

the office staff performs.

## THE OFFICE

## Redelegate Tasks When Installing an EHR

nstalling an electronic health record system at a primary care practice is a perfect time to also revamp the tasks

ciple: If a task does not require an MD or DO, then people with those degrees shouldn't do it. Instead, such a task should be delegated through a standing order to someone else on the staff, often a nurse or medical assistant.

top of their licensure is the guiding prin-

In a standing order, you extend your licensure to lower-level staff. Write the order in detail, sign it, put it into a folder, and keep it up-to-date for accreditation inspections.

With routine clinical tasks delegated to staff members using EHR tools, the typical duration of an office encounter between a primary care physician and patient can be shaved by a third, from 12 minutes to 8 minutes. The time saved can be used to see more patients or for leisure time.

EHRs allow these time savings in two main ways. First, they better enable a staff member, usually a nurse or medical assistant, to collect and enter into the record preliminary clinical information, such as past medical history, family history, vital signs, and reason for

When the physician enters the exam room to see the patient, reviewing what the staff has entered into the medical chart takes half as much time as entering the data initially and then they simply need to add the additional information that they gather.

The scope of preliminary information collected can be expanded and tailored to

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For example, a physician can teach staff members how to collect current information from patients with diabetes using a diabetes template, leaving fewer routine tasks for the physician to do. About a quarter of my patients have diabetes, so my nurse knows my diabetes template cold.

The second major time saving is through standing orders that a staff member can perform and that are automatically flagged for appropriate patients. These standing orders would include items such as administering immunizations, arranging for routine lab tests like prostate-specific antigen and hemoglobin  $A_{1c}$ , and reminding patients about and possibly scheduling routine screens like mammography and colonoscopy.

The key to making this sort of practice change work is to introduce new approaches in small steps and evaluate how each change works after a few weeks. The staff must be properly trained, and the changes should take into account how the staff currently works and how new tasks will affect them. Having a staff member on the development and implementation team is a big advantage.

Workflow changes following introduction of an electronic health record is an evolving and continuous process, not a static goal. Once an EHR system is in place and work assignments start to change, delegation of tasks will continue to evolve and the workflow will ideally continue to become more efficient over time.

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**HUMALOG®** 

Having all staff members work at the

INSULIN LISPRO INJECTION (rDNA ORIGIN)
BRIEF SUMMARY: Consult package insert for complete prescribing information.

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. However, in patients with type 2 diabetes, Humalog may be used without a longer-acting insulin when used in combination therapy with sulfonylurea agents.

Humalog may be used in an external insulin pump, but should not be diluted or mixed with any other insulin when used in the pump. Humalog administration in insulin pumps has not been studied in patients

CONTRAINDICATIONS: Humalog is contraindicated during episodes of hypoglycemia and in patients sensitive to

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 "PATIENT INFORMATION" leaflet 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 Pumps, and DOSAGE AND ADMINISTRATION).

Hypoglycemia is the most common adverse effect associated with the use of insulins, including Humalog.

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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 manufacturer 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—lovering 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 insulin preparations, 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 ohvisical activity.

Interest times in the same incurvous and is dependent on size or 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 induce 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.

Renal Impairment—The requirements for insulin may be reduced in patients with renal impairment. Hepatic Impairment—Hhough 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.

Allergy—Local Allergy—As with any insulin therapy, patients may experience redness, swelling, or itching at the site of injection. These minor reactions usually resolve 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 nijection technique.

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

Systemic Allergy—Less common, but potentially more serious, is generalized allergy to insulin, which 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 be life-threatening. Localized reactions and generalized myalgias 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 Humulin R\* (N=2969) and 30 patients receiving Humalog (N=2944) (P=.053).

Antibody Production—In large clinical trials, antibodies that cross-react with human insulin and insulin lispro were observed in both Humulin R+ and Humalog-treatment groups. As expected, the largest increase in the antibody levels during the 12-month clinical trials was observed with patients new to insulin therapy.

Usage of Humalog in External Insulin Pumps—The Infusion set (reservoir syringe, tubing, and catheter), Disetronic\* D-TRON®-2 or D-TRONplus®-2 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 pumps, the infusion set should be replaced and a new infusion site should be replaced every 48 hours or less.

When used in an external insulin pump, Humalog should not be diluted or mixed with any other insulin (see

selected every 48 hours or less.

When used in an external insulin pump, Humalog should not be diluted or mixed with any other insulin (see IMDICATIONS AND USAGE, WARNINGS, PRECAUTIONS, For Patients Using External Insulin Pumps, Mixing of Insulins, DOSAGE AND ADMINISTRATION, and Storage).

Information for Patients—Patients should be informed of the potential risks and advantages of Humalog and alternative therapies. Patients should use informed about the importance of proper insulin storage, injection technique, timing of dosage, adherence to meal planning, regular physical activity, regular blood glucose monitoring, periodic hemoglobin AIC testing, recognition and management of hypoglycemia and hyperglycemia, and periodic assessment for diabetes complications.

Patients should be advised to inform their physicial if the user across.

technique, liming or lossage, admerence to metal planning, regular physical activity, regular blood glucose monitoring, periodic hemoglobin ATC testing, recognition and management of hypoglycemia and hyperglycemia, and periodic assessment for diabetes complications.

