Smart drugs among students – multidimensional view on present ways of cognitive enhancement

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Abstract:

**Background:** Nootropic drugs, at the beginning of their career, were used in the treatment of diseases such as attention deficit hyperactivity disorder (ADHD), Alzheimer's disease and narcolepsy. Nowadays, they are becoming more and more popular among students, where they are often called smart drugs or cognitive enhancers. Their properties that improve
intellectual abilities and cognitive functions are used to, among other things, improve academic performance.

**Aim of the study:** The purpose of this article was to summarize the current state of knowledge relating to the use of smart drugs among students. This study discusses the multidimensional issues of using this group of drugs.

**Material and methods:** A cross-sectional search of relevant literature was performed in databases using accepted wording. Publications from 2003-2023 were included. The entire process has been supplemented with additional procedures to increase the reliability of this publication.

**Results:** In most sources, the most common representative of smart drugs turned out to be methylphenidate. Nootropics were most often obtained by obtaining a prescription from a doctor, online trading or from friends. The factors motivating the use of substances from this group are primarily the desire to improve academic results, competitiveness and the number of obligations at the university. There are many risks associated with their use, but students are more aware of the positive effects.

**Conclusion:** This paper presents various dimensions of the progressive spread of nootropic drugs. In addition to their undeniably positive effects, there are many risks, which are discussed in this article. Students' awareness should be formed through appropriate educational methods.

**Keywords:** nootropic agents; students; academic performance

**Introduction:**

Nootropic drugs have revolutionized the treatment of many disorders at the intersection of psychiatry and neurology. They were originally developed to treat a range of disorders, including attention deficit hyperactivity disorder (ADHD), Alzheimer's disease or narcolepsy [1]. Their multidirectional effects on improving cognitive brain function have been noticed and appreciated by many communities, thus gaining a lot of space for their off-label use. This had the foundation for defining a new group of drugs called smart drugs or cognitive
enhancers. To complete their definition, it was assumed that these are prescription drugs, taken without indication and in doses above the recommended one, which enhance intellectual abilities and cognitive functions, such as concentration, alertness or memory [2]. Research on this group of substances originated during World War II, when stimulants (mainly amphetamine and modafinil) were administered to soldiers to keep them alert and counteract the fatigue that accompanied them [3]. In modern times, college students have become the main consumers, and the scale of their prevalence is shown by estimates that the demand and subsequent profits from their sale will reach nearly $5 billion by 2025 [4].

This paper focuses on pharmacological nootropic agents and takes a closer look at the contemporary problem of their prevalence, the motivation for their use and the resulting risks among the student population. The mechanism of action of the three most popular substances belonging to the aforementioned group was also described.

**Material and methods:**

Initially, a cross-sectional search was performed in PubMed database. For this purpose, the following formulations were used: ((Nootropic drugs[Title/Abstract]) OR (Nootropics[Title/Abstract]) OR (Smart drugs[Title/Abstract]) OR (Brain enhancers[Title/Abstract]) OR (Study aids[Title/Abstract]) OR (Cognitive enhancers[Title/Abstract])) AND ((students[Title/Abstract]) OR (student[Title/Abstract])). Publications from 2003-2023 were included. A complementary hand search of the literature cited in the preselected articles was also performed to find additional publications meeting the objectives of this review. Reviewers conducted an initial qualification of articles based on titles and abstracts to assess their suitability to meet all the objectives of this work. Those that clearly did not meet the inclusion criteria were excluded. The remaining publications were meticulously analyzed in full text for final qualification for inclusion in the literature items of this work.

**Classification of smart drugs:**

The term neuroenhancement can be considered multidimensionally. Substances to which such a term is attributed belong to a numerous and heterogeneous group, which includes
methylphenidate, modafinil and piracetam, among others. Many authors complete the classification with stimulants such as amphetamine and its derivatives, explaining the enhancement of cognitive abilities through their effect of improving executive functions, as well as an increase in wakefulness or lack of feeling of fatigue.

On the other hand, the definition of neuroenhancement also includes drugs with sedative effects. For many researchers, it is clear that substances with sedative potential are not used directly to improve learning performance, but they can affect performance. A state of rest and relaxation can increase the likelihood of more effective learning. Thus, for example, taking cannabis to relax, stop recurring thoughts and facilitate falling asleep would allow one to remain more alert and focused the next day [5]. Mention should also be made of those natural and everyday counterparts to the aforementioned, also known by some as soft enhancers. There is no doubt that products such as coffee, energy drinks, dietary supplements and caffeine pills find many consumers in a much more representative group than the previously mentioned drugs. This is not surprising looking at their widespread availability and low addictive potential. Many authors include them in their publications and report data with them in mind. Soft enhancers are common among students in everyday life, and also their use seems to be socially acceptable [6].

