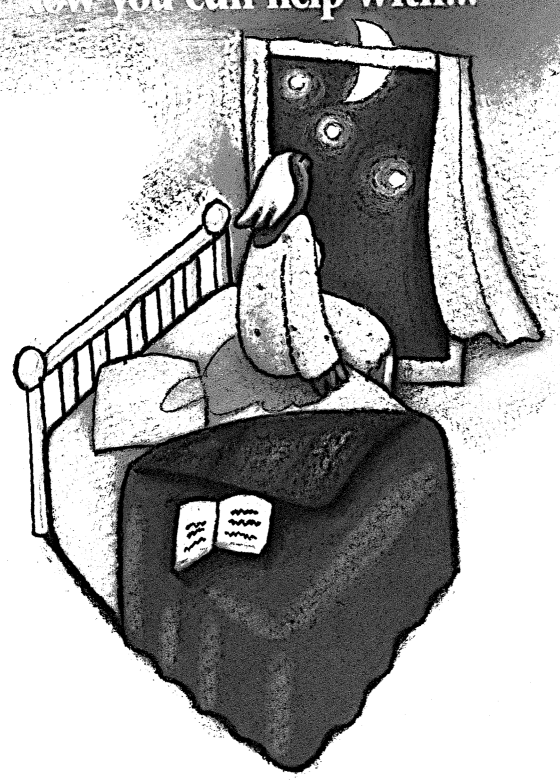
One in three adults say they occasionally have trouble sleeping.

How you can help with...



More sleep, better sleep, through the night



From a unique class of non-benzodiazepine sleep agents

More sleep

Total sleep time significantly increased compared with placebo. Patients fall asleep quickly; generally within 20 to 30 minutes.²⁴

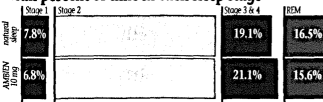
Better sleep

Awakenings were reduced, compared to placebo.

Through the night

No evidence of increased wakefulness during the last third of the night. Normal sleep stages are generally preserved² (clinical significance unknown).

Mean percent of time in each sleep stage²



In this multicenter, double-blind, randomized, controlled study of 631 healthy volunteers, no significant differences were found between the durations of sleep stages?

Short half-life

Mean 2.5-hour half-life, with no active metabolites.

With no objective evidence of tolerance or rebound insomnia

In studies of up to 35 consecutive nights at recommended doses.^{2,3}

Favorable safety and tolerability profile Adverse events with dosages of ≤ 10 mg that were statistically significant vs placebo

Short-term: ≤10 nights		Long-term: 28 to 35 nights	
drowsiness dizziness	2% 1%	dizziness drugged	5%
diarrhea	1%	feelings	3%

Recommended dosage

For adults:	ij,	one 10-mg tablet	Patients should take AMBIEN right before
For elderly/ debilitated patients:	ولاق	one 5-mg tablet	going to bed and when ready for sleep.

AMBIEN is indicated for the short-term treatment of insomnia. Prescriptions should not exceed a 1-month supply. Hypnotics should generally be limited to 7 to 10 days of use. Reevaluation of the patient is recommended if they are taken for more than 2 to 3 weeks.

In patients with hepatic dysfunction, treatment should be initiated with a 5-mg dose and patients closely monitored.



MORE SLEEP, BETTER SLEEP, THROUGH THE NIGHT

vs placebo

References: 1. National Commission on Sleep Disorders Research. Wake Up America. A National Sleep Alert. National Institute on Aging, National Institutes of Health, U.S. Department of Health and Human Services, Vol. I: January 1933. 2. Data on file, Searle. 3. Vogel G, Scharf M, Walsh J, et al. Effects of chronically administered zolpidem on the sleep of healthy insomniacs. Sleep Research. 1989;18:80. Abstract. 4. Walsh JK, Schweitzer PK, Sugerman JL, et al. Transient insomnia associated with a 3-hour phase advance of sleep time and treatment with zolpidem. J Clin Psychopharmacol, 1990:10:184-189.

BRIEF SUMMARY

INDICATIONS AND USAGE

Ambien (zolpidem tartrate) is indicated for the short-term treatment of insormia. Hypnotics should generally be limited to 7 to 10 days of use, and reevaluation of the patient is recommended if they are to be taken for more than 2 to 3 weeks.

Ambien should not be prescribed in quantities exceeding a 1-month supply (see Warnings).

