

POLICY & PRACTICE

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E-prescribing Controlled Substances

Starting June 1, physicians will have the option of electronically prescribing controlled substances. The U.S. Drug Enforcement Administration said it wants clinicians to take advantage of new technology while maintaining the current control system. The agency also said that eprescribing could reduce paperwork and forgery. The interim final rule outlines the security precautions, such as physician identity confirmation by an authorized third party.

Topamax Case Settled for \$81 Million

Two drug companies owned by Johnson & Johnson will pay more than \$81 million to settle criminal and civil charges that they illegally promoted the epilepsy drug Topamax for off-label psychiatric uses. Under the agreement reached with the Department of Justice, Ortho-McNeil Pharmaceutical LLC will plead guilty to a misdemeanor and pay a \$6 million criminal fine for promoting Topamax for psychiatric uses through its "Doctor-for-a-Day" program. Another Johnson & Johnson subsidiary, Ortho-McNeil-Janssen Pharmaceuticals, will pay more than \$75 million to settle the charge that it caused false claims to be submitted to government health care programs for uses of the drug that were not medically accepted indications. Ortho-McNeil-Janssen denies that it engaged in any wrongful conduct. Health and Human Services Inspector General Daniel R. Levinson said in a statement that the agreement requires Ortho-McNeil-Janssen to avoid future illegal drug promotion.

Gulf War Illnesses Need Attention

The U.S. government has committed more than \$400 million for research on Gulf War illnesses, but veterans with symptoms need more, according to a draft report from the Gulf War Veterans' Illnesses Task Force. The task force suggested that the VA do a better job of training primary care physicians, environmental health clinicians, pension examiners, mental health professionals, and social workers about Gulf War illnesses. It also recommended the VA work with the Centers for Disease Control and Prevention to include veteran-specific data collection in its national surveys.

Groups Push FDA Drug Enforcement

Two minority advocacy groups are urging the Food and Drug Administration to work harder to remove unapproved drugs from the market. The National Mi-

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Forest Laboratories, Inc. Heidelberg Engineering, Inc. UCB. Inc.

nority Quality Forum (NMQF) and MANA, a national Latina organization, both asked the FDA to push harder on its effort to remove unapproved drugs from the market, which was launched in 2006. MANA said that only 400 of what could be thousands of unapproved drugs have been removed from the market since 2006, and NMQF warned that patients

and physicians may not know some drugs are unapproved. "These unapproved drugs, which have not been evaluated by FDA's rigorous approval process, may compromise the health of patients and create increased liability for the physicians who prescribe them," the NMQF said in its letter.

Chemical Reforms Introduced

After several months of hearings, Sen. Frank Lautenberg (D-N.J.), who chairs a Senate environmental health subcommittee, has introduced a bill that would significantly strengthen federal enforcement powers over potentially toxic chemicals. The "Safe Chemicals Act of 2010" would grant the Environmental Protection Agency new powers to ask chemical manufacturers for safety information, to categorize chemicals by risks they pose, and to remove dangerous chemicals from the market. "The EPA has been able to require comprehensive testing on just 200 of the more than 80,000 chemicals produced and used in the U.S.," according to advocacy group Health Care Without Harm, which supports strong reforms.

-Mary Ellen Schneider



Brief Summary of Prescribing Information.

For complete details, please see full Prescribing Information for Namenda

INDICATIONS AND USAGE ine hydrochloride) is indicated for the treatment of

moderate to severe dementia of the Alzheimer's type

CONTRAINDICATIONS
Namenda (memantine hydrochloride) is contraindicated in patients v known hypersensitivity to memantine hydrochloride or to any excipie used in the formulation.

PRECAUTIONS

Information for Patients and Caregivers: Caregivers should be instructed in the recommended administration (twice per day for doses above 5 mg) and dose escalation (minimum interval of one week between dose increases). Neurological Conditions

Neurological Conditions
Seizures: Namenda has not been systematically evaluated in patients with a seizure disorder. In clinical trials of Namenda, seizures occurred in 0.2% of patients treated with Namenda and 0.5% of patients treated with placebo.

