44 PSYCHIATRY JULY 2010 • INTERNAL MEDICINE NEWS

Cataract Risk 15% Higher in Current SSRI Users

BY CALVIN GODFREY

FROM OPHTHALMOLOGY

Selective serotonin reuptake inhibitor use may lead to an increased risk of cataract development, a study by Canadian researchers suggests.

In a large nested case-control study, the researchers found that certain selective serotonin reuptake inhibitors (SSRIs) correlated with an increased likelihood of

cataract diagnosis and cataract surgery. "We don't want people to stop their medicine or switch," lead author Mahyar Etminan, Pharm.D., of the Vancouver Coastal Health Research Institute—said in an interview. "We don't want to stress the message of 'drug A increases risk over drug B' but rather the overall risk as a class effect. We want to shed light on cataracts as a possible side effect."

The study considered data from a co-

hort of 206,624 Quebec residents aged 65 or older. Researchers found that of the study's 18,784 cases diagnosed with cataracts, the 5.7% who were current SSRI users were 15% more likely to be diagnosed with cataracts or undergo cataract surgery than non-SSRI users.

Overall, the researchers found that it took an average of 656 days between the commencement of SSRI therapy and the diagnosis of cataracts

(Ophthalmology 2010;117:1251-5).

The study did not find a link between past SSRI use (individuals who had ceased taking SSRIs 30 days prior to diagnosis) and cataract development.

The findings come with some precedent. The use of amitriptyline, a tricyclic antidepressant, was linked to an increased risk of cortical cataracts in a study published in 2001 (Ophthalmology 2001;108:1670-4). Additionally, research conducted in animal models had shown an association between increased serotonin levels and cataract clouding.

Dr. Etminan and his colleagues were the first to specifically examine the catarogenic risks of SSRI use in humans, however.

While the study does not prove causation, its findings reveal an association between cataract complications and the use of the SSRIs fluvoxamine and paroxetine and venlafaxine, a serotonin and norepinephrine reuptake inhibitor. The results did not reveal a statistically significant association between cataract development and other commonly prescribed SSRIs.

In the study's findings, compared with those who never took SSRIs, current fluvoxamine users experienced the highest levels of cataract risk, increasing the likelihood of undergoing cataract surgery by 51% (rate ratio 1.51, after adjustment for variables such as gender, hypertension, and antihypertensive use). Users of paroxetine, according to the study, were 23% more likely to undergo surgery. Venlafaxine-which in 2007 was the 12th most-prescribed drug in the United States with 17 million prescriptions-increased likelihood of surgery 34%, the researchers wrote. In an interview, Dr. Etminan said that the variations between the individual drug results in this early study could be chalked up to random chance or measurement error.

Future studies are needed to determine the cataract risks of the individual agents, Dr. Etminan said.

The study's limitations, Dr. Etminan and his colleagues acknowledged, could not control for smoking histories or undiagnosed cataracts. The study investigators acknowledged that it "may have lacked adequate power to assess the risk of cataracts with all individual antidepressants."

Representatives from SSRI manufacturers Pfizer Inc. and GlaxoSmithKline responded by saying their companies would need more time to review the study before commenting.

Representatives from Jazz Pharmaceuticals Inc., which manufactures fluvoxamine, did not respond to requests for comment by press time.

Going forward, Dr. Etminan said he hoped that further research would help doctors identify an antidepressant that would offer a safe choice for, say, an elderly patient recovering from cataract surgery.

The researchers stated that they had no relevant conflicts of interest.

BYSTOLIC® (nebivolol) tablets Brief Summary of full Prescribing Information Initial U.S. Approval: 2007

INDICATIONS AND USAGE: Hypertension - BYSTOLIC is indicated for the treatment of hypertension *[see Clinical Studies (14.1)]*. BYSTOLIC may be used alone or in combination with other antihypertensive agents *[see Drug Interactions (7)]*.

CONTRAINDICATIONS: BYSTOLIC is contraindicated in the following conditions: Severe bradycardia; Heart block greater than first degree; Patients with cardiogenic shock; Decompensated cardiac failure; Sick sinus syndrome (unless a permanent pacemaker is in place); Patients with severe hepatic impairment (Child-Pugh >B); Patients who are hypersensitive to any component of this product.

