Confidentiality Is Critical for Teen Gyn. Care

BY DEEANNA FRANKLIN Senior Writer

BOSTON — A few adjustments might be needed to make your practice approachable and comfortable for adolescent patients, but the long-term payoffs can be worth it.

Why is it some people aren't so comfortable taking care of adolescents? They think they take more time in the office. They have varied issues. It's sometimes challenging, and a lot of ob.gyn. residency programs didn't address pediatric or adolescent gynecology specifically, Marc Laufer, M.D., said at an ob.gyn. meeting sponsored by Harvard Medical School.

"We're aware of that, and we're trying to address it through the American College of Obstetricians and Gynecologists and the North American Society for Pediatric and Adolescent Gynecology [NASPAG]," said Dr. Laufer of Harvard and chief of gynecology at Children's Hospital Boston.

Confidentiality is "one of the key issues" in making a practice more friendly for adolescents. One critical move is to make sure sound doesn't carry. If office walls aren't soundproof, Dr. Laufer suggested using sound machines, such as white noise machines or sound conditioners, which can help mask sounds between offices.

A lot of care for these patients can be conducted by their primary care providers. But when it comes to certain diseases, "if

References: 1. Schaff MB, Roth T, Vogel CW, Walsh JK. A multicenter, placebo-controlled study evaluating zolpidem in the treatment of chronic insomnia. *J Clin Psychiatry*. 1994;55:192-199. 2. Roth T, Roehrs T, Vogel G. Zolpidem in the treatment of the treatment of chronic insomnia. *J Clin Psychiatry*. 1994;55:192-199. 2. Roth T, Roehrs T, Vogel G. Zolpidem in the treatment of transient insomnia: a double-blind, randomized comparison with placebo. Sleep. 1995;18:246-251. 3. Elie R, Rüther E, Farr J, Emilien G, Salinas E, for the Zaleplon Clinical Study Group. Sleep latency is shortened during 4 weeks of treatment with zaleplon, a novel nonbenzodiazepine typnotic. *J Clin Psychiatry*. 1999;60:336-544. 4. AMBIEN Prescribing Information, Sandi-Synthelabo Inc. 5. Office of Applied Studies, Drug Abuse Varning Network (DAWN). Substance Abuse and Mental Health Services Administration, US Department of Health and Image Studies. 2002 Jane Jinge Studies 2002 Cline(Jinge/Jubae) 2402/CleyDubes 2402/CleyDube

Ambien[®] 🕅 (zolpidem tartrate)

BRIEF SUMMARY

INDICATIONS AND USAGE Ambien (zolpidem tartrate) is indicated for the short-term treatment of insomnia. Ambien has been shown to decrease sleep latency and increase the duration of sleep for up to 38 days in controlled dinical studies. 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 *Warning*)

CONTRAINDICATIONS

mediate evaluation. Following the rapid dose decrease or abrupt discontinuation of sedative/hyp-tics, there have been reports of signs and symptoms similar to those associ-ed with withdrawal from other CNS-depressant drugs (see *Drug Abuse and*

with withdrawal from other CNS-depressant drugs (see *Drug Abuse and* nadence). mbien, like other sedative/hypnotic drugs, has CNS-depressant effects. Due e rapid onset of action, Ambien should only be ingested immediately prior pations requiring complete mental alertness or motor coordination such as ating machinery or driving a motor vehicle after ingesting the drug, indud-otential impairment of the performance of such activities that may occur the following ingestion of Ambien, Ambien showed additive effects when com-d with achof and should not be taken with alcohol. Praients should also be oned about possible combined effects with other CNS-depressant drugs, use adjustments may be necessary when Ambien is administered with such ts because of the potentially additive effects.

PRECAUTIONS

General 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 debilitated patients. Therefore, the recommended Ambien dosage is 5 mg in such patients (see Dosage and Administration) to decrease the possibility of side effects. These patients should be closely monitored.

