'Young-Onset' Alzheimer's Is Increasing

BY BRUCE K. DIXON Chicago Bureau

CHICAGO — The increasing incidence of early-onset dementia represents both a national challenge and a future crisis, according to the Alzheimer's Association, which is about to change the designation to "young-onset."

Data from the national Health and Retirement Study indicate that in 2000, 480,000 Americans aged 55-64 had cognitive impairment at a level severe enough to be considered disabling.

Other research suggests that in 2004, Americans with young Alzheimer's and other dementias numbered from 220,000 to 640,000, according to an overview distributed at a conference on dementia sponsored by the Alzheimer's Association.

By the end of 2007, the Alzheimer's Association is expected to start using the term "young-onset" Alzheimer's because of the tendency of people to confuse "early-onset" with "early-stage" disease, said JoAnn Webster, early-onset coordinator for the Oklahoma and Arkansas chapters of the Alzheimer's Association in Tulsa.

"It's young people we're talking about, not early disease," she said in an interview.

However, treatment, support, and education often are delayed because of lack of awareness of early-onset dementia and physician reluctance to diagnose Alzheimer's disease in younger patients.

Patients often are told they are depressed or have other mental disorders, and the dementia goes unrecognized, Ms. Webster said during a seminar. "The person with early-onset dementia may indeed have mild depression or depressed mood because of uncertainty and fear, and often they are given diagnoses of anxiety or panic disorder, or they're given an antidepressant and sent home," she said.

When a woman begins to experience memory loss, she may be told by her physician that hormones are to blame and that her best option is to undergo a hysterectomy, she said.

When younger patients or their families complain of memory loss, physicians should launch the same battery of neurologic tests they would offer an elderly patient and, if dementia is diagnosed, they should immediately refer the patients to the Alzheimer's Association (800-272-3900) and its Web site (www.alz.org)

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for support and guidance, she added.

"I push for a proper diagnosis, because knowing what's wrong gives them the power and strength to fight back, whereas dodging the real diagnosis is extremely stressful for them," she added.

A proper diagnosis takes on additional importance when memory loss has led to job loss, Ms. Webster said, noting that having the diagnosis may confer eligibility for unemployment insurance.

'We've come to the point where we

have to make a diagnosis for these people who are young, proactive, and want to know what's wrong with them. They need to understand that Alzheimer's is a treatable illness, and the more all of us learn about it, and the more people we can engage in treatment and clinical trials of new drugs, the sooner we'll find a cure," Ms. Webster said.

She invited physicians to contact her by e-mail at joann.webster@alz.org or via the phone at 918-481-7746.





BRIEF SUMMARY OF FULL PRESCRIBING INFORMATION

INDICATIONS AND USAGE Rebif® (interferon-beta-1a) is indicated for the treatment of patients with relapsing forms of multiple sclerosis to decrease the frequency of clinical exacerbations and delay the accumulation of physical disability. The efficacy of Rebif® in chronic progressive multiple sclerosis has not been established.

Clinical Studies
Two multicenter studies evaluated the safety and efficacy of Rebif® in patients with relapsing-remitting multiple sclerosis. Study 1 demonstrated that Rebif® significantly reduced the number of relapses per patient compared to placebo at 2 years. Study 2 was a comparative trial comparing Rebif® 44 mcg sc tiw and Avonex® 30 mcg im qw. The results of this trial demonstrated that patients treated with Rebif® 44 mcg sc tiw were more likely to remain relapse-free at 24 and 84 weeks than were patients treated with Avonex® 30 mcg im qw. Adverse reactions over 48 weeks were generally similar between the two treatment groups. Exceptions included injection site disorders (83% of patients on Rebif® vs. 28% of patients on Avonex®), hepatic function disorders (18% on Rebif® vs. 10% on Avonex®), and leukopenia (6% on Rebif® vs. 10% on Avonex®), which were observed with greater frequency in the Rebif® group compared to the Avonex® group.