Genitourinary Conditions
Conditions that raise urine pH may decrease the urinary elimination of memantine resulting in increased plasma levels of memantine

Hepatic Impairment

Namenda undergoes partial hepatic metabolism, with about 48% of administered dose excreted in urine as unchanged drug or as the sum of parent drug and the N-glucuronide conjugate (74%). No dosage adjustment is needed in patients with mild or moderate hepatic impairment. Namenda should be administered with caution to patients with severe hepatic

No dosage adjustment is needed in patients with mild or moderate renal impairment. A dosage reduction is recommended in patients with severe renal impairment (see CLINICAL PHARMACOLOGY and DOSAGE AND ADMINISTRATION in Full Prescribing Information).

Drug-Drug Interactions *N-methyl-D-aspartate (NMDA) antagonists:* The combined use of Namenda with other NMDA antagonists (amantadine, ketamine, dextromethorphan) has not been systematically evaluated and such use should be approached with caution.

Should be approached with Caudon.

Effects of Namenda on substrates of microsomal enzymes: In vitro studies conducted with marker substrates of CYP450 enzymes (CYP1A2, -2A6, -2C9, -2D6, -2E1, -3A4) showed minimal inhibition of these enzymes by memantine. In addition, in vitro studies indicate that at concentrations exceeding those associated with efficacy, memantine does not induce the cytochrome P450 isoenzymes CYP1A2, CYP2C9, CYP2E1, and CYP3A4/5. No pharmacokinetic interactions with drugs metabolized by these enzymes are expected.

expected.

cts of inhibitors and/or substrates of microsomal enzymes on Namenda:
nantine is predominantly renally eliminated, and drugs that are
strates and/or inhibitors of the CYP450 system are not expected to alter
metabolism of memantine.

the metabolism of memantine. Acetylcholinesterase (AChE) inhibitors: Coadministration of Namenda with the AChE inhibitor donepezil HCl did not affect the pharmacokinetics of either compound. In a 24-week controlled clinical study in patients with moderate to severe Alzheimer's disease, the adverse event profile observed with a combination of memantine and donepezil was similar to that of donepezil alone.

Drugs eliminated via renal mechanisms: Because memantine is eliminated in part by tubular secretion, coadministration of drugs that use the same renal cationic system, including hydrochlorothiazide (HCTZ), triamterene (TA), cationic system, including hydrochlorothiazide (HCTZ), triamterene (TA), metformin, cimetidine, ranitidine, quinidine, and nicotine, could potentially result in altered plasma levels of both agents. However, coadministration of Namenda and HCTZ/TA did not affect the bioavailability of either memantine or TA, and the bioavailability of HCTZ decreased by 20%. In addition, coadministration of memantine with the antihyperglycemic drug Glucovance® (glyburide and metformin HCI) did not affect the pharmacokinetics of memantine, metformin and glyburide. Furthermore, memantine did not modify the serum glucose lowering effect of Glucovance®. Drugs that make the urine alkaline: The clearance of memantine was reduced by about 80% under alkaline urine conditions at pH 8. Therefore, alterations of urine pH towards the alkaline condition may lead to an accumulation of the drug with a possible increase in adverse effects. Urine pH is altered by diet, drugs (e.g. carbonic anhydrase inhibitors, sodium bicarbonate) and clinical state of the patient (e.g. renal tubular acidosis or severe infections of the urinary tract). Hence, memantine should be used with caution under these conditions.

Carcinogenesis, Mutagenesis and Impairment of Fertility

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Carcinogenesis, Mutagenesis and Impairment of Fertility

There was no evidence of carcinogenicity in a 113-week oral study in mice
at doses up to 40 mg/kg/day (10 times the maximum recommended human
dose [MRHD] on a mg/m² basis). There was also no evidence of
carcinogenicity in rats orally dosed at up to 40 mg/kg/day for 71 weeks
followed by 20 mg/kg/day (20 and 10 times the MRHD on a mg/m² basis,
respectively) through 128 weeks.

Memantine produced no evidence of genotoxic potential when evaluated in
the in vitro. Studyimurium of E. coli reverse mutation assay, an in vitro.

