FOCUS: PSYCHOSTIMULANTS

Mechanism of Action and Therapeutic Uses of Psychostimulants

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ABBREVIATIONS: ADHD = attention-deficit hyperactivity disorder; CNS = central nervous system; MDA = 3,4-methylenedioxyamphetamine; MDMA = 3,4-methylenedioxyamphetamine; NC = not controlled; OTC = over the counter.

INDEX TERMS: psychostimulants.


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The name stimulant can be given to any drug that increases the rate of a physiologic function. The term psychostimulant is more specific, referring to compounds that have direct neurological effects, typically: heightened alertness, increased energy, appetite suppression, and sometimes euphoria. Use of psychostimulants is widespread and occurs in both recreational and clinical settings. Therapeutic monitoring and screening for use and abuse of these drugs is common in clinical laboratories. Understanding the physiological effects and uses of stimulants will be of value to clinical and forensic laboratory scientists.

EXAMPLES OF PSYCHOSTIMULANTS

Table 1 lists several psychostimulants that are commonly prescribed, have widespread illicit use, or are well-known by the general public. Table 1 also identifies the control schedule of each compound. Many psychostimulants are listed as controlled substances under the Controlled Substances Act, Title II of the Comprehensive Drug Abuse Prevention and Control Act of 1970. Schedules I through IV are assigned to compounds based on their abuse potential and their medical utility. Schedules II through V contain drugs which have known medical uses whereas Schedule I compounds have no current, sanctioned medical use.

THERAPEUTIC USES OF PSYCHOSTIMULANTS

Amphetamine has long been known to be a mental stimulant. Because of its psychostimulant properties, amphetamine has been used successfully by U.S. fighter pilots because it enhances cockpit performance by reducing the effects of fatigue. Since amphetamine heightens alertness it has found use in the treatment of narcolepsy and in attention-deficit hyperactivity disorder. The drug Adderall® for example, is used widely in cases of ADHD and is merely a mixture of amphetamine's stereoisomers: dextro- and levoamphetamine.

Amphetamine also has appetite suppressant effects and thus is used in the treatment of obesity, although it is not FDA approved for this use. Although amphetamine can be considered the prototype psychostimulant, many psychostimulants have been identified or created that resemble amphetamine in their chemical structures. These compounds vary in their effects and utility as well as their abuse potential. Methamphetamine, like amphetamine has a history of illicit use and is known to be considerably more potent in vivo than the unmethylated amphetamine. Methamphetamine, like amphetamine, has been used with some success in ADHD patients. It is also approved for use in treating obesity.

Pseudoephedrine, the name given to the 1R2R enantiomer, and ephedrine, the 1R2S enantiomer, are common medications used primarily as nasal decongestants. Unlike amphetamine and methamphetamine they have little abuse potential allowing them to be obtained OTC. Ephedrine is also approved for treatment of bronchospasm, enuresis, hypotension, and narcolepsy.

Some psychostimulants have been found to have uses that are not typical for other drugs in the same class. Although bupropion is a phenylalkylamine, the structural dissimilarity with amphetamine is significant enough to produce considerably different physiologic effects. Bupropion is an...
example of a weak psychostimulant that has found use as an antidepressant (Wellbutrin®) and is also marketed as a smoking cessation aid (Zyban®). Although still a mild psychostimulant, the behavioral effects of bupropion are reported to be unlike amphetamine when used in humans. The abuse potential is also considerably less than with amphetamine.7 Off-label uses of bupropion include treatment of ADHD, diabetic neuropathy, and neuropathic pain.5 Another clinically-useful β-ketone psychostimulant is diethylpropion. Diethylpropion is currently used only in the treatment of obesity. This psychostimulant is commonly prescribed as an appetite suppressant under the name Tenuate®.

