Advance Directives Often Go Undiscussed

BY ROBERT FINN

SAN FRANCISCO — A minority of patients with relapsed leukemia or lymphoma discussed advance directives with their physicians, according to a survey of 75 such patients at Virginia Commonwealth University in Richmond.

Although 35 (47%) of the patients had an advance directive, just 5 (7%) had ever discussed advance directives with

HUMALOG®

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a O INJECTION (rDNA ORIGIN) RY: Consult package insert for complete prescribing information INSULIN LISPRO INJ BRIEF SUMMARY: C

INDICATIONS AND USAGE: Humalog is an insulin analog that is indicated in the treatment of patients with diabetes mellitus for the control of hyperglycemia. Humalog has a more rapid onset and a shorter duration of action than regular human insulin. Therefore, in patients with type 1 diabetes, Humalog should be used in regimens that include a longer-acting insulin. However, in patients with type 2 diabetes, Humalog may be used without a longer-acting insulin when used in combination therapy with sulforylurea agents. Humalog may be used in an external insulin pump, but should not be difuted or mixed with any other insulin when used in the pump. Humalog administration in insulin pumps has not been studied in patients with type 2 diabetes.

CONTRAINDICATIONS: Humalog is contraindicated during episodes of hypoglycemia and in patients sensitive to Humalog or any of its excipients.

- Numany of any of its excipients. WARNINGS: This human insulin analog differs from regular human insulin by its rapid onset of action as well as a shorter duration of activity. When used as a mealtime insulin, the dose of Humalog should be given within 15 minutes before or immediately after the meal. Because of the short duration of action of Humalog, patients with type 1 diabetes also require a longer-acting insulin to maintain glucose control (except when using an external insulin pump). External Insulin Pumps: When used in an external insulin pump, Humalog should not be diluted or mixed with any other insulin, Patients should carefully read and follow the external insulin pump manufacturer's instructions and the "PATLENT INFORMATION" leafter before using Humalog. Physicians should carefully evaluate information on external insulin pump use in the Humalog physician package insert and in the external insulin pump manufacturer's instructions. If unexplained hyperglycemia or ketosis occurs during external insulin pump use, prompt identification and correction of the cause is necessary. The patient may require interim therapy with subcutaneous insulin injections (see PRECAUTIONS, For Patients Using External Insulin Fumpi, and DOSAGE AND ADMINISTRATION). Hypoglycemia is the most common adverse effect associated with the use of insulins, including Humalog. As with all insulins, the timping of hypoglycemia may differ among various insulin formulations. Glucose

Hypoglycemia is the most common adverse effect associated with the use of insulins, including Humalog As with all insulins, the timing of hypoglycemia may differ among various insulin formulations. Glucose monitoring is recommended for all patients with diabetes and is particularly important for patients using an external insulin pump. Any change of insulin should be made cautiously and only under medical supervision. Changes in insulin strength, manufacturer, type (eg, regular, NPH, analog), species, or method of manufacture may result in the need for a change in dosage.

PRECAUTIONS: General—Hypoglycemia and hypokalemia are among the potential clinical adverse effects associated with the use of all insulins. Because of differences in the action of Humalog and other insulins, care should be taken in patients in whom such potential side effects might be clinically relevant (eg, patients who are fasting, have autonomic neuropathy, or are using potassium-lowering drugs or patients taking drugs sensitive to serum potassium level). Lipodystrophy and hypersensitivity are among other potential clinical adverse effects associated with the use of all insulins. As with all insulins, the time course of Humalog 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.

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. Insulin requirements may be altered during illness, emotional disturbances, or other stress. **Hypoglycemia** — As with all insulin preparations, hypoglycemic reactions may be associated with the administration of Humalog. Rapid changes in serum glucose concentrations may be associated with the administration of Humalog. Rapid changes in serum glucose concentrations may induce symptoms of hypoglycemia in persons with diabetes, regardless of the glucose value. Early warning symptoms of adaetic nerve disease, use of medications such as beta-hockers, or intensified diabetes control. **Renal Impairment**—The requirements for insulin may be reduced in patients with renal impairment. **Hepatic Impairment**—Although impaired hepatic function does not affect the absorption or disposition of Humalog, careful glucose monitoring and dose adjustments of insulin, including Humalog, may be necessary. **Altergy**—Lacial.Altergy—As with any insulin threapy, patients may experience refness, swelling, or riching at the site of injection. These minor reactions usually resolve in a terv days to a few weeks. In some instances, these reactions may be related to factors other than insulin, is as kin cleansing agent or poor injection technique.

