Sibutramine's Weight Maintenance Lasts 2 Years

BY JANE SALODOF MACNEIL

Senior Editor

PHOENIX — Sibutramine helped people who had lost substantial amounts of weight maintain much of their weight loss for 2 years in a multicenter, doubleblind, randomized trial. By 3 years, however, they had regained about as much as a control group on placebo.

In all, 466 participants started the maintenance study after losing an average of 30 kg in nonpharmacologic weight-loss programs. Although everyone had begun to put the pounds back on before entering the trial, they had to have maintained at least half of their peak weight loss for 6 months, said Dr. James W. Anderson, the lead investigator, reported in an analysis at the annual meeting of The Obesity Society.

Indeed, the net weight gain for the two groups was significantly less with sibutramine (Meridia) for the first 2 years-0.6% vs. 3.1% at 6 months, 2.8% vs. 5.6% at 1 year, and 7.1% vs. 9.8% at 24 months. At 3 years, the sibutramine group still had less net weight gain (9.6% vs. 11.7%), but the difference was no longer significant, and both groups appeared to plateau, said Dr. Anderson of the University of Kentucky, Lexington.

Abbott Laboratories became the study sponsor after acquiring sibutramine and decided to curtail follow-up at 36 months, according to Dr. Anderson, who emphasized that he has not received support from Abbott for more than 5 years.

The trial randomized 236 subjects to 15 mg of sibutramine per day and 230 to placebo at 26 sites. Of these, 150 placebo patients and 144 sibutramine patients completed 36 months. Blood pressure and pulse rate were significantly higher with sibutramine. Serious adverse events were more common with placebo (19% vs. 17%), but more people discontinued because of adverse events on sibutramine (15% vs. 7%).



insulin detemir (rDNA origin) injection

Rx ONLY BRIEF SUMMARY. Please see package insert for

INDICATIONS AND USAGE

VEVENIR is indicated for once- or twice-daily subcutaneous dministration for the treatment of adult and pediatric patients with type 1 diabetes mellitus or adult patients with type 2 iabetes mellitus who require basal (long acting) insulin for the patient of the meaning of the control of the programming. control of hyperglycemia

WARNINGS
Hypoglycemia is the most common adverse effect of insulin therapy, including LEVEMIR. As with all insulins, the timing of hypoglycemia may differ among various insulin formulations.

Glucose monitoring is recommended for all patients with diabetes.

Any change of insulin dose should be made cautiously and only under medical supervision. Changes in insulin strength, timing of dosing, manufacturer, type (e.g., regular, NPH, or insulin analogs), species (animal, human), or method of manufacture (rDNA versus animal-source insulin) may result in the need for a change in dosage. Concomitant oral antidiabetic treatment may need to be adjusted.

PRECAUTIONS

Inadequate dosing or discontinuation of treatment may lead to hyperglycemia and, in patients with type 1 diabetes, diabetic ketoacidosis. The first symptoms of hyperglycemia usually occur gradually over a period of hours or days. They include nausea, vomiting, drowsiness, flushed dry skin, dry mouth, increased urination, thirst and loss of appetite as well as acetone breath. Untreated hyperglycemic events are potentially fatal.

Untreated nypergiycemic events are potentially tatal. LEVEMIR is not intended for intravenous or intramuscular administration. The prolonged duration of activity of insulin detemir is dependent on injection into subcutaneous tissue. Intravenous administration of the usual subcutaneous dose could result in severe hypoglycemia. Absorption after intramuscular administration is both faster and more extensive than absorption after subcutaneous administration.

LEVEMIR should not be diluted or mixed with any other insulin preparations (see PRECAUTIONS, Mixing of Insulins)

Insulin may cause sodium retention and edema, particularly if previously poor metabolic control is improved by intensified

As with all insulin preparations, the time course of LEVEMIR action may vary in different individuals or at different times in the same individual and is dependent on site of injection, blood supply, temperature, and physical activity.

Adjustment of dosage of any insulin may be necessary if patients change their physical activity or their usual meal plan.

