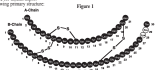
Lilly DESCRIPTION man insulin analog that is a rapid-acting, parenteral blood glucose-lowering agent nalog, created when the amino acids at positions 28 and 29 on the insulin B-chain are hogenic laboratory strain of Escherichia coli bacteria that has been genetically alterec Humalog* (insulin lispro, rDNA origin) is a human Chemically, it is Lys(B28), Pro(B29) human insulin analo reversed. Humalog is synthesized in a special non-path by the addition of the gene for insulin lispro. Humalog has the following primary structure:



Insulin lispro has the empirical formula C_2 :H₂N₂O₂S₃, and a molecular weight of \$808, both identical to that of human insulin. The viak, cartridges, and Pers contain a sterile solution of Humaling force as an injection. Humaling injection consists of rain-risualing the state of t

CLINICAL PHARMACOLOGY

Antidiahetic Activity
The primary activity of insulin, including Humalog, is the regulation of glucose metalosism. In addition, all insulins have several anabolic and microathorism con many insoes in the body, in muscle and other insues (except the brain), insufin causes inpid transport of glucose and anti-catabolic actions on many insoes in the body. In muscle and other insues (except the brain), insufin causes inpid transport of glucose glucose in the form of glycogen, inhibits glucoseogeneosis, and promotes the conversion of excess glucose into fat.

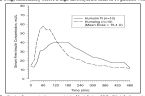
Humalog has been shown to be equipote to lumain insulin on analor basis. One unit of Humalog has been same glucose-lowering effect as one unit of human regular insulin, but its effect is more rapid and of shorter duration. The glucose-lowering activity of Humalog and human regular insulin is comparable when administered to normal voluntees by the intervenous route.

human regular insulin is comparable when administered to normal volunteres sy me turns-courso-to-use.

Pharmacokinetic.

Absorption and Biomendability—Humalog is as biosenaliable as human regular insulin, with absolute biosenaliability ranging between 55% to 75% with dossolute biomens. Humalog is as biosenaliable as human regular insulin, with type 1 (insulin-dependent) diabetes. The stress of the properties of the propert

Figure 2: Serum Humalog and insulin levels after subcutaneous injection of human regular insulin or Humalog (0.2 U/kg) immediately before a high carbohydrate meal in 10 patients with type 1 diabetes.*



Distribution — The volume of distribution for Humalog is identical to that of human regular insulin, with a range of 0.26 to 0.36 L/kg. Metabolism — Human metabolism studies have not been conducted. However, animal studies indicate that the metabolism of Humalog is given bushteneously, its lays be about the metabolism of humanogue and the metabolism of the studies of the metabolism of the studies of the stu

and 1 ong and 02 ong encycleverey.

Pharmodynamic
Shudies in normal volunteers and patients with diabetes demonstrated that Hamalog has a more rapid onset of glucose-lowering activity.

Shudies in normal volunteers and patients with diabetes demonstrated that Hamalog has a more rapid onset of glucose-lowering activity.

a carlier onset of activity of Hamalog is directly related to its more rapid rate of absorption. The time course of action to resolute and install time of the control of activity is known to be affected by the site of injection, exercise, and other variables (see PRECALTIONS, General).

Figure 3: Blood glucose levels after subcutaneous injection of human regular insulin or Humalog (0.2 U/kg) immediately before a high carbohydrate meal in 10 patients with type 1 diabetes.*



Special Populations

Age and Gender — Information on the effect of age and gender on the planmacokinetics of Humalog is unavailable. However, in large and fender — Information on the effect of age and gender dat to thindicate any difference in prosponding altonous parameters between Humalog and human regular insulin.

Smoking — The effect of smoking on the pharmacokinetics and pharmacokynamics of Humalog has not been studied. Programacy — The effect of programacy on the pharmacokinetics and pharmacokynamics of Humalog has not been studied. Obesity — The effect of solvenid and substantances fat the thickness on the pharmacokinetics and pharmacokynamics of Humalog has not been studied. Obesity — The effect of solvenid and substantances in the thickness on the pharmacokynamics of Humalog has not been studied. In large clinical trials, which included patients with Body Maks index up to and including 35 kgm²r, no consistent differences have been studied. The pharmacokinetic differences between Humalog and pharmacokynamics of Humalog has not been studied. The pharmacokinetic differences between Humalog and been as they of 25 patients with type 2 diabetes and a wide range of renal function, the pharmacokinetic differences between Humalog and patients with real displacements of the pharmacokinetic differences between Humalog and patients with real displacements of the pharmacokinetic differences between Humalog and patients with required by distinction.

