CLINICAL

HF Rate Has Held Steady for 20 Years The incidence of heart failure has not improved over the past 20 years, according to Veronique L. Roger, M.D., of the Mayo Clinic, Rochester, Minn., and her associates.

The researchers conducted a community-based study to examine trends in heart failure (HF), hoping to avoid the selection biases shown to be present in previous studies of the issue. They studied a cohort of 4,537 cases that were first diagnosed in one Minnesota county between 1979 and 2000, a population with characteristics similar to those of whites across

CAPSULES

the United States but not to those of other racial groups.

The incidence of HF did not decrease appreciably over these 2 decades, ranging from 360/100,000 men in 1979-1984 to 383/100,000 men in 1996-2000 and from 284/100,000 women to 315/100,000 women in those same periods (JAMA 2004;292:344-50).

Overall HF mortality improved somewhat, from 57% in 1979-1984 to 48% in 1996-2000. However, improvements were not equal across different age groups and genders. Only the youngest men showed improved survival, while women and elderly people of both sexes had steady mortality over the 20-year study period.

A full 42% of cases were diagnosed as outpatients, and 26% of these patients were never hospitalized for their HF. Thus, hospital-based surveillance programs and studies based on inpatient data fail to identify a significant portion of HF cases, the investigators noted.

Pericardial Valves Superior to Porcine

Second-generation pericardial valves are more durable than traditional porcine valves as aortic valve replacements, said Guangqiang Gao, M.D., and associates at

