Hypoglycemia Prevention Initiative: Frequently Asked Questions

Table of Contents
Hypoglycemia Prevention Initiative: Frequently Asked Questions..............................1
A. Overview ..................................................................................................................1
  1. What is the Hypoglycemia Prevention Initiative?..............................................1
  2. Who is the Endocrine Society?.............................................................................2
  3. Who is Avalere Health?.......................................................................................2
  4. What is the Hypoglycemia Prevention Study (HypoPrevent)?..........................2
  5. Who should participate in the Hypoglycemia Prevention Study?.....................3
B. Benefits and Resources .........................................................................................4
  4. What are the key benefits of participating in HypoPrevent?..............................4
  5. What support will I receive as a participating practice?....................................4
  6. What is the HypoPrevent Toolkit?......................................................................5
C. Expectations and Responsibilities of Participation ..............................................5
  7. How will HypoPrevent’s Intervention and resources be integrated into my practice’s existing clinical workflow?..............................................................5
  8. How can I determine if my practice is ready?.....................................................5
  9. What are the expectations of participants in HypoPrevent?..............................5
10. What is the process for Internal Review Board (IRB) Approval and patient informed consent for HypoPrevent?.................................................................6
11. What are the data collection and submission requirements for HypoPrevent?6

A. Overview
1. What is the Hypoglycemia Prevention Initiative?
As an ongoing effort to reduce the frequency of diabetes-related hypoglycemia in patients with Type 2 diabetes (T2D), Endocrine Society and Avalere Health have been working collaboratively to develop and implement the Hypoglycemia Prevention Initiative. The Initiative engages in various opportunities to support research, advocacy, education, and quality of care. The Hypoglycemia Prevention Study, HypoPrevent, is a central component to the Initiative.

The goals of the Initiative are to:
- Support providers in adopting clinical guidance in practice
- Promote patients’ confidence in managing hypoglycemia
- Test interventions for decreasing the risk of hypoglycemia in clinical settings
2. Who is the Endocrine Society?
The Endocrine Society is the oldest and largest global professional membership organization representing the field of endocrinology with over 18,000 members in 122 countries. Our members care for patients and are dedicated to advancing hormone research and excellence in the clinical practice of endocrinology. These expert scientists and physicians shape the genesis and delivery of patient care from bench to bedside to well-being.

Since 2014, the Endocrine Society has made hypoglycemia a strategic priority, conducting a series of roundtables with key stakeholders from the federal government, provider community, patient advocates, and payers to determine a comprehensive approach to this problem. Endocrine Society, in collaboration with Avalere Health, has now embarked on a multi-year quality improvement project, the Hypoglycemia Prevention Initiative.

3. Who is Avalere Health?
Avalere Health is a healthcare advisory firm headquartered in Washington, D.C., specializing in strategy, policy, and data analysis across health sectors. Avalere delivers a comprehensive perspective, compelling substance, and creative solutions to help organizations make better business decisions. In 2016, Avalere partnered with the Endocrine Society to develop and initiate a multi-year quality improvement project, the Hypoglycemia Prevention Initiative.

4. What is the Hypoglycemia Prevention Study (HypoPrevent)?
HypoPrevent, a central component of the Initiative, is a multi-site quality improvement study designed to test a three-pronged intervention in primary care practices to identify patients at risk of hypoglycemia and assess methods to decrease their risk through the individualization of their A1c goals and/or changes to their medications. HypoPrevent will be implemented in primary care practices in 2 phases, the Feasibility Phase and the Impact Phase. The target population is patients 65 years of age and older who are treated with insulin or sulfonylureas with a most recent A1c level of less than 7%. HypoPrevent will also serve as a vehicle for testing patient safety quality measures intended for eventual use in value-based payment programs.

a. What is the intervention being tested in HypoPrevent?
HypoPrevent is testing a dual-pronged intervention, containing two primary components:

i. **Hypoglycemia Risk Screening Tool**: an evidence-based algorithm to enable providers to identify patients with T2D aged 65 and older within their patient panel who are at risk for medication-related hypoglycemia based on risk factors: prescription for hypoglycemic agents including insulin and/or sulfonylureas and a most recent hemoglobin A1c (A1c) value of < 7.0%

ii. **Provider Hypoglycemia Reduction Clinical Decision Support Tool**: an evidence-based, point-of-care, decision support tool to guide providers in patient assessment, care planning, and monitoring and ongoing evaluation of patients through shared decision-making with a focus on reducing risk of
hypoglycemia through the individualization of A1c goals and/or changes in medication management

b. What is the difference between the Feasibility Phase and the Impact Phase of the HypoPrevent?

