Non Benzodiazepines Hypnotics: Another Way to Induce Sleep

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Non Benzodiazepines Hypnotics: Another Way to Induce Sleep

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Summary:

The third generation hypnotics, zolpidem and zopiclone can be used to optimize rest periods during sustained operations and help the military personnel. This is possible because these drugs provide a good quality of sleep, without residual effects in the morning after administration. A literature review of the possible limitations of use of these drugs in pilots has been presented.

1. Introduction

Technological advances have improved the ability of military Air Force crews to withstand the operational demands of increased workload in a hostile environment (60). The 1986 air raids in Libya (59) and the Falklands war have shown, however, that physical and psychological fatigue of the pilots and ground support crews can reduce operational success. This is especially the case when operations take place from aircraft carriers in extreme conditions of frequent flights and little rest, including flights 24 hours a day, for long periods of time and lack of soundproof buildings.

Mc Couley (39) has claimed that crew tiredness is a common denominator of human errors responsible for accidents during the flight and that 60% of accidents happen during approach and landing procedures. One study (53) examined the relationship between alertness and successful landing. The results have shown that reduced alertness represents an important factor in several accidents during approach and landing. A number of studies (10) have shown that aircrews taking part in mid and long range flights have anomalies of the sleep-wakefulness cycle, which are caused by disruption of circadian rhythm. Another factor that disrupts circadian rhythms is the conduct of nocturnal operations, a current problem as the majority of NATO attacks on Yugoslavia take place in the night-time. Dealing with crew tiredness is therefore vital in accident prevention. To achieve this, it is necessary to overcome the sleep difficulties caused by operational stress, hostile environment and disruption of circadian rhythms.

Hypnotic drugs can help to deal with these difficult situations. The perfect hypnotic should act rapidly, have a short duration of action, not lead to dependence and have no residual effect so that any impairment is not carried into the next day. It is also important to preserve the architectural structure of normal sleep. Drugs should not reduce sleep stages III and IV, which are essential for rest. They should also not interfere with R.E.M. sleep during which dreams occur, as dreams play an important role in psychological well being and are vital for effective rest.

The non-benzodiazepine 3rd generation hypnotics (zolpidem, zopiclone) are close to the above characteristics of the perfect hypnotics as existing studies prove that they preserve good sleep quality and no suppression of higher cognitive functions 6 hours after their administration.

Before reviewing the relevant literature in more detail, let us see first what exactly sleep and insomnia are. It should be noted that it is not uncommon for individuals to complain about lack of sleep and impaired daytime performance even though there is no objective evidence of insomnia from sleep recordings. This is known as pseudoinsomnia, or sleep state misperception (56).

2. Sleep and Insomnia

Even though we spend more time sleeping than in any other activity, it is difficult to define exactly what sleep is and what precise purpose it serves. We experience sleep as a loss of consciousness of our environment and we feel intuitively that it must fulfill some restorative function. Many circadian rhythms are linked tightly to our sleep/wake cycle. An adequate period of sleep at night is necessary in order to feel refreshed and effective the next day. It has been demonstrated that, while we sleep, bones grow (61), cells divide (12), wounds heal (51) and that hormones, such as growth hormones, are released (67). Sleep seems to facilitate the reinforcement of long-term memory traces (18), but is associated with a general slowing of brain activity and metabolism (38).

Arousal and wakefulness seem to be related to a high level of electrical activity within the cerebral cortex (27). When this electrical activity wanes, drowsiness and sleep set in. It seems that cortical activity is maintained by stimulatory input from a structure in the brainstem called the Reticular Activating System (RAS) (27, 63). Activation of the cortex by the RAS elicits awakening and vigilance. The RAS consists of a diffuse closed...
network of excitatory and inhibitory interneurons lying along the rostrocaudal axis of the brain system. The excitatory interneurons of the RAS are believed to use glutamic acid as their neurotransmitter, and the inhibitory ones to use γ-aminobutyric acid (GABA). GABA is the principal inhibitory neurotransmitter in the central nervous system (62) and many hypnotic, sedative, anxiolytic, anti-epileptic, anti-spastic and anaesthetic drugs owe their activity to their effects on GABA-mediated neurotransmission (16).