Interestive, use recommended Ambien dosage is 5 mg in such patients (see Dosage 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 illness: Clinical experience with Ambien in industry of the desist at those in patients with diseases or conditions that could affect metabolism or hemodynamic responses. Although studies did not reveal respiratory main the Total Arousal Index together with a reduction in lowest oxygen saturation and increase in the times of oxygen desaturation below 80% and 90% was observed in patients with mild to-moderate below. However, precautions should be observed if Ambien 10 mg when compared to placebob. However, precautions should be observed if Ambien is prescribed to the theorem of the time of which in the Total Arousal Index together with a reduction in lowest oxygen desaturation below 80% and 90% was observed if Ambien is prescribed to the times of the time of the times of oxygen desaturation below 80% and 90% was observed if Ambien is prescribed to the times of the times the time of the times the time of the times the times of the times the times of the times the times of the times the times of the times of the times the time

Laboratory tests: There are no specific laboratory tests recommended.

interactions active drugs: Ambien was evaluated in healthy volunteers in single-dose action studies for several CNS drugs. A study involving haloperidol and dem revealed no effect of haloperidol on the pharmacolinetics or pharma-mamics of zolpidem. Imiparanine in combination with zolpidem produced no macokinetic interaction other than a 20% decrease in peak levels of ramine, but there was an additive effect of decreased alertness. Similarly, promazine in combination with zolpidem produced no pharmacokinetic action, but there was an additive effect of decreased alertness and psy-notor performance. The lack of a drug interaction following single-dose inistration does not predict a lack following chronic administration. additive effect on psychomotor performance between alcohol and zolpi-was demonstrated.

An additive effect on psychomotor performance between alcohol and zolpi-lem was demonstrated. A single-dose interaction study with zolpidem 10 mg and fluoxetine 20 mg at teady-state levels in male volunteers did not demonstrate any dinically signifi-ant pharmacokinetic or pharmacodynamic interactions. When multiple doses of colpidem and fluoxetine at steady-state concentrations were evaluated in healthy emales, the only significant change was a 17% increase in the zolpidem half-life. There was no evidence of an additive effect in psychomotor performance. Following five consecutive inghithy doses of zolpidem 10 mg in the presence of entraline 50 mg (17 consecutive daily doses, at 7:00 am, in healthy female vol-inteers), zolpidem C_{sw} was significanthy higher (43%) and T_{sw} was significantly decreased (53%). Pharmacokinetics of sertraline and N-desmethylsertraline were unaffected by zolpidem. Since the systematic evaluations of Ambien in combination with other CNS-active drugs have been limited, careful consideration should be given to the barmacology of any CNS-active drug to be used with zolpidem. Any drug with NS-depressant effects could potentially enhance the CNS-depressant effects of olpidem.

Drugs that affect drug metabolism via cytochrome P450: A randomized, double-blind, crossover interaction study in ten healthy volunteers between itraconazide (200 mg once daily for 4 days) and a single dose of zolpidem (10 mg) given 5 hours after the last dose of itraconazide resulted in a 34% increase in AUC_{barce} of zolpidem. There were no significant pharmacodynamic effects of zolpidem con subjective drowsiness, posturel sway, or psychomotor performance. A randomized, placebo-controlled, crossover interaction study in eight healthy female volunteers between 5 consecutive daily doses of rifampin (600 mg) and a single dose of zolpidem to the 20 mg interact term of the AUC - 20 mg interact terms of the study - (-36%), and T₁₂(-36%) of zolpidem together with significant reductions in the pharmacodynamic effects of zolpidem.

zolpidem.
Other drugs: A study involving cimetidine/zolpidem and ranitidine/zolpidem combinations revealed no effect of either drug on the pharmacokinetics or pharmacodynamics of zolpidem. Zolpidem had no effect on digoxin kinetics and dig not affect prothrombin time when given with warfarin in normal subjects Zolpidem's sedative/hypnotic effect was reversed by furmazenii, however, no sig nificant alterations in zolpidem pharmacokinetics were found.

Torug/Laboratory test interactions: Zolpidem is not known to interfere with com-monly employed clinical laboratory tests. In addition, clinical data indicate that zolpidem does not cross-react with benzodiazepines, polates, barbiturates, cocaine, cannabinoids, or amphetamines in two standard urine drug screens.

cocaine, cannabinoids, or amphetamines in two standard urine drug screens. Carcinogenesis, mutagenesis, impairment of fortility Carcinogenesis: Zolpidem was administered to rats and mice for 2 years at dietary dosages of 4, 18, and 80 mg/kg/day. In mice, these doses are 26 to 520 times or 2 to 35 times the maximum 10-mg human dose on a mg/kg or mg/m² basis, respectively. In rats these doses are 43 to 876 times or 6 to 115 times the maximum 10-mg human dose on a mg/kg or mg/m² basis, respectively. No evi-dence of carcinogenic potential was observed in mice. Renal liposarcomas were seen in 4/100 rats (3 males, 1 female) receiving 80 mg/kg/day and a renal lipoma was observed in one male rat at the 18 mg/kg/day dose. Incidence rates of lipo-ma and liposarcoma for zolpidem were comparable to those seen in historical controls and the tumor findings are thought to be a spontaneous ocurrence.