In a study of 22 patients with type 2 diabetes, impaired bepatic function, the pharmacokinetic differences between Humalog and the pharmacokinetic difference the section of the substances of the pharmacokinetic

CLINICAL STUDIES
In open-label, cross-over studies of 1008 patients with type 1 diabetes and 722 patients with type 2 (non-insulin-dependent amalog reduced postprandial glucose compared with human regular insulin (see Table 1). The clinical significance of important properties of the properties of t

Table 1: Comparison of Means of Glycemic Parameters at the End of Combined Treatment Periods.
All Randomized Patients in Cross-Over Studies (3 months for each treatment)

An Randomized Fadents in Cross-Over Studies (5 months for each treatment)				
Type 1, N=1008 Glycemic Parameter, (mg/dL)	Humaloga	Humulin Ra*		
Fasting Blood Glucose 1-Hour Postprandial 2-Hour Postprandial HbA _{1c} (%)	209.5 ± 91.6 232.4 ± 97.7 200.9 ± 95.4 8.2 ± 1.5	204.1 ± 89.3 250.0 ± 96.7 231.7 ± 103.9 8.2 ± 1.5		
Type 2, N=722 Glycemic Parameter, (mg/dL)	Humalog ^a	Humulin R ^a		
Fasting Blood Glucose 1-Hour Postprandial 2-Hour Postprandial HbA _{1c} (%)	192.1 ± 67.9 238.1 ± 79.7 217.4 ± 83.2 8.2 ± 1.3	183.1 ± 66.1 250.0 ± 75.2 236.5 ± 80.6 8.2 ± 1.4		

In 12-month parallel studies in patients with type 1 and type 2 diabetes, HbA₁₁, did not differ between patients treated with human regular insulin and those treated with Humalog.

Hypoglycomia — White the overall rate of hypoglycemia did not differ between patients with type 1 and type 2 diabetes treated with Humalog compared with human regular insulin, patients with type 1 diabetes treated with Humalog and 6-me in Polycomic episodes between midnight and 6-m. The lower rate of hypoglycemia in the Humalog-treated group may have been related to higher nocturnal blood glucose levels, as reflected by a small increase in mean fasting blood glucose levels.

Humalog in Combinations with Sidoglyriure Agents— In a two-month study in patients with fasting hyperplycemia despite maximal dosing with sallonylureas (SU), patients were randomized to one of three treatment regimens: Humalins 'NPI' at beditine plus SU. Humalog teric times a day before measal path SU or Humalog three times a day before measal and Humalin NPII at beditine. The combination of Humalog and SU resulted in an improvement in HbA₁₀ accompanied by a weight gain (see Table 2).

Table 2: Results of a Two-Month Study in Which Humalog Was Added to Sulfonylurea Therapy in Patients Not Adequately Controlled on Sulfonylurea Alone

	Humulin N h.s. + SU ^a	Humalog a.c. + SU	Humalog a.c. + Humulin N h.s.		
Randomized (n)	135	139	149		
HbA _{1c} (%) at baseline	9.9	10.0	10.0		
HbA _{1c} (%) at 2-months	8.7	8.4	8.5		
HbA _{1s} (%) change from baseline	-1.2	-1.6	-1.4		
Weight gain at 2-months (kg)	0.6	1.2	1.5		
Hypoglycemia* (events/mo)	0.11	0.03	0.09		
Number of injections	1	3	4		
Total insulin dose (II/kg) at 2-months	0.23	0.33	0.52		

a.c.-three times a day before meals. h.s.-at bedrime. SU-oral sulfonylurea agent blood glucose \$36mg/dl. or needing assistance from third party.

Humalog in External Insulin Pamps — To evaluate the administration of Humalog via external insulin pumps, two open-label cross-over design studies were performed in patients with type I dubetes. One study involved 39 patients treated for 24 weeks with Humalog or regular human insulin. After 12 weeks of treatment, the mean HbAs, values descreased from 78 is 70 78% in the Immalog-tested patients and from 78% to 75% in the regular insulin-treated patients. Another study involved 60 patients treated for 24 weeks with either Humalog croid patients and remained unchanged from 7.7% in 6 the Humalog croid patients and remained unchanged from 7.7% in the Humalog croid patients and remained unchanged from 7.7% in the buffered regular insulin-treated patients. Alse of hypoglycease were comparable between treatment groups in 60st studies. Humalog administration in insulin pumps has not been studied in patients with type 2 dubetes.

INDICATIONS AND USAGE

Humalog is an insulin analog that is indicated in the returned or platests with diabetes mellitus for the control of hyperglycemia. Humalog has a more rapid onest and a shorter duration of action than human regular insulin. Therefore, in patients with type I diabetes, the malog ghould be used in regiments that include a longer-acting insulined particular between the majories that where the control is the state of the

CONTRAINDICATIONS Humalog is contraindicated during episodes of hypoglyce sensitive to Humalog or one of its excipients

This human issulin analog differs from human regular insulin by its rapid onset of action as well as a shorter duration of activity. When used as a meditime insulin, the dose of Humalog should be given within 15 minutes before or immediately after the med. Because of the short duration of action of Humalog, patients with type 1 diabetes also require a longe-setting insulin to maintain glooses control (except the control of action of Humalog, patients with type 1 diabetes also require a longe-setting insulin to maintain glooses control (except for patients using an external insulin jumps maniforing it recommended for all patients with diabetes and is particularly important for patients using an external insulin jumps maniforing its recommended for all patients with diabetes and is particularly important of the patients with diabetes and is particularly important of the patients with diabetes. Any change of insulin should be made causinosity and only under medical supervision. Changes in insulin strength, manufacturer, type and the patient with diabetes.