Methcathinone is a Schedule I psychostimulant that is easily synthesized from ephedrine. Methcathinone is the N-methylated version of cathinone, a natural psychostimulant obtained from the Catha edulis plant and known as “qat” or “khar”. Methcathinone has no sanctioned medical use and is only used recreationally. Its effects are those of a classic psychostimulant, causing ‘flying euphoria’ followed by a five-to-eight hour period of feeling tough, invincible with increased libido, and desire to be physical.8,9

The next two compounds listed in Table 1 are also Schedule I compounds. 3,4-methylenedioxymethamphetamine (MDMA) and 3,4-methylenedioxyamphetamine (MDA) are popular recreational drugs known respectively as ‘the love (or hug) drug’ and ‘ecstasy’. MDMA use is growing rapidly and in some European countries is the second most frequently used illegal drug after marijuana.10 Interestingly, both MDMA and MDA were used therapeutically before being classified Schedule I in 1985. These compounds were used by patients in constructive psychotherapy sessions.8 MDMA has received much negative attention in recent years as claims of its neurotoxicity mounted. However data concerning the neurotoxicity of MDMA have been inconsistent and one major study was even retracted after it was found that the toxicity in the study was brought about by methamphetamine rather than MDMA.11,12 The widespread use and popularity of MDMA coupled with the lack of definitive reports on neurotoxicity have encouraged some to reconsider the utility of this drug.13-15 Because of MDMA’s unique effects, a new study investigating the therapeutic use of MDMA has recently earned IRB and FDA approval. This research will assess the use of MDMA in the treatment of post-traumatic stress disorder.16,17

Perhaps the most well-known psychostimulant after amphetamine is methylphenidate, made popular under the trade name Ritalin®. Methylphenidate has pharmacological effects similar

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**Table 1. Some common psychostimulants**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Trade or common name</th>
<th>Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphetamine</td>
<td>Dextedrine®, Adderall®</td>
<td>C-II</td>
</tr>
<tr>
<td>Methamphetamine</td>
<td>Desoxyn®, Methamphetamine® ‘speed’, ‘crystal meth’</td>
<td>C-II</td>
</tr>
<tr>
<td>Ephedrine</td>
<td>Herbal sources: ma huang, ephedra, ephedra sinica, epitonin, Pretz-D® OTC</td>
<td></td>
</tr>
<tr>
<td>Bupropion</td>
<td>Wellbutrin®, Zyban®, Buproban™</td>
<td>NC</td>
</tr>
<tr>
<td>Diethylpropion</td>
<td>Tenuate®, Dipro™ Durad™ Radtue™</td>
<td>C-IV</td>
</tr>
<tr>
<td>Methcathinone</td>
<td>‘cat’</td>
<td>C-I</td>
</tr>
<tr>
<td>3,4-Methylenedioxymethamphetamine</td>
<td>MDA, ‘love drug’</td>
<td>C-I</td>
</tr>
<tr>
<td>3,4-Methylenedioxyamphetamine</td>
<td>MDMA, ecstasy, XTC</td>
<td>C-I</td>
</tr>
<tr>
<td>Methylphenidate</td>
<td>Concerta™, Metadate®, Methylin™, Ritalin®</td>
<td>C-II</td>
</tr>
<tr>
<td>Phenetermine</td>
<td>Adipex-P®, Fastin®, Obenix®, Phentamine®, Supramine™</td>
<td>C-IV</td>
</tr>
<tr>
<td>Fenfluramine</td>
<td>Pondimin® (discontinued, removed from the U.S. market)</td>
<td>C-IV</td>
</tr>
<tr>
<td>Cocaine</td>
<td>Methyl benzoyllecgonine, coke, crack</td>
<td>C-II</td>
</tr>
</tbody>
</table>

C-I = Schedule I: Compounds with high potential for abuse having no currently accepted medical use in treatment in the United States.
C-II = Schedule II: Compounds with high potential for abuse having currently accepted medical use(s) in treatment in the United States or a currently accepted medical use with severe restrictions.
C-IV = Schedule IV: Compounds having a low potential for abuse relative to drugs scheduled as C-III.
NC = not controlled (unscheduled).
OTC = over the counter.
to amphetamine; however, its abuse potential is somewhat lower, although there are many conflicting reports regarding methylphenidate abuse potential (see reference 18 for review). Methylphenidate has become the drug of choice for treating ADHD but has also found off-label use as an antidepressant.5