Sleep is not a uniform process but consists of different phases, which are called sleep stages. Five are traditionally identified: I & II (light sleep), III & IV (deep sleep) and REM. During the night, several recurring sleep cycles occur, going from awake down through the first four sleep stages and back again. The deep stages III and IV, characterized by slow waves on the electroencephalogram (EEG), are considered to be the most restorative. The fifth sleep stage, REM sleep, is characterized by paradoxical EEG activity, vivid dreams, inhibited muscle tone, rapid eye movements, bursts of sympathetic activity and penile erections in males. Although most people sleep for around seven hours each night, it is important to realize that some people function perfectly well on four hours sleep a night, whilst others require ten hours of sleep. This means that there is no normal or correct duration of sleep and thus that insomnia cannot be defined simply by a certain sleep duration. Instead, we can identify insomnia when an individual complains of not sleeping sufficiently long or well to function properly in the day.

Insomnia should be considered as a symptom of an underlying pathology, or of a change in life-style, rather than a disease in itself. Treatment should thus be directed primarily at resolving the underlying cause. Insomnia may be transient or chronic and may be characterized by difficulties in initiating or maintaining sleep. Impaired daytime functioning is the primary consequence of insomnia.

Different types of insomnia can be identified, dominated by difficulty in falling asleep, by early awakening or by multiple nocturnal awakenings.

The causes of insomnia are numerous and can be physical, such as pain or respiratory disease, psychological, such as stress or bereavement, psychiatric, such as depression or anxiety, or pharmacological, such as alcoholism or benzodiazepine withdrawal (13, 24). Often, transient insomnia can be triggered by, for example, stress or illness, which then evolves into chronic insomnia, which persists even when the original trigger is removed.

Insomnia can be related to severe consequences including elevated mortality (31, 76), increased probability of having traffic accidents (1) and comororbidity with psychiatric (20) and cardiovascular (22) disease. Several studies have demonstrated impaired quality of life in insomniacs (29) and there is generally a deterioration in social and professional relationships (26, 64, 11).

A major determinant of sleep quality is sleep hygiene (15). Poor sleep hygiene can trigger transient insomnia and favour the evolution of transient insomnia into chronic insomnia. The first-line treatment of insomnia should aim to improve sleep hygiene. This can be attempted by imposing regular bedtime and rising time, optimizing the sleeping environment and evening food intake, avoiding alcohol, caffeine and sugar also in the evening, as well as naps in the daytime and by taking physical exercise during the day.

3. Treatment of insomnia with hypnotics.

3.1 History

Hypnotic medication has been available since the introduction of chloral hydrate in the early nineteenth century. This was followed by the discovery and introduction of the barbiturates (1903), chloromethiazole (1958), the benzodiazepines (1961) and finally by zopiclone and zolpidem (1987/1988). All these drugs act by amplifying GABA-mediated neurotransmission in the brain.

The modern hypnotics in use today are the benzodiazepines such as (flunitrazepam, temazepam and triazolam) the cyclopyrolone zopiclone and the imidazopyridine zolpidem. These drugs are all extremely well tolerated compared to the previous generations of drugs, which had significant toxicity. In the vast majority of cases, they are the only hypnotic drugs that should be prescribed for the treatment of insomnia today. Benzodiazepines, zopiclone and zolpidem all interact with the same receptor in the brain, the α subunit of the GABA_A receptor. In binding to this protein, they facilitate receptor activation by GABA (16).

3.2 The GABA_A receptor: site of action of hypnotic drugs.

GABA (γ-aminobutyric acid) is the major inhibitory neurotransmitter in the mammalian central nervous system (62). It inhibits neurons by interacting with an oligomeric protein in the cell membrane, the GABA_A receptor. This protein is composed of five transmembrane peptide subunits arranged in a rosette. When the GABA_A receptor is activated by GABA, the peptide subunits move apart to form an ion-permeable pore across the membrane. This pore allows the passage of chloride ions across the membrane in response to the electrical gradient, which provides a short-circuit preventing the neuron from raising an action potential (62). The GABA_A receptor is thought to be composed of two α subunits, two β subunits and one γ subunit (34).

