Ecstasy-Journey from Consumption to Death

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Submission: September 10, 2017; Published: December 11, 2017

Abstract
Ecstasy is an illicit compound, use of which is very famous amongst ravers and dance clubbers. It stimulates the over-release of serotonin and other neurotransmitters which psycho-physiologically cause thermal stress, elevated breathing and heartbeat, sharpened visio-audio senses and high blood pressure. User experiences the euphoric, light-minded and optimistic feelings which are exaggerated under loud, crowded and physically wearing environment. Although these feelings are short term and after frequent consumption, wondrous, care-free sensations are superseded by depression, pessimism, sleeplessness and loss of appetite. Current review describes the molecular reasons behind the effects ecstasy poses on naïve users and highlights how addiction for this drug begins and if not controlled how it eventually brings the demise of its users.

Introduction
3,4-methylenedioxymethamphetamine (MDMA) which is commonly known as “ecstasy”, is an amphetamine derived illicit compound which stimulates the release of serotonin and other neuro-hormones like cortisol, oxytocin and prolactin [1]. It is highly consumed by youngsters after cannabis. Its peak consumption was reported between early and mid-2000s [2]. But most recently, the use of its highly purified, crystalline prepared form has brought up some serious concerns [1].

Around three decades ago, it was consumed as a recreational drug. Since that time, ecstasy is widely used by youngsters in dance clubs and raves. Initially, college or high school students walking in groups and holding hands or laughing or singing together were recognized as the potential candidates and approached by researchers for MDMA related surveys [2]. MDMA is preferentially used among the youngsters because of its peculiar characteristics like mild hallucination, increased sociability, heightened perception of colour and sound and feeling of empathy [3]. Contrarily, its frequent use or high doses leads to dose dependency which is associated with depressive symptoms, increased anxiety and persistent deficits in cognition [4]. So, the purpose of the current literature is to shed light on the potential mechanism behind imposing its effects and bring into spotlight factors that could lead to its addiction.

Different Routes of MDMA Consumption
According to global drug survey, the pleasure and risk of any drug is directly linked to the method it is administered by the user. Globally, MDMA is administered either orally in the form of capsule or is inhaled accompanied by physical exertion (e.g. dancing). Around 90% MDMA users consume it via oral route (among which 48.9% swallow a capsule, 15.3% bomb it, 15.1% use it as dab powder and 9.7% drink it by dissolving it in any liquid) and 10% of its users snort it (GDS, 2017).

Ecstasy is easily absorbed from intestinal tract and within 2 hours of its oral consumption, it reaches at its peak plasma concentration. While its administration through nasal route brings about the quicker effect as finely ground drug easily gets absorbed in the porous nasal mucus lining and directly reaches bloodstream. That’s why its users are more attracted to snorting because of its rapid onset. The duration of MDMA effect and the duration of its onset after administration are indirectly proportional to each other. Due to quicker drug response, the duration of its effect is shorter which leads to frequent redoes and increases the risk of MDMA resistance. Moreover, rapid increase in its blood concentration puts the user at elevated risk of anxiety, palpitation and perspiration [5,6].

Symptoms and Lethal Effects of MDMA
MDMA has a potential of showing various psychological and adverse effects along with immune dysfunctioning. The diverse psychological and adverse effects are caused by plasma-MDMA induced elevated secretion levels of cortisol, prolactin, ACTH, dehydroepiandrosterone (DHEA), and antidiuretic hormone (ADH, vasopressin). As the plasma concentration of MDMA changes, it induces various immune dysfunctionalities. These might include CD4 T-helper lymphocytes decreased circulation, increase in NK cells and impaired lymphocyte mitogen-induced proliferation [7].
These effects can be categorized into: acute, short-term and chronic effects.

Short-term effects are those that last 24 hours after drug consumption. Depressed mood, fatigue, insomnia, dry mouth, heavy legs, loss of appetite, drowsiness, lack of energy, weakness, muscular tension, lack of concentration and head ache are the short term effects of ecstasy. Late short-term effects of ecstasy include anxiety, drowsiness, lack of energy, fatigue, and muscular tension which last for about 7 days [3,4].