Two other drugs worth mentioning as psychostimulants are phentermine and fenfluramine. Both of these drugs have been used as appetite suppressants and both have amphetamine-like sympathomimetic effects. Fenfluramine contains a chiral center and fenfluramine is the name given to the racemic mixture of D- and L-isomers. The D-isomer (dexfenfluramine) was purportedly responsible for the anorectic actions of fenfluramine and was also associated with fewer side effects than the racemic fenfluramine. Dexfenfluramine was marketed as Redux®.19 By 1997 both fenfluramine and dexfenfluramine were removed from the U.S. market at the request of the FDA when cases of cardiac valvulopathy were reported.5,20,21 Fenfluramine is perhaps best known by its off-label use with phentermine, known as “Fen-phen”. Fen-phen was widely used for the long-term management of obesity. The fenfluramine-phentermine combination was also associated with valvulopathies and a bulletin was issued by the FDA to cease off-label use of the Fen-Phen combination.5,21 Of the two drugs, only phentermine is still used as an appetite suppressant although tolerance to the anorexiant effects of phentermine usually develops a few weeks after starting therapy.

When considering the clinically-used drugs listed in Table 1, cocaine is the only one that is not used for its psychostimulant effects. Cocaine is used clinically only as a local anesthetic, usually in mucosal or ophthalmic procedures.

MECHANISM OF ACTION

It is easy to see the resemblance between the chemical structures of common psychostimulants and endogenous monoamine neurotransmitters (Figure 1). The prototype psychostimulant amphetamine closely resembles the catecholamine neurotransmitters norepinephrine, epinephrine, and dopamine. Since many psychostimulants share the features of a phenyl ring, a nitrogen group, and carbon side chains of varying lengths, many stimulants fall into the category of phenylalkylamines. With the possible exception of cocaine, all the compounds listed in Table 1 can be classified as phenylalkylamines. Because amphetamine is considered the prototype stimulant, other compounds that have similar chemical structures and similar physiologic effects are often termed ‘amphetamine’ (Figure 2).

Given their structural similarity to endogenous neurotransmitters, it is not surprising that many phenylalkylamines have autonomic nervous system activity, i.e., sympathomimetics, as well as mood-altering effects. Amphetamine and other closely related phenylalkylamines can activate receptors that normally bind catecholamines or serotonin. In addition, amphetamine and related compounds can cause the release of catecholamines and serotonin from nerve endings.22-26 Once released, the endogenous neurotransmitters are free to act on their extracellular receptors. When catecholamine receptors in the brain are activated, myriad effects can result. Neuropsychological effects of catecholamine receptor activation or potentiation by psychostimulants can include: increased alertness, insomnia, euphoria, decreased appetite, and at higher doses, psychosis. Indeed, amphetamine can correctly be called a “psychotomimetic” since high doses can bring about a psychosis very similar to that seen in schizophrenic patients.27,28

In the periphery, activation or potentiation of catecholamine receptors by psychostimulants can result in: vasoconstriction and subsequent hypertension, mydriasis, tachycardia, and other general sympathomimetic effects. The mechanism of action of amphetamine and amphetamine-like psychostimulants involves four major effects:

1. Binding to extracellular catecholamine receptors
2. Inhibition of monoamine neurotransmitter uptake
3. Release of catecholamines from neurons
4. Inhibition of monoamine oxidase

Figure 1. Structure of amphetamine and catecholamine neurotransmitters
Psychostimulants alter neurotransmitter release. Psychostimulants can effectively alter the amount or the rate of neurotransmitter released by a monoaminergic neuron. The resulting change in mood or behavior is the summation of these neuromodulations and is quite complex. At the cellular level this neuromodulation is due to one or more of the effects listed above. Amphetamine-like compounds have a wide range of affinities to catecholamine and serotonin receptors. The overall pattern of receptor binding for a psychostimulant is unique for a given stimulant and contributes to the distinctive behavioral effects of a given compound.