Hypoglycemia
As with all insulin preparations, hypoglycemic reactions may be associated with the administration of LEVEMIR. Hypoglycemia is the most common adverse effect of insulins. Early warning symptoms of hypoglycemia may be different or less pronounced under certain conditions, such as long duration of diabetes, diabetic nerve disease, use of medications such as beta-blockers, or intensified diabetes control (see PRECAUTIONS, Drug Interactions). Such situations may result in severe hypoglycemia. Interactions). Such situations may result in severe hypoglycemia (and, possibly, loss of consciousness) prior to patients' awareness of hypoglycemia.

of hypoglycemia.

The time of occurrence of hypoglycemia depends on the action profile of the insulins used and may, therefore, change when the treatment regimen or timing of dosing is changed. In patients being switched from other intermediate or long-acting insulin preparations to once- or twice-daily LEVEMIR, dosages can be prescribed on a unit-to-unit basis; however, as with all insulin preparations, dose and timing of administration may need to be adjusted to reduce the risk of hypoglycemia.

Renal Impairment

Renal Impairment
As with other insulins, the requirements for LEVEMIR may need to be adjusted in patients with renal impairment.

Hepatic ImpairmentAs with other insulins, the requirements for LEVEMIR may need to be adjusted in patients with hepatic impairment.

to be adjusted in patients with nepatic impairment.

Injection Site and Allergic Reactions
As with any insulin therapy, lipodystrophy may occur at the injection site and delay insulin absorption. Other injection site reactions with insulin therapy may include redness, pain, itching, hives, swelling, and inflammation. Continuous rotation of the injection site within a given area may help to reduce or prevent these reactions. Reactions usually resolve in a few days to a few

weeks. On rare occasions, injection site reactions may require discontinuation of LEVEMIR.

In some instances, these reactions may be related to factors other than insulin, such as irritants in a skin cleansing agent or poor injection technique.

Systemic allergy: Generalized allergy to insulin, which is less common but potentially more serious, may cause rash (including pruritus) over the whole body, shortness of breath, wheezing, reduction in blood pressure, rapid pulse, or sweating. Severe cases of generalized allergy, including anaphylactic reaction, may he life-threatening.

Intercurrent Conditions
Insulin requirements may be altered during intercurrent conditions such as illness, emotional disturbances, or other stresses.

stresses.

Information for Patients
LEVEMIR must only be used if the solution appears clear and colorless with no visible particles. Patients should be informed about potential risks and advantages of LEVEMIR therapy, including the possible side effects. Patients should be offered continued education and advice on insulin therapies, injection technique, life-style management, regular glucose monitoring, periodic glycosylated hemoglobin testing, recognition and management of hypo- and hyperglycemia, adherence to meal planning, complications of insulin therapy, timing of dosage, instruction for use of injection devices and proper storage of insulin. Patients should be informed that frequent, patient-performed blood glucose measurements are needed to achieve effective glycemic control to avoid both hyperglycemia and hypoglycemia. Patients must be instructed on handling of special situations such as intercurrent conditions (Illness, stress, hypoglycemia. Patients must be instructed on handling of special situations such as intercurrent conditions (illness, stress, or emotional disturbances), an inadequate or skipped insulin dose, inadvertent administration of an increased insulin dose, inadequate food intake, or skipped meals. Refer patients to the LEVEMIR "Patient Information" circular for additional informat

As with all patients who have diabetes, the ability to concentrate and/or react may be impaired as a result of hypoglycemia or hyperglycemia Patients with diabetes should be advised to inform their health care professional if they are pregnant or are contemplating pregnancy (see PRECAUTIONS, Pregnancy).

Laboratory Tests
As with all insulin therapy, the therapeutic response to LEVEMIR
should be monitored by periodic blood glucose tests. Periodic
measurement of HbA_t, is recommended for the monitoring of
long-term glycemic control.

Drug Interactions

A number of substances affect glucose metabolism and may require insulin dose adjustment and particularly close monitoring.

The following are examples of substances that may reduce the blood-glucose-lowering effect of insulin: corticosteroids, danazol, diuretics, sympathomimetic agents (e.g., epinephrine, albuterol, terbutaline), isoniazid, phenothiazine derivatives, somatropin, thyroid hormones, estrogens, progestogens (e.g., in oral contraceptives).