ARICEPT® (Donepe	ezil Hydrochlorid	le Tablets)		Table 3. Adverse Events Reported in Controlled Clinical Trials in at Least 2% of Patients Receiving ARICEPT® and at a Higher Frequency than Placebo-treated Patients			
treatment of mild to moderate demen	seri for full prescribing inform tia of the Alzheimer's type. CON ochloride or to piperidine dei	TRAINDICATIONS AND USAGE ITRAINDICATIONS ARICEPT® is contu rivatives WARNINGS Anesthesia:	aindicated in patients with known	Body System/Adverse Event	Placebo (n=355)	ARICEPT® (n=747)	
inhibitor, is likely to exaggerate suc	ccinvlcholine-type muscle rel	axation during anesthesia. <i>Cardiovas</i>	cular Conditions: Because of	Percent of Patients with any Adverse Event	(11=000) 72	74	
their pharmacological action, cholir	nesterase inhibitors may have	vagotonic effects on the sinoatrial and a	trioventricular nodes. This effect	Body as a Whole		14	
may manifest as bradycardia or hear	rt block in patients both with an	d without known underlying cardiac con	duction abnormalities. Syncopal	Headache	9	10	
episodes have been reported in as	sociation with the use of ARIC	CEPT [®] . Gastrointestinal Condition	Through their primary action,	Pain, various locations	8	9	
cholinesterase inhibitors may be exp	pected to increase gastric acid	secretion due to increased cholinergic a	stivity. Therefore, patients should	Accident	6	7	
e a those with a history of ulcer dis	s of active of occult gastronne sease or those receiving conc	urrent nonsteroidal anti-inflammatory d	ased lisk for developing diders,	Fatigue Cardiousseular Sustem	3	5	
of ABICEPT® have shown no incr	rease, relative to placebo, in th	he incidence of either peptic ulcer dise	se or astrointestinal bleeding.	Svincone	1	2	
ARICEPT®, as a predictable consec	quence of its pharmacological	I properties, has been shown to produce	diarrhea, nausea and vomiting.	Digestive System	1	-	
These effects, when they occur, app	ear more frequently with the 1	0 mg/day dose than with the 5 mg/day d	ose. In most cases, these effects	Nausea	6	11	
have been mild and transient, so	ometimes lasting one to thr	ee weeks, and have resolved during	continued use of ARICEPT®.	Diarrhea	5	10	
Genitourinary: Although not ob	oserved in clinical trials of A	RICEP1 [®] , cholinomimetics may caus	e bladder outflow obstruction.	Vomiting	3	5	
Neurological Conditions: Sel20	Jres: Unolinomimetics are bell fostation of Alzbeimer's Diseas	leved to nave some potential to cause ge	ef their ehelinemimetic actions	Anorexia	2	4	
cholinesterase inhibitors should l	he prescribed with care to r	nationts with a history of asthma or o	hstructive nulmonary disease	Feedwaren in Experimentation System	2	4	
PRECAUTIONS Drug-Drug Inte	eractions (see Clinical Pharm	nacoloov: Clinical Pharmacokinetics: Di	ug-drug Interactions) <i>Effect of</i>	Metaholic and Nutritional Systems	5	4	
ARICEPT® on the Metabolisn	n of Other Drugs: No in vivo	clinical trials have investigated the effect	t of ARICEPT® on the clearance	Weight Decrease	1	3	
of drugs metabolized by CYP 3A4 ((e.g. cisapride, terfenadine) or	by CYP 2D6 (e.g. imipramine). Howeve	; <i>in vitro</i> studies show a low rate	Musculoskeletal System			
of binding to these enzymes (mea	an K _i about 50-130 µM), that	, given the therapeutic plasma concent	rations of donepezil (164 nM),	Muscle Cramps	2	6	
indicates little likelihood of interferer	nce. Whether ARICEP I® has a	any potential for enzyme induction is not	rnown. Formal pharmacokinetic	Arthritis	1	2	
Studies evaluated the potential of AF	RICEP I © TOR INTERACTION WITH TR	neophylline, cimetidine, warrarin, digoxii	and ketoconazole. No effects of	Nervous System	0	0	
Ketoconazole and quinidine inhibito	ts of these drugs were observers of CVP/50, 34/ and 206, r	veu. <i>Ellect of other brugs on the l</i> espectively inhibit donenezil metabolism	in vitro Whether there is a clinical	INSOMNIA	6	9	
effect of quinidine is not known. In a	17-day crossover study in 18 h	espectively, in hibit donepezi metabolisi realthy volunteers, ketoconazole (200 mc	n d) increased mean donenezil	Dizziness	0	8	
(5 mg q.d.) concentrations (AUCo	and C) by 36%. The clir	nical relevance of this increase in concer	tration is unknown. Inducers of	Abnormal Dreams	0	3	
CYP 2D6 and CYP 3A4 (e.g., pheny	toin, carbamazepine, dexamet	hasone, rifampin, and phenobarbital) cou	ld increase the rate of elimination	Somnolence	<1	2	
of ARICEPT®. Formal pharmacokin	netic studies demonstrated that	t the metabolism of ARICEPT® is not sig	nificantly affected by concurrent	Urogenital System			
administration of digoxin or cimetion	dine. Use with Anticholine	ergics: Because of their mechanism of	action, cholinesterase inhibitors	Frequent Urination	1	2	
have the potential to interfere with the	e activity of anticholinergic me	dications. <i>Use with Cholinomimetic</i>	s and Uther Cholinesterase	Other Adverse Events Observed During Clinical Trials	ARICEPT® has been admini	stered to over 1700 individuals	during clinical
nnibitors: A synergistic ellect m	ay de expected when chointe r cholineraic agonists such a	esterase inflibitors are given concurrent	iy with succinyicholine, similar	trials worldwide. Approximately 1200 of these patients have been	treated for at least 3 months	and more than 1000 patients ha	ve been treated
Fertility No evidence of a carcinon	enic notential was obtained in	an 88-week carcinonenicity study of dor	enezil hydrochloride conducted	for at least 6 months. Controlled and uncontrolled trials in the U	nited States included approx	cimately 900 patients. In regards	s to the highest
in CD-1 mice at doses up to 180 m	o/ko/dav (approximately 90 ti	mes the maximum recommended huma	n dose on a mo/m² basis), or in	dose of 10 mg/day, this population includes 650 patients treate	d for 3 months, 475 patients	treated for 6 months and 116 p	patients treated
a 104-week carcinogenicity study in Sprague-Dawley rats at doses up to 30 mg/kg/day (approximately 30 times the maximum				for over 1 year. The range of patient exposure is from 1 to 121-	4 days. Treatment emergent States were recorded as adv	signs and symptoms that occ	urrea auring 3 stigstore uning
recommended human dose on a mg/m ² basis). Donepezil was not mutagenic in the Ames reverse mutation assay in bacteria, or in a				terminology of their own chonsing. To provide an overall estimate of the proportion of individuals having similar types of events the events.			
mouse lymphoma forward mutation assay <i>in vitro</i> . In the chromosome aberration test in cultures of Chinese hamster lung (CHL) cells, some				were grouped into a smaller number of standardized categories using a modified COSTART dictionary and event frequencies were calculated			
ciastogenic effects were observed. Donepezil was not clastogenic in the <i>in vivo</i> mouse micronucleus test and was not genotoxic in an <i>in</i>				across all studies. These categories are used in the listing below. The frequencies represent the proportion of 900 patients from these trials			
the maximum recommended human dose on a mo/m ² hasis) Prennancy Prennancy Category C *Teratology studies conducted in				who experienced that event while receiving ARICEPT® All adverse events occurring at least twice are included, except for those already listed			