Memantine produced no evidence of genotoxic potential when evaluated in the *in vitro S. typhimurium or E. coli* reverse mutation assay, an *in vitro* chromosomal aberration test in human lymphocytes, an *in vivo* cytogenetics assay for chromosome damage in rats, and the *in vivo* mouse micronucleus assay. The results were equivocal in an *in vitro* gene mutation assay using Chinese hamster V79 cells.

No impairment of fertility or reproductive performance was seen in rats administered up to 18 mg/kg/day (9 times the MRHD on a mg/m² basis) orally from 14 days prior to mating through gestation and lactation in females, or for 60 days prior to mating in males.

Pregnancy Pregnancy Category B: Memantine given orally to pregnant rats and pregnant rabbits during the period of organogenesis was not teratogenic up to the highest doses tested (18 mg/kg/day in rats and 30 mg/kg/day in rabbits, which are 9 and 30 times, respectively, the maximum recommended human dose [MRHD] on a mg/m² basis).

Slight maternal toxicity, decreased pup weights and an increased incidence of non-ossified cervical vertebrae were seen at an oral dose of 18 mg/kg/day in a study in which rats were given oral memantine beginning pre-mating and continuing through the postpartum period. Slight maternal toxicity and decreased pup weights were also seen at this dose in a study in which rats were treated from day 15 of gestation through the post-partum period. The no-effect dose for these effects was 6 mg/kg, which is 3 times the MRHD

on a mg/m² basis.

There are no adequate and well-controlled studies of memantine in pregnant women. Memantine should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Nursing Mothers
It is not known whether memantine is excreted in human breast milk.
Because many drugs are excreted in human milk, caution should be
exercised when memantine is administered to a nursing mother.

ADVERSE REACTIONS

The experience described in this section derives from studies in patients with Alzheimer's disease and vascular dementia.

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Adverse Events Leading to Discontinuation: In placebo-controlled trials in which dementia patients received doses of Namenda up to 20 mg/day, the likelihood of discontinuation because of an adverse event was the same in the Namenda group as in the placebo group. No individual adverse event was associated with the discontinuation of treatment in 1% or more of Namenda-treated patients and at a rate greater than placebo

Adverse Events Reported in Controlled Trials: The reported adverse events in Namenda (memantine hydrochloride) trials reflect experience gained under closely monitored conditions in a highly selected patient population. In actual practice or in other clinical trials, these frequency estimates may not apply, as the conditions of use, reporting behavior and the types of patitreated may differ. Table 1 lists treatment-emergent signs and symptoms were reported in at least 2% of patients in placebo-controlled dementia trials and for which the rate of occurrence was greater for patients treated with Namenda than for those treated with placebo. No adverse event occurred at a frequency of at least 5% and twice the placebo rate.

Table 1: Adverse Events Reported in Controlled Clinical Trials in at Least 2% of Patients Receiving Namenda and at a Higher Frequency than Placebo-

Body System	Piacedo	warnenda
Adverse Event	(N = 922)	(N = 940)
	%	%
Body as a Whole		
Fatigue	1	2
Pain	1	3
Cardiovascular System		
Hypertension	2	4
Central and Peripheral Nervous System		
Dizziness	5	7
Headache	3	6
Gastrointestinal System	-	0
Constipation	3	5
Vomiting	2	3
Musculoskeletal System		
Back pain	2	3
Psychiatric Disorders		
Confusion	5	6
Somnolence	2	3
Hallucination	2	3
Respiratory System		
Coughing	3	4
Dyspnea	1	2

Other adverse events occurring with an incidence of at least 2% in Namenda-treated patients but at a greater or equal rate on placebo were agitation, fall, inflicted injury, urinary incontinence, diarrhea, bronchitis, insomnia, urinary tract infection, influenza-like symptoms, abnormal gait, depression, upper respiratory tract infection, anxiety, peripheral edema, nausea, anorexia, and arthralgia.

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The overall profile of adverse events and the incidence rates for individual adverse events in the subpopulation of patients with moderate to severe Alzheimer's disease were not different from the profile and incidence rates described above for the overall dementia population.

described adover on the overall demental population.