CONTRAINDICATIONS
Rebiff (interferon beta-1a) is contraindicated in patients with a history of hypersensitivity to natural or recombinant interferon, human albumin, mannitol USP, sodium acetate, or Water for Injection USP.

WARNINGS
Rebif* (interferon beta-1a) should be used with caution in patients with depression, a condition that is common in people with multiple sclerosis. Depression, suicidal ideation, and suicide attempts have been reported to occur with increased frequency in patients receiving interferon compounds, including Rebif*. Patients should be advised to report immediately any symptoms of depression and/or suicidal ideation to the prescribing physician. If a patient develops depression, cessation of treatment with Rebif*should be considered.

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Severe liver injury, including some cases of hepatic failure requiring liver transplantation has been reported rarely in patients taking Rebif®. Symptoms of liver dysfunction began from one to six months following the initiation of Rebif®. If jaundice or other symptoms of liver dysfunction appear, treatment with Rebif® should be discontinued immediately due to the potential for rapid progression to liver failure. Asymptomatic elevation of hepatic transaminases (particularly SGPT) is common with interferon therapy (see ADVERSE REACTIONS). Rebif® should be initiated with caution in patients with active liver disease, alcohol abuse, increased serum SGPT (>2.5 times ULN), or a history of significant liver disease. Also, the potential risk of Rebif® used in combination with known hepatotoxic products should be considered prior to Rebif® administration, or when adding new agents to the regimen of patients already on Rebif®. Reduction of Rebif® dose should be considered if SGPT rises above 5 times the upper limit of normal. The dose may be gradually re-escalated when enzyme levels have normalized.

Anaphylaxis and other allergic reactions (some severe) have been reported as a rare complication of Rebif*. Other allergic reactions have included skin rash and urticaria, and have ranged from mild to severe without a clear relationship to doscor duration of exposure. Several allergic reactions, some severe, have occurred after prolonged use.

PRECAUTIONS
General: Caution should be exercised when administering Rebif® to patients with pre-existing seizure disorders. Seizures have been associated with the use of beta interferons. A relationship between occurrence of seizures and the use of Rebif® has not been established. Leukopenia and new or worsening thyroid abnormalities have developed in some patients treated with Rebif®. Regular monitoring for these conditions is recommended.

Information for Patients: All patients should be instructed to read the Rebif® Medication Guide supplied to them. Patients should be cautioned not to change the dosage or the schedule of administration without medical consultation. Patients should be informed of the most common and the most severe adverse reactions associated with the use of Rebif®. Patients should be advised of the symptoms associated with these conditions, and to report them to their physician.

Female patients should be cautioned about the abortifacient potential of Rebif®

Patients should be instructed in the use of aseptic technique when administering Rebif®. Appropriate instruction for self-injection or injection by another person should be provided, including careful review of the Rebif® Medication Guide. If a patient is to self-administer Rebif® the physical and cognitive ability of that patient to self-administer and properly dispose of syringes should be assessed. The initial injection should be performed under the supervision of an appropriately qualified health care professional. Patients should be advised of the importance of rotating sites of injection with each dose, to minimize the likelihood of severe injection site reactions or necrosis.

Laboratory Tests: In addition to those laboratory tests normally required for monitoring patients with multiple sclerosis, blood cell counts and liver function tests are recommended at regular intervals (1, 3, and 6 months) following introduction of Rebif* therapy and then periodically thereafter in the absence of clinical symptoms. Thyroid function tests are recommended every 6 months in patients with a history of thyroid dysfunction or as clinically indicated. Patients with myelosuppression may require more intensive monitoring of complete blood cell counts, with differential and platelet counts.

Immunization: Patients taking Rebif® may receive concomitant influenza vaccination and achieve similar positive antibody response to the vaccination as patients not receiving Rebif®. The exact relationship of antibody titers to vaccine efficacy is unknown in patients taking Rebif®.

