# **Calcium Scoring Useful in Asymptomatic Patients**

BY KERRI WACHTER Senior Writer

oronary artery calcium scoring by CT can be a useful tool in the evaluation of asymptomatic patients with an intermediate risk of coronary heart disease but not those with low or high risk, according to expert consensus.

On Jan. 23, the American College of Cardiology Foundation and the American Heart Association jointly released an expert consensus document that updates information and opinion on coronary artery calcium (CAC) scoring by CT, particularly with regard to global cardiovascular risk assessment and evaluation of patients with chest pain (J. Am. Coll. Cardiol. 2007; 49:378-402). The last consensus document on the use of electron-beam CT for the diagnosis and prognosis of coronary artery disease (CAD) was published in 2000.

On the basis of data available since that time, the committee concluded that CAC measurement using CT scanning is a reasonable tool for evaluating asymptomatic patients with a 10-year risk of estimated CHD events between 10% and 20%.

'The test does what it does very well it detects calcium. It's a marker of atherosclerosis and ergo a marker of higher risk," said Dr. Robert O. Bonow, a member of the writing committee and chief of the division of cardiology at Northwestern Memorial Hospital in Chicago in an interview. With the intermediate group,

her than the therapeutic exposure to ramelteon and M-II, resp MRHD based on AUC).

So interpolary Labor and Delivery The potential effects of ROZEREM on the duration of labor and/or delivery, for either the mother or the fetus, have not been studied. ROZEREM has no established use in labor and delivery.

Nursing Mohers Ramelteon is secreted into the milk of lactating rats. It is not known wi this drug is excreted in human milk. No clinical studies in nursing mot have been performed. The use of ROZEREM in nursing mothers is not recommended.

Pediatric Use Safety and effectiveness of ROZEREM in pediatric patients have not been established. Further study is needed prior to determining that this product may be used safely in pre-pubescent and pubescent patients.

may be used safely in pre-publescent and publescent patients. **Certairt: Use** A total of 654 subjects in double-blind, placebo-controlled, efficacy trials received ROZEREM were at least 65 years of age, of these, 199 were 75 of age or older. No overall differences in safety or efficacy were observed between elderly and younger adult subjects. **ADVERSE REACTIONS** 

Dverview The data described in this section reflect exposure to ROZEREM in 4251 sub-ects, including 346 exposed for 6 months or longer, and 473 subjects for

percent of the 3594 individual subjects exposed to ROZEREM in clinica dies discontinued treatment owing to an adverse event, compared with of the 1370 subjects receiving placebo. The most frequent adverse even ding to discontinuation in subjects receiving ROZEREM were somnolence 3%), dizziness (0.5%), nausea (0.3%), fatigue (0.3%), headache (0.3%), insomnia (0.3%).

se Reactions Resulting in Discontinuation of Treatment ercent of the 3594 individual subjects exposed to ROZEREM in

(u x%), drzzness (0.5%), nausea (0.3%), fatigue (0.3%), headache (0.3%), and insomnia (0.3%). **ROZEREM Most Commonly Diserved Adverse Events in Phase 1-3 trials** The incidence of adverse events during the Phase 1 through 3 trials (% placebo, n=1370; % ramelton [8 mg], n=1250) were; headache NOS (7%, 7%), somolence (3%, 5%), latigue (2%, 4%), dupler sepiratory tract infection NOS (2%, 3%), diarrhea NOS (2%, 4%), upper reparatory tract depression (1%, 2%), disyonali exacettated (2%, 5%), upper reparatory tract infection NOS (2%, 3%), diarrhea NOS (2%, 2%), wupper sepiratory tract depression (1%, 2%), dysquesi (1%, 2%), arthratiga (1%, 2%), influenza (0, 1%), blood cortisol decreased (0, 1%) Because clinical trials are conducted under widely varying conditions, adver reaction rates observed in the clinical trials of a drug cannot be directly con pared to rates in clinical trials of other drugs, and may not reflect the rates observed in practice. The adverse reaction information from clinical trials does, however, provide a basis for identifying the adverse events that appea to be related to drug use and for approximating rates. **DRUG ABUSE AND DEFENDENCIE** 