These differ in their amino acid composition and in their function. The β subunits are responsible for binding GABA, and thus initiating receptor activation (9). The α subunits bind benzodiazepines, zopiclone and zolpidem (34). Binding of hypnotic drugs to the α subunit increases the probability of channel opening in response.
to GABA, and thus facilitates GABA-mediated inhibition (65).

There are several different isoforms of each of α, β and γ subunits whose presence in the receptor determines its pharmacological and biophysical properties (77). There are six different isoforms of the α subunit, α1 to α6. Receptors containing any of these isoforms recognize benzodiazepines and zopiclone, but only those containing α1 have high affinity for zolpidem (49). GABA_A receptors containing α1 subunits have been referred to as BZ, or δ subtypes, and those containing other α subunit isoforms BZ2 or δ2. The predominant form of native GABA_A receptor in the mammalian brain has α subunit composition of α1 β2 γ2 (40). The functional significance of GABA_A receptor heterogeneity and subtype selectivity remains unknown (17).

3.3 Absorption and elimination of hypnotic drugs.

The pharmacokinetic properties of hypnotic drugs are an extremely important determinant of therapeutic efficacy (23). A good hypnotic needs to be absorbed rapidly, to penetrate the brain and trigger sleep within thirty minutes of oral absorption of the drug. Plasma levels should remain sufficiently high to have pharmacological activity throughout the night and thus prevent nocturnal awakenings. Drug levels should have fallen below pharmacologically active levels when it is time to wake up, so that there are no residual sedative effects during the day. Drugs with very short half-life do not maintain their activity throughout the night (28), and may not prevent early awakenings. It has also been suggested (mainly from experience with triazolam) that such drugs may be more likely to produce dependence (28) and behavioral side effects (58, 42) during the day than longer-acting hypnotic drugs. Hypnotics with long half-life, on the other hand, will continue to be active the next day, producing sedation and impairment of cognitive performance (25). Difficulty waking up, and a muzzy feeling on waking (benzodiazepine hangover) may be experienced.

3.4 New generation hypnotic drugs:

A. Zolpidem

Zolpidem is an hypnotic belonging to a new chemical class, the imidazopyridines, which are structurally unrelated to benzodiazepines. Zolpidem is a selective ligand for the central omega-1 receptor subtype with a high intrinsic activity and a potent hypnotic effect. It has a low affinity for ω1 (including δ1 or δ3 subunits) and no affinity for the peripheral BZD subtype. The ω1 and ω3 subtypes have specific central nervous system distribution; the ω1 subtype being predominant in the cerebellum whilst both subtypes are present in the cerebral cortex and only the ω2 subtype is present in the spinal cord. This specific affinity of zolpidem for the W1 subtype receptor results in marked hypnotic properties with only minor anxiolytic, myorelaxant and anticonvulsant properties. Finally, zolpidem has a short elimination half life (2.5 hours) and no active metabolites, allowing a good hypnotic activity without residual effects in the morning (48).

Clinical material, involving both healthy volunteers and insomniac populations, has clearly demonstrated the hypnotic efficacy of zolpidem in daily doses ranging from 5mg to 20mg. Moreover, the sleep architecture generally remains unaffected at these dosages, apart from an enhancing effect upon slow-wave sleep (36, 41).

In two studies (69, 5) polysomnographic recordings showed that zolpidem and flunitrazepam (BZD) significantly shortened sleep onset latency. Zolpidem respected the overall sleep architecture, without disturbance of N-REM III and IV stages and REM sleep. Flunitrazepam significantly decreased REM sleep, slow wave sleep and increased stage II.

