G9a are seen to be directly opposing the inductions of ΔFosB in the nucleus accumbens (i.e., they confine increases in its expression) [75]. Sufficient overexpression of ΔJunD in the nucleus accumbens with viral vectors can entirely block many of the chronic drug abuse associated with behavioral and neural alterations (i.e., the alterations mediated by ΔFosB). ΔFosB also have significant functions in the regulation of behavioral responses to natural rewards including exercise, sex and palatable food. As these addictive drugs and natural rewards can induce expression of ΔFosB (i.e., they cause the brain to produce more of it), thus, long-term acquisition of these natural rewards can lead to similar pathological state of addiction. Therefore, ΔFosB is considered as one of the most significant factors found to be involved in both addiction of amphetamine and compulsive sex addictions induced by amphetamine [76,77]. These sex addictions are linked with a dopamine dysregulation syndrome (DDS) which arises in some patients who are taking dopaminergic drugs [78].

Dose and route of administration largely determine the amphetamine effects on the regulation of gene. Most of the studies on addiction and gene regulation are based on intravenous administration of large doses of amphetamine in animal models. These studies confirmed that therapeutic uses of amphetamine do not significantly alter gene regulation [73]. Amphetamine dependence generally indicates that addiction has occurred and the user does not feel “normal” without the drug [79]. This dependence is associated with a range of physical and mental health problems. As stated by the DSM-IV-TR, in the diagnostic features of amphetamine dependence are presented [69].

Amphetamine withdrawal

Withdrawal is this period of readjustment [80]. Amphetamine dependence is also characterized by withdrawal symptoms, once the drug usage is stopped. Some of the common withdrawal symptoms are offered in Table 8 [81].

A previous study on people who used amphetamine and methamphetamine compulsively, reported that abrupt cessation of

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### Table 4: Symptoms of amphetamine abuse [67].

| Most Common Symptoms     | Less Desirable Symptoms                  |
|--------------------------|-----------------------------------------|
| • Increased body temperature | • Hostility                           |
| • Euphoria               | • Paranoia                              |
| • Increased blood pressure | • Aggressiveness                       |
| • Dry mouth              | • Cardiovascular system failure         |
| • Faster breathing       | • Irregular heart beat                  |
| • Dilated pupils         | • Nausea                                |
| • Increased energy and alertness | • Headache                           |
| • Decreased fatigue      | • Reduction of social inhibitions       |
| • Decreased appetite     | • Altered sexual behavior               |
|                          | • Blurred vision                        |
|                          | • Chest pain                            |
|                          | • Hallucinations                        |
|                          | • Unrealistic ideas of personal ability and power |
|                          | • Convulsions                           |
|                          | • Malnutrition                          |
|                          | • Skin disorders                        |
|                          | • Amphetamine-induced psychosis         |

### Table 5: Mood, behavioral, physical and psychological symptoms of amphetamine abuse [68].

| Mood Symptoms     | Behavioral Symptoms | Physical Symptoms | Psychological Symptoms |
|------------------|---------------------|-------------------|-----------------------|
| • Euphoria       | • Hostility         | • Increased body temperature | • Paranoia            |
| • Increased alertness | • Aggressiveness   | • Increased blood pressure | • Psychosis           |
| • Irritability   | • Altered sexual behavior | • Dilated pupils   | • Hallucinations       |
| • Mood swings    | • Not sleeping for prolonged period of time | • Faster breathing  | • Clear and focused feeling |
| • Depression     | • Not eating for prolonged period of time | • Decreased fatigue |                       |
| • Anxiety        | • Unrealistic ideas of personal ability and power | • Dry mouth         |                       |
|                  | • Fast and excessive talking |                       |                       |
|                  | • Grinding of teeth    |                       |                       |
|                  | • Increased confidence |                       |                       |
|                  | • Rectangular feeling  |                       |                       |