Drug Interactions: Drug interaction studies have not been conducted with Rebif[®]. Due to its potential to cause neutropenia and lymphopenia, proper monitoring of patients is required if Rebif[®] is given in combination with myelosuppressive agents. Also, the potential for hepatic injury should be considered when Rebif[®] is used in combination with other products associated with hepatic injury, or when new agents are added to the regimen of patients already on Rebif[®] (see WARNINGS).

Carcinogenesis, Mutagenesis, Impairment of Fertility: No carcinogenicity data for Rebif® are available in animals or humans. Rebif® was not mutagenic when tested in the Ames bacterial test and in an in vitro cytogenetic assay in human lymphocytes in the presence and absence of metabolic activation. No studies have been conducted to evaluate the effects of Rebif® of fertility in humans. In studies in normally cycling female cynomolgus monkeys given daily sc injections of Rebif® for six months at doses of up to 9 times the recommended weekly human dose (based on body surface area), no effects were observed on either menstrual cycling or serum estradiol levels. The validity of extrapolating doses used in animal studies to human doses is not established. In male monkeys, the same doses of Rebif® had no demonstrable adverse effects on sperm count, motility, morphology, or function.

Pregnancy Category C: Rebif® treatment has been associated with significant increases in embryolethal or abortifacient effects in cynomolgus monkeys administered doses approximately 2 times the cumulative weekly human dose (based on either body weight or surface area) either during the period of organogenesis (gestation day 21-89) or later in pregnancy. There were no fetal malformations or other evidence of teratogenesis noted in these studies. These effects are consistent with the abortifacient effects of other type I interferons. There are no adequate and well-controlled studies of Rebif® in pregnant women. However, in Studies 1 and 2, there were 2 spontaneous abortions observed and 5 fetuses carried to term among 7 women in the Rebif® groups. If a woman becomes pregnant or plans to become pregnant while taking Rebif®, should be informed about the potential hazards to the fetus and discontinuation of Rebif® should be considered. A pregnancy registry has been established to monitor pregnancy outcomes of women exposed to Rebif® while pregnant. Register patients online at www.RebifPregnancyRegistry.com or call MS LifeLines™ at 1-877-447-3243.

Nursing Mothers: It is not known whether Rebif® is excreted in human milk.

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Pediatric Use: The safety and effectiveness of Rebif® in pediatric patients have not been studied. Geriatric Use: Clinical studies of Rebif® did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently than younger subjects.

ADVERSE REACTIONS

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The most frequently reported serious adverse reactions with Rebif® were psychiatric disorders including depression and suicidal ideation or attempt (see WARNINGS). The incidence of depression of any severity in the Rebif®-treated groups and placebo-treated group was approximately 25%. In post-marketing experience, Rebif® administration has been rarely associated with severe liver dysfunction, including hepatic failure requiring liver transplantation (see WARNINGS). The most commonly reported adverse reactions were injection site disorders, influenza-like symptoms (headache, fatique, fever, rigors, chest pain, backpain, myalgia), abdominal pain, depression, elevation of liver enzymes and hematologic abnormalities. The most frequently reported adverse reactions resulting in clinical intervention (e.g., discontinuation of Rebif®, adjustment in dosage, or the need for concomitant medication to treat an adverse reaction symptom) were injectionsite disorders, influenza-like symptoms, depression and elevation of liver enzymes (See WARNINGS). Injection site necross was rare.