pregnant rats at doses up to 16 mg/	/ko/dav (approximately 13 tim	the maximum recommended humar	in Tables 2 or 3, COSTART terms too general to be informative, or events less likely to be drug caused. Events are classified by body				
pregnant rabbits at doses up to 10 mg/kg/day (approximately 16 times the maximum recommended human dose on a mg/m ² basis) did				system and listed using the following definitions: <i>Trequent adve</i>	<i>rse events</i> —tnose occurrin	g in at least 1/100 patients; <i>intre</i> silv related to A DICEDT® treatme	quentaaverse
not disclose any evidence for a teratogenic potential of donepezil. However, in a study in which pregnant rats were given up to 10 mg/kg/day				events—those occurring in 1/100 to 1/1000 patients. These adverse events are not necessarily related to AKICEP1® treatment and in most			
(approximately 8 times the maximum recommended human dose on a mg/m ² basis) from day 17 of gestation through day 20 postpartum,				seen in studies conducted outside the United States. Rody as a Whole: Frequent: influenza chest pain toothache: Infrequent: feuere edema			
there was a slight increase in still births and a slight decrease in pup survival through day 4 postpartum at this dose; the next lower dose				face, periorbital edema, herria hiatal, abscess, cellulitis, chills, ceneralized coldness, head fullness, listlessness, Cardiovascular System:			
only if the notantial benefit justifier	the potential rick to the fature	Suules In pley I all wonten. Anicer 1° s	Frequent: hypertension, vasodilation, atrial fibrillation, hot flashes, hypotension; Infrequent: angina pectoris, postural hypotension,				
human hreast milk ABICEPT® has	s no indication for use in nursi	ing mothers Pediatric Use There are i	n adequate and well-controlled	myocardial infarction, AV block (first degree), congestive heart	myocardial infarction, AV block (first degree), congestive heart failure, arteritis, bradycardia, peripheral vascular disease, supraventricular		
trials to document the safety and effic	cacy of ARICEPT® in any illne	ss occurring in children. Geriatric Use	Alzheimer's disease is a disorder	tachycardia, deep vein thrombosis. Digestive System: Frequent: fecal incontinence, gastrointestinal bleeding, bloating, epigastric pain;			
occurring primarily in individuals o	ver 55 years of age. The mean	age of the patients enrolled in the clinica	studies with ARICEPT® was 73	Intrequent: eructation, gingivitis, increased appetite, flatuience, p	periodontal abscess, choielit	nasis, diverticulitis, drooling, di	y mouth, tever
years; 80% of these patients were b	between 65 and 84 years old a	and 49% of the patients were at or abov	e the age of 75. The efficacy and	sole, yasulius, iiiliabie colori, iongue edenia, epiyasulic distress thirst jaundice, melena, polydinsia, duodenal ulcar stomach ulc	or Endocrine System: Intersection	idi iSdi i ili idSeS, i ieni uni inuita, il raguant: diabatas mallitus, gaita	r Hemic and
safety data presented in the clinical t	trials section were obtained fro	om these patients. There were no clinical	y significant differences in most	I vmnhatic System: Infrequent: anemia thrombocythemia	a thrombocytonenia eosir	iophilia erythrocytonenia Me	taholic and
adverse events reported by patient (groups ≥65 years old and <6	b years old. ADVERSE REACTIONS	Reading to the ADIOFDER C	Nutritional Disorders: Frequent: dehydration; Infrequent: dol	ut, hypokalemia, increased ci	eatine kinase, hyperglycemia. w	eight increase.
ma/day treatment aroune were con	mnarable to those of nlacebo	 united trials of AntiGEMT® due to adve treatment around at anonovimately 5% 	The rate of discontinuation of	increased lactate dehydrogenase. Musculoskeletal Syst	em: Frequent: bone fract	ure; Infrequent: muscle weak	ness, muscle
natients who received 7-day escala	itions from 5 ma/day to 10 m	1/day was higher at 13%. The most cor	nmon adverse events leading to	fasciculation. Nervous System: Frequent: delusions, trem	ior, irritability, paresthesia,	aggression, vertigo, ataxia, inc	reased libido
discontinuation, defined as those or	curring in at least 2% of patient	ts and at twice the incidence seen in place	po patients, are shown in Table 1.	restlessness, abnormal crying, nervousness, aphasia; Infreque	ant: cerebrovascular acciden	t, intracranial hemorrhage, tran	sient ischemic
Table 4 Mart Francis Adv		Withdrawal from Controlled Oliv	aal Triala hu Daaa Orana	allack, emotional lability, neuralgia, coluness (localized), n	Nuscie spasini, dysprioria, luenhacia, hostility decrease	gait abriormanty, riypertorna, id libido, melancholia, emotior	nypokinesia,
iable 1. Wost Frequent Adv	erse events Leading to t	williurawai from controlled Clin	cal mais by Dose Group	nystaomus, nacino, Respiratory System: Frequent: dysnue	a sore throat, bronchitis: <i>Infr</i>	e <i>ouent:</i> enistaxis, post nasal dri	n. pneumonia
Dose Group	Placebo	5 mg/day ARICEPT®	10 mg/day ARICEPT®	hyperventilation, pulmonary congestion, wheezing, hypoxia, pl	haryngitis, pleurisy, pulmon	ary collapse, sleep apnea, snor	ing. Skin and
Patients Randomized	355	350	315	Appendages: Frequent: pruritus, diaphoresis, urticaria; Infrequent	<i>ent:</i> dermatitis, erythema, skir	discoloration, hyperkeratosis, a	lopecia, fungal
Event/% Discontinuing				dermatitis, herpes zoster, hirsutism, skin striae, night sweats, ski	n ulcer. Special Senses: /	requent: cataract, eye irritation,	vision blurred
Nausea	1%	1%	3%	Intrequent: dry eyes, glaucoma, earache, tinnitus, blepharitis, de	ecreased hearing, retinal hen	norrhage, otitis externa, otitis m	edia, bad taste,
Diarrhea	0%	<1%	3%	conjunctival nemorriage, ear ouzzing, motion sickness, spots be	ovetitie enurgenital Sys	em. <i>Frequent:</i> Urinary incontin	ence, nocturia, bility to empty
Vomiting	<1%	<1%	2%	hladder breast fibroadenosis, fibroevetie breast maetitie, pourie	renal failure vaninitie Dre	tintroduction Reports Volume	unity to critiply stary reports of
Most Frequent Adverse Clini	ical Events Seen in Asso	ciation with the Use of ARICEPT	® The most common adverse	adverse events temporally associated with ARICFPT® that have	been received since market	introduction that are not listed a	above, and that
events, defined as those occurring	at a frequency of at least 5%	in patients receiving 10 mg/day and tw	ice the placebo rate. are laroely	there is inadequate data to determine the causal relationship wi	there is inadequate data to determine the causal relationship with the drun include the following: andominal nain, anitation, cholecustitis		
predicted by ARICEPT®'s choling	omimetic effects. These inclu	ide nausea, diarrhea, insomnia, vomiti	ng, muscle cramp, fatigue and	confusion, convulsions, hallucinations, heart block (all types), he	confusion, convulsions, hallucinations, heart block (all types), hemolytic anemia, hepatitis, hyponatremia, neuroleptic malignant syndrome,		
anorexia. These adverse events we	ere often of mild intensity an	d transient, resolving during continued	ARICEPT® treatment without	pancreatitis, and rash. OVERDOSAGE Because strategies	for the management of	overdose are continually e	volving, it is
the need for dose modification. The	here is evidence to suggest	that the frequency of these common a	dverse events may be affected	advisable to contact a Poison Control Center to deter	rmine the latest recomi	nendations for the manag	ement of an