More information about HypoPrevent, based on the Phase, can be found in the table below.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Phase 1: Feasibility Phase</th>
<th>Phase 2: Impact Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Goal/Purpose</td>
<td>To assess the feasibility of implementing the intervention in real-world clinical practice, and to inform the final design and implementation of Phase 2</td>
<td>To explore the impact of the interventions on patient outcomes and provider behaviors</td>
</tr>
<tr>
<td>Expected Duration</td>
<td>2 months (enrollment period + baseline visit) + follow-up interviews</td>
<td>9 months (baseline visit + two quarterly follow-up visits) + follow-up surveys</td>
</tr>
<tr>
<td>Endpoints</td>
<td>• Facilitators and barriers to implementation of the intervention • Facilitators and barriers to implementation of the provider self-efficacy</td>
<td>• Impact of hypoglycemia on the patient before and after the intervention • Size of the at-risk population before and after the intervention is implemented, • Change in the both patient and provider self-efficacy in addressing hypoglycemia</td>
</tr>
<tr>
<td>Number of participating practices</td>
<td>5-10 providers across 2 practices</td>
<td>45 providers across 6-10 practices</td>
</tr>
<tr>
<td>Number of enrolled patients per provider</td>
<td>Minimum of 2 patients</td>
<td>15-30 patients</td>
</tr>
<tr>
<td>Number of visits per patient</td>
<td>1 baseline visit</td>
<td>1 baseline visit; up to 2 follow up visits</td>
</tr>
</tbody>
</table>

3. Who should participate in the Hypoglycemia Prevention Study?
HypoPrevent is designed for primary care practices with experience in quality improvement that have an interest in a population management approach to identify patients at risk of hypoglycemia and to mitigate that risk on an individual patient basis. Practices with a large number of older (65+) patients with T2D and without a formal mechanism to identify those with hypoglycemia will have the greatest opportunity to make an impact on their patient population through participation in this study.
B. Benefits and Resources

4. What are the key benefits of participating in HypoPrevent?
There are numerous ways that you and your practice will benefit from HypoPrevent, including improved quality of care and safety for your patients, provider and patient focused resources related to hypoglycemia and shared decision-making, and the chance to have input on developing new quality measures. Some of these benefits include:

For Practices…
- Opportunity to be recognized as a leader and early adopter in hypoglycemia quality improvement
- Opportunities to publish and present on your practice’s experience and results
- Obtain technical assistance with collection of performance data to track and monitor implementation progress

For Providers…
- Improve the safety of your patients with T2D in your busy clinical practice with an intervention that helps you better identify individuals at risk of hypoglycemia and help them mitigate their risk
- Improve your ability to effectively engage patients on the topic of hypoglycemia through easy to use resources
- Provider-focused resources to help facilitate shared-decision making and medication changes designed to maximize patient engagement and reduce the risk of hypoglycemia
- Patient-focused resources to help patients recognize, treat, and avoid hypoglycemia
- Access to educational resources and training to implement guidelines into practice related to setting individualized A1c goals for older patients
- Opportunities for educational activities that may include CME, MOC Part IV, and help meet MIPS Improvement Activities reporting requirements
- Have input on developing new quality measures for diabetes to improve patient safety

5. What support will I receive as a participating practice?
Practices will receiving ongoing support before, during, and after HypoPrevent Intervention period.

Before participating in HypoPrevent, you will have the opportunity to engage in a series of onboarding and training webinars with experts in the field.