at the site of injection. These minor reactions usually résolve in a few days to a few weeks. 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. <u>Systemic Alleyce</u>—Less common, but potentially more serious, is generalized allergy to insulin, which may cause rash (including puritus) 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 be life-threatening. Localized reactions and generalized malaging have been reported with the use of cresol as an injectable excipient. In Humalog-controlled clinical trials, pruritus (with or without rash) was seen in 17 patients receiving Humalog. The starting Humalog (Hz 2944) (*P*=.053). <u>Antibody Production</u>—In large clinical trials, antibodies that cross-react with human insulin and insulin lispor were observed in both Humalog. Teatment groups. As expected, the largest increase in the antibody levels during the 12-month clinical trials was observed with patients new to insulin therapy. <u>Usage of Humalog in External Insulin Pumps</u>—The infusion set (reservoir syringe, tubing, and cathteer), Disetronic® D-TRON®²³ or D-TRONPlus²⁸²: cartridge adapter, and Humalog in the external insulin pump reservoir should be replaced and a new infusion site selected every 48 hours or Less. Humalog in the external insulin pump, the infusion set should be replaced and a new infusion site should

using Disetronic Rapid⁴⁹² infusion sets. The infusion set (reservoir syringe, tubing, catheter), D-TRON^{82,3} or D-TRONplus^{42,3} cartridge adapter, and Humalog in the external insulin pump reservoir should be replaced, and a new infusion site selected every 48 hours or less. Humalog in the external pump should not be exposed to temperatures above **37°C (98.6°F).** A Humalog 3 mL cartridoe used in the D-TRON^{62,3} or D-TRON^{62,4} or D-TRON^{64,4} or D-TRON⁶

C (98.6°F). A Humalog 3 mL cartridge used in the D-TRON®2.2 or D-TRONplus®2.3 pump should be discarded after 7 days, in if it still contains Humalog, Infusion sites that are erythematous, pruritic, or thickened should be reported to dical personnel, and a new site selected. Humalog should not be diluted or mixed with any other insulin when used in an external insulin pump. Laboratory Tests—As with all insulins, the therapeutic response to Humalog should be monitored by periodic od glucose tests. Periodic measurement of hemoglobin A1C is recommended for the monitoring of long-term comic control.

block glucose tests. Periodic Ineastientent of hemoglucin ArC is recommended to the monitoring of long-term *Drug Interactions*—Insulin requirements may be increased by medications with hyperglycemic activity, such as corticosteroids, sionitad, certain ligid-towering drugs (e.g., niacin), estrogens, oral contraceptives, phenothazines, and thyroid replacement therapy (see CLINICAL PHARMACOLOGY). Insulin requirements may be decreased in the presence of drugs that increase insulin sensitivity or have hypoglycemic activity, such as oral antidiabetic agents, salicylates, sulfa antibiotics, certain antidepressants (monoamine oxidase inhibitors), angiotensin-converting-enzyme inhibitors, angiotensin Il receptor blocking agents, beta-adrenergic blockers, inhibitors of pancreatic function (e.g. octrovide), and alcohol. Beta-adrenergic blockers may mask the symptoms of hypoglycemia in some patients. **Mixing of Insulins**—Care should be taken when mixing all insulins as a change in peak action may occur. The American Diabetes Association warns in its Position Statement on Insulin Administration, "On mixing, physiochemical changes in the mixture may occur (either immediately or over time). As a result, the physiological with Humulin® N or Humulim® U does not decrease the absorption rate or the total bioavailability of Humalog.

their oncologist, said Dr. Thomas J. Smith at the annual Oncology Congress

presented by Reed Medical Education. You would think that most of them would have had a discussion about advance directives, about code status with their doctor," said Dr. Smith, the Massey Endowed Professor of Palliative Care Research and Medicine at VCU. But "in only 2 out of 75 cases had the doctor ever brought it up."

In initial questioning, only 23% of the patients expressed a desire to discuss advance directives with their oncologist, according to the study. Yet 86% were willing to discuss advance directives with their admitting hospitalist—a physician they had never met before—and 95% thought it would be important to have this discussion.