WARNINGS AND PRECAUTIONS: Abrupt Cessation of Therapy - Do not abruptly discontinue BYSTOLIC therapy in patients with coronary artery disease. Severe exacerbation of angina, myocardial infarction and ventricular arrhythmias have been reported in patients with coronary artery disease following the abrupt discontinuation of therapy with β-blockers. Myocardial infarction and ventricular arrhythmias may occur with or without preceding exacerbation of the angina pectoris. Caution patients without overt coronary artery disease against interruption or abrupt discontinuation of therapy. As with other β-blockers, when discontinuation of BYSTOLIC is planned, carefully observe and advise patients to minimize physical activity. Taper BYSTOLIC over 1 to 2 weeks when possible. If the angina worsens or acute coronary insufficiency develops, restart BYSTOLIC promptly, at least temporarily. Angina and Acute Myocardial Infarction - BYSTOLIC was not studied in patients with angina pectoris or who had a recent MI. Bron-chospastic Diseases - In general, patients with bronchospastic diseases should not receive β-blockers. Anesthesia and Major Surgery - Because beta-blocker withdrawal has been associated with an increased risk of MI and chest pain, patients already on beta-blockers should generally continue treatment throughout the perioperative period. If BYSTOLIC is to be continued perioperatively, monitor patients closely when anesthetic agents which depress myocardial function, such as ether, cyclopropane, and trichloroethylene, are used. If β-blocking therapy is withdrawn prior to major surgery, the impaired ability of the heart to respond to reflex adrenergic stimuli may augment the risks of general anesthesia and surgical procedures. The β-blocking effects of BYSTOLIC can be reversed by β-agonists, e.g., dobutamine or isoproterenol. However, such patients may be subject to protracted severe hypotension. Additionally, difficulty in restarting and maintaining the heartbeat has been reported with β-blockers. The β-blockers may m

in patients with hypertension and in patients with heart failure. The observed adverse reaction profile was consistent with the pharmacology of the drug and the health status of the patients in the clinical trials. Adverse reactions reported for each of these patient populations are provided below. Excluded are adverse reactions considered too general to be informative and those not reasonably associated with the use of the drug because they were associated with the condition being treated or are very common in the treated population. The data described below reflect worldwide clinical trial exposure to BYSTOLIC in 6545 patients, including 5038 patients treated for hypertension and the remaining 1507 subjects treated for other cardiovascular diseases. Doses ranged from 0.5 mg to 40 mg. Patients received BYSTOLIC for up to 24 months, with over 1900 patients treated for at least 6 months, and approximately 1300 patients for more than one year. HYPERTENSION: In placebo-controlled clinical trials comparing BYSTOLIC with placebo, discontinuation of therapy due to adverse reactions was reported in 2.8% of patients treated with nebivolol and 2.2% of patients given placebo. The most common adverse reactions that led to discontinuation of BYSTOLIC were headache (0.4%), nausea (0.2%) and bradycardia (0.2%). Table 1 list treatment-emergent adverse reactions that were reported in three 12-week, placebo-controlled monotherapy trials involving 1597 hypertensive patients treated with either 5 mg, 10 mg, or 20-40 mg of BYSTOLIC and 205 patients given placebo and for which the rate of occurrence was at least 1% of patients treated with nebivolol and greater than the rate for those treated with placebo in at least one dose group. Table 1. Treatment-Emergent Adverse Reactions with an Incidence (over 6 weeks) = 1% in BYSTOLIC-treated Patients and at a Higher Frequency than Placebo in at least one dose group. Table 1. Treatment-Emergent Adverse Reactions with an Incidence of the patients and the product of the patients and the pati

These adverse reactions have been chosen for inclusion due to a combination of seriousness, frequency of reporting or potential causal connection to BYSTOLIC. Adverse reactions common in the population have generally been omitted. Because these adverse reactions were reported voluntarily from a population of uncertain size, it is not possible to estimate their frequency or establish a causal relationship to BYSTOLIC exposure: abnormal hepatic function (including increased AST, ALT and bilirubin), acute pulmonary edema, acute renal failure, atrioventricular block (both second- and third-degree), bronchospasm, erectile dysfunction, hypersensitivity (including urticaria, allergic vasculitis and rare reports of angioedema), myocardial infarction, pruritus, psoriasis, Raynaud's phenomenon, peripheral ischemia/claudication, somnolence, syncope, thrombocytopenia, various rashes and skin disorders, vertigo, and vomiting.