Mechanism of action:

The group of substances collectively referred to as nootropics is not homogeneous. They differ in chemical structure, mechanism of action and effects on the body, producing different clinical effects. Considering social aspects, they are characterized by varying prevalence and popularity, the greatest of which is attributed to the drugs presented below, where their mechanism of action is discussed.

Methylphenidate

Methylphenidate is a representative of compounds in the phenylethylamine group. It stimulates the noradrenergic and dopaminergic systems in the central nervous system. Its action takes place through the release, sensitive to reserpine, of the contents of synaptic vesicles [7]. Methylphenidate increases dopamine and norepinephrine concentrations in the synaptic space by blocking the dopamine transporter (DAT) and the norepinephrine transporter (NET). It has been shown to improve cognitive function, enhance planning skills
and improve working memory. However, it worsens local blood flow in various brain regions [8]. Methylphenidate's effects are also based on its agonist function toward serotonergic receptor 1A (HTR1A) [9,10]. In addition, it affects neurotrophic effects on neural tissue via brain-derived neutrophic factor (BDNF), which influences neuronal survival and plasticity. The evidence for the above actions is unclear, as some studies suggest inhibition [11], while others show activation and an increase in plasma BDNF levels in children with ADHD [12]. It has also been reported that methylphenidate affects the regulation of apoptosis-related proteins, which may have a role in neuronal survival [13]. Moreover, studies suggest the immunomodulatory properties of methylphenidate by triggering inflammation via the release of interleukin 1 (IL-1) and tumor necrosis factor (TNF) [14]. Methylphenidate's range of action also includes regulation of mediators and transcription factors such as adenylyl kinase isoenzyme 1 (AK-1), mitogen-activated protein kinase 3 (MAPK-3), c-Fos protooncogene (FOS), calcium and calmodulin-dependent kinase type IV (CAMK-4) [15].

**Modafinil**

Modafinil stimulates the noradrenergic system of the thalamus, frontal cortex and hypothalamus. It also activates dopaminergic receptors of the nucleus accumbens, striatum, frontal cortex and locus coeruleus [16]. It causes an increase in the concentration of norepinephrine and dopamine in the synaptic gap [17,18]. Modafinil also exhibits neuroprotective and antioxidant effects, which may affect excitatory properties [19,20]. In other studies, modafinil has been shown to reduce the release of gamma-aminobutyric acid. Unlike amphetamine and its derivatives, modafinil exhibits weaker dopaminergic activity, which has a tidal effect on its weaker addictive potential. An additional mechanism of action of modafinil involves the hypocretin system. Hypocretin, also known as orexin, regulates appetite. One study found reduced levels of hypocretin in dogs with narcolepsy, so it is speculated that it may be responsible for wakefulness. Hypocretin peptides stimulate histamine release in the tuberomammillary nucleus, affecting the sleep-wake cycle. In one study, hypocretin administered to rats caused a prolongation of the wakefulness phase [21].

**Piracetam**

Piracetam is another nootropic drug used to improve cognitive abilities. It belongs to the pyrrolidone group and is a derivative of gamma-aminobutyric acid (GABA). It improves memory, concentration and psychomotor performance. It has also been proven to enhance memory and thinking processes in patients after a stroke or head injury. It also improves
cognitive function in patients who have experienced memory loss as a result of coronary artery bypass graft (CABG) surgery. [22]. Piracetam improves brain nerve cell metabolism, increases oxygen and glucose utilization, and improves cerebral blood supply [23]. Moreover, it protects mitochondria from oxidative stress and aging. Although the protective properties of piracetam in in vitro studies are modest, which can be explained by unnatural and overly demanding conditions. Leuner et al. report that in the pathophysiological view of the slowly progressive changes that occur during aging, these processes are sometimes completely inhibited by treatment with piracetam. This is likely to happen through an increase in the fluidity of mitochondrial membranes, which may be the most important factor regulating their function [24]. It should be added that piracetam does not exhibit free radical scavenging activity [25], so it can be assumed that its action is limited directly to the mitochondria.