In attempting to learn why many patients would not want to discuss this top-

Given alone or mixed with Humulin N, Humalog results in a more rapid absorption and glucose-lowering effect

Given alone or mixed with Humulin N, Humalog results in a more rapid absorption and glucose-lowering effect compared with regular human insulin.
 Pregnancy—Teratogenic Effects—Pregnancy Category B—Reproduction studies with insulin lispro have been performed in pregnant rats and rabits at parenterial doses up to 4 and 0.3 times, respectively, the average thrang that more that shall be average thrange that the transmitter of the start start and shall ogg. There are, however, no adequate and well-controlled studies with Humalog in pregnant women. Because animal reproduction studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed.
 Athough there are limited clinical studies of the use of Humalog in pregnancy, published studies with human insulins suggest that optimizing overall glycemic control, including postprandial control, before conception and during pregnancy improves fetal outcome. Although the fetal complications of maternal hyperglycemia have been well documented, fetal toxicity also has been reported with maternal hyperglycemia. Insulin requirements usually all during the first trimesters during the second and third timesters. Careful monitoring of the patient is required throughout pregnancy. During the perinatal period, careful monitoring of infants born to mothers with diabetes is warranted.
 Mursing Muthers—I is unknown whether Humalog is excreted in significant amounts in human milk. Many fungs, including human nisulin, are excreted in human milk. For this reason, caution should be exercised when future algoses. mer plan, or both.
 Prediatric Use—In a 9-month, crossover study of prepubescent children (n=60), aged 3 to 11 years, comparable glycemic control as measured by AIC was achieved regardless of treatment group: regular human insulin 30 minutes before meals 8.4%, Humalog immediately before meals 8.7%. The incidence of hypoglycemia distributes of the distributes of the dist

ADVERSE REACTIONS: Clinical studies comparing Humalog with regular human insulin did not demonstrate a difference in frequency of adverse events between the 2 treatments. Adverse events commonly associated with human insulin therapy include the following: Body as a Whole—allergic reactions (see PRECAUTIONS). Skia and Anonedrase...injection site reaction. Involvetophy purpties rach

Body as a Whole—allergic reactions (*see* PRECAUTIUNS). **Skin and Appendages**—injection site reaction, lipodystrophy, pruritus, rash. **Other**—hypoglycemia (*see* WARNINGS *and* PRECAUTIONS).

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 neurolo impairment may be treated with intramuscular/subcutaneous glucogon or concentrated intravenous glucose. Sustained carbohydrate intake and observation may be necessary because hypoglycemia may recur after ponarote objected processor. apparent clinical recovery

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Sustained carbohydrate intake and observation may be necessary because hypoglycemia may recur after apparent clinical recovery.
DOSAGE AND ADMINISTRATION: Humalog is intended for subcutaneous administration, including use in select external insulin pumps (see DOSAGE AND ADMINISTRATION, *External Insulin Pumps*). Dosage regimens of Humalog will vary among patients and should be determined by the healthcare provider familiar with the patient's metabolic needs, eating habits, and other lifestyle variables. Pharmacokinetic and pharmacodynamic studies showed Humalog to be equipotent to regular human insulin, but with more rapid activity. The quicker glucose-lowering effect as one unit of regular human insulin, but with more rapid activity. The quicker glucose-lowering effect as one unit of regular human insulin, but with more rapid activity. The quicker glucose-lowering effect as all insulin may be needed when a patient changes from other insulins to Humalog, patient's med as a meditime insulin, Humalog should be given within 15 minutes before or immediately after a meal. Regular human insulin given may need to be adjusted when using Humalog. The rate of insulin absorption and consequently the onset of activity are known to be affected by the site of function in longer-acting insulin being given may need to be adjusted when using Humalog. The rate of insulin absorption and consequently the onset of activity are known to be affected by the ther insulins. Humalog maintains its rapid onset of activity are known to be affected by the site of functions are higher train those following debtoid or has less variability in its onset of action amorg injection sites compared with regular human insulin (see PRECAUTIONS). After abdominal administration, Humalog is olightly shorter following abdominal injection. Compared with debtoid and fenoral injections. As with all insulin my be used at a metal that be serverably in different individuals or within the same simple with offer the solution and the inspected visually before use w