Any change of insulin should be made causinosity and only under medical supervision. Changes in insulin strength, manufacturer, type for a change in dosage.

L, regular, FPH, analoga, specces (attitums, munuary, or memory of a clausing indicase), a clausing indicase, a clausing indicase.

External Insulin Pumper When used in an external insulin pump, Humalog should not be diluted or mixed with any other insulin, its should carefully read and follow the external insulin pump manufacturer's instructions and the "INFORMATION FOR THE insuling the control of the contr

Physicians should carefully evaluate information on external insulin pump use in this Humalog physician package insert and in the external insulin pump manufacturer's instructions. If unexplained hyperglycema or ketosis occurs during external insulin pump use, prompt identification and correction of the cause is necessary. The patient user premaring this subcutaneous insulin injections (see PRECAUTIONS, For Patients Using External Insulin Pumps, and DOSAGE AND ADMINISTRATION). 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 (e.g., patients who a rating, have autonomic neuropathy, or are using potassuma-bowering days or printers shain of discovering the control of the property of the control of the

Information for Patients
Patients should be informed of the potential risks and advantage

Indicates the Patients
Design of the process of the

Laboratory Tests As with all insulins, the therapeutic response to Humalog should be monitored by periodic blood glucose tests. Periodic measurement of glycosylated hemoglobin is recommended for the monitoring of long-term glycemic control.

insulin requirements may be increased by medications with hyperplycemic activity such as corticosteroids, isoniazid, certain lipid-lowering drugs (e.g., niacin), estrogens, oral contraceptives, phenothiazines, and thyroid replacement therapy (see CLINICAL PHARMACOLOGY).

PHARMACOLOGY), use to decrease in the presence of drugs with hypoglycenic activity, such as only hypoglycenic activity. In the presence of the presence of drugs with hypoglycenic activity, such as only hypoglycenic activity, and as only hypoglycenic activity, such as only hypoglycenic activity, and activity activity

studied (see WARNING) with a longer-setting insulin, such as Humulin V, Humulin U, Humulog should be drawn into the syringe first to prevent clouding of the Humulog by the longer-acting insulin, such as Humulin V, Humulog should be drawn into the syringe first to prevent clouding of the Humulog by the longer-acting insulin. Injection should be made immediately after mixing. Mixtures should not be

Carcinogenesis, Mataguenis, Impairment of Fertility

Long-term studies in animals have not been performed to evaluate the carcinogenic potential of Humalog, Humalog was not matagenic in a naturey of in vitor and in vivo genetic trockivity assays (bacterial mutation tests, unecheduled DNA synthesis, mouse lymphoma assay, chromosomal aberariston tests, and a micromoticus test). There is no evidence from animal studies of Humalog-induced impairment of rettility.

Pregancy
Troutogonic Effects — Pregancy Category B — Reproduction studies have been performed in pregnant rats and rabbits at parenteral doses up to
A and 0.3 times, respectively, the average human dose (40 units/day) based on body surface area. The results have revealed no evidence of
impaired fertility or harm to the felus date to Humalog. There are, however, no adequate and well-controlled studies in pregnant women. Because
animal reproduction studies are not always preductive or human response, this draps abund to use of unitge greganney only of electry needed.
Although there are limited clinical studies of the use of Humalog in pregnancy, published studies with human insulins suggest that
Although there are limited clinical studies of the use of Humalog in pregnancy, published studies with human insulins suggest that
Although there fair domipalications of natural by-perdy term lane been need tho cummend, feel altonicity about host base reported with maternal
bypodypeensia. Insulin requirements usually fail during the first trimester and increase during the second and third trimesters. Careful
monitoring of the patient is required throughout pregnancy. During the perinatal period, careful monitoring of infants born to mothers with
diabetes is warranted.

Nursing Mothers It is unknown Nating abotters

It is unknown whether Humalog is excreted in significant amounts in human milk. Many drugs, including human insulin, are excreted in human milk. For this reason, caution should be exercised when Humalog is administered to a nursing woman. Patients with diabetes who are leathing may require adjustments in Humalog does, med plan, or both.

are lacturing may require adjustments in Humalog dose, meal plan, or both.