Vital Sign Changes: Namenda and placebo groups were compared with respect to (1) mean change from baseline in vital signs (pulse, systolic blood pressure, diastolic blood pressure, and weight) and (2) the incidence of patients meeting criteria for potentially clinically significant changes from baseline in these variables. There were no clinically important changes in vital signs in patients treated with Namenda. A comparison of supine and standing vital sign measures for Namenda and placebo in elderly normal subjects indicated that Namenda treatment is not associated with orthostatic changes.

with orthostatic changes.

Laboratory Changes: Namenda and placebo groups were compared with respect to (1) mean change from baseline in various serum chemistry, hematology, and urinalysis variables and (2) the incidence of patients meeting criteria for potentially clinically significant changes from baseline in these variables. These analyses revealed no clinically important changes in laboratory test parameters associated with Namenda treatment.

In laboratory test parameters associated with Namenda treatment.

EGG Changes: Namenda and placebo groups were compared with respect to (1) mean change from baseline in various ECG parameters and (2) the incidence of patients meeting criteria for potentially clinically significant changes from baseline in these variables. These analyses revealed no clinically important changes in ECG parameters associated with Namenda treatment.

treatment.

Other Adverse Events Observed During Clinical Trials

Namenda has been administered to approximately 1350 patients with dementia, of whom more than 1200 received the maximum recommended dose of 20 mg/day. Patients received Namenda treatment for periods of up to 884 days, with 862 patients receiving at least 24 weeks of treatment and 387 patients receiving 48 weeks or more of treatment.

Treatment emergent signs and symptoms that occurred during 8 controlled clinical trials and 4 open-label trials were recorded as adverse events by the clinical investigators using terminology of their own choosing. To provide an overall estimate of the proportion of individuals having similar types of

overall estimate of the proportion of individuals having similar types of events, the events were grouped into a smaller number of standardized

categories using WHO terminology, and event frequencies were calculated

across all studies.
All adverse events occurring in at least two patients are included, except for All adverse events occurring in at least two patients are included, except for those already listed in Table 1, WHO terms too general to be informative, minor symptoms or events unlikely to be drug-caused, e.g., because they are common in the study population. Events are classified by body system and listed using the following definitions: frequent adverse events - those occurring in at least 1/100 patients; infrequent adverse events - those occurring in 1/100 to 1/1/000 patients. These adverse events are not necessarily related to Namenda treatment and in most cases were observed at a similar frequency in placebo-treated patients in the controlled studies. Body as a Whole: Frequent: syncope. Infrequent: hypothermia, allergic reaction.

Cardiovascular System: Frequent: cardiac failure. Infrequent: angina pectoris, bradycardia, myocardial infarction, thrombophlebitis, atrial fibrillation, hypotension, cardiac arrest, postural hypotension, pulmonary

emooism, puimonary edema. Central and Peripheral Nervous System: Frequent: transient ischemic attack, cerebrovascular accident, vertigo, ataxia, hypokinesia. Infrequent: paresthesia, convulsions, extrapyramidal disorder, hypertonia, tremor, aphasia, hypoesthesia, abnormal coordination, hemiplejai, hyperkinesia, involuntary muscle contractions, stupor, cerebral hemorrhage, neuralgia, particular augments.

Gastrointestinal System: Infrequent: gastroenteritis, diverticulitis, gastro-intestinal hemorrhage, melena, esophageal ulceration.

Hemic and Lymphatic Disorders: Frequent: anemia. Infrequent: leukopenia. Metabolic and Nutritional Disorders: Frequent: increased alkaline phosphatase, decreased weight. Infrequent: dehydration, hyponatremia, aggravated diabetes mellitus.

aggi avateu utauetes meintus. Psychiatric Disorders: Frequent: aggressive reaction. Infrequent: delus personality disorder, emotional lability, nervousness, sleep disorder, lili increased, psychosis, amnesia, apathy, paranoid reaction, thinking abnor crying abnormal, appetite increased, paroniria, delirium, depersonalizat neurosis, suicide attempt.