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Amphetamine use in the chronic heavy users may lead to time-limited withdrawal syndrome that may take place as early as 24 h of their last dose [82]. This study also concluded that as high as 87.6% of the cases withdrawal symptoms are most commonly seen amongst the chronic users who use amphetamine in larger doses on regular basis [82]. Symptoms of amphetamine withdrawal may include drowsiness, lack of motivation, anxiety, depression, craving for amphetamine, increased appetite, fatigue, hypoactivity or hyperactivity and lucid dreams [82]. This study also indicated that the extents of withdrawal symptoms are directly related to the degree of dependence and these symptoms are less likely to take place along with the therapeutic doses and uses [82]. Unfortunately, often manufacturer prescribing information does not indicate the risk of developing withdrawal symptoms following discontinuation of amphetamine use even after an extended period at therapeutic doses [46].

Amphetamine tolerance

The first signs of amphetamine addiction tend to be tolerance. Tolerance is the reaction that the body has been given a drug repeatedly [65]. Over time, sometimes faster if taken with other drugs, the body will have less of a reaction to the drug and the user will have to take more of the drug in order to feel the same effects [83]. As tolerance builds, the user will no longer feel the same effects of amphetamines and may resort to using more or to using the drugs more often [65].

Regular amphetamine use can quickly lead to tolerance of the drug.
Figure 3: Signaling cascade in the nucleus accumbens of brain’s reward center that lead to amphetamine addiction. Co-release of presynaptic dopamine and glutamate through psychostimulants (i.e., amphetamine), post-synaptic receptors for these neurotransmitters cause internal signaling events by a cAMP pathway then calcium-dependent path that finally increased the phosphorylation of CREB. Phosphorylated CREB raises the level of ΔFosB that represses the c-Fos gene by means of co-repressors, c-Fos repression that serves as a molecular switch which enables the accumulation of ΔFosB in the neuron. Phosphorylated ΔFosB once persists in neurons for the period of one or two months, gradually accumulates following repeated high-dose exposure to stimulants through this process. ΔFosB operates is liable for addiction-related structural changes in the brain and afterward sufficient accrual, using its downstream targets (e.g. NF-κB), it encourages an addictive state [Adapted from 20,21].

| Time Since Last Amphetamine Use | Amphetamine Withdrawal Symptoms |
|-------------------------------|--------------------------------|
| 1–3 Days                      | Comedown:                      |
|                               | • Exhaustion                   |
|                               | • Increased Sleep              |
|                               | • Depression                   |
| 2–10 Days                     | Withdrawal:                    |
|                               | • Strong urges (i.e., cravings) to use amphetamines |
|                               | • Mood swings (i.e., feeling anxious, irritable or agitated or feeling flat and lacking energy) |
|                               | • Poor sleep                   |
|                               | • Poor concentration           |
|                               | • General aches and pains, headaches |
|                               | • Increased appetite (very hungry) |
|                               | • Strange thoughts             |
|                               | • Misunderstanding things around surrounding (e.g. seeing things that are not there) |
| 7–28 Days                     | Most symptoms start to settle down, although common symptoms include: |
|                               | • Mood swings (i.e., feeling anxious, irritable or agitated or feeling flat and lacking energy) |
|                               | • Poor sleep                   |
|                               | • Cravings                     |
| 1–3 Months                    | • Return of normal sleep and levels of activity and mood |
|                               | • Major improvements in general health and mood |

Table 8: Common symptoms associated with amphetamine withdrawal [81].
Though tolerance does build rapidly, taking a break from the drug use can quickly cause the tolerance to diminish [65,84]. Unfortunately, this is one of the greatest dangers associated with amphetamine addiction treatment; users who stop taking amphetamines for a period of time have a lowered tolerance and when they relapse are more likely to overdose because they think that they can return to previous patterns of drug abuse which could prove to be too much for their newly reduced levels of tolerance [65].

Tolerance is also expected to develop with regular substituted amphetamine use [85]. When substituted amphetamines are abused, drug tolerance develops rapidly [72].