HOW SUPPLIED:

log (insulin lispro injection, USP [rDNA origin]) is available in t	he following package size:	s (with each
tion containing 100 units insulin lispro per mL [U-100]):		
0 mL vials	NDC 0002-7510-01	(VL-7510)
x 3 mL cartridges ³	NDC 0002-7516-59	(VL-7516)
x 3 mL disposable insulin delivery devices (Pen)	NDC 0002-8725-59	(HP-8725)
x 3 mL disposable insulin delivery devices (KwikPen™)	NDC 0002-8799-59	(HP-8799)

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. *Storage*—Unopened Humalog should be stored in a refrigerator (2° to 8°C (36° to 46°F)), but not in the zer. Do not use Humalog if it has been frozen. Unrefrigerated (below 30°C (86°F)) 12 vials, cartridges, Pens, I KwikPens must be used within 28 days or be discarded, even if they still contain Humalog. Protect from

and Knikerens must be used within 25 days or be discarded, even in they suit contain Humaiog. Protect from direct heat and light. *Use in an External insulin Pump*—A Humalog 3mL cartridge used in the D-TRON®2.3 or D-TRONplus®2.3 should be discarded after 7 days, even if it still contains Humalog. Infusion sets, D-TRON®2. and D-TRONplus®2. cartridge adapters, and Humalog in the external insulin pump reservoir should be discarded every 48 hours or less. Literature revised May 27, 2009

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Lucrature revised May 27, 2009 KwikPens manufactured by Eli Lilly and Company, Indianapolis, IN 46285, USA. Pens manufactured by Eli Lilly and Company, Indianapolis, IN 46285, USA or Lilly France, F-67640 Fegersheim, France. Vials manufactured by Eli Lilly and Company, Indianapolis, IN 46285, USA or Hospira, Inc., Lake Forest, IL 60045, USA or Lilly France, F-67640 Fegersheim, France. Cartridges manufactured by Lilly France, F-67640 Fegersheim, France for Eli Lilly and Company, Indianapolis, IN 46285, USA.

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ic with their oncologist, Dr. Smith and his colleagues questioned them closely. During this additional questioning, 48% of the patients said they would, in fact, prefer to discuss advance directives with their oncologist, and 36% said they would prefer to discuss this topic with their primary care physician. "They want to discuss it with ... the doctors who know them best," Dr. Smith said. "They're waiting for us to talk about it first."

But physicians are often reluctant to initiate this discussion for fear of upsetting the patient or causing the patient to give up hope. In fact, advance directives have never been associated with worse survival in any study, Dr. Smith said. One study even showed that advance directives were associated with a 2.2-fold better chance of survival in patients undergoing bone marrow transplants (J. Clin. Oncol. 2007;25:5643-8).

Other studies have shown that patients who are overly optimistic about survival receive worse end-of-life care. They are more likely to be resuscitated and to die in an ICU or on a ventilator, and less likely to die at home, where most patients would prefer to die. Excessive optimism was not associated with improved survival in this study (JAMA 2008;299:2667-78).

Dr. Smith serves on the speakers bureau for Medtronic Inc., and has received honoraria from UnitedHealthcare and grant support from the National Cancer Institute and the National Library of Medicine. Reed Medical Education and this news organization are owned by Elsevier.

INDEX OF ADVERTISERS

AstraZeneca LP. Symbicort	59-62
Boehringer Ingelheim Pharmacet Micardis Twynsta	uticals, Inc. 19-20 29-31
Bristol-Myers Squibb Onglyza	72-74
Covidien AG TussiCaps	5-6
Cephalon, Inc. Amrix	17-18, 47-48
Eisai Inc. and Pfizer Inc. Aricept	36a-36b
Forest Laboratories, Inc. Namenda Savella Bystolic Lexapro	8a-8b 11-15 39-42 67-71
King Pharmaceuticals, Inc. Embeda	77-81
Eli Lilly and Company Evista Humalog Merck & Co., Inc.	24-27 34-36, 82-84
Januvia P&G Priolosec	52a-52b 43
Pfizer Inc. Lipitor Corporate Toviaz Lyrica Caduet	3-4 7, 32-33, 56a-56d, 57-58 21-22 44a-44d, 45-46 49-51
Stryker IVS Corporate	55
Wyeth Pharmaceuticals Inc. Pristiq	87-88
Xanodyne Pharmaceuticals, Inc. Zipsor	64-66