hv the rate of titration. An open-label study was conducted with 269 patients who received placebo in the 15 and 30-week studies These patients were titrated to a dose of 10 mg/day over a 6-week period. The rates of common adverse events were lower than those seen in patients titrated to 10 mg/day over one week in the controlled clinical trials and were comparable to those seen in patients on 5 mg/day. See Table 2 for a comparison of the most common adverse events following one and six week titration regimens.

lable 2. Comparison of Rates of Adverse Events in Patients litrated to 10 mg/day Over 1 and 6 weeks								
Adverse Event	No tit Placebo (n=315)	ration 5 mg/day (n=311)	One week titration 10 mg/day (n=315)	Six week titration 10 mg/day (n=269)				
Nausea	6%	5%	19%	6%				
Diarrhea	5%	8%	15%	9%				
Insomnia	6%	6%	14%	6%				
Fatigue	3%	4%	8%	3%				
Vomiting	3%	3%	8%	5%				
Muscle cramps	2%	6%	8%	3%				
Anorexia	2%	3%	7%	3%				
Adverse Events Report of clinical trials in a highly may not apply, as the condi signs and symptoms that w the rate of occurrence was frequently in formale action	ted in Controlled Tri selected patient popula tions of use, reporting t vere reported in at least greater for ARICEPT® a	als The events cited ation. In actual clinica behavior, and the kind 2% of patients in pla assigned than placeb	reflect experience gained under clc al practice or in other clinical trials, ls of patients treated may differ. Table cebo-controlled trials who received o assigned patients. In general, adv	sely monitored conditions these frequency estimates e 3 lists treatment emergent d ARICEPT [®] and for which rerse events occurred more				

