she added.

Sulfonylureas May Elevate Risk in MI Survivors

BY BRUCE JANCIN

BARCELONA — Four widely prescribed oral sulfonylurea drugs are associated with significantly increased risk of all-cause mortality compared with metformin in type 2 diabetic patients having a history of MI, according to a comprehensive Danish national cohort study.

The study included all Danish adults with a prior MI who started on oral glu-

cose-lowering monotherapy during 1997-2006. The conclusion: Glimepiride, glyburide, glipizide, and tolbutamide were associated with 33%-43% higher mortality risk than was metformin, Dr. Tina Ken Schramm said at the annual congress of the European Society of Cardiology.

In contrast, single-agent gliclazide and repaglinide had all-cause mortality risks similar to metformin.

"We believe that metformin in general should be part of the treatment of type 2 diabetes to reduce mortality, but gliclazide and repaglinide may be good alternatives," said Dr. Schramm of the Heart Center at Copenhagen University National Hospital.

Metformin deserves the nod as the first-line agent on the basis of the results of the landmark United Kingdom Prospective Diabetes Study, which con-

Table 2. Treatment-Emergent Adverse Reaction Incidence in Placebo-Controlled Trials in Fibromyalgia Patients (Events Occurring in at Least 2% of All Savella-Treated Patients and Occurring More Frequently in Either Savella Treatment Group Than in the Placebo Treatment

aroup/(common/				
System Organ Class- Preferred Term	Savella 100 mg/day (n = 623) %	Savella 200 mg/day (n = 934) %	All Savella (n = 1557) %	Placebo (n = 652) %
Vascular Disorders				
Hot flush	11	12	12	2
Hypertension	7	4	5	2
Flushing	2	3	3	1

Weight Changes-In placebo-controlled fibromyalgia clinical trials, patients treated with Savella for up to 3 months experienced a mean weight loss of approximately 0.8 kg in both the Savella 100 mg/day and the Savella 200 mg/day treatment groups, compared with a mean weight loss of approximately 0.2 kg 3 months experienced a mean weight loss of approximately 0.8 kg in both the Savella 100 mg/day and the Savella 200 mg/day treatment groups, compared with a mean weight loss of approximately 0.2 kg in placebo-treated patients. **Genilourinary Adverse Reactions in Males**-In the placebo-controlled fibromyalgia studies, the following treatment-emergent adverse reactions related to the genitourinary system were observed in at least 2% of male patients treated with Savella, and occurred at a rate greater than in placebo-treated male patients: dysuria, ejaculation disorder, erectile dysfunction, ejaculation failure, libido decreased, prostatitis, scrotal pain, testicular pain, testicular swelling, urinary hesitation, urinary retention, urethral pain, and urine flow decreased. **Other Adverse Reactions Observed During Clinical Trials of Savella in Fibromyalgia**-Following is a list of frequent (those occurring on one or more occasions in at least 1/100 patients) treatment-emergent adverse reactions reported from 1824 fibromyalgia patients treated with Savella for periods up to 68 weeks. The listing does not include those events already listed in Table 2, those events for which a drug cause was remote, those events which were so general as to be uninformative, and those events reported only once which did not have a substantial probability of being acutely life threatening. Adverse reactions are categorized by body system and listed in order of decreasing frequency. Adverse reactions for alarrhea, dyspepsia, gastro-esophageal reflux disease, flatulence, abdominal distension; General Disorders – fatigue, peripheral edema, irritability, pyrexia, Infections – urinary tract infection, cystitis; Injury, Poisoning, and Procedural Complications – contusion, fall; Investigations – weight decreased or increased; Metabolism and Nutrition Disorders – depression, stress; Skin Disorders – night sweats **Postmarketing Spontaneous Reports**. The following additional adverse reactions have been chosen for inclusion because of a combination combination of seriousness, frequency of reporting, or potential causal connection to Savella. However, because these adverse reactions were reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure. These events include: Blood and Lymphatic System Disorders – leukopenia, neutropenia, thrombocy-topenia; Cardiac Disorders – supraventricular tachycardia; Eye Disorders – accommodation disorder; Endocrine Disorders – nyperprolactinemia; Hepatobiliary Disorders – hepatitis; Metabolism and Nutri-tion Disorders – anorexia, hyponatremia; Musculoskeletal and Connective Tissue Disorders – rhabdomyolysis; Nervous System Disorders – convulsions (including grand mal). loss of consciousness, Parkinsonism; Psychiatric Disorders – delirium, hallucination; Renal and Urinary Disorders – acute renal failure, urinary retention; Reproductive System and Breast Disorders – hypertensive crisis **DBUG INTERACTIONS**; Minacioran undergoes minimal CVP450 related metabolism, with the majority of