Human Data: See the CLINICAL TRIALS section, Studies Pertinent to Safety Concerns for Sleep-Promoting Agents in the Complete Prescri

on in animals or in humans after chronic adm thdrawal signs. Ramelteon does not appear to

ent. JZEREM was administered in single doses up to 160 mg in an abuse liabil-trial. No safety or tolerability concerns were seen. **ccommended Treatment** meral symptomatic and supportive measures should be used, along with mediate gastric lavage where appropriate. Intravenous fluids should be ministered as needed. As in all cases of drug overdose, respiration, pulse, od pressure, and other appropriate vital signs should be monitored, and neral supportive measures employed.

general supportive measures employed. Hemodialysis does not effectively reduce exposure to ROZEREM. Therefore, the use of dialysis in the treatment of overdosage is not appropriate. **Poison Control Center** As with the management of all overdosage, the possibility of multiple drug ingestion should be considered. The physician may contact a poison control center for current information on the management of overdosage.

continuation of ramelteon i on did not produce withdr duce physical dependence.

Rx only

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a novel hypnotic lacking abuse Arch Gen Psychiatry. In press.

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CAC measurement could help cardiologists decide how aggressive to be with treatment.

However, the authors advised against the use of CAC measurement in patients with low CHD risk (below 10% 10-year risk of estimated CHD events). They noted that CAC measurements in this patient group would be similar to using the technique for population screening, which the committee also counseled against.

Likewise, the authors advised against the use of CAC measurements in asymptomatic patients with high CHD risk (greater than 20% 10-year risk of estimated CHD events or established coronary disease, or other high-risk diagnoses). Patients in this



'If you have someone at low risk and a positive calcium scan doubles your risk from 1% to 2%, it's still low risk.'

DR. BONOW

category "should be treated aggressively consistent with secondary prevention goals based upon the current National Cholesterol Education Program III guidelines and thus should not require additional testing, including CAC scoring, to establish this risk evaluation," they wrote.

"If you have someone at low risk and a positive calcium scan doubles your risk from 1% to 2%, it's still low risk," said Dr. Bonow. "If you're very high risk, it's high risk no matter what."

While the recommendations give the thumbs up to the use of CAC to evaluate patients with intermediate risk, the authors noted that there have been no headto-head comparisons of CAC with other assessment tools. Some, such as ankle/ brachial index or carotid ultrasound, may be less expensive.

There have also been no randomized trials that demonstrate that CAC measurement improves outcomes. "This created a lot of discussion in the committee," said Dr. Bonow. "The problem is that it's not clear that the trial will ever be done. Meanwhile, we have data that [CAC] might be a useful test in certain subsets of patients,' said Dr. Bonow.

The committee also noted that the strongest CAC data are for white men. Until additional data in other groups are available, the authors recommended caution in extrapolating CAC data derived from these studies in women and ethnic minorities.

The committee also reviewed the use of CAC measurement in patients with diabetes. It has been noted in several cross-sectional studies that patients with diabetes have a higher prevalence and degree of coronary calcium than nondiabetic patients.

The authors noted that there is some evidence to suggest "that coronary calcium might be useful to further stratify shortterm risk in diabetic patients." However, they cautioned that additional studies from nonreferral populations with longer follow-up are needed. 

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### mary of Prescribing Information

ROZEREM™

# INDICATIONS AND USAGE ROZEREM is indicated for the treatment of insomnia characterized by diffi

CONTRAINDICATIONS ROZEREM is contraindicated in patients with a hypersensitivity to rameltec or any components of the ROZEREM formulation.

If Unproteins of the treatment of the presenting manifestation of a physis sleep disturbances may be the presenting manifestation of a physis re psychiatric disorder, symptomatic treatment of insomnia should be ted only after a careful evaluation of the patient. The failure of insom mit after a reasonable period of treatment may indicate the presence primary psychiatric and/or medical illness that should be evaluated arening of insomnia, or the emergence of new cognitive or behaviora mailties, may be the result of an unrecognized underlying psychiatric ical disorder and requires further evaluation of the patient. As with olics, exacerbation of insomnia and emergence of cognitive and bel abnormalities were seen with ROZEREM during the clinical develop ram ROZEREM should not be used by patients with severe hepatic impai

Invocence shown once used by patients with severe hepatic impairment. ROZEREM should not be used in combination with fluvoxamine (see **PRE-CUTIONS: Drug Interactions)**. A variety of cognitive and behavior changes have been reported to occur in association with the use of hypotolcs. In primarily depressed patients, worsening of depression, including suicidal ideation, has been reported in association with the use of hypotolcs.