Toxicity of amphetamine and associated psychosis

Toxic central effects of amphetamine use include psychosis, hyperthermia, seizures and rhabdomyolysis, while cardiovascular toxicity includes ventricular arrhythmias, acute myocardial infarction and cardiomyopathies [86]. Neurotoxicity refers to neurological changes that persist after cessation of use and evidence suggests that chronic methamphetamine use leads to dopamine depletion and possibly also changes in serotonergic function [87]. Amphetamine toxicity produces a wide range of clinical effects displayed in Table 9 [62].

There is overwhelming evidence that patients with psychotic disorders have an increased vulnerability to compulsive use of drugs of abuse [88], including psycho stimulants like amphetamines [89]. The DSM-5 describes the following amphetamine-related psychiatric disorders presented in Table 10 [70]. The symptoms of psychosis induced by amphetamines are very similar to those of acute schizophrenia spectrum psychosis and include: lack of concentration, delusions of persecution, increased motor activity, disorganization of thoughts, lack of insight, anxiety, suspicion and auditory hallucinations [90-91]. Some studies have suggested differences with more pronounced grandiosity and visual hallucinations [92].

Clinical Signs Indicating Amphetamine Toxicity

• Seizure
• Focal neurological deficit
• Reduced level of consciousness
• Abnormal motor movements
• Hyperthermia
• Dysrhythmias
• Hypotension
• Hypertension
• Coronary artery spasm and related sequelae
• Autonomic instability and other major disturbance of homeostasis

Table 9: Clinical signs representative significant amphetamine toxicity [62].

Amphetamine-Related Psychiatric Disorders

• Amphetamine-induced anxiety disorder
• Amphetamine-induced bipolar disorder
• Amphetamine-induced psychotic disorder with delusions
• Amphetamine-induced psychotic disorder with hallucinations
• Amphetamine-induced sexual dysfunction
• Amphetamine-induced sleep disorder
• Amphetamine intoxication
• Amphetamine intoxication delirium
• Amphetamine withdrawal
• Amphetamine-induced obsessive-compulsive and related disorders
• Unspecified amphetamine-related disorder

Table 10: Psychiatric disorders associated with amphetamine [70].

There may be several explanations for this increased comorbidity, but there is convincing evidence from animal studies that this may be due to shared vulnerabilities for both psychosis and drug use disorders [93]. These animal studies also point to possible neural mechanisms explaining the increased comorbidity [94]. In rodents and primates, sufficiently high doses of amphetamine cause dopaminergic neurotoxicity, or damage to dopamine neurons, this is characterized by dopamine terminal degeneration and reduced transporter and receptor function [20,95]. There is no evidence that amphetamine is directly neurotoxic in humans [39]. However, large doses of amphetamine may indirectly cause dopaminergic neurotoxicity as a result of hyperpyrexia, the excessive formation of reactive oxygen species and increased autoxidation of dopamine [20,96]. Animal models of neurotoxicity from high-dose amphetamine exposure indicate that the occurrence of hyperpyrexia (i.e., core body temperature ≥ 40°C) is necessary for the development of amphetamine-induced neurotoxicity [97]. Prolonged elevations of brain temperature above 40°C likely promote the development of amphetamine-induced neurotoxicity in laboratory animals by facilitating the production of reactive oxygen species, disrupting cellular protein function and transiently increasing blood-brain barrier permeability [20,97].

A severe amphetamine overdose can result in a stimulant psychosis that may involve a variety of symptoms, such as delusions and paranoia [20,22]. A previous report on treatment for amphetamine, dextroamphetamine and methamphetamine psychosis states that about 5 to 15% of users fail to recover completely [20,98]. According to the same review, there is at least one trial that shows antipsychotic medications effectively resolve the symptoms of acute amphetamine psychosis [22]. Psychosis very rarely arises from therapeutic use [20,24].