erythema multiforme, Stevens Johnson syndrome; Vascular Disorders – hypertensive crisis DRUG INTERACTIONS: Milnacipran undergoes minimal CYP450 related metabolism, with the majority of the dose excreted unchanged in urine (55%), and has a low binding to plasma proteins (13%). In vitro and in vivo studies showed that Savella is unlikely to be involved in clinically significant pharmacokinetic drug interactions [see Pharmacokinetics in Special Populations]. Clinically Important Interactions with Other Drugs-Lithium: Serotonin syndrome may occur when lithium is co-administered with Savella and with other drugs that impair metabolism of serotonin [see Warnings and Precautions – Serotonin Syndrome or Neuroleptic Malignant Syndrome (MMS)-Like Reactions]. Epinephrine and norepinephrine: Savella inhibits the reuptake of norepinephrine. Therefore concomitant use of Savella with epinephrine and norepinephrine may be associated with paroxysmal hypertension and possible arrhythmia [see Warnings and Precautions – Effects on Blood Pressure and Effects on Heart Rate] Serotoneria [clinically clinically in hypertension and administration of Savella with other inhibitors of serotonin re-uptake may result in hypertension and and Precautions – Effects on Blood Pressure and Effects on Heart Rate] Serotonergic Drugs: Co-administration of Savella with other inhibitors of serotonin re-uptake may result in hypertension and coronary artery vasoconstriction, through additive serotonergic effects [see Warnings and Precautions]. Digoxin: Use of Savella concomitantly with digoxin may be associated with potentiation of adverse hemodynamic effects. Postural hypotension and tachycardia have been reported in combination therapy with intravenously administered digoxin (1 mg). Co-administration of Savella and intravenous digoxin should be avoided [see Warnings and Precautions] Clonidine: Because Savella inhibits norepinephrine reuptake, co-administration with clonidine may inhibit clonidine's anti-hypertensive effect. Clomipramine: In a drug-drug interaction study, an increase in euphoria and postural hypotension was observed in patients who switched from clomipramine to Savella. CNS-active drugs: Given the primary CNS effects of Savella, caution should be used when it is taken in combination with other centrally acting drugs, including those with a similar mechanism of action. Monoamine Oxidase Inhibitors (MAOIs): [see Contraindications]. Contraindicationel

Contraindications). USE IN SPECIFIC POPULATIONS: Pregnancy-Pregnancy Category C. Milnacipran increased the incidence of dead fetuses in utero in rats at doses of 5 mg/kg/day (0.25 times the MRHD on a mg/m² basis). Administration of milnacipran to mice and rabbits during the period of organogenesis did not result in embryotoxicity or teratogenicity at doses up to 125 mg/kg/day in mice (3 times the maximum recom-mended human dose [MRHD] of 200 mg/day on a mg/m² basis). In rabbits, the incidence of the skeletal variation (6 times the MRHD of 200 mg/day on a mg/m² basis). In rabbits, the incidence of the skeletal variation actra signal in was increased following administration of milacore at 15 mg/kg/day during the period Actra single rib, was increased following administration of milinacipran at 15 mg/kg/day during the period of organogenesis. There are no adequate and well-controlled studies in pregnant women. Savella should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. <u>Nonterato-genic Effects</u>; Neonates exposed to dual reuptake inhibitors of serotonin and norepinephrine, or selective serotonin reuptake inhibitors late in the third trimester have developed complications requiring prolonged hospitalization, respiratory support, and tube feeding. Such complications can arise immediately upon delivery. Reported clinical findings have included respiratory distress, cyanosis, apnee, seizures, temperature instability, feeding difficulty, vomiting, hypoglycemia, hypotonia, hypertonia, hyperreflexia, tremor, jitteriness, irritability, and constant crying. These features are consistent with either a direct toxic effect of these classes of drugs or, possibly, a drug discontinuation syndrome. It should be noted that, in some cases, the clinical picture is consistent with serotonin syndrome [see Warnings and Precautions]. In rats, a decrease in pup body weight and viability on postpartum day 4 were observed when milnacipran, at a dose of 5 mg/kg/day (approximately 0.2 times the MRHD on a mg/m² basis), was administered orally to rats during late gestation. The no effect dose for maternal and offspring toxicity was 2.5 mg/kg/day (approximately 0.1 times the MRHD on a mg/m² basis). Labor and Delivery-The effect of milacipran on labor and delivery is unknown. The use of Savella during labor and delivery is not recommended. **Nursing Mothers**. There are no adequate and well-controlled studies in nursing mothers. It is not known fin linacipran is excreted in hreast milk. Because many drugs are excreted in human milk and because of known if milnacipran is excreted in human milk. Studies in animals have shown that milnacipran or its metabolites are excreted in breast milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from milnacipran, a decision should be made whether to discontinue the drug, taking into account the importance of the drug to the mother. Because the safety of Savella in infants is not known, nursing while on Savella is not recommended. **Pediatric Use**-Safety and effectiveness of Savella in a fibromyalgia pediatric population below the age of 17 have not been established [see Box Warning and Warnings and Precautions]. The use of Savella is not recommended in pediatric patients. **Geriatric Use**-in controlled clinical studies of Savella, 402 patients were 60 years or older, and no overall differences in safety and efficacy were observed between these patients and younger patients. In view of the predominant excretion of unchanged milnacipran via kidneys and the expected decrease in renal function with age renal function should be considered prior to use of Savella in he elderly [see Dosage and Administration]. SNRIs, SSRIs, and Savella in have been to use of Savella in the elderly [see Dosage and Administration]. SNRs, SSRs, sna Savella, have been associated with cases of clinically significant hyponatremia in elderly patients, who may be at greater risk for this adverse event [see Warnings and Precautions].