In a study with 1772 hospitalized insomniac patients, only 2.2% of them reported adverse events and less than 1% of them withdrew due to intolerance. 1.5% had their nightly dose reduced to 5mg/day due to adverse drug reactions. The most frequently reported were drowsiness, hangover, dizziness and nausea and nightmares and hallucinations (33). In healthy volunteers, zolpidem 10 or 20mg/day did not significantly alter respiratory parameters (35).

In a recent article (54) the author reviewed the literature to determine whether the behavioral pharmacologic profile of zolpidem also differs from that of benzodiazepines. The specific topics that are reviewed include abuse potential, tolerance producing effects and physiological dependence-producing effects. The most parsimonious conclusion is that despite its unique neuropharmacological profile, the behavioral effects of zolpidem are generally similar to those of benzodiazepines.

Another study (71) examined behavioral effects and development of physical dependence after once-daily increasing doses of zolpidem in three baboons. The conclusion was that zolpidem produced physical dependence and the severity of the withdrawal syndrome can be characterized as intermediate.

Other studies have shown that zolpidem, at a dose of 10mg, does not lead to dependence and that the side effects are rare and comparable to placebo (57, 74). It has also been shown that its pharmacokinetics are not altered by food, alcohol, caffeine and timing of administration (6).

Disturbances in psychomotor and cognitive functioning and memory have long been recognized to be associated with the use of hypnotic drugs, especially those that are eliminated slowly. Several classes of hypnotic drugs can affect next-day functioning and memory. Barbiturates, benzodiazepines and various over-the-counter drugs have all been shown to possess such properties, to varying degrees.

Next-day drowsiness or incoordination can be dangerous when performing tasks such as car driving or the operation of machinery and psychomotor impairment...
may contribute to accidents. It is therefore important that hypnotic drugs should, as much as possible, be free from such undesirable "hangover" effects (69).

Morselli (43) reviewed double-blind studies involving observations made on the day after a single dose of 20mg zolpidem. Two studies included flunitrazepam as a positive control and two others triazolam. Despite using a higher dose than that currently recommended (10mg), measurements of saccadic eye movement peak velocity and repeated testing of daytime sleep latency showed that zolpidem did not affect alertness on the day after administration, although such an effect was clearly detected with the reference benzodiazepine. Next-day psychomotor function testing did not show any impairment of simple tasks. Creation time tests, digit symbol substitution tests etc. after either zolpidem or triazolam, although a performance decrement was seen after flunitrazepam. In a more complex simulated driving test, zolpidem did not differ significantly from placebo, while triazolam measurably affected performance.

In another study (18) flunitrazepam significantly impaired attention and memory compared with zolpidem and placebo, while zolpidem did not differ from placebo the day after administration.

The more recent studies (14, 68) also reach the conclusion that, at the clinically recommended dose (10mg), zolpidem treatment appears to give rise to no, or minimal next-day effects on psychomotor and cognitive functions.

Many studies showed that zolpidem is capable of impairing both memory and psychomotor functions near the time of peak plasma concentration. These effects, behavioral and subject-rated, are similar to those of benzodiazepine hypnotics and zopiclone (55, 2).

Another study (75) concluded that there was no significant interaction between zolpidem and alcohol, which did not potentiate the performance-impairing effects of zolpidem in the time of peak plasma concentration.

The literature on this is, however, inadequate. Further research is needed to determine whether consumption of alcohol is safe in combination with zolpidem (3rd generation hypnotics).

The sedative effects of 10mg of zolpidem are not antagonized by 150-300mg of caffeine in pharmacodynamic or pharmacokinetic terms (37).

Another study (72) showed that flumazenil (1.0mg) a benzodiazepine receptor antagonist, can reverse memory impairment caused by agonists of the benzodiazepine receptor, like triazolam and zolpidem.

B. Zopiclone

Zopiclone is an hypnotic with a short half-life (5 hours) which was demonstrated efficacy in sleep quality as assessed by questionnaires (46).