Patients should be advised to inform their physician if they are pregnant or intend to become pregnant. Refer patients to the "PATIENT INFORMATION" leaflet for timing of Humalog dosing (≤15 minutes before or immediately after a meal), storing insulin, and common adverse effects.

For Patients Using Insulin Pen Delivery Devices: Before starting therapy, patients should read the "PATIENT INFORMATION" leaflet that accompanies the drug product and the User Manual that accompanies the delivery device. They should also reread these materials each time the prescription is renewed. Patients should be instructed on how to properly use the delivery device, prime the Pen, and properly dispose of needles. Patients should be advised not to share their Pens with others.

For Patients Using External Insulin Pumps: Patients using an external infusion pump should be trained in intensive insulin therapy and in the function of their external insulin pump and pump accessories. Humalog was tested in the MiniMed<sup>®</sup> Models 506, 507, and 508 insulin pumps using MiniMed<sup>®</sup> Polyfine³ infusion sets. Humalog was also tested in the Distertonic ®+1+RONJ®\*23 and D−1RONJPuls®\*23 insulin pumps (with Humalog 3 mL cartridges) using Disetronic Rapid®\* infusion sets.

The Infusion set (reservoir syringe, tubing, catheter), D−1RONJPuls®\*23 and a new infusion site selected every 48 hours or less. Humalog in the external lump phould not be exposed to temperatures above 37°C (98.6°F).

A Humalog 3 mL cartridge used in the D−1RON®\*23 or D−1RONJPuls®\*23 pump should be discarded after 7 days, even if it still contains Humalog, Infusion sites that are erythematous, pruritic, or thickened should be reported to medical personnel, and a new site selected.

Humalog should not be diluted or mixed

blood glucose tests. Periodic measurement of hemogroum ATU is recommended to the monitoring of nongream glycemic control.

\*\*Drug Interactions\*\*—Insulin requirements may be increased by medications with hyperglycemic activity, such as corticosteroids, isoniazid, certain lipid-lowering drugs (eg., niacin), estrogens, oral contraceptives, phenothiazines, 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 (eg., octreotide), 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 response to the insulin mixture may differ from that of the injection of the insulins separately." Mixing Humalog with Humulin® N or Humulin® U does not decrease the absorption rate or the total bioavailability of Humalog.

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 licera beauty.

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).

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 neurologic impairment may be treated with intramuscular/subcutaneous glucagon or concentrated intravenous glucose. 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 (ie, one unit of Humalog has the same glucose-lowering effect as one unit of regular human insulin), but with more rapid activity. The quicker glucose-lowering effect of basal insulin may be needed when a patient changes from other insulins to Humalog, assistantly in revenent premial themselvermia.

owering effect of Humalog is related to the more rapid absorption rate from subcutaneous tissue. An adjustment of dose or schedule of basal insulin may be needed when a patient changes from other insulins to Humalog, particularly to prevent premeal hyperglycemia.

When used as a mealtime insulin, Humalog should be given within 15 minutes before or immediately after a meal. Regular human insulin is best given 30 to 60 minutes before a meal. To achieve optimal glucose control, the amount of longer-acting insulin being given may need to be adjusted when using Humalog.

The rate of insulin absorption and consequently the onest of activity are known to be affected by the site of injection, exercise, and other variables. Humalog was absorbed at a consistently faster rate than regular human insulin or Humalog at abdominal, deltoid, or femoral sites, the 3 sites often used by patients with diabetes. When not mixed in the same syringe with other insulins, Humalog maintains its rapid onset of action and has less variability in its onset of action and has less variability in its onset of action and has less variability in its onset of action and has less variability in its onset of action and has less variability in its onset of action and has less variability in its onset of action and has less variability in its onset of action and many concentrations. Also, the duration of action of Humalog is slightly shorter following abdominal injection, compared with deltoid and femoral injections. As with all insulin preparations, the time course of action of Humalog may vary considerably in different individuals or within the same individual. Patients must be educated to use proper injection techniques.

Humalog in a vail may be diluted with STERILE DILUENT for Humalog, Humulin N, Humulin R, Humulin 70/30, and Humuline® RJ-50 to a concentration of 1:10 (equivalent to U-10) or 1:2 (equivalent to U-50). Diluted Humalog may remain in patient use for 28 days when stored at 5°C (41°P) and for 14 days when stored at 30°C (86°P). Do not

HUMBUSUPFLIED:
Humalog (insulin lispro injection, USP [rDNA origin]) is available in the following package sizes (with each presentation containing 100 units insulin lispro per mL [U-100]):
10 mL vials
5 x 3 mL cartridges³
5 x 3 mL cartridges³
5 x 3 mL disposable insulin delivery devices (Pen)
NDC 0002-7516-59 (VL-7516)
NDC 0002-7516-59 (VL-7516)
NDC 0002-8799-59 (HP-8725)
NDC 0002-8799-59 (HP-8799)

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Disetronic®, H-TRONplus®, D-TRON®, and Rapid® are registered trademarks of Roche Diagnostics GMBH.

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Other product and company names may be the trademarks of their respective owners.

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

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\*\*Use in an External Insulin Pump—A Humalog 3mL cartridge used in the D-TRON®23 or D-TRONplus®23

\*\*should be discarded after 7 days, even if it still contains Humalog, Infusion sets, D-TRON®23 and D-TRONplus

\*\*cartridge adapters, and Humalog in the external insulin pump reservoir should be discarded every 48 hours

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.
www.humalog.com

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