Pharmaceutical products of amphetamine

Commonly prescribed amphetamine preparations like Adderall, Evekeo, Dyanavel XR, contain both the enantiomers. Among them Eveko is the racemic amphetamine sulfate [99]. Amphetamine is also available in the form of enantiopure and prodrug as dextroamphetamine and lisdexamfetamine correspondingly [46,100]. The prodrug form, lisdexamfetamine is structurally different from amphetamine and remains inactive until it is metabolized into dextroamphetamine [101]. Benzedrine, Symptamedrine and Psychodrine are the previously available free base of racemic amphetamine. On the other hand, Cylpril is the previously available form of levoamphetamine. Most of the current amphetamine preparations are available in the salt form rather than the free base form, due to the high volatility of the free bases [1]. Nevertheless, dosage forms like oral suspension and orally disintegrating tablet were introduced in the market in 2015 and 2016, respectively [102]. Some of the current brands and their generic equivalents are listed in Table 11 [102-104].

Regulatory status of amphetamine in society and culture

From the 1960s onward, amphetamine has been popular with many youth cultures in Britain and other parts of the world as a recreational drug [103]. It has been commonly used by novelists, mathematicians, film procurer and punks, gangsters and casuals in all night soul and ska dances, punk concerts, basement shows, etc. In Table 12, slang names of frequently abused amphetamines and substituted amphetamines are presented [62]. As a consequence of potential abuse, distribution of amphetamines and their analogues are controlled by the respective authorities.

According to United Nations 1971 Convention on Psychotropic Substances, amphetamine has been classified into schedule II controlled substance (Table 13) [104,105]. Therefore, use of amphetamine is
comprehensively controlled in most countries. Countries like South Korea and Japan have even banned the therapeutic uses of substituted amphetamines [106,107]. In contrast, some countries, including the Netherlands (list I drug), the US (schedule II drug), Canada (schedule I drug), Australia (schedule 8), United Kingdom (class B drug) and Thailand (category 1 narcotic) allow medical uses of amphetamine [108].

However, in these countries the medical uses of amphetamine are on a restrictive national drug schedule. Unfortunately, the illegal synthesis of amphetamine is still taking place in underground labs and is sold illegally, predominantly in European countries [108]. In 2013, about 1.2 million young adults of European countries used amphetamine or methamphetamine illegally. Whereas, the midst of 2012 around 5.9 metric tons of illegal amphetamine were detained within EU member countries. However, outside Europe, illegal market for amphetamine is much lesser as compared to MDMA and methamphetamine [109].

**Conclusion**

Even though amphetamine can be abused, these substances are additionally utilized for therapeutic purposes. Nevertheless, addiction to amphetamine particularly develops when prescription amphetamines are taken at much higher doses as compared to the doses used for therapeutic purposes. Consumption of gradually higher dosages due to “binge and crash” cycle and due to tolerance, physical dependence and psychological dependence are the possible phenomena with the abuse of amphetamines. In binge-crash cycle, following binge episodes, the amphetamine abuser crashes and faces anxiety, extreme fatigue, severe depression and a desire for more drugs. So, for the benefit of the nation, especially younger generation, caution must be taken to ensure rational use of amphetamines.

**Authors’ Contributions**

This work was carried out in collaboration between all authors. Author MSU designed the study, wrote the protocol, managed the analyses of the study and prepared the draft of the manuscript. Authors MAS, MTK, MFH, MN, II, AAM and MTI managed the literature searches and also participated in manuscript preparation. Author SK reviewed the scientific contents of the manuscript. All the authors read and approved the final manuscript.

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**Competing Interests**

The authors proclaim that they have no competing interests.
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5. European monitoring centre for drugs and drug addiction (2016) Table GPS-2: Last 12 months prevalence of drug use by age and country, most recent national general population survey available since 2000.
6. European monitoring centre for drugs and drug addiction (2016) Table EYE-20: ESPAD 2011 school surveys: Lifetime prevalence (percentages) of psychoactive substance use among students 15-16 years.
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