advisable to contact a Poison Control Center to determine the latest recommendations for the mmangement of an overdose of any drug. As in any case of overdose, general supportive measures should be utilized. Overdosage with cholinesterase inhibitors can result in cholinergic crisis characterized by severe nausea, vomiting, salivation, sweating, bradycardia, hypotension, respiratory depression, collapse and comvulsions. Increasing muscle weakness is a possibility and may result in chash if respiratory muscles are involved. Tetrary anticholinergics such as atropine may be used as an antichole for ARICCPT® overdosage. Intravenous atropine sultate titrated to effect is recommended: an initial dose of 1.0 to 2.0 mg IV with subsequent doses based upon clinical response. Adpical responses in blood pressure and heart rate have been reported with other cholinomimetics when co-administerd with qualarnary articholinergics such as glycogruptial. It is not hornow whether ARICEPT® and/or its metabolities can be removed by dialysis (hornoida) sis, performal dialysis, or hernotilitation). Dose-related signs of toxicity in animals included reduced spontaneous movement, prone position, extension and lacerization consultione deversed respirations estimation extension and lacer body and being theory and the article respiration extension consistence and heart rate brave bear reported with other cholinomimetics when co-administer with qualarnary anticholinergics such as glycogruptials. It is not known whether ARICEPT® and/or its metabolities can be removed by dialysis (hornoida) extension and the utilized to effect any extension of the state of the order based and the order of the order based and the order based and the transformation extension and the interview of the order and the transformation of the order based and the order and the article to any extension order state and the order based and the order based and the order based and the article to any extensis order torder and the order and theory based and the or Benotice utarysis, and the second sec is likely to be associated with a higher incidence of cholinergic adverse events than the 5 mg dose. In open label trials using a 6 week titration, the frequency of these same adverse events was similar between the 5 mg and 10 mg dose groups. Therefore, because steady state is not ecause the incidence of untoward effects may nfluenced by th of dose escalation, treatmen of 10 mg should not be contemplated until patients have been on a daily dose of 5 mg for 4 to 6 weeks. ARICEPT® should be taken in the of 10 mg should not be contemplated uniting attents have been on a daily does or 5 mg for 4 to 6 weeks. Artic P1° exhold be taken in the verify that of the first provide the provided by the provided of the provided by the 15°C to 30°C (59°F to 86°F).