controls and the turnor intenings are thought to be a spontaneous occurrence. Mutagenesis: Colpidem did not have mutagenic activity in several tests includ-ing the Ames test, genotoxicity in mouse lymphoma cells in vitro, chromosomal aberrations in cultured human Hymphocytes, unscheduled DNA synthesis in rat hepatocytes in vitro, and the micronucleus test in mice.

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

mg/m². No effects on any other tertainty parameters were noted. Pregnacy Pregnacy Teratogenic effects: Category B. Studies to assess the effects of zolpidem on human reproduction and development have not been conducted. Teratology studies were conducted in rats and rabbits. In rats, adverse maternal and fetal effects occurred at 20 and 100 mg base/kg and included obse-related maternal lettargy and ataxia 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, three was an increase in postimplantation fetal loss and underossification of sternebrae in viable fetuses. This drug should be used during pregnancy only if clearly needed. Maternancesic affects: Studies to assess the effects on children whose mothers

This drug should be used during pregnancy only if dearly needed. Nonteratogenic 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/hypotic drugs may be at some risk for with-drawal symptoms from the drug during the postnatal period. In addition, neona-tal flacacidity has been reported in infants born of mothers who received sedative/ hypotic drugs during pregnancy. Labor and delivery: Ambien has no established use in labor and delivery. Nursing mothers: Studies in lacating mothers indicate that between 0.004 and 0.019% of the total administered dose is excreted into milk, but the effect of zolpi-dem on the infant is unknown. The use of Ambien in nursing mothers is not recommended.

Pediatric use: Safety and effectiveness in pediatric patients below the age of 18 have not been established.

have not been established. Geriatric use: A total of 154 patients in U.S. controlled clinical trials and 897 patients in non-U.S. clinical trials who received zolpidem were ≥60 years of age. For a pool of U.S. patients receiving zolpidem at doses of ≤10 mg or placebo, there were three adverse events occurring at an incidence of at least 3% for zolpi-dem and for which the zolpidem incidence was at least twice the placebo inci-dence (ie, they could be considered drug related).

Adverse Event	Zolpidem	Placebo
Dizziness	3%	0%
Drowsiness	5%	2%
Diarrhea	3%	1%

A total of 30/1,959 (1.5%) non-U.S. patients receiving zolpidem reported falls, including 28/30 (93%) who were \geq 70 years of age. Of these 28 patients, 23 (82%) were receiving zolpidem doess >10 mg. A total of 24/1,95% (1.2%) non-U.S. patients receiving zolpidem reported confusion, including 18/24 (75%) who were \geq 70 years of age. Of these 18 patients, 14 (78%) were receiving zolpidem doess >10 mg.

ADVERSE REACTIONS

Associated with discontinuation of treatments Approximately 4% of 1,701 patients who received xalpidem at all doses (1.25 to 90 mg) in U.S. premarketing chinal trials discontinued treatment because of an adverse dinical event. Events most commonly associated with discontinuation from U.S. trials were dayline drowsiness (0.5%), dizziness (0.4%), headache (0.5%), nausee (0.6%), and vomit-

drowshiess (0.2%), ruliness (0.4%), headache (0.5%), hausea (0.5%), and vorhiting (0.5%). Approximately 4% of 1,959 patients who received zolpidem at all doses (1 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 (1.1%), dizziness/vertigo (0.8%), amnesia (0.5%), nau-sea (0.5%), headache (0.4%), and falls (0.4%). Data from a dinical study in which selective serotonin reuptake inhibitor-(SSRI) treated patients were given zolpidem revealed that four of the seven dis-continuations during double-blind treatment with zolpidem (n-95) were associ-ated with impaired concentration, continuing or aggravated depression, and manic reaction; one patient treated with placebo (n-97) was discontinued after an attempted suicide.