Patients should avoid engaging in hazardous activities that require concentra-ion (such as operating a motor vehicle or heavy machinery) after taking 302EREM.

After taking ROZEREM, patients should confine their activities to those neces-sary to prepare for bed.

PRECAUTIONS

Jeneral OZEREM has not been studied in subjects with severe sleep apnea or severe COPD and is not recommended for use in those populations. Patients should be advised to exercise caution if they consume alcohol in combination with ROZEREM.

COMDINATION WITH PLACETERN. Use in Adolescents and Children ROZEREM has been associated with an effect on reproductive hormone adults, e.g. decreased testosterone levels and increased prolactin levels not known what effect chronic or even chronic intermittent use of ROZE may have on the reproductive axis in developing humans (see Pediatric

mation for Patients the should be advised to take ROZEREM within 30 minutes prior to to bed and should confine their activities to those necessary to pre-

tor bed. Patients should be advised to avoid engaging in hazardous activities (such as operating a motor vehicle or heavy machinery) after taking ROZEREM. Patients should be advised that they should not take ROZEREM with or immediately after a high fat meal.

Patients should be advised to consult their health care provider if they experience worsening of insomnia or any new behavioral signs or symptoms of

concern. Patients should consult their health care provider if they experience one of the following: cessation of menses or galactorrhea in females, decreased libido, or problems with fertility.

Laboratory Tests No standard monitoring is required.

ents presenting with unexplained amenorrhea, galactorrhea, decrea or problems with fertility, assessment of prolactin levels and testos levels should be considered as appropriate.

Drug Interactions R0ZERAM has a highly variable inter-subject pharmacokinetic profile (approximately 100% coefficient of variation in C<sub>max</sub> and AUC). As noted above, CYP1A2 is the major isozyme involved in the metabolism of R0ZEREM; the CYP2C subfamily and CYP3A4 isozymes are also involve to a minor degree.

minor degree. cts of Other Drugs on ROZEREM Metabolism

Effects of Other Drugs on ROZEREM Metabolism Fluvoxamine (strong CYP142 Inhibitor): When fluvoxamine 100 mg twice daily was administered for 3 days prior to single-dose co-administration of ROZEREM 16 mg and fluvoxamine, the AU06<sub>961</sub> for ramelteon increased approximately 190-101d, and the Gma. Increased approximately 70-101d, compared to ROZEREM administered alone. ROZEREM should not be used in combination with fluvoxamine (See WARNINGS). Other less potent CYP142 inhibitors have not been adequately studied. ROZEREM should not be administered with caution to patients taking less strong CYP142, inhibitors. Ri/ampin (strong CYP enzyme induce):/ Administration of rifampin 600 mg once daily for 11 days resulted in a mean decrease of approximately 80% /40% to 90%) in total exposure to rameteon and metabolite M-11, looth AU0<sub>6941</sub> and G<sub>max</sub>) after a single 32 mg dose of ROZEREM. Efficacy may be reduced when ROZEREM is used in combination with strong CYP enzyme inducers such as rifampin.

Invuouers such as Intamplin. Ketoconazole (strong CVP3A4 inhibitor): The AUC<sub>0-bit</sub> and C<sub>max</sub> of ramelteon increased by approximately 84% and 36%, respectively, when a single 16 mg dose of ROZEREM was administered on the fourth day of ketoconazole 200 mg twice daily administration, compared to administration of ROZEREM alone. Similar increases were seen in M-II pharmacokinetic variables. ROZEREM should be administered with caution in subjects taking strong CVP3A4 inhibitors such as ketoconazole.

