

About This Guide

This Guide provides a high-level overview of Health Management Reminders in Greenway Intergy. The overview is meant to provide guidance for you, your practice electronic health record (EHR) champion, or IT staff.

Experienced users are the main focus of this Guide; not every step is included in the instructions, and this is not a replacement for training from Greenway Intergy. There are several ways to approach each workflow in Greenway Intergy. This Guide highlights one workflow; however, you may be familiar with an alternative approach. Please note that this Guide was created based upon Greenway Intergy version 10.10. Screens and features may change as new software versions are released.

INDICATIONS AND USAGE

- PRALUENT is a PCSK9 (Proprotein Convertase Subtilisin/Kexin Type 9) inhibitor antibody indicated as adjunct to diet and maximally tolerated statin therapy for the treatment of adults with heterozygous familial hypercholesterolemia or clinical atherosclerotic cardiovascular disease, who require additional lowering of LDL-C
- The effect of PRALUENT on cardiovascular morbidity and mortality has not been determined

IMPORTANT SAFETY INFORMATION

- PRALUENT is contraindicated in patients with a history of a serious hypersensitivity reaction to PRALUENT. Reactions have included hypersensitivity vasculitis and hypersensitivity reactions requiring hospitalization

Please see accompanying full [Prescribing Information](#)



Using Health Management Reminders

Clinical decision support (CDS) tools such as Health Management Reminders provide clinicians and staff with patient-specific information, intelligently filtered and presented at appropriate times to help improve the delivery of care. Health Management Reminders can assist with identifying gaps-in-care and may enhance patient outcomes and improve care consistency—a benefit to both providers and patients.

Health Management Reminders can be created to alert health care professionals (HCPs) to consider PRALUENT therapy for appropriate patients during the visit.

Factors That May Impact Health Management Reminders

The display of Health Management Reminders may be impacted by the clinical data available in the EHR; for example, if lab results are saved in the EHR as a PDF file and not available for use as “data” to be queried. Or, if lab results are received from multiple laboratories, it may be necessary to map each lab’s order codes to LOINC codes to enable all appropriate patients to appear on the patient list. Also, medications prescribed before the EHR was implemented might not be included in a patient’s medications list. These information gaps can limit the number of patients on which reminders display.

Health Management Reminders can help identify whether the patient is a candidate for PRALUENT treatment based on clinical appropriateness and payer utilization management criteria via:

- Reminders during the patient exam workflow, such as ordering lab tests or other diagnostics, adjusting or adding medications to a patient’s treatment plan, or providing patient education on LDL-C goals
- Identification of patients who, for example, have clinical atherosclerotic cardiovascular disease (ASCVD) and are on maximally tolerated statin therapy atorvastatin 40 mg/day and having elevated LDL-C ≥ 70 mg/dL or ≥ 100 mg/dL, depending on insurance

IMPORTANT SAFETY INFORMATION

- Hypersensitivity reactions (e.g., pruritus, rash, urticaria), including some serious events (e.g., hypersensitivity vasculitis and hypersensitivity reactions requiring hospitalization), have been reported with PRALUENT treatment. If signs or symptoms of serious allergic reactions occur, discontinue treatment with PRALUENT, treat according to the standard of care, and monitor until signs and symptoms resolve

Please see accompanying full [Prescribing Information](#)



Reminders: Using Health Management in Greenway Intergy

Greenway Intergy enables the setup of reminders based on criteria in the Health Management Reminders feature. Available criteria include diagnosis, current and prior medications, lab values, and other clinical or patient demographic information. The Health Management Reminder is displayed when the data in the patient chart meet the criteria.

For example, a Health Management Reminder may be created to alert the HCP of patients with uncontrolled LDL-C who are clinically appropriate and approvable for PRALUENT therapy based upon the approved Indication.

The following steps illustrate how to create reminders to identify patients who may be candidates for treatment with PRALUENT.

To Create Reminders in Health Management:

NOTE: To see a list of patients for whom this reminder will display, use the EHR patient identification tip sheet to create a patient list. The list will identify patients who may be appropriate for PRALUENT using specific criteria (eg, diagnosis of ASCVD, atorvastatin 40 mg/day, LDL-C \geq 70 mg/dL).

1. Navigate to **Health Management Setup**

2. Select **Tools**, Select **New**

3. Add items to the Care Condition Set.

For example: Medications: atorvastatin, rosuvastatin. Diagnosis: (Clinical ASCVD [eg, stroke, transient ischemic attack, acute coronary syndromes, history of myocardial infarction, stable or unstable angina, coronary or other arterial revascularization, peripheral arterial disease] plus hypercholesterolemia OR HeFH). Lab Results: ie, LDL-C \geq 70 mg/dL or \geq 100 mg/dL, depending on insurance

Health Management Reminder
Guideline type: Care Condition
Gender: Both Default Diag:
Age Category: All
Recall Reason: <Do Not Generate Recall>
Knowledge Links:
Quality Reporting:

Care Condition Set	Type	Code
<input type="checkbox"/>		
Statin	drugClass	
Diagnosis	Diag	
Lab - LDL	Lab	

Reminders	Frequency	Tier	Conditional Inclusion	Warm Schedule	Recall	Link
Patient is candidate for treatment intensification with PRALUENT	Once	2	None			

IMPORTANT SAFETY INFORMATION

- The most commonly occurring adverse reactions (\geq 5% of patients treated with PRALUENT and occurring more frequently than with placebo) are nasopharyngitis, injection site reactions, and influenza