Publistric Use
In a 9-month, rons-over study of pre-purbescent children (resp. daged 3 to 11 years, comparable glycemic control as measured by HbA₁, was achieved regardless of treatment group; human regular modili 30 minutes before meals 8.4%, furnalise immediately before meals glycemic control as measured by HbA₂, was achieved regardless of treatment group; human regular insulin 30 to 45 minutes before meals 8.7% and Humanog immediately before meals 8.7%. The furnal regular insulin 30 to 45 minutes before meals 8.7% and Humanog insulin sides of the regular control as measured by HbA₂, was achieved regardless of treatment group; human regular insulin 30 to 45 minutes before meals 8.7% and Humanog in saw similar for all three treatments. Adjustment of basel insulin may be required. To improve accuracy in dossing in podatitre patients, a dilutert may be used. If the dilutent is added detectly to the Humanog vail, the side-life may be reduced, or DoSAGE, ADA DAMINISTRATION.

Geriarie Use
Of the total number of subjects (n=2834) in eight clinical studies of Humalog, twelve percent (n=338) were 65 years of age or over. The
mojerity of these were type? Indicats, HoA-, values and hypoglycemia rates did not differ by age. Pharmacokinetic/pharmacokynamic studies
to assess the effect of age on the onset of Humalog action have not been performed.

ADVERSE REACTIONS
Clinical studies comparing Humalog with human regular insulin did not demonstrate rate a difference in frequency of adverse events between

the two teatments.

Adverse events commonly associated with human insulin therapy include the following:
Body as a Whole —allerige reactions (see PRECAUTIONS).
Skin and Appendages — injection sile reaction, lipodytophy, pruritus, rash.
Other — hypoglycemia (see WARNINGS and PRECAUTIONS).

OVERDOSAGE

Hypoglycenia may occur as a result of an excess of insulin relative to food intake, energy expenditure, or both. Mild quisodes of hypoglycenia usually can be rearded with onal glacose. Adjustments in drug dosage, medi patterns, or esercise, may be needed. More severe quisodes with coma, searner, or neurologic impattement may be treated with intaminestal-relativalisationers glacusgic or contentional interactions. Decease the contention of the content of the contention to the content of the content of

DOSAGE AND ADMINISTRATION

Humalog is intended for subcutaneous administration, including use in select external insulin pumps (see DOSAGE AND ADMINISTRATION)

Humalog is intended for subcutaneous administration, including use in select external insulin pumps (see DOSAGE AND ADMINISTRATION). Attental finalin pumps (see DOSAGE AND ADMINISTRATION). Attental finalin pumps (see DOSAGE AND ADMINISTRATION). The pump (see DOSAGE AND ADMINISTRATION). The pump (see DOSAGE AND ADMINISTRATION) are subcutanteed by the pharmacodynamic studies showed Humalog to be equipotent to human regular insulin, but with more rapid activity. The quicker gloseco-lowering gredability as one unit of human regular insulin, but with more rapid activity. The quicker gloseco-lowering gredability is one unit of human regular insulin, but with more rapid activity. The quicker gloseco-lowering gredability are pump attent changes from other insulins in fulmane, particularly to prevent pre-mail perpendicular and appropriate appropriate appropriate and appropriate appropriate and appropriate app

when stored at S°C (41°F) and for 14 days when stored at 30°C (80°F). Do not cutture transmage common us a non-special material mainting purposes. The production of the produ How SUPPLED

Humalog (insulin lispro injection, fDNA origin) value are available in the following package size:

100 units per ml. (L'100)

NDC 0002-7516-91 (VL-7510)

NDC 0002-7516-91 (VL-7510)

SA' 3ml. cartridges'

NDC 0002-7516-99 (VL-7516)

Humalog (insulin lispro injection, fDNA origin) eartridges are available in the following package size:

NDC 0002-7516-99 (VL-7516)

SA' 3ml. distribution of DNA origin) eartridges are available in the following package size:

NDC 0002-7516-99 (VL-7516)

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Stronge — Unopened Humalog should be stored in a refrigerator $(2^b \text{ to } 8^b\text{C} [36^b \text{ to } 46^b\text{F}])$, but not in the freezer. Do not use Humalog if it has been frozen. Unrefrigerated (below $30^b\text{C} [86^b\text{F}])$ vials, cartridges, and Pens must be used within 28 days or be discarded, even if they still contain Humalog. Frotest from direct beat and light, See table below:

	Not in-use (unopened) Room Temperature below 86°F (30°C)	Not in-use (unopened) Refrigerated	In-use (opened) Room Temperature, below 86°F (30°C)
10 mL Vial	28 days	Until expiration date	28 days, refrigerated/room temperature.
3 mL Cartridge	28 days	Until expiration date	28 days, Do not refrigerate.
3 ml Pen	28 days	Until expiration date	28 days Do not refrigerate

Use in an External Insulin Pump — A Humalog 3 mL cartridge used in the D-TRON823 or D-TRON823 plus should be discarded after 7 days, even if it still contains Humalog. Infusion sets, D-TRON823 and D-TRON823 plus cartridge adapters, and Humalog in the external insulin pump reservoir should be discarded every 48 hours or less.