CONTRAINDICATIONS

WARNINGS

WARNINGS

Since sleep disturbances may be the presenting manifestation of a physical and/or psychiatric disorder, symptomatic treatment of insomina should be initiated only after a careful evaluation of the patient. The failure of insomnia to renti after 7 to 10 days of treatment may indicate the presence of a primary psychiatric and/or medical iliness which should be evaluated. Worsening of insomnia or the emergence of new thinking or behavior shnormalities may be the consequence of an unrecognized psychiatric or physical disorder. Such findings have emerged during the course of treatment with seditive/hypnotic drugs, including Ambien. Because some of the important adverse effects of Ambien appear to be dose related (see Precautions and Dosage and Administration), it is important to use the smallest possible effective dose, especially in the elderive been A variety of abnormal entring. A variety of abnormal entring in the second of the second of the consequence of the control of t

depressed patients, who earn is association with the use of sedetive/ hyphotics. It can rarely be determined with certainty whether a perticular instance of the abnormal behaviors listed above are drug induced appointment of the programment of the programment

PRECAUTIONS

General PRECAUTIONS

Use in the elderly and/or debilitated patients: impaired motor and/or cognitive performance after repeated exposure or unusual sensitivity to sedative/hypnotic drugs is a concern in the treatment of elderly and/or debirated patients. Therefore, the recommended Ambien dosage is 5 mg in such patients (see Posage and Administration) to decrease the possibility of side effects. These patients should be closely monitored.

Use in patients with concomitant illness: Clinical experience with Ambien in patients with concomitant systemic illness is limited. Caution is advisable in using Ambien in patients with diseases or conditions that could affect metabolism or hemodynamic responses. Although preliminary studies did not reveal respiratory depressant effects at hypnotic doses of Ambien in normals, precautions should be observed if Ambien is prescribed to patients with compromised respiratory function, since sedative/hypnotics have the capacity to depress respiratory divine. Post-marketing reports of respiratory insufficiency, most of which involved patients with pre-assing respiratory vinction or efferations in pharmacolimatic patients surface and an activities of the patients with repatients with pre-assing respiratory wind and activities of the patients and activities and patients surface and patients and patients surface and patients and patients and patients and patients and patients and patients with hepatic compromise, and they should be closely monitored.

monitored. Use in depression: As with other sedetive/hypnotic drugs, Ambien should be administered with caution to patients exhibiting signs or symptoms of depression. Suicidal tendencies may be present in such patients and protective measures may be required. Intentional overdosage is more common in this group of patients; therefore, the least amount of drug that is feasible should be prescribed for the patient at any one time.

Information for patients: Patient information is printed in the complete prescribing information and is available in pads for distribution to patients.

ratory tests: There are no specific laboratory tests recon

plete prescribing information and is available in pade for distribution to patients. Laboratory tests: There are no specific laboratory tests recommended. Drug interactions CNS-active drugs: Ambien was evaluated in healthy volunteers in single-dose interaction studies for several CNS drugs. A study involving haloperidol and zolpidem revealed no effect of haloperidol on the pharmacokinertics or pharmacodynamics of zolpidem. Impramine in combination with zolpidem produced no pharmacokinertic interaction other than a 20% decrease in peak levels of impramine, but there was an additive effect of decreased alertness. Similarly, chloripromazine in combination with zolpidem produced no pharmacokineric interaction, but there was an additive effect of decreased alertness and psychomotor performance. The lack of a drug interaction following angle-dose administration does not predict a lack following chronic administration.

An additive effect on psychomotor performance between alcohol and zolpidem was demonstrated.

Since the systematic evaluations of Ambien in combination with other CNS-active drugs have been limited, careful consideration should be given to the pharmacokyor of any CNS-active drug to be used with zolpidem. Any drug with CNS-depressant effects could potentially enhance the CNS-depressant effects could potentially enhances the CNS-depressant effects of active drug on the pharmacokinetics or pharmacokyntemics. Zolpidem. Zolpidem. A study interactive and administration in a zolpidem pharmacokinetics or pharmacokyntemics of a pharmacokyntemic section to a section and active and act

kg/day dose, incidence rates of lipoms and liposarcoms for zolpidem were comparable to those seen in historical controls and the tumor findings are thought to be a spontaneous occurrence.

Mutagenesis: Zolpidem did not have mutagenic activity in several tests including the Ames test, genotoxicity in mouse lymphoma cells in vitro, chromosomal aberrations in cultured human lymphocytes, unscheduled DNA synthesis in rat hepatocytes in vitro, and the

unscheduled DNA synthesis in rat nepatocytes in vitro, and the micronucleus test in mice.