**Commonness of cognitive enhancement:**

There are conflicting data on the prevalence of smart drugs among students. Esposito et al. report that it ranges from 2-80%, with an average of 22.81% [26]. In the United States, on the other hand, Teter et al. found that 24.5% of students included in the study confirmed taking methylphenidate [27]. In another American study, this percentage is as high as 96% [28]. The number of users of brain enhancers is steadily increasing, and men are more likely to admit to using them [26]. In contrast, Miranda et al. published the results of their study, in which women were more likely to use coffee, tea and dietary supplements [29]. According to some sources, the largest number of students used smart drugs without a prescription or medical consultation. The most common ways to obtain them were through friends and e-commerce [26]. Data on the prevalence of student use of nootropics varies depending on the source. This is influenced by the nationality of the study group, the year of the study, the field of study and the country in which the study was conducted. Among the French population of medical and pharmacy students, Franchi et al. proved that 5.8% of the study group admitted to having used substances from the aforementioned group. Methylphenidate was the most popular (3.6%), followed closely by amphetamine and piracetam, whose use was confirmed by 2.9% of respondents each, respectively [30]. The popularity of methylphenidate was also reflected among the Portuguese academic community, where, of the drugs used, methylphenidate again had the highest percentage (35.1%). The next choices were modafinil (10.4%) and idebenone (2.6%), indicating a different preference than noted by Franchi et al [29].
Despite the greatest popularity of methylphenidate among smart drugs, which was demonstrated in most of the studies included in the literature of this paper, there are substances that legitimize greater prevalence in some academic communities. Among Western Australian students, dextroamphetamine was the most common pharmaceutical of choice, with 5.3% of respondents admitting to using it. This is more than three times the rate for methylphenidate, whose prevalence was put at 1.5% [31]. The vast majority of users of these drugs did not have a prescription or medical recommendation for their use, and all of them began consumption during their first year of school [30]. In comparison, other sources found no statistically significant relationship between the frequency of use of enhancers and the year of study [29]. In one large study involving 6725 students, 13.8% of respondents indicated that they had used prescription drugs or narcotics at least once for cognitive enhancement. The most common drugs were again methylphenidate (4.1%), and surprisingly sedatives (2.7%) and beta blockers (1.2%). The above results were more applicable to those students who reported higher levels of perceived stress at the same time. Consumption of soft enhancers, which included coffee, energy drinks and vitamins, was found to be much more common. These were consumed daily in the month leading up to the exam, while their aforementioned pharmacological counterparts were used relatively infrequently and more sporadically, mainly immediately before a pass. The highest predilection for the use of smart drugs was among architecture students (19.6%), and the lowest in mathematics (8.6%).

A very important aspect is the source from which young people obtain the mentioned substances. It turns out that they most often get them from a doctor who writes a prescription [6]. This is also confirmed by Miranda et al. where they detailed the exact medical professions where students realize their needs. It was proven that 33% of the respondents, despite the lack of a diagnosis, manage to get a prescription from a family doctor, another 22% from a psychiatrist. On the other hand, they mainly get their knowledge and information from websites and medical books [32].

**Motivation for using smart drugs:**

Finding a common denominator to comprehensively explain what motivates students to decide to take the above measures seems impossible, due to the complexity of the issue at hand. Esposito et al. in their systematic review showed that smart drugs are most often taken for their effects on better concentration, greater cognitive abilities, reduced stress levels, more
efficient time optimization and curiosity [26]. What is more, students who reported feeling more pressure to perform in their studies, at work or among their families were more likely to admit to experiencing nootropics [6]. One study on a group of UK students aimed to examine the relationship between attitudes toward using aids. They proved that the most important positive predictors for their use were competitiveness and superficial motives, understood as the need to work enough to meet only minimum requirements.

In contrast, among the negative predictors, students' academic performance was found to be the most significant [33]. Among the French population of medical and pharmacy students, the desire to improve academic performance was the most common motivation for their use, as confirmed by all respondents. Increasing arousal and concentration, as well as reducing body weight and achieving a state of euphoria were found to be the next most common reasons [30]. Students who admitted to using pharmacological cognitive enhancers presented higher levels of perceived stress and lower coping abilities. As a strategy to cope with the demands of academic life, they chose the one based on coping with emotions, and the closer they got to their deadlines, they focused directly on their chosen goals [34]. In turn, Merwidi-Ląd et al. report that among the Polish population of medical and dental students, the most common reasons for reaching for reinforcers were feeling overwhelmed by the number of tasks (68% of respondents), followed by stress during studies (50%). What is more, nearly 20% of respondents answered that the mentioned means were recommended by their friends, who suggested the effectiveness of these substances, although it is questionable according to some studies. One of these is an experiment on the potential of methylphenidate at different doses (10 mg, 20 mg, 40 mg and placebo) in affecting a wide range of cognitive functions in healthy and young adults. Participants were subjected to tests of attention and episodic and working memory. No differences in performance were observed in any case, thus negating the cognitive effects of methylphenidate in this group of subjects [35].