Literature revised February 6, 2007

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PA 9089 FSAMP

INFORMATION FOR THE PATIENT CARTRIDGE



(rDNA ORIGIN)

(100 UNITS PER ML (U-100)



3 ML CARTRIDGE

For use in Eli Lilly and Company's HumaPen® MEMOIRTM and HumaPen® LUXURATM HD insulin delivery devices, Owen Mumford, Ltd.'s Autopen®² 3 mL insulin delivery device (reusable insulin Pen), Disetronic®³ D-TRON®³ or D-TRON®³ plus insulin pumps.

WARNINGS

THIS LILLY HUMAN INSULIN ANALOG IS DIFFERENT FROM OTHER INSULINS BECAUSE IT HAS A RAPID ONSET AND SHORTER DURATION OF ACTION. THE RAPID ONSET OF ACTION MEANS THAT YOU SHOULD TAKE YOUR DOSE OF HUMALOG* (INSULIN LISPRO INJECTION, rDNA ORIGIN) WITHIN 15 MINUTES BEFORE OR IMMEDIATELY AFTER EATING A MEAL. THE SHORT DURATION OF ACTION OF HUMALOG MEANS THAT IF YOU HAVE TYPE 1 DIABETES, YOU ALSO NEED TO USE A LONGER-ACTING INSULIN TO GIVE THE BEST GLUCOSE CONTROL (EXCEPT WHEN USING AN EXTERNAL INSULIN PUMP). IF YOU HAVE TYPE 2 DIABETES, HUMALOG MAY BE USED WITHOUT A LONGERACTING INSULIN WHEN USED IN COMBINATION THERAPY WITH SULFONYLUIREA ACENTS.

ANY CHANGE OF INSULIN SHOULD BE MADE CAUTIOUSLY AND ONLY UNDER MEDICAL SUPERVISION. CHANGES IN STRENGTH, MANUFACTURER, TYPE (E.G., REGULAR, NPH, LENTE), SPECIES (BEEF, PORK, BEEF-PORK, HUMAN), OR METHOD OF MANUFACTURE (rDNA VERSUS ANIMAL-SOURCE INSULIN) MAY RESULT IN THE NEED FOR A CHANGE IN THE TIMING OR DOSAGE OF HUMALOG OR THE LONGERACTING INSULIN, OR BOTH.

PATIENTS TAKING HUMALOG MAY REQUIRE A CHANGE IN DOSAGE FROM THAT USED WITH OTHER INSULINS. IF AN ADJUSTMENT IS NEEDED, IT MAY OCCUR WITH THE FIRST DOSE OR DURING THE FIRST SEVERAL WEEKS OR MONTHS.

USE IN REUSABLE INSULIN PEN: TO OBTAIN AN ACCURATE DOSE, CAREFULLY READ AND FOLLOW THE INSULIN DELIVERY DEVICE MANUFACTURER'S INSTRUCTIONS AND THIS "INFORMATION FOR THE PATIENT" INSERT BEFORE USING THIS PRODUCT IN AN INSULIN PEN (see INSTRUCTIONS FOR USE section).

USE IN AN EXTERNAL INSULIN PUMP: CAREFULLY READ AND FOLLOW THE EXTERNAL INSULIN PUMP MANUFACTURER'S INSTRUCTIONS AND THIS "INFORMATION FOR THE PATIENT" INSERT BEFORE USING THIS PRODUCT IN AN EXTERNAL INSULIN PUMP (see INSTRUCTIONS FOR USE section).

DIABETES

Insulin is a hormone produced by the pancreas, a large gland that lies near the stomach. This hormone is necessary for the body's correct use of food, especially sugar. Diabetes occurs when the pancreas does not make enough insulin to meet your body's needs.

To control your diabetes, your doctor has prescribed injections of insulin products to keep your blood glucose at a near-normal level. You have been instructed to test your blood and/or your urine regularly for glucose. Studies have shown that some chronic complications of diabetes such as eye disease, kidney disease, and nerve disease can be significantly reduced if the blood sugar is maintained as close to normal as possible. The American Diabetes Association recommends that if your pre-meal glucose levels are consistently above 130 mg/dL or your hemoglobin A_{1c} (HbA_{1c}) is more than 7%, consult your doctor. A change in your diabetes therapy may be needed. If your blood tests consistently show below-normal glucose levels, you should also let your doctor know. Proper control of your diabetes requires close and constant cooperation with your doctor. Despite diabetes, you can lead an active and healthy life if you eat a balanced diet, exercise repularly and take your insulin injections as prescribed

Always keep an extra supply of Humalog as well as a spare syringe and needle on hand. Always wear diabetic identification so that appropriate treatment can be given if complications occur away from home.