A number of comparative studies with benzodiazepine hypnotics, including flunitrazepam, flurazepam, nitrazepam, triazolam and temazepam, in insomniac patients have not identified significant differences in perceived sleep quality between zopiclone and the benzodiazepines (44, 70). In EEG studies, differences between zopiclone and certain benzodiazepine hypnotics have been observed. The most important of these is that zopiclone does not appear to decrease the time spent in slow-wave sleep and in REM sleep (44, 70). In some (but by no means all) of these studies, zopiclone actually increases the time spent in these sleep phases.

Morning-after effects of zopiclone are mild compared to most BZD hypnotics (70). Subjective assessment of alertness the next morning in insomniacs generally has shown zopiclone (7.5mg) to be superior to nitrazepam (5 or 10mg), flurazepam (30mg) and flunitrazepam (1 or 2mg). Impairment of psychomotor performance after zopiclone, however, has been demonstrated objectively in some, but not all, studies in healthy volunteers (32, 7).

In a comparative study (45) with zopiclone, flunitrazepam and nitrazepam, all three drugs induced some impairment of memory, but effects were more pronounced with the two benzodiazepines and especially flunitrazepam.

The adverse events seen with zopiclone are bitter taste, dry mouth, drowsiness, dizziness, tiredness, lack of coordination, depression and headache.

We conclude that zolpidem, because of its very short half-life, has better results in measures of next-day psychomotor performance and memory than zopiclone. A drawback of the short half-life is that the hypnotic effect may wane during the night, leading to a deterioration of sleep quality towards the end of the night. A similar phenomenon has been reported for short-acting benzodiazepine hypnotics (28). In contrast, zopiclone provide significantly fewer spontaneous awakenings (52).

Caffeine moderately antagonizes the effects of triazolam and zopiclone on the psychomotor performance of healthy subjects (45). But zopiclone counteracted the effects of caffeine more easily than zopiclone counteracted the decremental effects of zopiclone.

In another study (30) the authors studied the co-administration of alcohol with zopiclone and triazolam (BZD). Alcohol enhanced and prolonged the effects of both hypnotics without modifying their plasma concentrations. Drug-alcohol interactions were mainly additive though more obvious with TRZ. The hypnotics were free from residual psychomotor and cognitive effects at 8h even after the co-administration of alcohol.

In another study (4) the authors interpreted their findings as suggesting that the users of anxiolytic benzodiazepines and zopiclone were at increased risk of road-traffic accidents.

Finally there is a report (3) of physical dependence on zopiclone in individuals with depended personalities.
In this table we can see the main properties of hypnotic drugs:

<table>
<thead>
<tr>
<th></th>
<th>ZOLPIDEM</th>
<th>ZOPICLONE</th>
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<tbody>
<tr>
<td></td>
<td>Short</td>
<td>Long</td>
</tr>
<tr>
<td>1. Efficacy to induce sleep</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2. Sleep quality</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3. Anxiolytic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Elimination longlife</td>
<td>2.5 h</td>
<td>5 h</td>
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<tr>
<td>5. Adverse events</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>6. Spontaneous awakenings</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>7. Residual effects in the morning</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>8. Disturbances of sleep architecture</td>
<td>-</td>
<td>+/-</td>
</tr>
<tr>
<td>9. Next day psychomotor performance</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>10. Caffeine antagonizes</td>
<td>+/-</td>
<td>+</td>
</tr>
<tr>
<td>11. Alcohol interaction</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>12. Dependence/Tolerance effects</td>
<td>+/-</td>
<td>++</td>
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<tr>
<td>Abuse potential</td>
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**C. Antihistamines and anxiolytics.**

The antihistamines do not increase total sleep time, but offer a shorter sleep latency period. Side effects of residual daytime sedation are well documented. The anxiolytics have a long half-life and residual daytime effects.

**D. Melatonin.**

Several studies suggest that melatonin may produce behavioral sedation or facilitate objectively measured sleep, others have found no appreciable effect. There are theoretical reasons to be concerned about its use: melatonin is vasoconstrictive and has unwanted disruptive effects on reproductive endocrine function.

**3.5 Hypnotic drug use in military operations.**

A French study (60) examined the action of 10mg of zolpidem and 1mg of flunitrazepam compared to placebo in 12 pilots and 12 ground personnel. The subjects underwent psychomotor testing in a flight simulator and spectroscopic EEG analysis.