Table 1. Adverse Reactions and Laboratory Abnormalities in Study 1			
BODY SYSTEM Preferred Term	Rebif® Placebo tiw (n=187)	Rebif® 22 mcg tiw (n=189)	44mcg tiw (n=184)
BODY AS A WHOLE			
Influenza-like symptoms	51%	56%	59%
Headache	63%	65%	70%
Fatigue	36%	33%	41%
Fever	16%	25%	28%
Rigors	5%	6%	13%
Chest Pain	5%	6%	8%
Malaise	1%	4%	5%
	1 /0	470	370
INJECTION SITE DISORDERS	200/	000/	000/
Injection Site Reaction	39%	89%	92%
Injection Site Necrosis	0%	1%	3%
CENTRAL & PERIPH NERVOU	5		
SYSTEM DISORDERS	E0/	70/	C0/
Hypertonia	5%	7% 5%	6% 4%
Coordination Abnormal Convulsions	2% 2%	5% 5%	4% 4%
	2%	5%	4%
ENDOCRINE DISORDERS			
Thyroid Disorder	3%	4%	6%
GASTROINTESTINAL SYSTEM DISORDERS	1		
Abdominal Pain	17%	22%	20%
Dry Mouth	1%	1%	5%
LIVER AND BILIARY SYSTEM DISORDERS			
SGPT Increased	4%	20%	27%
SGOT Increased	4%	10%	17%
Hepatic Function Abnormal	2%	4%	9%
Bilirubinaemia	1%	3%	2%
MUSCULO-SKELETAL SYSTEI DISORDERS	М		
Myalgia	20%	25%	25%
Back Pain	20%	23%	25%
Skeletal Pain	10%	15%	10%
HEMATOLOGIC DISORDERS			
Leukopenia	14%	28%	36%
Lymphadenopathy	8%	11%	12%
Thrombocytopenia	2%	2%	8%
Anemia	3%	3%	5%
PSYCHIATRIC DISORDERS	5,0	570	370
Somnolence	1%	4%	5%
	1 70	470	370
SKIN DISORDERS			
Rash Erythematous	3%	7%	5%
Rash Maculo-Papular	2%	5%	4%
URINARY SYSTEM DISORDEI	RS		
Micturition Frequency	4%	2%	7%
Urinary Incontinence	2%	4%	2%
VISION DISORDERS			
Vision Abnormal	7%	7%	13%
Xerophthalmia	0%	3%	1%
Acropridiania	0 /0	3/0	1 70

The safety of Rebif® (22 mcg Ine safety of Rebri[®] (22 mg and 44 mg) vs placebo was studied in 560 patients with RRMS who were treated for 24 months (Study 1). Table 1 enumerates ackerse events and laboratory abnormalities that occurred at an incidence that was at least 2% more in either Rebif®-treated group than was observed in the placebo group.

Immunogenicity:
As with all therapeutic proteins, there is a potential for immunogenicity. Serum Nab were detected in 31% and 24% of Rebiff*-treated patients at the 22 mcg and 44 mcg tiw dose respectively at one or more times during Study 1. The clinical significance of the presence of NAb to Rebiff sunknown. Comparison of the incidence of antibodies to other products maybe misleading.

DOSAGE AND ADMINISTRATION

ADMINISTRATION
Dosages of Rebil® shown to
be safe and effective are 22
mcg and 44 mcg sc tiv.
Rebil® should be
administered, if possible, at
the same time (preferably in
the late afternoon or
evening) on the same three
days (e.g. Monday,
Wednesday, and Friday) at
least 48 hours apart each
week. Generally, patients least 48 hours apart each week. Generally, patients should be started at 20% of the prescribed dose and increased over a 4-week period to the targeted dose, either 22 mcg or 44 mcg sc tiw. Leukopenia or elevated liver function tests may necessitate dose reduction or discontinuation of Rebiffordministration until toxicity is resolved.

Rebif® is intended for use under the guidance and supervision of a physician. It is recommended that physicians or qualified medical personnel train patients in the proper technique for self-administering subcutaneous injections using the pre-filled syringe. Patients should be advised to rotate sites for sc injections. Concurrent use of analgesics and/or antipyretics may help ameliorate flu-like symptoms on treatment days. Rebif® should be inspected visually for particulate matter and discoloration prior to administration.

Rx only. Manufacturer: EMD Serono, Inc., Rockland, MA 02370

Co-marketed by: EMD Serono, Inc., Rockland, MA 02370 Pfizer, Inc., New York, NY 10017

Rebif is a registered trademark of EMD Serono, Inc. NS LifeLines is a service mark of EMD Serono, Inc. Avonex is a registered trademark of Biogen Idec. Revised: April 2007 07-19234