DRUG INTERACTIONS: CYP2D6 Inhibitors - Use caution when BYSTOLIC is co-administered with CYP2D6 inhibitors (quinidine, propafenone, fluoxetine, paroxetine, etc.) [see Clinical Pharmacology (12.5)]. Hypotensive Agents - Do not use BYSTOLIC with other β-blockers. Closely monitor patients receiving catecholamine-depleting drugs, such as reserpine or guanethidine, because the added β-blocking action of BYSTOLIC may produce excessive reduction of sympathetic activity. In patients who are receiving BYSTOLIC and clonidine, discontinue BYSTOLIC several days before the gradual tapering of clonidine. Digitalis Glycosides - Both digitalis glycosides and β-blockers slow atrioventricular conduction and decrease heart rate. Concomitant use can increase the risk of bradycardia. Calcium Channel Blockers - BYSTOLIC can exacerbate the effects of myocardial depressants or inhibitors of AV conduction, such as certain calcium antagonists (particularly of the phenylalkylamine [verapamil] and benzothiazepine [diltiazem] classes), or antiarrhythmic agents, such as disopyramide.

classes), or antiarrhythmic agents, such as disopyramide.

USE IN SPECIFIC POPULATIONS: Pregnancy: Teratogenic Effects, Category C - Decreased pup body weights occurred at 1.25 and 2.5 mg/kg in rats, when exposed during the perinatal period (late gestation, parturition and lactation). At 5 mg/kg and higher doses (1.2 times the MRHD), prolonged gestation, dystocia and reduced maternal care were produced with corresponding increases in late fetal deaths and stillbirths and decreased birth weight, live litter size and pup survival. Insufficient numbers of pups survived at 5 mg/kg to evaluate the offspring for reproductive performance. In studies in which pregnant rats were given nebivolol during organogenesis, reduced fetal body weights were observed at maternally toxic doses of 20 and 40 mg/kg/day (5 and 10 times the MRHD), and small reversible delays in sternal and thoracic ossification associated with the reduced fetal body weights and a small increase in resorption occurred at 0 mg/kg/day (10 times the MRHD). No adverse effects on embryo-fetal viability, sex, weight or morphology were observed in studies in which nebivolol was given to pregnant rabbits at doses as high as 20 mg/kg/day (10 times the MRHD). Labor and Delivery - Nebivolol caused prolonged gestation and dystocia at doses ≥ 5 mg/kg in rats (1.2 times the MRHD). These effects were associated with increased fetal deaths and stillborn pups, and decreased birth weight, live litter size and pup survival rate, events that occurred only when nebivolol was given during the perinatal period (late gestation, parturition and lactation). No studies of nebivolol were conducted in pregnant women. Use BYSTOLIC during pregnancy only if the potential benefit justifies the potential risk to the fetus. Nursing Mothers - Studies in rats have shown that nebivolol or its metabolities cross the placental barrier and are excreted in breast milk. It is not known whether this drug is excreted in human milk. Because of the potential for β-blockers to produce serious adverse

OVERDOSAGE: In clinical trials and worldwide postmarketing experience there were reports of BYSTOLIC overdose. The most common signs and symptoms associated with BYSTOLIC overdosage are bradycardia and hypotension. Other important adverse reactions reported with BYSTOLIC overdosage are bradycardia and hypotension. Other important adverse reactions reported with BYSTOLIC overdose include cardiac failure, dizziness, hypoglycemia, fatigue and vomiting. Other adverse reactions associated with β-blocker overdose include bronchospasm and heart block. The largest known ingestion of BYSTOLIC worldwide involved a patient who ingested up to 500 mg of BYSTOLIC along with several 100 mg tablets of acetylsalicylic acid in a suicide attempt. The patient experienced hyperhidrosis, pallor, depressed level of consciousness, hypokinesia, hypotension, sinus bradycardia, hypoglycemia, hypokalemia, respiratory failure, and vomiting. The patient recovered. Because of extensive drug binding to plasma proteins, hemodialysis is not expected to enhance nebivolol clearance. If overdose occurs, provide general supportive and specific symptomatic treatment. Based on expected pharmacologic actions and recommendations for other β-blockers, consider the following general measures, including stopping BYSTOLIC, when clinically warranted: Bradycardia: Administer IV atropine. If the response is inadequate, isoproterenol or another agent with positive chronotropic properties may be given cautiously. Under some circumstances, transthoracic or transvenous pacemaker placement may be necessary. Hypotension: Administer IV fluids and vasopressors. Intravenous glucagon may be useful. Heart Block (second- or third-degree): Monitor and treat with isoproterenol infusion. Under some circumstances, transthoracic or transvenous pacemaker placement may be necessary. Congestive Heart Failure: Initiate therapy with digitalis glycosides and diuretics. In certain cases, consider the use of inotropic and vasodilating agents. Bronchospasm: Administer Iv office. Pro

Forest Pharmaceuticals, Inc. Subsidiary of Forest Laboratories, Inc. St. Louis, MO 63045, USA Licensed from Mylan Laboratories, Inc. Under license from Janssen Pharmaceutica N.V., Beerse, Belgium

Rev. 02/10 © 2010 Forest Laboratories, Inc.