HUMALOG

Description

Humalog (insulin lispro [rDNA origin]) is made by a special non-disease-producing laboratory strain of Escherichia coli bacteria that has been genetically altered by the addition of the gene for this human insulin analog. Humalog consists of zinc-insulin lispro crystals dissolved in a clear fluid. Humalog is a sterile solution and is for subcutaneous injection. It should not be used intramuscularly. The concentration of Humalog is 100 units/mL (U-100). Humalog starts lowering blood glucose more quickly and has a shorter duration of action compared to regular human insulin. This means that your dose of Humalog should be given within 15 minutes before or immediately after eating a meal (regular insulin works best when given 30 to 60 minutes before eating a meal). The short duration of action of Humalog means that if you have type 1 diabetes, you need to use a longer-acting insulin to give the best glucose control (except when using an external insulin pump). If you have type 2 diabetes, Humalog may be used without a longer-acting insulin when used in combination therapy with sulfonylurea agents. The time course of Humalog action, like that of other insulins, may vary in different individuals or at different times in the same individual, based on dose. site of injection, blood supply, temperature, and physical activity. Identification

Cartridges of insulin lispro injection (rDNA origin), by Eli Lilly and Company, have the trademark Humalog. Your doctor has prescribed the type of insulin that

nesine beneves is best for you.

DO NOT USE ANY OTHER INSULIN EXCEPT ON YOUR DOCTOR'S ADVICE AND DIRECTION.

Cartridges of Humalog 3 mL are available in boxes of 5.

3 mL Cartridge

Humalog[®] 3 mL cartridges are for use in Eli Lilly and Company's HumaPen* MEMOIR™ and HumaPen* LUXURA™ HD¹ insulin delivery devices, Owen Mumford, Ltd.'s Autopen®'3 mL insulin delivery device (reusable insulin Pen) and in Disetronic D-TRON®' or D-TRON®'plus insulin pumps using Disetronic Rapid®¹ infusion sets

The cartridge containing Humalog is not designed to allow any other insulin to be mixed in the cartridge or for the cartridge to be refilled with insulin.

Always examine the appearance of Humalog solution in a cartridge before administering a dose. When using a Humalog cartridge in an external insulin pump, inspect the cartridge before inserting it in the external insulin pump and periodically during use. Humalog is a clear and colorless liquid with a water-like appearance and consistency. Do not use if it appears cloudy, thickened, or slightly colored, or if solid particles are visible. If you note anything unusual in its appearance or notice your insulin requirements changing markedly, consult your doctor.

Storage

When used in Reusable Insulin Pen

Not in-use (unopened): Unopened Humalog cartridges should be stored in a refrigerator but not in the freezer. Do not use a Humalog cartridge if it has been frozen

In-use: Humalog cartridges in-use should NOT be refrigerated but should be kept at room temperature (below 86°F [30°C]) away from direct heat and light. Humalog cartridge that you are using must be discarded 28 days after the first use.

Do not use Humalog after the expiration date stamped on the label.

When used in an External Insulin Pump

Infusion sets (tubing and catheters) and D-TRON® or D-TRON® plus cartridge adapter should be discarded every 48 hours or less. Humalog in an external insulin pump should not be exposed to temperatures above 98.6°F (37°C) such as in sauna or hot tub, hot showers, direct sunlight, or radiant heater. A Humalog 3 mL cartridge used in the D-TRON® or D-TRON® plus pump should be discarded after 7 days, even if it still contains Humalog.

INSTRUCTIONS FOR USE

Reusable insulin Pens and external insulin pumps differ in their operation. It is important to read, understand, and follow the instructions for use of the reusable insulin Pen or external insulin pump you are using.

NEVER SHARE INSULIN PENS, EXTERNAL INSULIN PUMPS, INFUSION SETS, CARTRIDGES, OR NEEDLES.

PREPARING FOR AN INJECTION USING REUSABLE INSULIN PEN OR EXTERNAL INSULIN PUMP

- 1. Inspect the appearance of Humalog solution before you insert the cartridge into the reusable insulin Pen or external insulin pump. Humalog should look clear and colorless. Do not use Humalog if it appears cloudy, thickened, slightly colored, or if solid particles are visible. Once the cartridge is in use, inspect the insulin in the insulin Pen before each injection. When using a Humalog cartridge in an external insulin pump, inspect the cartridge before inserting it in the external insulin pump and periodically during use.
- Use in Reusable Insulin Pen Follow the reusable insulin Pen manufacturer's
 instructions carefully for loading the cartridge into the insulin Pen and for use
 of the insulin Pen.
- a. Use an alcohol swab to wipe the exposed rubber surface on the metal cap end
 of the cartridge.
- Follow the insulin needle manufacturer's instructions for attaching and changing the needle.
- Use in an External Insulin Pump Follow the external insulin pump manufacturer's instructions carefully for use of Humalog 3 mL cartridges in the D-TRON®3 or D-TRON®-blus insulin pump.

GENERAL INSTRUCTIONS

- For use in Reusable Insulin Pen
- Wash your hands.
- To avoid tissue damage, choose a site for each injection that is at least 1/2 inch from the previous injection site. The usual sites of injection are abdomen, thighs, and arms.
- 3. Cleanse the skin with alcohol where the injection is to be made
- 4. With one hand, stabilize the skin by spreading it or pinching up a large area.
- Inject the dose as instructed by your doctor. Hold the needle under the skin for at least 5 seconds after injecting.
- After injecting a dose, pull the needle out and apply gentle pressure over the injection site for several seconds. Do not rub the area.
- 7. Immediately after an injection, remove the needle from the insulin Pen. Doing so will guard against contamination, and prevent leakage of Humalog, reentry of air, and needle clogs. Do not reuse needles. Place the used needle in a puncture-resistant disposable container and properly dispose of it as directed by your Health Care Professional.