Sleep quality was similar after administration of zolpidem and flunitrazepam. The ground personnel who received benzodiazepines showed difficulty in walking, somnolence and lack of energy compared to those receiving zolpidem or placebo. On the contrary, no differences were found, either in self-assessments or in psychomotor testing, in pilots. This can be explained by the fact that pilots are highly motivated and therefore able to overcome certain potentially adverse factors.

The EEG analysis showed no effect of zolpidem, while those receiving benzodiazepines had a greater prevalence of slow waves, reduction of the a rhythm, and development of fast rhythms.

Other studies (66) have shown that zopiclone, zolpidem and temazepam do not influence athletic performance. A U.S. army Aeromedical Research Laboratory study (8) compared the effects of zolpidem-induced prophylactic naps and forced rest periods in prolonged work schedules. The conclusion was that zolpidem-induced naps were more effective because the subjects obtained more sleep after zolpidem administration. Post-nap grogginess persisted for about 2 hours after either the zolpidem or placebo nap, a fact which, despite the overall benefits from prophylactic naps, could compromise performance under operational conditions if insufficient time for awakening was planned.

Another study (50) concluded that zolpidem is a hypnotic which appears to cause less global impairment than benzodiazepine during peak effect, and is free of persistent performance decrement or hangover effect.

A study of the influence of daytime administration of zolpidem and triazolam on performance showed that there is no advantage of zolpidem over triazolam (63) in performance-imparing effects.

Finally, it was shown that zolpidem improved sleep quality at high altitudes (4,000 meters) without affecting respiration (5).

**CONCLUSIONS**

It is evident from the above that 3rd generation hypnotics (zolpidem and zopiclone) do have some advantages over the older benzodiazepines, primarily because they do not significantly influence higher cognitive functions the day after the administration. Specifically, zolpidem is free of any such influence 6 hours after administration, while the same may be true of zopiclone a few hours later due to its longer half-life.

These new hypnotics also facilitate the easy achievement of good quality sleep, as they do not disrupt its architecture and allow both deep stages III and IV sleep, which is necessary for adequate rest, and REM sleep, which is necessary for dreaming, a process fundamental for the psychological well being.

The side effects are very few but the possibilities of some form of dependence and of synergistic effect with alcohol have not been adequately studied yet. We do not have many studies concerning these issues but the few which exist, give encouraging results.

Most studies, about the use of drugs in military operations, are performed in simulated conditions, which do not fully reproduce the stress of an actual operation (the presence of enemy, loss of companions, unfamiliar and/or hostile environment). Moreover, they do not take into account the pilot's psychological profile and any human weaknesses. Despite the progress of psychopharmacology, chemically induced sleep cannot match the quality of normal sleep and a pilot who needs a drug to get an adequate rest may induce the psychological perception of a dependent person.
Observations from military Air Forces have shown that pilots may occasionally try to confront their stress with alcohol. What could then be the effect of co-administration of a drug and how can the complete absence of a synergistic effect be guaranteed? Could the flight surgeon be seen as co-responsible of an accident? Another possibility is that the pilot may, depending on the time and composition of the last meal be at risk of relative hypoglycemia or he may not be at the peak of his health during the operation. What would be the effect of drug administration then?

Also, there should be an urgent operational need, when the drug is near its peak concentration, when the pilot will clearly be unsuitable.

Frequent nocturnal operations disrupt circadian rhythms. In order to deal with this an adequate number of pilots must be required to allow sufficient time for rest and to alternate participation in operations.

Finally, let us not forget that a real combat situation brings to the surface all the aspects of somebody’s personality. We are not dealing with a car engine, which regularly needs lubrication for optimum functionality, but with human beings that (may) have ethical dilemmas regarding their mission. Consciences are not determined by propaganda and may revolt particularly when risks of civilian casualties are high. Let us not forget that one of the three Enola-Gay pilots became insane after the bombing of Hiroshima.

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