Frequently reported acute side effects reported in the users are lack of appetite, jaw clenching, dry mouth, thirst and restlessness. Some other reported effects include palpitations, impaired balance, sensitivity to cold, drowsiness and insomnia. Muscular tension, anxiety, trismus, nystagmus and tremor are also found to be occurring in frequent users. MDMA also induces panic attacks, delirium and psychotic episodes which are resolved immediately after ceased drug activity [8,9].

Chronic effects of MDMA can be described as mild, moderate or severe. Nausea, vomiting, mydriasis, dry mouth, sweating, restlessness, tremor, hyper-reflexia, irritability, pallor, bruxism, trismus and palpitations are its mild effects. While moderate effects are hyperactivity, aggression, panic attacks, psychosis, muscle tension, tachycardia and increase in body temperature. Severe intoxication leads to hypotension, hyperthermia, seizures, delirium, renal failure and coma. Heat stroke and hyponatraemia are another severe complication linked with ecstasy toxicity which ultimately leads to renal failure and hepatic necrosis, respectively [10,11].

**Mechanism of Action**

The intensity and time of MDMA effect is largely dependent upon its route of consumption but through whatever route it is consumed it eventually finds its way to liver and there cytochrome P450 (CYP) CYP2D6, 1A2, 2B6, and 3A4 0-demethylate it and forms 3,4-dihydroxymethamphetamine (HHMA) . If its pharmacokinetics adopt minor pathway, then it is N-demethylated by the actions of CYP2B6, 1A2, and 2D6 and forms 3,4-methylenedioxyamphetamine (MDA). But before it is metabolized, it binds with CYP2D6 and inhibits its function and accumulates in plasma [12,13]. So, this plasma-MDMA actually interacts with serotonin reuptake transporters (SERT) and triggers the release of serotonin (5-HT) [14]. The elevated level of serotonin leads to serotonin syndrome which along with inflammatory environment (in most cases raves which includes loud music, light shows and are overcrowded) leads to mental confusion, hyperactivity, trismus and hyperthermia [15]. All this in psychophysiology term, causes faster breathing, tachycardia and high blood pressure in ecstasy user [16]. Moreover, MDMA induces the release of 80% of available serotonin in synaptic cleft [17] which along with reduction in left amygdale activity [18], brings about the initiation of dopamine and NO signaling pathway take a recreational user in a euphoric condition [19]. In this condition, user feels stress free, relax, light minded and lose all judgments for risks.

Meanwhile rest of the CYP enzymes act on MDA and convert it into its metabolites. Around 10% of MDA is converted into MDA (20) which is a agonist for 5-HT receptor and is a potent monoamine releaser [21]. Like MDA, it is also accumulate in plasma. Plasma-MDA then induces the over-production of cortisol [22] which leads to hyperthermia [14]. This thermal stress makes user feel hot along with pronounced perspiration and dehydration [23]. Some MDMA users then drink water in excessive amount which dilutes blood sodium electrolytes and bring about hyponatraemia [24].

This burst of energy, light-mindedness and joyous sensation feels good initially. But after frequent encounters with MDMA, the levels of serotonin and DA remain stable [25,26]. This creates the bodily urge to increase its dose and the phenomenon for addiction begins. NO which is a free radical gas plays very crucial role in addictive behavior. It acts as secondary messenger in central nerves system (CNS) and is produced by NO synthase from L-arginie [25]. The exact mechanism of how MDMA induces its production is still obscure but blocking NO signaling pathway is proven effective in relapse from MDMA abuse [19].