In addition to having direct receptor-binding effects, many psychostimulants inhibit monoamine transporters. Monoamine transporter proteins serve to recycle neurotransmitters after they are released from the neuron and in so doing, terminate the neurotransmitter signal. These same monoamine transporters are also the targets for antidepressant medications including the popular drugs fluoxetine (Prozac®), paroxetine (Paxil®) and sertraline (Zoloft®). Inhibitors of monoamine transporters block the uptake, or ‘reuptake’, of neurotransmitters by neurons (Figure 3). This blockade effectively increases the concentration of neurotransmitter in the synapse, resulting in increased binding of neurotransmitters to their receptors.

**Figure 2. Structures of some common psychostimulants**

Structural similarity can be seen among many of the compounds containing the phenylalkylamine structure (compounds shown are from Table 1).
Psychostimulants can also elevate the concentration of neurotransmitter in the synapse by evoking release of neurotransmitters. The release of stored neurotransmitters can be triggered directly when psychostimulants bind receptors present on neurons or can release neurotransmitters indirectly via an exchange mechanism occurring through monoamine transporter proteins. The exchange mechanism or efflux process which can occur via monoamine transporters has been observed with many amphetamine derivatives. This efflux is thought to be a type of ‘reverse-transport’ mediated by monoamine transporters. Finally, amphetamine and amphetamine-like psychostimulants often act as competitive inhibitors of the enzyme monoamine oxidase (MAO). MAO is a mitochondrial enzyme that breaks down monoamine neurotransmitters. Products of catecholamine breakdown include 3-methoxy-4-hydroxymandelic acid (VMA), homovanillic acid (HVA), and dihydroxyphenylacetic acid (DOPAC). These compounds are commonly-measured metabolites of catecholamines whose formation is due in part or in full to MAO. Inhibition of MAO would thus bring about an expected increase in monoamine neurotransmitters since their breakdown would be impeded.

In summary, psychostimulants can affect neurotransmitter release in at least four ways; by binding extracellular receptors, by blocking monoamine transporters, by evoking neurotransmitter release, or by inhibition of monoamine oxidase.
and these systems are also affected by psychostimulants like amphetamine. In addition to appetite control, amphetamine is also used for its ability to focus attention. The basis for this effect is not well understood. The seemingly paradoxical fact that psychostimulants can act as calming agents in humans seems enigmatic. The effects of amphetamine on attention and alertness are ultimately due to the modulation of normal patterns of central activity. These modulations are brought about by modulating serotonergic, dopaminergic, and noradrenergic pathways but the precise mechanism of action of amphetamine in ADHD is unknown (see references 2 and 40 for reviews).

Euphoria, of course, is an important, high-dose effect of amphetamine. The euphoric effects of amphetamine and methamphetamine are very similar to cocaine however unlike cocaine, amphetamine and methamphetamine are taken orally and are metabolized more slowly than cocaine, making it the psychostimulant of choice for many drug users. Centrally, amphetamine and cocaine cause an acute dopamine release in addition to inhibiting neuron dopamine uptake. This release of dopamine in the nucleus accumbens and prefrontal cortex is thought to cause the prominent euphoric and reinforcing effects associated with amphetamine and cocaine. These psychostimulants can produce a significant euphoria, locomotor stimulation, reduced fatigue, sexual stimulation, increased mental attention, and increased sociality. Higher doses can produce tremors and vomiting, as well as tonic-clonic convulsions.

CONCLUSION
Psychostimulants are important pharmacological agents used in a variety of conditions including the treatment of obesity, ADHD, narcolepsy, and as decongestants. Psychostimulants typically contain a phenylalkylamine structure that closely resembles that of endogenous monoamines. The actions of psychostimulants are mediated through their ability to bind monoamine receptors and/or modulate monoamine transmitter release. Although psychostimulants have a history of abuse and misuse their therapeutic uses are numerous and significant.

REFERENCES
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