During HypoPrevent, participants will have access to individual technical assistance calls with HypoPrevent leaders, opportunities to informally engage with implementation experts, data collection guides, templates, and other resources to facilitate implementation of the Intervention, and feedback reports to monitor improvement and evaluate impact of the Intervention. Finally, resources to support uptake of the interventions with providers and patients will be made available in the Hypoglycemia Prevention Study Toolkit.

After HypoPrevent’s conclusion, participants will be supported as they look to disseminate their results and expand and sustain their efforts to identify patients at risk for hypoglycemia.
6. **What is the HypoPrevent Toolkit?**

The Toolkit is an exclusive resource for participating practices that contains provider and patient resources that are: an evidence-based, easily accessible, supportive of practice-based quality improvement, and aligned with the goals of the Hypoglycemia Prevention Initiative. The Toolkit provides both quality and clinical resources for members of the practice to use in implementing the intervention at every stage of the clinical workflow, in addition to providing resources aimed at patients and helping them better understand and manage their hypoglycemia risk.

**C. Expectations and Responsibilities of Participation**

7. **How will HypoPrevent’s Intervention and resources be integrated into my practice’s existing clinical workflow?**

HypoPrevent is designed to be integrated into the practice’s existing workflow, occurring during regular recurring visits standard for the T2D population and requiring data that is for the most part already being collected. The Intervention is intended to assist primary care providers, who often face competing priorities and limited time with complex patients, to effectively manage this at-risk population by focusing their encounters on the areas in which they can have the greatest impact in reducing risk of hypoglycemia.

8. **How can I determine if my practice is ready?**

We will work with practices that are interested in participating in the study to determine whether they are ready to participate. We will use a combination of surveys and pre-enrollment screening calls to gather information about practices to ensure they fit our eligibility criteria and that our study population is nationally representative.

In order to successfully participate in the study, practices must:

- Have experience in protocol-driven initiatives
- Systematically assign patients to a primary care provider’s panel
- Be able to pull customized patient panel reports from their EHR based on specific patient information such as lab results, age, diagnoses, medication use, etc.

9. **What are the expectations of participants in HypoPrevent?**

Once selected, practice participants will be asked to complete the following tasks:

- Sign a Practice Participation Agreement and Business Associate Agreement (BAA) with HypoPrevent’s Project Team
- Participate in practice onboarding activities, specifically, educational webinars, practice-specific training, and receiving the Practice Welcome Announcement
- Provide HypoPrevent’s Project Team with a report of all patients in their panel, including characteristics used to determine risk
- Conduct outreach to patients identified as at risk for hypoglycemia regarding HypoPrevent participation
- Enroll patients into HypoPrevent and collect informed consent
- Commit to collecting and supporting analysis of data on all study endpoints (different for the Phase in which the practice is participating; see **Question 3c** above for differentiation)
• Complete pre- and post-study assessments including, but not limited to, a provider self-efficacy survey to determine change in providers’ perceptions of their ability to identify and manage patients at risk of hypoglycemia

10. What is the process for Internal Review Board (IRB) Approval and patient informed consent for HypoPrevent?

**IRB Submission:** HypoPrevent protocol will be submitted and approved by Quorum Review IRB, in anticipation that many of the practices may not have access to their own IRB. Quorum is a nationally recognized central IRB whose boards comply fully with relevant U.S. and Canadian regulations and has full accreditation from the Association for the Accreditation of Human Research Protection Programs. All practices will receive information necessary to register their site under Quorum. Practices who are required by their organization to obtain local IRB approval will receive materials to support their application to the local IRB.

**Patient Informed Consent:** Each patient will consent in writing to participation in HypoPrevent prior to any assessments and will be obtained by the provider or a designated member of the practice. Practices should safely store signed informed consent documents.

11. What are the data collection and submission requirements for HypoPrevent?

• Practices will be required to collect data from a variety of different sources for purposes of HypoPrevent, including: patient demographic information, A1c lab test results, patient-reported surveys, provider surveys, and documentation of actions taken during the visit.

• Submission requirements will vary based on the Phase of the study in which the practice is participating. Phase 1 will utilize a combination of EHR data extraction, patient-reported surveys, and Adobe PDF forms; Phase 2 will involve EHR data extraction and submission of data via a web-based application.