Once addiction for MDMA begins, the user either goes through frequent low dose encounters or sudden exposure to heavy a dose which eventually leads to neural injury at cellular level. Serotonin toxicity is common phenomenon observed [27]. First, its exposure induces the production of reactive oxygen species (ROS) in mitochondria and then initiates caspase dependent apoptosis pathway [28]. Further, MDMA also disrupts the homeostasis of Ca2+ ions by activating calpains and intensify mitochondrial fragmentation [29]. The fragmentation of mitochondria causes accumulation of phagocytes and leads to neurotoxicity [27]. It also induces the expression of Atg5 autophagy gene [30]. This injury consequently results into neuron depletion and chronic tolerance to MDMA. At this stage, all the pleasurable and light-minded sensations first experienced by user are replaced by episodes of depression, insomnia, reduced appetite and adverse health conditions [31,32].

If still the user does not stop and keeps on ingesting heavy doses of ecstasy, then as it activates sympathetic autonomic nervous system, so, user suffers from fever, vasconstriction and organ failure accompanied by cardiac failure. Later, sufferer also suffers from barotraumas [33].

**MDMA to Death**

MDMA or ecstasy is a ring substituted amphetamine which induces the euphoric feelings. Its half life is of 7 hours and reaches its peak levels after 1.5 to 2 hours of consumption. If its concentration elevates to 0.5 to 10mg/L in blood then it can pose serious toxic effects and in certain conditions can be life threatening [34,35].

A case study reported the death of 39 years old woman who after oral intake of MDMA collapsed and died after 7 days. On ingesting MDMA, she felt discomfort in anterior chest area and went through cardiopulmonary arrest. Her serum samples were taken on admission in hospital and revealed 1.2mg/L MDMA
concentration. Her autopsy results revealed myocytes necrosis along with macrophage inflammatory response and calcification. Liver necrosis, myoglobinuria, bronchopneumonia and neuron degeneration in whole brain is also reported. All these happenings clearly suggests that the demise of this lady was by MDMA intoxication [34].

The use of MDMA is itself very lethal but its consumption with other substances increases the likelihood of death. A recent study reported the death of 19 year old woman who two days prior to her death took MDMA along with alcohol consumption. The death was caused by diabetic ketoacidosis, a condition provoked by combined ingestion of MDMA and alcohol. Her biochemical tests revealed the vitreous glucose concentration of 6.5mmol/L and 13.86mmol/L b-OH butyrate. Further, microhemorrhages and swelling in her midbrain and corpus callosum is also reported [36].

The Therapeutic Role of MDMA

Recently the abilities of MDMA has been tested to cure posttraumatic stress disorder (PTSD), a psychologically depressive condition associated with psychological fragmentation, diminished sense of self-worth, trust and safety and loss of sense of self-coherence. In extreme conditions, PTSD sufferers go through severe personality changes including withdrawal, hopelessness, feeling of emptiness, negative self-perception, low-self esteem and disturbed emotions [37,38].

In such patients, MDMA assisted psychotherapy has shown recuperating outcomes. Positive alteration in the personality structure is reported which accounts for reduction in neuroticism and increase in openness to new social experiences and self-examination. Such changes ultimately lead to decrease in PTSD symptoms and are long-term [39]. Although, the hallucinogenic property of MDMA [3] is proven beneficial to improve depressive symptoms, still, more clinic experimentation is requisite to further understand its curative properties.

Conclusion

MDMA is a popular choice of recreational drug amongst youngsters and is usually used in raves and dance club. Naïve users enjoy its euphoric properties under loud and physically extorting environment. At molecular level, such users experience bust of serotonin and electrolyte imbalance which in long run leads to depression, anti-social behavior, organ failure and cardiac failure. So, ecstasy is actually a method of slow-suicide adopted by young generations which after 5-10 years of frequent use irreversibly damage neural system of users. Although anti-narcotic agencies are trying to control its trafficking worldwide and awareness programs are also being run. Still, there is a strong urge to monitor its use in colleges and universities. Administration should periodically conduct seminars and workshops to aware youngsters and their parents about the dark side of drug use and should also spread knowledge on avoiding the impulse for its use. Through the effort of many researchers, today we know most about the mechanism of MDMA addiction and also know potential ways to heal from its addiction. Despite all that, the room for more confirmatory experiments and discovery of novel recovery treatments is still available.

References