Impairment of fertility: In a rat reproduction study, the high dose (100 mg base/kg) of zolpidem resulted in irregular estrus cycles and prolonged precottal intervals, but there was no effect on male or female fertility after daily oral doses of 4 to 100 mg base/kg or 5 to 130 times the recommended human dose in mg/m². No effects on any other fertility parameters were noted.

to 130 times the recommended human dose in mg/m². No effects on any other fertility parameters were noted.
Pregnancy Category B. Studies to assess the effects of zolpidem on human reproduction and development have not been conducted.
Teratology studies were conducted in rets and rabbits.
In rats, adverse maternal and fetal effects occurred at 20 and 100 mg base/kg and included dose-related maternal lethargy and ataxis and a dose-related trend to incomplete ossification of fetal skull bones.
In rabbits, dose-related maternal sedation and decreased weight gain occurred at all doses tested. At the high dose, 16 mg base/kg, there was an increase in postimplantation total loss and underossification of sternebrae in viable fetuses.
This drug should be used during pregnancy only if clearly needed.
Montrearbogenic effects: Studies to assess the effects on children whose mothers took zolpidem during pregnancy have not been conducted. However, children born of mothers taking sedative/hypnotic drugs during pregnancy.

Labor and delivery: Ambien has no established use in labor and

Versing mothers: Studies in lactating mothers indicate that between 0.004 and 0.013% of the total administered does is excerted into milk, but the effect of zoipidem on the inflant is unknown.

The use of Ambien in nursing mothers is not recommended. Safety and effectiveness in children below the age of 18 have not

ADVERSE REACTIONS

Associated with discontinuation of treatment: Approximately 4% of 1,701 patients who received zolpidem at all doses (1.25 to 90 mg) in U.S. premarketing clinical trials discontinued treatment because of an adverse clinical event. Events most commonly associated with discontinuation from U.S. trials were daytime drowsiness (0.5%), dizziness (0.4%), headache (0.5%), nauses (0.6%), and vomiting (0.5%).

Approximately 8% of 1.320 naisas (1.320 n

(0.5%). Approximately 6% of 1,320 patients who received zolpidem at all doses (5 to 50 mg) in similar foreign trials discontinued treatment because of an adverse event. Events most commonly associated with discontinuation from these trials were daytime drowsiness (1.6%), armesus (0.6%), dizziness (0.6%), headache (0.6%), and nauses (0.6%).

nauses (0.6%). Incidence in controlled clinical trials. Most commonly observed adverse events in controlled trials: Most commonly observed adverse events in controlled trials: Dung short-term treatment (up to 10 nights) with Ambien at doses up to 10 mg, the most commonly observed adverse events associated with the use of zolipdem and seen at statistically significant differences from placebo-treated patients were drowsiness (reported by 2% of zolipdem patients), dizzness (1%), and diarrhae (1%). During longer-term treatment (28 to 35 nights) with zolipidem at doses up to 10 mg, the most commonly observed adverse events associated with the use of zolipidem and seen at statistically significant differences from placebo-treated patients were dizzness (5%) and drugged feelings (3%).

Incidence of Treatment-Emergent Adverse Experiences in Short-term Placebo-Controlled Clinical Trials (Percentage of patients reporting)

Body System/ Adverse Event*	Zolpidem (≤ 10 mg) (N=685)	Placebo (N=473)
Central and Peripheral Nervous System Headache	7	6
Drowsiness	2	-
Dizziness	1	-
Gastrointestinal System Nausea	•	3
Diarrhea	î	-
Musculoskeletal System		
Myalgia	1	2

*Events reported by at least 1% of Ambien patients are included.

lence of Treatment-Emergent Adverse Experiences in Long-term Placebo-Controlled Clinical Trials (Percentage of patients reporting)

Body System/ Adverse Event*	Zolpidem (≤ 10 mg) (N≃152)	Placebo (N≈161)
Autonomic Nervous System	3	•
Dry mouth	•	•
Body as a Whole Allergy	4	
Back pain	3	ż
Influenza-like symptoms	4 3 2 1	_
Chest pain	ī	_
Fatigue	i	2
Cardiovascular System	-	_
Palpitation	2	-
Central and Peripheral Nervous System	_	
Headache	19	22
Drowsiness	8	
Dizziness	8 5 3 2 2 1 1	5 1 1 1
Lethargy	3	1
Drugged feeling	3	-
Lightheadedness	2	1
Depression	2	1
Abnormal dreams	1	-
Amnesia		1 3
Anxiety	!	1
Nervousness	3	3
Sleep disorder	T	-
Gastrointestinal System	_	_
Nausea	6	
Dyspepsia	5	6
Diarrhea	6 3 2 2 1	6 2 2 1 1
Abdominal pain	2	2
Constipation	2	
Anorexia		
Vorniting	1	,
Immunologic System	•	
Infection	•	•
Musculoskeletal System	7	٠.
Myalgia Arthralois	4	'
ADDRAIGH	•	•

Incidence of Treatment-Emergent Adverse Experiences in Long-term Placebo-Controlled Clinical Trials (Cont'd) (Percentage of patients reporting)

Body System/ Adverse Event*	Zolpidem (≤ 10 mg) (N= 152)	Placebo (N=161)
Respiratory System Upper respiratory infection Sinusitis Pharyngitis Rhimitis	5 4 3 1	6 2 1 3
Skin and Appendages Rash Urogenital System Urinary tract infection	2	1