The following are examples of substances that may increase the blood-glucose-lowering effect of insulin and susceptibility to hypoglycemia: oral antidiabetic drugs, ACE inhibitors, disopyramide, fibrates, fluoxetine, MAO inhibitors, propoxyphe salicylates, somatostatin analog (e.g., octreotide), and sulfonamide antibiotics.

Beta-blockers, clonidine, lithium salts, and alcohol managementiate or weaken the blood-glucose-lowering eff potentiate or weaken the blood-glucose-lowering effect of insulin. Pentamidine may cause hypoglycemia, which may sometimes be followed by hyperglycemia. In addition, under the influence of sympatholytic medicinal products such as beta-blockers, clonidine, guanethidine, and reserpine, the signs of hypoglycemia may be reduced or absent.

The results of *in-vitro* and *in-vivo* protein binding studies demonstrate that there is no clinically relevant interaction bet insulin detemir and fatty acids or other protein bound drugs.

Mixing of Insulins

If LEVEMIR is mixed with other insulin preparations, the profi
of action of one or both individual components may change.
Mixing LEVEMIR with insulin aspart, a rapid acting insulin
analog, resulted in about 40% reduction in AUC_(0.2h) and C_{max} for insulin aspart compared to separate injections when the ratio of insulin aspart to LEVEMIR was less than 50%.

LEVEMIR should NOT be mixed or diluted with any other

Carcinogenicity, Mutagenicity, Impairment of Fertility
Standard 2-year carcinogenicity studies in animals have not
been performed. Insulin determir tested negative for genotoxic
potential in the *in-vitro* reverse mutation study in bacteria,
human peripheral blood lymphocyte chromosome aberration
test, and the *in-vivo* mouse micronucleus test.

Pregnancy: Teratogenic Effects: Pregnancy Category C

Pregnancy: Teratogenic Effects: Pregnancy Category C In a fertility and embryonic development study, insulin detemir was administered to female rats before mating, during mating, and throughout pregnancy at doses up to 300 nmol/kg/day (3 times the recommended human dose, based on plasma Area Under the Curve (AUC) ratio). Doses of 150 and 300 nmol/kg/da produced numbers of litters with visceral anomalies. Doses up to 900 nmol/kg/day (approximately 135 times the recommended human dose based on AUC ratio) were given to rabbits during organogenesis. Drug-dose related increases in the incidence of fetuses with gall bladder abnormalities such as small, bilobed, bifurcated and missing gall bladders were observed at a dose of 900 nmol/kg/day. The rat and rabbit embryofetal development studies that included concurrent human insulin control groups

indicated that insulin determir and human insulin had similar effects regarding embryotoxicity and teratogenicity.

Nursing mothers It is unknown whether LEVEMIR is excreted in significant amounts in human milk. For this reason, caution should be exercised when LEVEMIR is administered to a nursing mother. Patients with diabetes who are lactating may require adjustments in insulin dose, meal plan, or both.

Pediatric useIn a controlled clinical study, HbA_{1c} concentrations and rates of hypoglycemia were similar among patients treated with LEVEMIR and patients treated with NPH human insulin.

Geriatric use
Of the total number of subjects in intermediate and long-tern clinical studies of LEVENIR, 85 (type 1 studies) and 363 (type 2 studies) were 65 years and older. No overall differences in safety or effectiveness were observed between these subjects safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out. In elderly patients with diabetes, the initial dosing, dose increments, and maintenand dosage should be conservative to avoid hypoglycemic reactio Hypoglycemia may be difficult to recognize in the elderly.

Body as Whole: allergic reactions (see PRECAUTIONS, Allergy) Skin and Appendages: lipodystrophy, pruritus, rash, Mild injection site reactions occurred more frequently with LEVEMIR than with NPH human insulin and usually resolved in a few days to a few weeks (see PRECAUTIONS, Allergy).

Hypoglycemia: (see WARNINGS and PRECAUTIONS).