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fluconazole. praction studies of concomitant administration of ROZEREM with flu (CVP206 inhibitor), omeprazole (CVP1A2 inducer/CVP2C19 inhibit ophylline (CVP1A2 substrate), and dextromethorphan (CVP206 subs not produce clinically meaningful changes in either peak or total exp es to ramelteon or the M-II metabolite.

sures to rameteon or the M-II metabolite. Effects of ROZEREM on Metabolism of Other Drugs Concomitant administration of ROZEREM with omeprazole (CYP2C19 strate), dextromethorphan (CYP2O6 substrate), midazolam (CYP3A4 substrate), did warfarin (CYP2O5 Substrate), digoxin (p-glycoprotein strate), and warfarin (CYP2O5 [S]/CYP1A2 [R] substrate) did not pro-clinically meaningful changes in peak and total exposures to these dri Effect of Algobia on Romania

*f Alcohol on Rozerem* With single-dose, daytime co-administration of ROZEREM 32 mg oblo (0.6 g/kg), there were no clinically meaningful or statistically s

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nificant effects on peak or total exposure to ROZEREM. However, an additive effect was seen on some measures of psychomotor performance (i.e., the Digit Symbol Substitution Test, the Psychomotor Vigilance Task Test, and a Visual Analog Scale of sedation) at some post-dose time points. No additive effect was seen on the Delayed Word Recognition Test. Because alcohol by itself impairs performance, and the intended effect of ROZEREM is to pro-mote sleep, patients should be cautioned not to consume alcohol when usin higher than the therapeutic exposure to ramelteon and M-II, respective the MRHD based on AUC). The effects of ramelteon on pre- and post-natal development in the r studied by administration of ramelteon to the pregnant rat by oral gar doess of 0, 30, 100, or 300 mg/kg/day from day 6 of gestation throug turrition to postnatal (lactation) day 21, at which time offspring were v Maternai toxicity was noted at doess of 100 mg/kg/day or greater ans sisted of reduced body weight gain and increased adrenal gland weig Reduced body weight during the post-weating period was also notic offspring of the groups given 100 mg/kg/day and higher. Offspring in 300 mg/kg/day group demonstrated physical and developmental dela including delayed eruption of the lower incisors, a delayed acquisition offspring in the 300 mg/kg/day group was likely due to altered ma behavior and function observed at this dose level. Offspring of the 300 mg/kg/day group also showed evidence of diaphragmatic hernia, ging observed in the emproductive capacity of forspring and the ensuiting progregory were not different from those of vehicle-treated off The no-effect level for pre- and postnati development in this study of mg/kg/day (39-lines higher than the MRHD on a mg/m<sup>2</sup> basis). Labor and Delivery

ROZEREM. Drug/Laboratory Test Interactions ROZEREM is not known to interfere with commonly used clinical il tests. In addition, *in vitro* data indicate that ramelteon does not can positive results for henzodiazepines, opiates, barbiturates, cocaine noids, or amphetamines in two standard urine drug screening met

ogenesis, Mutagenesis, and Impairment of Fertility

rcinogenesis a two-year carcinogenicity study, B6C3F, mice were administered namelte a two-year carcinogenicity study, B6C3F, mice were administered namelte obsess of 0.30, 100, 300, or 1000 mg/kg/day by oral gavage. Male mice bibited a dose-related increase in the incidence of hepatic tumors at doss test > 200 mg/kg/day including hepatic admons, hepatic carcinoma, and atoblastoma. Female mice developed a dose-related increase in the inci-ce of hepatic adenomas at dose levels > 300 mg/kg/day and hepatic cinoma at the 1000 mg/kg/day (dose level. The no-effect level for hepatic tors in male mice was 30 mg/kg/day (103-times and 3-times and 3-times and exposure to ramelteon and the active metabolite M-II, respectively, at the strumm recommended human dress MBHD hepatic admonsed and the structure and the