*Events reported by at least 1% of patients treated with Ambien

There is evidence from dose comparison triels suggesting a dose relationship for many of the adverse events associated with zolpidem use, particularly for certain CNS and gastrointestinal adverse events. Adverse events are further classified and enumerated in order of decreasing frequency using the following definitions: frequent adverse events are defined as those occurring in greater then 1/100 subjects infrequent adverse events are those occurring in 1/100 to 1/1,000 patients; rare events are those occurring in less than 1/1,000 patients.

patients, all events are tioned booking in his set and 17,000 patients. Frequent: abdominal pain, amnesis, ataxis, confusion, depression, darrhea, diplopa, dizaness, dreaming abnormal, drowsiness, drugged feeling, dry mouth, dyspepsia, euphonia, fatigue, heedache, insominis, ethargy, kipritheadedness, myelgia, nausea, upper respiratory infection, vertigo, vision abnormal, vomining. Infrequent: signation, allergy, anoraxis, anxiety, arthreigia, arthritis, asthemia, back pain, bronchitis, cerebrovascular disorder, chest pain, constipation, coughing, cystilis, addressed cignition, detached, difficulty concentrating, dysarthris, dysphagia, dyspines, edema, endough the control to the constitution, falling, lever, flatulence, gastroentisch, hallucination, floroup, hyperglycenia, hypertension, hyporadien, miraction, influenza-like symptome, milasse stresses paramygistic, paramyg

sweating increased, tachycardia, taste perversion, tinnitus, tootti disorder, traums, tremor, urinary incontinence, urinary tract infection, veginitis.

Rare: abdominel body sensation, abscess, acne, acute renel feiture, aggressive reaction, allergic reaction, allergy aggrevated, anaphylactic shock, anemia, spetite increased, arrhythmia, arteritis, erhosis, birlubinemia, breast fibroadenosis, breast neoplasm, breast pain female, bronchospasm, bulious eruption, BUN increased, circulatory failure, comeal ucceration, deluson, dementia, depersonalizations dermatitis, dysphasia, dysuria, edema periorbital, enteritis, epistaxis, face derma, feeling strange, flushing, trunculosis, gastrille, glatinis, social control, and the stranger flushing, trunculosis, gastrille, glatinis, periorbital enteritis, political, herpes zoster, hot flashes, hypercholesteremie, hyperhemolophomia, hyperploemia, hyperploemia, hyperpension aggravated, hypotension, hypotension, hyperpension, signature, and the stranger of the stranger of

ORUG ABUSE AND DEPENDENCE
Controlled substance: Schedule IV.
Abuse and dependence: Studies of abuse potential in former drug abusers found that the effects of single doses of zolpidem tertrate 40 mg were similer, but not identical, to diszapam 20 mg, while zolpidem tertrate 10 mg was difficult to distinguish from placebo-Sedetive-hyponotics have produced withdrawal signs and symptoms following abrupt discontinuation. These reported symptoms rangle from mild dysphora sand insomnia to a withdrawal syndrome that may include abdominal and muscle cramps, vomiting, sweating-termors, and convulsions. The US-clinical thial experience from zolpidem doses not reveal any clear evidence for withdrawal syndrome Nevertheless, the following adverse events included in DSM-fill-criteria for uncomplicated sedative/hypnotic withdrawal were reported at an incidence of ≤ 15 during US-clinical thial following placebo-substitution occurring within 48 hours following last zolpidem treatment: fatigue, nauses fit salming, fightlesdedness, uncontrolled crying, smessis, stomach cramps, panic attack, nervoueness, and abdominal discomfort.

Individuals with a history of addiction to, or abuse of drugs of andividuals with a history of addiction to, or shuse of fings of individuals with a history of addiction to, or shuse of fings of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individuals with a history of addiction to, or shuse of drugs of individ

under cereia surveillance when receiving any hypnotic.

OVERDOSAGE

Signs and symptoms: In European postmarketing reports of overdose with zolpidem alone, impairment of consciousness has ranged from sommolence to light come, with one case such of cardiovasculer and respiratory compromise. Individuals heve fully recovered from zolpidem tertrate overdoses up to 400 mg (40 times the maximum recommended dose). Overdose cases involving multiple CNS-depressent sgents, including solpidem, have resulted in more severe symptomatology, including fatal outcomes. Recommended treatment: General symptomatic and supportive measures should be used along with immediate gastric lavage where appropriate. Intravenous fluids should be administered as needed-flumazemi may be useful. Respiration, pulse, blood pressure, and other appropriate signs should be monitored and general supportive measures employed. Sedating drugs should be withheld following colpidem overdossge. Zolpidem is not dialyzable.

The possibility of multiple drug ingestion should be considered.

Caution: Federal law prohibits dispensing without prescription.

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