Respiratory System: Frequent: pneumonia. Infrequent: apnea.

Skin and Appendages: Frequent: rash. Infrequent: skin ulceration, pruritus, cellulitis, eczema, dermatitis, erythematous rash, alopecia, urticaria.

Special Senses: Frequent: cataract, conjunctivitis. Infrequent: macula lutea degeneration, decreased visual acuity, decreased hearing, tinnitus, blepharitis, blurred vision, corneal opacity, glaucoma, conjunctival hemorrhage, eye pain, retinal hemorrhage, exerophthalmia, diplopia, abnormal lacrimation, myopia, retinal detachment.

Urinary System: Frequent: frequent micturition. Infrequent: dysuria, hematuria, urinary retention.

Events Reported Subsequent to the Marketing of Namenda, both US and

EX-US Although no causal relationship to memantine treatment has been found, Although no causal relationship to memantine treatment has been found, the following adverse events have been reported to be temporally associated with memantine treatment and are not described elsewhere in labeling: aspiration pneumonia, asthenia, atrioventricular block, bone fracture, carpal tunnel syndrome, cerebral infarction, chest pain, cholelithiasis, claudication, collitis, deep venous thrombosis, depressed level of consciousness (including loss of consciousness and rare reports of coma), dyskinesia, dysphagia, encephalopathy, gastritis, gastroesophageal reflux, grand mal convulsions, intracranial hemorrhage, hepatitis (including increased ALT and AST and hepatic failure), hyperglycemia, hyperipidemia, hypoglycemia, ileus, increased INF, impotence, lethargy, malaise, myoclonus, neuroleptic malignant syndrome, acute pancreatitis, Parkinsonism, acute renal failure (including increased creatinine and renal insufficiency), prolonged OT interval, restlessness, sepsis, Stevens-Johnson syndrome, suicidal ideation, sudden death, supraventricular tachycardia, tachycardia, tardive dyskinesia, thrombocytopenia, and hallucinations (both visual and auditory). hallucinations (both visual and auditory).

ANIMAL TOXICOLOGY

ANIMAL TOXICOLOGY

Memantine induced neuronal lesions (vacuolation and necrosis) in the multipolar and pyramidal cells in cortical layers III and IV of the posterior cingulate and retrosplenial neocortices in rats, similar to those which are known to occur in rodents administered other NMDA receptor antagonists. Lesions were seen after a single dose of memantine. In a study in which rats were given daily oral doses of memantine for 14 days, the no-effect dose for neuronal necrosis was 6 times the maximum recommended human dose na months. human dose on a mg/m^2 basis. The potential for induction of central neuronal vacuolation and necrosis by NMDA receptor antagonists in humans is

DRUG ARUSE AND DEPENDENCE

DRUG ABUSE AND DEPENDENCE
Controlled Substance Class: Memantine HCl is not a controlled substance.
Physical and Psychological Dependence: Memantine HCl is a low to moderate affinity uncompetitive NMDA antagonist that did not produce any evidence of drug-seeking behavior or withdrawal symptoms upon discontinuation in 2,504 patients who participated in clinical trials at therapeutic doses. Post marketing data, outside the U.S., retrospectively collected, has provided no evidence of drug abuse or dependence.

OVERDINASE*

OVERDOSAGE

OVERDOSAGE
Signs and symptoms associated with memantine overdosage in clinical trials and from worldwide marketing experience include agitation, confusion, ECG changes, loss of consciousness, psychosis, restlessness, slowed movement, somnolence, stupor, unsteady gait, visual hallucinations, vertigo, vomiting, and weakness. The largest known ingestion of memantine worldwide was 2.0 grams in a patient who took memantine in conjunction with unspecified antidiabetic medications. The patient experienced coma, diplopia, and agitation, but subsequently recovered.

Because strategies for the management of overdose are continually evolving, it is advisable to contact a poison control center to determine the latest recommendations for the management an overdose of any drug. As in any cases of overdose, general supportive measures should be utilized, and treatment should be symptomatic. Elimination of memantine can be enhanced by acidification of urine.



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