DRUG ABUSE AND DEPENDENCE: Controlled Substance - Milnacipran is not a controlled substance Abuse-Milnacipran did not produce behavioral signs indicative of abuse potential in animal or human studies. Dependence-Milnacipran produces physical dependence, as evidenced by the emergence of withdrawal symptoms following drug discontinuation, similar to other SNRIs and SSRIs. These withdrawal symptoms can be severe. Thus, Savella should be tapered and not abruptly discontinued after extended use [see Discontinuation of Treatment with Savella].

OVERDOSAGE: There is limited clinical experience with Savella overdose in humans. In clinical trials cases of acute ingestions up to 1000 mg, alone or in combination with other drugs, were reported with cases of actue ingestions up to 1000 mg, alone of in combination with other drugs, were reported with none being fatal. In postmarketing experience, fatal outcomes have been reported for acute overdoses primarily involving multiple drugs but also with Savella only. The most common signs and symptoms included increased blood pressure, cardio-respiratory arrest, changes in the level of consciousness (ranging from somnolence to coma), confusional state, dizziness, and increased hepatic enzymes. Management of Overdose-There is no specific antidote to Savella, but if serotonin syndrome ensues, available to the with expected diagender of the temperature acorton way hopencient (and the unit of the temperature acorton) way hopencient (and the unit of the temperature acorton). Management of overouse inter is no specific antidote to Savelia, out in serotioning and the ensues, specific treatment (such as with cyproheptadine and/or temperature control) may be considered. In case of acute overdose, treatment should consist of those general measures employed in the management of overdose with any drug. An adequate airway, oxygenation, and ventilation should be assured and cardiac rhythm and vital signs should be monitored. Induction of emesis is not recommended. Gastric lavage with a large-bore orogastric tube with appropriate airway protection, if needed, may be indicated if performed soon after ingestion or in symptomatic patients. Because there is no specific antidote for Savella, symptometic patients. soon and ingestion of in symptomatic patients. Because there is no specific and other of savena, symptomatic care and treatment with gastric lavage and activated charcoal should be considered as soon as possible for patients who experience a Savella overdose. Due to the large volume of distribution of this drug, forced diuresis, dialysis, hemoperfusion, and exchange transfusion are unlikely to be beneficial. In managing overdose, the possibility of multiple drug involvement should be considered. The physician should consider contacting a poison control center for additional information on the treatment of any overdose. Telephone numbers for certified poison control centers are listed in the Physicians' Desk Reference (PDR).

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'We believe that metformin in general should be part of the treatment of type 2 diabetes to reduce mortality.

DR. SCHRAMM

glyburide (13%), glipizide and gliclazide (7% each), tolbutamide (6%), and repaglinide (2%). Acarbose was prescribed as monotherapy in only 44 patients nationwide-far too small a number to allow meaningful results. Similarly, the thiazolidinediones were used too seldom to draw any conclusions, Dr. Schramm explained.

vincingly established the drug as the

safest glucose-lowering agent available,

Out of the total Danish population of

roughly 4.1 million, 107,870 type 2 dia-

betic individuals initiated monotherapy

with a glucose-lowering agent during

the 9-year study period. Among them

were 9,135 with a prior MI, who formed

Glimepiride was the most widely prescribed of the glucose-lowering medications in Denmark, being used by 43% of

subjects. Next came metformin (32%),

the population for this study.

Metformin served as the comparator in determining all-cause mortality risks for the other oral glucose-lowering agents in a multivariate analysis. Dr. Schramm reported having no financial conflicts of interest.

History of Foot **Ulcers** Predictive In the Elderly

history of foot ulcers is associated Awith an increased risk of mortality among community-dwelling adults and elderly people with diabetes, according to a 10-year follow-up of the Nord-Trøndelag Health Study (HUNT 2).

The risk persisted after adjustment for comorbidity and depression scores, suggesting the need for close monitoring, said Marjolein M. Iversen of Bergen (Norway) University, and associates.

The study included 155 diabetic persons with a history of foot ulcers, 1,339 diabetic persons without a history of foot ulcers, and 63,632 nondiabetic persons, followed for 10 years with mortality as the end point (Diabetes Care 2009 Sept. 3 [doi: 10.2337/dc09-0651]).

In a Cox regression analysis, having a history of foot ulcers was associated with more than a twofold (2.29 [95% confidence interval 1.82-2.88]) hazard risk for mortality, versus those in the nondiabetic group, and a 47% increase in mortality, versus diabetic persons without a history of foot ulcers.

-Caroline Helwick