AB214802F

Providence Health System, Portland, Ore.

The first generation of pericardial valves was taken off the market because of the high rate of structural failure. The second generation was marketed in 1991 in the United States but has not been as widely used as porcine valves, largely because of the poor history of its predecessors. Dr. Gao and associates compared the two types of valve by reviewing the experience at their facility with all 518 aortic valve replacements using porcine valves, which were done between 1974 and 1996, and all 1,021 aortic valve replacements using second-generation pericardial valves, which were done between 1991 and 2002 (J. Am. Coll. Cardiol. 2004;44:384-8).

Overall late survival rates were similar between the two groups, as was freedom from valve-related complications such as thromboembolic events and endocarditis. But the estimated 10-year rate of explantation was much higher for porcine (10%)than for pericardial valves (3%). Only four pericardial valves required replacement because of structural deterioration.

When Thrombolysis Is the Sole Option

For acute MI patients who either can't afford invasive intervention or don't have ready access to a hospital with a cath lab, medical thrombolysis is the only viable option. Fortunately, it has proved beneficial to most such patients, either as a definitive treatment or as a bridge to later intervention, according to Rami Khouzam, M.D., of the Tucson Hospitals Medical Education Program, and his associates.

The investigators reviewed the experience of 42 MI patients treated between 1999 and 2002 at a small county hospital in southern Arizona that offered only thrombolysis. MI patients in whom thrombolysis produced resolution of ST-segment changes within 90 minutes had a very high likelihood of a patent infarct artery and a very low mortality. Further PCI would be unlikely to improve their outcome. In contrast, those in whom thrombolysis did not produce ST-segment resolution within 90 minutes had a 40% probability for an occluded artery and required transfer for "rescue" PCI, the researchers said (Chest 2004;126:457-60).

Gemfibrozil Does Not Slow Renal Decline

Even though gemfibrozil cuts triglyceride and total cholesterol levels while raising HDL cholesterol, it does not slow renal decline in patients with mild renal insufficiency and concomitant coronary disease, reported Marcello Tonelli, M.D., of the University of Alberta, Edmonton, and his associates.

The researchers analyzed data from a trial conducted at 20 Veterans Affairs medical centers involving 399 subjects with both coronary heart disease and renal insufficiency (Am. I. Kidney Dis. 2004:44:832-9).

Compared with placebo, gemfibrozil (Lopid) cut triglycerides by 55.0 mg/dL, cut total cholesterol by 6.7 mg/dL, and raised HDL cholesterol by 2.3 mg/dL. But it did not slow the decline of renal function or improve subjects' chances of progressing to moderate or severe renal insufficiency.

Both gemfibrozil and statins improved cardiovascular outcomes in this population, but it appears that only statins exert renal benefits as well.