tumors in male mice are young uses much, the no-effect level for hepatic tumors in male mice was 30 mg/kg/day (131 times and 3-times the thrapp tic exposure to ramelteon and the active metabolite M-II, respectively, at the maximum recommended human dose [MRHD] based on an are-under-the curve [AUC] comparison). The no-effect level for hepatic tumors in female mice was 100 mg/kg/day (327 -times and 12-times the thrapeutic exposure to orametteon and M-II, respectively, at the MRHD based on AUC). In a two-year carcinogenicity study conducted in the Sprague-Dawley rat, male and female rats were administered rametteon at doses 0 o, 15, 60, 250 or 1000 mg/kg/day dose level. Fenale tast active administered ramet and benigin Levgid cell tumors in the incidence of hepatic adenoma and benigin Levgid cell tumors in the 1000 mg/kg/day dose level. Female rats % mice admines the dose related increase in the 1000 mg/kg/day dose level. Female rats % mice admines the dose mg/kg/day (342 + 142-times and 12-times the therapeutic exposure to rametteon and M-II, respectively, at the MRHD based on AUC). The no-effect level for hepatic umors and benign Levgid cell tumors in male rats was 60 mg/kg/day = 1,429-times and 12-times the therapeutic exposure to rametteon and M-II, respectively, at the MRHD based on AUC). The no-effect level for hepatic umors in fmale rats was 15 mg/kg/day (342 - times and 15-times the therapeutic exposure to rametteon and M-II, respectively, at the MRHD based on AUC). The no-effect level for hepatic admines rats was 15 mg/kg/day (342 - times and 15-times the therapeutic exposure to rametteon and M-II), respectively, at the MRHD based on AUC). The development of hepatic tumors in rodents following characteristic exposure to rametteon and M-II, respectively, at the MRHD based on AUC).

therapeutic exposure to ramelteon and M-II, respectively, as the INTEL based on AUC). The development of hepatic tumors in rodents following chronic treatme with non-genotoxic compounds may be secondary to microsonal enzym induction is mechanism for tumor generation not thought to occur in humans. Leydig cell tumor development following treatment with non-genotoxic compounds in rodents has been linked to reductions in circula testistic and the evel with compensatory increases in luteinizing hormone release, which is a known proliferative stimulus to Leydig cells in the rat testis. Rat Leydig cells are more sensitive to the stimulatory effects of luteinizing hormone than human Leydig cells. In mechanistic studies com ducted in the rat, daily ramelteno administration at 250 and 1000 mg/kg/ for 4 weeks was associated with a reduction in plasma testosterone level in the same study, luteinizing hormone inversion at 250 and 1000 mg/kg/ for 4 ueeks was associated with a reduction in plasma testosterone level in the same study, luteinizing ontrono levels were elevated over a 24 ho period after the last ramelton treatment however, the durability of this cuteinizing hormone finding and its support for the proposed mechanistic explanation was not clearly established.

unauou was not cearly established. ugh the rodent tumors observed following ramelteon treatment occu sama levels of ramelteon and M-II in excess of mean clinical plasma a tations at the MHRD, the relevance of both rodent hepatic tumors and pr rat Leydig cell tumors to humans is not known.

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Information. Animal Data. Ramelteon did not produces any signals from animal behavioral studies indicating that the drug produces rewarding effects. Monkeys did not self-administer ramelteon and the drug did not induce a conditioned place preference in rats. There was no generalization between ramelteon and midazolam. Ramelteon did not affect rotorod performance, an indicator of disruption of motor function, and it did not potentiate the ability of diazepam to interfere with rotorod performance. studies. Impairment of Fertility Ramelteon was administered to male and female Sprague-Davley rats in an initial fertility and early embryonic development study at dose levels of 6. 60, or 600 mg/kg/day. No effects on male or female mating or fertility were observed with a ramelteon dose up to 600 mg/kg/day. (786-times higher than the MRHD on a mg/m² basis). Irregular estrus cycles, reduction in the num-ber of implant a ramelteon dose up to 800 mg/kg/day. (786-times higher than the MRHD on a mg/m² basis). Irregular estrus cycles, reduction in the num-ber of implants, and reduction in the number of corpora lutea occurred at the 600 mg/kg/day dose level. Administration of ramelteon up to 600 mg/kg/day to male rats tor 7 weeks had no effect on sperm quality and when the treated male rats were mated with untreated female rats there was no effect on implants or embryos. In a repeat of this study using oral administration of ramettoon at 2,00 or 200 mg/kg/day (rthe say Study), the say of effect one effects were seen on implantation or embryor viability. The *n*-offect dose for fertility endpoints was 20 mg/kg/day in themales (26-times the MRHD on a mg/m² basis) whon considering all studies. **Pregnancy: Pregnancy: Category C** produce physical dependence. OVERDOSAGE Signs and Symptoms No cases of ROZEREM overdose have been reported during clinical develop ment

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