8. 3 mL cartridge — Use the gauge on the side of the cartridge to help you judge how much insulin remains. The distance between each mark on the 3 mL cartridge is about 20 units.

For use in an External Insulin Pump

Your doctor should train you on intensive insulin therapy including sterile techniques. You should also be trained on the use of your external insulin pump and pump accessories.

You should replace the infusion set (tubing and catheter) and D-TRON®3 or D-TRON®3plus cartridge adapter every 48 hours or less. You should also choose a new infusion site every 48 hours or less. A Humalog 3 mL cartridge used in the pump should be discarded after 7 days, even if it still contains Humalog. Contact your doctor if your infusion sites are red, itching, or thickened, and then choose a new infusion site.

Follow the external insulin pump manufacturer's instructions carefully for use of Humalog 3 mL cartridges in Disetronic D-TRON®3 or D-TRON®3plus insulin pump.

DOSAGE

Your doctor has told you which insulin to use, how much, and when and how often to inject it. Because each patient's case of diabetes is different, this schedule has been individualized for you. Your usual Humalog dose may be affected by changes in your food, activity, or work schedule. Carefully follow your doctor's instructions to allow for these changes. Other things that may affect your Humalog dose are:

Illness, especially with nausea and vomiting, may cause your insulin requirements to change. Even if you are not eating, you will still require insulin. You and your doctor should establish a sick day plan for you to use in case of illness. When you are sick, test your blood glucose/urine glucose and ketones frequently and call your doctor as instructed.

Pregnancy

Good control of diabetes is especially important for you and your unborn baby. Pregnancy may make managing your diabetes more difficult. If you are planning to have a baby, are pregnant, or are nursing a baby, consult your doctor. Humalog has not been tested in pregnant or nursing women.

Geriatric Use

Elderly patients using Humalog had HbA₁₆ values and hypoglycemia rates similar to those observed in younger patients. The onset of action of Humalog may be different in elderly patients.

Medication

Insulin requirements may be increased if you are taking other drugs with hyperglycemic activity, such as oral contraceptives, corticosteroids, or thyroid replacement therapy. Insulin requirements may be reduced in the presence of drugs with blood-glucose-lowering activity, such as oral antidiabetic agents, salicylates (for example, aspirin), sulfa antibiotics, alcohol, and certain antidepressants. Your Health Care Professional is aware of other medications that may affect your diabetes control. Therefore, always discuss any medications you are taking with your doctor.

Exercise

Exercise may lower your body's need for insulin products during and for some time after the physical activity. Exercise may also speed up the effect of a dose of Humalog, especially if the exercise involves the area of injection site. Discuss with your doctor how you should adjust your regimen to accommodate exercise.

Persons traveling across more than 2 time zones should consult their doctor concerning adjustments in their insulin schedule

COMMON PROBLEMS OF DIABETES

Hypoglycemia (Low Blood Sugar)

Hypoglycemia (too little glucose in the blood) is one of the most frequent adverse events experienced by insulin users. It can be brought about by:

- Missing or delaying meals.
- Taking too much insulin.
- Exercising or working more than usual.
- An infection or illness (especially with diarrhea or vomiting).
- A change in the body's need for insulin.
- 6. Diseases of the adrenal, pituitary, or thyroid gland, or progression of kidney or
- 7. Interactions with other drugs that lower blood glucose, such as oral antidiabetic agents, salicylates (for example, aspirin), sulfa antibiotics, and certain antidepressants.

8. Consumption of alcoholic beverages.

Symptoms of mild to moderate hypoglycemia may occur suddenly and can include:

drowsiness

· blurred vision

· slurred speech

· irritability

· seizures

· depressed mood

· abnormal behavior

· unsteady movement

· personality changes

anxiety

· sleep disturbances

- sweating
- dizziness
- palpitation · tremor
- hunger
- · restlessness
- · tingling in the hands, feet, lips, or tongue
- · lightheadedness
- · inability to concentrate
- · headache

Signs of severe hypoglycemia can include: disorientation

· unconsciousness death Therefore, it is important that assistance be obtained immediately.

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, changing insulin preparations, or intensified control (3 or more injections per day) of diabetes. A few patients who have experienced hypoglycemic reactions after transfer from animal-source insulin to human insulin have reported that the early warning symptoms of hypoglycemia were less pronounced or different from those experienced with their previous insulin.

Without recognition of early warning symptoms, you may not be able to take steps to avoid more serious hypoglycemia. Be alert for all of the various types of symptoms that may indicate hypoglycemia. Patients who experience hypoglycemia without early warning symptoms should monitor their blood glucose frequently, especially prior to activities such as driving. If the blood glucose is below your normal fasting glucose, you should consider eating or drinking sugar-containing foods to treat your hypoglycemia.