Iaffini teauton, on semi-liafini teauton, on semi-natempted suicide. Icidence in controlled clinical trials fost commonly observed adverse events in controlled trials: During short-term eatment (up to 10 nights) with Ambien at doses up to 10 mg, the most com-nohy observed adverse events associated with the use of coljoidem and seen at tatistically significant differences from placebo-treated patients were drowsi-ess (reported by 2% of zoljoidem patients), dizziness (1%), and diarrhea (1%). Juring longer-term treatment (28 to 35 nights) with zolpidem at doses up to 10 mg, the most commonly observed adverse events associated with the use of objectem and seen at statistically significant differences from placebo-treated valents were dizziness (5%) and drugged feelings (3%).

patients were dizziness (5%) and drugged feelings (3%). Treatment-emergent adverse experiences in placebo-controlled clinical trials: The following are treatment-emergent adverse events from U.S. placebo-con-rolled dinical trials. Data are limited to data from doses up to and including 10 mg. In short-term trials, events seen in zolpidem patients (m=686) at an incidence equal to 1% or greater compared to placebo (m=473) were: headache (7% vs 6%) for placebo, drowsiness (2% vs 0%), dizzines (1% vs 0%), nausea (2% vs 3%), diarrhea (1% vs 0%), and myalgia (1% vs 2%). In long-term clinical trials, events seen in zolpidem patients (n=152) at an incidence of 1% or greater compared to placebo (n=161) were: dry mouth (3% vs 1% for placebo), **allergy** (4% vs 1%),

back pain (3% vs 2%), influenza-like symptoms (2% vs 0%), chest pain (1% vs 0%), fatigue (1% vs 2%), palpitation (2% vs 0%), headache (19% vs 22%), drowsiness (3% vs 5%), dizziness (5% vs 1%), depression (2% vs 1%), aboromal dreams (1% vs 0%), ammesia (1% vs 0%), anxiety (1% vs 1%), arousenss (1% vs 3%), sleep disorder (1% vs 0%), nauesa (3% vs 2%), downess disorder (1% vs 0%), anxiety (12% vs 1%), arousenss (1% vs 3%), sleep disorder (1% vs 0%), inaresta (1% vs 1%), arousenss (1% vs 3%), sleep disorder (1% vs 0%), inaesto (1% vs 1%), arousenss (1% vs 3%), dishomial pain (12% vs 2%), constitution (2% vs 1%), arousens (1% vs 2%), rash (2% vs 1%), and urinary tratification (2% vs 2%).

Intection (2% vs 2%). **Dose relationship for adverse events:** There is evidence from dose comparisor trials suggesting a dose relationship for many of the adverse events associatec 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 than 1/100 subjects; infrequent adverse events are those occurring in f1/100 to 1/1/000 patients; rare events are those occurring in less than 1/1,000 patients.

tess than 1/1,000 patients. Frequent: abdominal pain, abnormal dreams, allergy, amnesia, anoxie ety, arthralgia, asthenia, ataxia, back pain, chest pain, confusion, constipation depression, diarrhea, diplopia, dizzinaes, drowsiness, drugged feeling, dr, mouth, dyspepsia, euphoria, fatigue, headache, hiccup, infection, influenza-like symptoms, insomnia, lethargy, lightheadedness, myalgia, nausea, nervousness papitation, sleep disorder, vertigo, vision abnormal, vomiting.

papitation, steep disorder, vertigo, vision abnormal, vormiting. Infrequent: abnormal hepatic function, agitation, arthritis, bronchitis, cere-brovascular disorder, coughing, cystitis, decreased cognition, detached, difficul-ty concentrating, dysarthria, dysphagia, dysphae, edema, emotional lability, eye irritation, eye gain, falling, ever, flatulence gastroenteritis, hallucination, hyper-glycemia, hypertension, hypoesthesia, illusion, increased SGPT, increased sweating, leg cramps, malaise, menstrual disorder, migraine, pallor, paresthesia, postural hypotension, puruitus, scleritis, sleeping (after daytime dosing), speech disorder, stupor, syncope, tachyacria, taste perversion, thirst, tinnitus, trauma, tremor, urinary incontinence, vaginitis.

disorder, stupor, syncöpe, tachycardia, taste perversion, thirst, tinnitus, trauma, tremor, urinary incontinence, vaginitis.
Rare: abdominal body sensation, abnormal accommodation, abnormal gait, abnormal thinking, abscess, acne, acute renal failure, aggressive reaction, allering, and the start of the star

Abuse and dependence: Studies of abuse potential in former drug abusers found that the effects of single doses of zolpidem tartrate 40 mg were similar, but not identical, to diazepam 20 mg, while zolpidem tartrate 10 mg was difficult to dis-tionuish form alreade