In trials of up to 6 months duration in patients with type 1 and type 2 diabetes, the incidence of severe hypoglycemia with LEVEMIR was comparable to the incidence with NPH, and, as expected, greater overall in patients with type 1 diabetes (Table 4)

weight gain:
In trials of up to 6 months duration in patients with type 1 and type 2 diabetes, LEVEMIR was associated with somewhat less weight gain than NPH (Table 4). Whether these observed differences represent true differences in the effects of LEVEMIR and NPH insulin is not known, since these trials were not blinded and the protocols (e.g., diet and exercise instructions and monitoring) were not specifically directed at exploring hypotheses related to weight effects of the treatments compared. The clinical significance of the observed differences has not been established.

Safetv Information on Clinical Studies

	Treatment	# of subjects	Weight (kg)		<u>Hypoglycemia</u> (events/subject/month)	
			Baseline	End of treatment	Major*	Minor**
Type 1						
Study A	LEVEMIR	N=276	75.0	75.1	0.045	2.184
	NPH	N=133	75.7	76.4	0.035	3.063
Study C	LEVEMIR	N=492	76.5	76.3	0.029	2.397
	NPH	N=257	76.1	76.5	0.027	2.564
Study D	LEVEMIR	N=232	N/A	N/A	0.076	2.677
Pediatric	NPH	N=115	N/A	N/A	0.083	3.203
Type 2						
Study E	LEVEMIR	N=237	82.7	83.7	0.001	0.306
	NPH	N=239	82.4	85.2	0.006	0.595
Study F	LEVEMIR	N=195	81.8	82.3	0.003	0.193
	NPH	N=200	79.6	80.9	0.006	0.235

Major = requires assistance of another individual because of neurologic impairment
 Minor = plasma glucose <56 mg/dl, subject able to deal with the episode him/herself

OVERDOSAGE
Hypoglycemia may occur as a result of an excess of insulin relative to food intake, energy expenditure, or both. Mild episodes of hypoglycemia usually can be treated with oral glucose. Adjustments in drug dosage, meal patterns, or exercise may be needed. More severe episodes with coma, seizure, or neurologic impairment may be treated with intramuscular/ subcutaneous glucagon or concentrated intravenous glucose. After apparent clinical recovery from hypoglycemia, continued observation and additional carbohydrate intake may be necessary to avoid reoccurrence of hypoglycemia.

More detailed information is available on request.

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Liraglutide Beat Orlistat for Losing Weight

PHOENIX — Liraglutide, an investigational drug given once a day, produced significantly more weight loss than orlistat in a randomized, 20-week, placebo-controlled trial conducted in obese patients who for the most part were not diabetic.

Participants on four different doses of liraglutide—1.2, 1.8, 2.4, and 3.0 mg—tested in the study lost significantly more weight, compared with a control group on placebo. Those treated at the two highest doses (2.4 and 3.0 mg per day) lost significantly more weight than did those given 120 mg of orlistat (Xenical) three times a day.

The mean weight loss ranged from 4.8 kg with the lowest 1.2-mg dose of liraglutide to 7.2 kg with the 3.0-mg dose, according to the investigators. The mean weight loss for placebo was little more than 2 kg and about 4 kg with orlistat.

Dr. Arne Astrup, the lead author and head of the department of human nutrition at the University of Copenhagen, described results of the six-arm, 564-patient study in an oral presentation at the annual scientific meeting of NAASO, the Obesity Society. Novo Nordisk A/S sponsored the trial.

Nearly two-thirds of the participants in the current study did not have diabetes; most of the rest were classified with prediabetes, leaving about 3% with the disease. About three-quarters of the population were women, and the average age was in the mid-40s (range 18-65 years). Body mass index ranged from 30 kg/m² to 40 kg/m² and weight from about 96 kg to 98 kg. The protocol encouraged all of the participants to go on a reduced-calorie diet and engage in physical activity.

The proportion of participants who lost 5% or more of body weight was 44% with orlistat but ranged from 54% to 76% with the liraglutide doses; and 28% of those on the highest dose lost more than 10% of their body weight. The most common adverse events were nausea and vomiting.

Dr. Astrup disclosed being a consultant to Novo Nordisk and receiving support for serving on advisory boards relative to liraglutide. The investigators also included Novo Nordisk employees, one of whom was a shareholder in the company, and other scientists who had received financial support and/or served on advisory boards.

-Jane Salodof MacNeil