Mild to moderate hypoglycemia may be treated by eating foods or drinks that contain sugar. Patients should always carry a quick source of sugar, such as candy mints or glucose tablets. More severe hypoglycemia may require the assistance of another person. Patients who are unable to take sugar orally or who are unconscious require an injection of glucagon or should be treated with intravenous administration of glucose at a medical facility

You should learn to recognize your own symptoms of hypoglycemia. If you are uncertain about these symptoms, you should monitor your blood glucose frequently to help you learn to recognize the symptoms that you experience with hypoglycemia.

If you have frequent episodes of hypoglycemia or experience difficulty in recognizing the symptoms, you should consult your doctor to discuss possible changes in therapy, meal plans, and/or exercise programs to help you avoid hypoglycemia.

Hyperglycemia and Diabetic Ketoacidosis (DKA)

Hyperglycemia (too much glucose in the blood) may develop if your body has too little insulin. Hyperglycemia can be brought about by any of the following:

- Omitting your insulin or taking less than the doctor has prescribed.
- Eating significantly more than your meal plan suggests
- Developing a fever, infection, or other significant stressful situation.

In patients with type 1 or insulin-dependent diabetes, prolonged hyperglycemia can result in DKA. The first symptoms of DKA usually come on gradually, over a period of hours or days, and include a drowsy feeling, flushed face, thirst, loss of appetite,

and fruity odor on the breath. With DKA, urine tests show large amounts of glucose and ketones. Heavy breathing and a rapid pulse are more severe symptoms. If uncorrected, prolonged hyperglycemia or DKA can lead to nausea, vomiting, stomach pains, dehydration, loss of consciousness, or death. Therefore, it is important that you obtain medical assistance immediately.

Lipodystrophy

Rarely, administration of insulin subcutaneously can result in lipoatrophy (depression in the skin) or lipohypertrophy (enlargement or thickening of tissue). If you notice either of these conditions, consult your doctor. A change in your injection technique may help alleviate the problem.

Local Allergy - Patients occasionally experience redness, swelling, and itching at the site of injection of insulin. This condition, called local allergy, usually clears up in a few days to a few weeks. In some instances, this condition may be related to factors other than insulin, such as irritants in the skin cleansing agent or poor injection technique. If you have local reactions, contact your doctor.

Systemic Allergy — Less common, but potentially more serious, is generalized allergy to insulin, which may cause rash over the whole body, shortness of breath, wheezing, reduction in blood pressure, fast pulse, or sweating. Severe cases of generalized allergy may be life threatening. If you think you are having a generalized allergic reaction to insulin, notify a doctor immediately.

ADDITIONAL INFORMATION

Additional information about diabetes may be obtained from your diabetes

DIABETES FORECAST is a magazine designed especially for people with diabetes and their families. It is available by subscription from the American Diabetes Association (ADA), P.O. Box 363, Mt. Morris, IL 61054-0363, 1-800-DIABETES (1-800-342-2383)

Another publication, COUNTDOWN, is available from the Juvenile Diabetes Research Foundation International (JDRFI), 120 Wall Street, 19th Floor, New York, NY 10005, 1-800-533-CURE (1-800-533-2873).

Additional information about Humalog can be obtained by calling The Lilly Answers Center at 1-800-LillyRx (1-800-545-5979).

Patient Information revised February 6, 2007

Manufactured by Lilly France F-67640 Fegersheim, France for Eli Lilly and Company Indianapolis, IN 46285, USA

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5 Cartridges 3 mL

NDC 0002-7516-59 VL - 7516 100 units per mL





insulin lispro injection (rDNA origin)

Rx only

U-100

For use in Eli Lilly and Company's HumaPen® MEMOIR™ and HumaPen® LUXURA™ HD insulin delivery devices and Owen Mumford, Ltd.'s Autopen® 3 mL insulin delivery device.

SH 8662 FSAMS

Humalog® insulin lispro injection (rDNA origin)

U-100

5 x 3 mL cartridges 100 units per mL

Humalog® insulin lispro injection (rDNA origin)

001-U

5 x 3 mL cartridges

IMPORTANT-SEE WARNINGS ON ENCLOSED INSERT

Keep in a cold place. Avoid freezing. Warning: Any change of insulin should be made cautiously and only under medical supervision.

For subcutaneous use.

See enclosed insert for dosage.

Each mL contains 100 Units of insulin lispro; glycerin, 16 mg; dibasic sodium phosphate, 1.88 mg; Metacresol, 3.15 mg; zinc oxide content adjusted to provide 0.0197 mg zinc ion; trace amounts of phenol, and water for injection. Hydrochloric acid 10% and/or sodium hydroxide 10% may be added to adjust pH.

Neutral

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3 mL 444, DASHIN TOURIS PER ML U-100 UNIS PER ML