Despite the increasing percentage of students who choose to supplement with the above drugs, there is still a group that believes the benefits of their use are overestimated. Students who have never used smart drugs are more likely to argue that their colleagues' reasons for reaching for them are due to insufficiently well-planned work. They also add that the above pharmaceuticals even hinder the formation of planning and learning skills. Some work argues that users of brain enhancers overestimate the benefits associated with their potential intellectual enhancement effects [36]. Nevertheless, participants in one study agreed that the decision to use them should be individual and autonomous. Only those who had no experience
with these substances supported the finding of unregulated access to them, indicating that they may cause more pressure among the student population and that it may result in inequality in educational and professional development. Interesting findings also came from those students who were in the process of preparing their presentations for scientific conferences. A significant percentage (13.41%) responded that speaking and communication skills were most important to them, and the desire to improve them was the main motivator for using smart drugs [29].

In addition to the reasons, understood as the goals that individuals want to achieve with the help of pharmacology, it has been proven that there are certain personality traits that predispose to this. It turned out that low extroversion, low agreeableness, high conscientiousness are conducive to reaching for them. In addition, circumstances such as alcohol consumption and anxiety intensify the consumption of nootropic drugs [37].

**Risks and side effects:**

Many studies demonstrate the lack of knowledge regarding the side effects of smart drugs and their addictive potential [38]. Many authors agree that this awareness can be built through educational campaigns [39]. In one study, the most commonly reported side effects of this group of agents include nervousness (27.1%), sleep disturbances (26.4%) and headaches (25%). Participants also reported experiencing depression (18.1%), loss of appetite (17.9%) and tachycardia (15.8%). A less numerous but socially important group of side effects included problems with family (6.3%) or friends (3.8%). Some students experienced anxiety attacks (7.4%) or fits of aggression (6.6%). Several respondents admitted to financial problems (2.4%), problems with the police (2.3%), accidents (0.9%), and even unconsciousness (2.5%) [6]. Students were generally more knowledgeable about the benefits and possibilities of using nootropic drugs than about their potential side effects, which is a significant predictor of abuse. The potential for addiction was the most frequently cited problem resulting from their consumption. In contrast, respondents confirmed the lack of available information on health effects as a source of the aforementioned risks [40]. In another study, respondents who were not interested in enhancing their intellectual abilities with this group of drugs unanimously answered that the main factors that kept them from doing so were lack of need to do so (73.23% of respondents) and fear of their health consequences (37%).
The dangers of using enhancers are generally well understood and widely reported in the literature. Among both students who have used and those who have not had experience with them, there is a belief that such substances cannot be considered without negative consequences. The frequency and type of side effects that accompanied those who used drugs from the group in question depended, among other things, on the type of pharmaceutical. For the aforementioned dextroamphetamine, popular among Australians, almost two-thirds of students experienced negative effects, where, in contrast to modafinil, their frequency was less than half. Respondents mainly complained of insomnia, anxiety and loss of appetite [31].

Conclusions:

This paper presents various dimensions of the progressive spread of nootropic drugs used as pharmacological cognitive enhancement. It may entail many risks due to the lack of sufficient knowledge and awareness of how smart drugs can harm health. Unfortunately, the measures being taken to curb these problems are not keeping pace with the rate at which brain enhancers are gaining popularity among students. What is needed, therefore, are targeted strategies and the dissemination of information about the effects of smart drugs and their risks. Universities should be responsible in educating students in this area. Promising results have been noted using an education methodology referred to as "peer to peer," in which people with similar levels of experience learn from each other and exchange knowledge [40]. The aspects considered in this article focus on off-label use of the aforementioned substances. It is important to remember that the treatment of many neurological diseases would not be possible without them, which thus puts them in a completely different light.

Author Contributions:

Conceptualization, Piotr Pisera and Aleksandra Kielkowicz; methodology, Daniel Ślusarczyk and Bartłomiej Źmuda; software, Filip Pactwa and Zuzanna Popińska; check, Wiktoria Jakubowska and Michał Żuberek; formal analysis, Piotr Pisera and Daniel Ślusarczyk; investigation, Aleksandra Kielkowicz and Zuzanna Popińska; resources, Bartłomiej Żmuda, Wiktoria Jakubowska and Michał Żuberek; data curation, Filip Pactwa, Daniel Ślusarczyk and Wiktoria Jakubowska; writing - rough preparation, Piotr Pisera; writing - review and editing, Piotr Pisera and Aleksandra Kielkowicz; visualization, Piotr Pisera, Bartłomiej Żmuda and
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