Please see accompanying full [Prescribing Information](#)



Reporting: Using Health Management Reminders

4. Add Clinical Event(s)

For example:

- Patient visit
- LDL result ≥ 70 mg/dL

5. Create Reminder

- Define reminder text
- Define satisfying event(s)
- For example:
 - LDL result ≥ 70 mg/dL

View Reminder [?] [X]

Reminder Text:
Patient candidate for Treatment with PRALUENT Tier: 2

Knowledge Link: []
Default Diag for Posting Orders: []

Guideline Defaults:
Frequency: Once []
Warm Schedule: []
Recall Schedule: []

Satisfying Events and Orders

Description	Type	Code	Satisfies Reminders	Display as Order
Praluent	Order		<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>

[Edit] [Close]

To View Health Management Reminders:

Health Management Reminders are viewable in the patient chart via flyover buttons in Flowsheets, Visit Notes, and Orders/Charges screens.

Chart Flowsheets Note Order/Charges Patient Summary

NO NARCOTICS LETTER SENT [dismiss X]

Test, Patient
04/27/1995 21 y M ■■■ xxx-xx-6789 ■■ Care Program None [Add a Sticky Note]

[Patient Encounter] with Allen, Christopher C MD on 12/07/2016
[Print Summary] [Send Summary] [More...]

Today's Vitals
No vitals entered today

Health Reminders Due
Cholesterol Care Due
Immunizations Due
None

Problems: atherosclerosis Medications: atorvastatin [Print Summary]

IMPORTANT SAFETY INFORMATION

- Local injection site reactions including erythema/redness, itching, swelling, and pain/tenderness were reported more frequently in patients treated with PRALUENT 75 mg and/or 150 mg every 2 weeks (7.2% versus 5.1% for PRALUENT and placebo, respectively). Few patients discontinued treatment because of these reactions (0.2% versus 0.4% for PRALUENT and placebo, respectively), but patients receiving PRALUENT had a greater number of injection site reactions, had more reports of associated symptoms, and had reactions of longer average duration than patients receiving placebo

Please see accompanying full [Prescribing Information](#)



INDICATIONS AND USAGE

- PRALUENT® (alirocumab) is a PCSK9 (Proprotein Convertase Subtilisin/Kexin Type 9) inhibitor antibody indicated as adjunct to diet and maximally tolerated statin therapy for the treatment of adults with heterozygous familial hypercholesterolemia or clinical atherosclerotic cardiovascular disease, who require additional lowering of LDL-C
- The effect of PRALUENT on cardiovascular morbidity and mortality has not been determined

IMPORTANT SAFETY INFORMATION

- PRALUENT is contraindicated in patients with a history of a serious hypersensitivity reaction to PRALUENT. Reactions have included hypersensitivity vasculitis and hypersensitivity reactions requiring hospitalization
- Hypersensitivity reactions (e.g., pruritus, rash, urticaria), including some serious events (e.g., hypersensitivity vasculitis and hypersensitivity reactions requiring hospitalization), have been reported with PRALUENT treatment. If signs or symptoms of serious allergic reactions occur, discontinue treatment with PRALUENT, treat according to the standard of care, and monitor until signs and symptoms resolve
- The most commonly occurring adverse reactions ($\geq 5\%$ of patients treated with PRALUENT and occurring more frequently than with placebo) are nasopharyngitis, injection site reactions, and influenza
- Local injection site reactions including erythema/redness, itching, swelling, and pain/tenderness were reported more frequently in patients treated with PRALUENT 75 mg and/or 150 mg every 2 weeks (7.2% versus 5.1% for PRALUENT and placebo, respectively). Few patients discontinued treatment because of these reactions (0.2% versus 0.4% for PRALUENT and placebo, respectively), but patients receiving PRALUENT had a greater number of injection site reactions, had more reports of associated symptoms, and had reactions of longer average duration than patients receiving placebo
- The once-monthly (Q4W) 300 mg dosing regimen had a higher rate of local injection site reactions as compared to PRALUENT 75 mg Q2W or placebo (16.6%, 9.6%, and 7.9%, respectively) in a trial in which all patients received an injection of drug or placebo every 2 weeks to maintain the blind. The discontinuation rate due to injection site reactions was 0.7% in the 300 mg Q4W arm and 0% in the other 2 arms
- Neurocognitive events were reported in 0.8% of patients treated with PRALUENT and 0.7% of patients treated with placebo. Confusion or memory impairment were reported more frequently by those treated with PRALUENT (0.2% for each) than in those treated with placebo ($<0.1\%$ for each)
- Liver-related disorders (primarily related to abnormalities in liver enzymes) were reported in 2.5% of patients treated with PRALUENT and 1.8% of patients treated with placebo, leading to treatment discontinuation in 0.4% and 0.2% of patients, respectively. Increases in serum transaminases to greater than 3 times the upper limit of normal occurred in 1.7% of patients treated with PRALUENT and 1.4% of patients treated with placebo
- The most common adverse reactions leading to treatment discontinuation in patients treated with PRALUENT were allergic reactions (0.6% versus 0.2% for PRALUENT and placebo, respectively) and elevated liver enzymes (0.3% versus $<0.1\%$)
- PRALUENT is a human monoclonal antibody. As with all therapeutic proteins, there is a potential for immunogenicity with PRALUENT

Please see accompanying full [Prescribing Information](#)



©2017 Sanofi and Regeneron Pharmaceuticals, Inc.
All rights reserved. November 2017 SAUS.PRL.17.11.9044