Autise and dependence: Studies of abuse potentiaritate 40 mg words subsets found that the effects of single doese of abpidem tartrate 40 mg words subsets found identical, to diazepan 20 mg, while zolpidem tartrate 40 mg were similar, but not identical, to diazepan 20 mg, while zolpidem tartrate 40 mg were similar, but not identical, to diazepan 20 mg, while zolpidem tartrate 40 mg were similar, but not identical, to diazepan 20 mg, while zolpidem tartrate 40 mg were similar, but not detriced to the similar target of the similar target of the similar target and insomnia to a withdrawal syndrome that may include abdominal and mus-de cramps, vomiting, sweating, tremors, and convulsions. The U.S. clinical trial experience from zolpidem does not reveal any dear evidence for withdrawal syndrome. Nevertheless, the following adverse events included in DSMIHR cri-teria for uncomplicated sedative/hypnotic withdrawal were reported at an inci-dence of 51 Mg during U.S. clinical trial 510 Minung placebo substitution occurring within 48 hours following last zolpidem treatment; fatigue, nausea, flushing, lightheaddendess, uncontrolled crying, emesis, stomach cramps, panic attack, nervousness, and abdominal discomfort. Rare post-marketing reports of abuse, dependence and withdrawal have been received. Mutividuals with a history of addiction to, or abuse of, drugs or alcohol are at increased risk of habituation and dependence; they should be under careful sur-vellance when receiving any hypnotic. **OVENOSACE Signs and symptoms:** In European postmarketing reports of overdose with zolpi-dem alone, impairment of consciousness has ranged from somonlence to light during the maximum recommended does). Overdose cases involving multiple CN3-depressant agents, including zolpidem, have resulted in more severe symptomatology, including fatel outcomes. **Recommended treatment:** General symptomatic and supportive measures

ymptomatology, including tatal outcomes. ecommended treatment: General symptomatic and supportive mea hould be used along with immediate gastric lavage where approp itravenous fluids should be administered as needed. Flumazenil may be u sepiration, pulse, blood pressure, and other appropriate signs should be ored and general supportive measures employed. Sedating drugs shou ithheld following zopidem overdosage. Zopidem is not diatyzahe. The possibility of multiple drug ingestion should be considered.

sanofi~synthelabo

Distributed by: Sanofi-Synthelabo Inc. New York, NY 10016

Revised August 2002

we treat and diagnose them when people are younger, we may improve their longterm health care," Dr. Laufer said at the meeting cosponsored by Brigham and Women's Hospital. Dr. Laufer said if polycystic ovary syndrome were diagnosed and treated during adolescence, there would be a greater chance of decreasing rates of obesity and diabetes. The same holds for endometriosis. An early diagnosis likely would result in less pelvic pain over the patient's lifetime and lead to improved long-term fertility.

Since most adolescents are "Internet savvy," Dr. Laufer encouraged physicians to direct young patients to online resources such as the Web site by the Center for Young Women's Health at Children's Hospital Boston (www.young womenshealth.org). The site offers education information in English and Spanish and an online chat room where teens can ask questions and get answers from health professionals.

The NASPAG's Web site (www.naspag.org) also offers information for teens and physicians and includes links to other adolescent care Web sites.

FDA Warns About Imported Test Kits

The Food and Drug Administration is Twarning consumers about possible false results from several unapproved home-use diagnostic test kits marketed in the United States via the Internet by Globus Media of Montreal.

The test kits are not approved for sale in the United States. There is concern that the use of these products could lead to false results that could contribute to significant adverse health consequences, but there are no confirmed instances of false results, according to the FDA.

The kits are labeled as Rapid HIV Test Kit, Rapid Syphilis Test Kit, One Step Cas-

Use of these tests could lead to false results that could contribute to adverse health consequences.

sette Style Cocaine Test, One Step Cassette Style Marijuana (THC) Test, One Step Cassette Style Amphetamine Rapid Test, Dengue Fever Test, and One

Step Midstream Style HCG Urine (Home) Pregnancy Test.

The tests, sold through Web sites such as www.htkit.com, have been distributed nationwide. The name of the kit appears on the instructions, but the envelope, instructions, and packaging may not accurately identify the manufacturer, packer, or distributor. Anyone who has used one of these products should be retested using valid test methods.

No home-use test kits intended for diagnosing HIV, syphilis, and dengue fever have been approved for sale in the United States.