Broadening Access to Continuous Glucose Monitoring for Patients With Type 2 Diabetes

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Persons from racial and ethnic minority populations, those in low-income groups, and other socially marginalized groups are disproportionately affected by type 2 diabetes and experience higher disease prevalence, poorer glycemic control, higher rates of diabetes complications, and higher prevalence of comorbid conditions.1,2 Achieving glucose targets that will reduce the risk of diabetes complications, particularly among high-risk groups, is critical to improve the health and well-being of those with diabetes and to reduce health care utilization and expenditures. Yet, diabetes control remains elusive. Self-monitoring of blood glucose, while still a standard part of diabetes self-management, has not been shown to result in self-adjustments to insulin in primary care settings. This represents a significant opportunity gap because 30% of patients with type 2 diabetes are treated with some form of insulin.3

Real-time continuous glucose monitoring (CGM), which measures glucose levels in subcutaneous interstitial fluid as frequently as every 5 minutes, has been shown to improve diabetes control, reduce hypoglycemia, and be cost-effective for patients with type 1 diabetes.4,5 Less research has been conducted among patients with type 2 diabetes, but clinical trials involving patients using intensive insulin regimens (eg, basal/bolus insulin) have shown reductions in hemoglobin A1c (HbA1c) levels and shorter intervals of hypoglycemia.6,7 Several questions remain: Can the results of clinical trials of patients with type 2 diabetes be translated into usual care settings? Can patients with type 2 diabetes who use less intensive insulin regimens benefit from CGM? Can CGM be feasibly implemented in primary care settings, where most of type 2 diabetes management occurs? In this issue of JAMA, the randomized controlled trial (RCT) reported by Martens et al8 and the observational study reported by Karter et al9 provide new data that help provide answers to these questions.

Martens et al8 conducted an RCT of CGM (n = 116) vs blood glucose meter (BGM) monitoring (n = 59) among adults with type 2 diabetes who were taking basal insulin without prandial insulin and were recruited from primary care practices. At 8 months, the mean HbA1c level improved from 9.1% to 8.0% in the CGM group and from 9.0% to 8.4% in the control group (adjusted difference, −0.4% [95% CI, −0.8% to −0.1%]). This effect size may have been greater if the control group had received usual care rather than instructions on how to self-titrate insulin based on BGM data. Compared with the BGM group, the time in range, or the amount of time spent in the target blood glucose range (70-180 mg/dL), was 3.6 hours per day higher, the mean glucose level was 26 mg/dL lower (95% CI, −41 to −12), and the time with glucose levels greater than 250 mg/dL was 3.8 hours per day less in the CGM group (all P < .001). There were also high rates of satisfaction among CGM users.

Karter et al9 conducted a retrospective cohort study of 41,753 adult patients (36,080 with type 2 diabetes, 5,673 with type 1 diabetes) who were treated with insulin and were receiving care at Kaiser Permanente.9 The authors followed the outcomes of those who initiated CGM (3,806 patients) compared with those who did not; the CGM group primarily used basal/bolus insulin regimens, whereas the control group was treated with various forms of insulin. Over the 4-year study period (which ended in December 2018), the authors reported a difference-in-difference reduction in HbA1c level of −0.40% (95% CI, −0.48% to −0.32%) and in rates of emergency department visits and hospitalization for hypoglycemia of 2.7% (95% CI, −4.4% to −1.1%). The net change in HbA1c level was greater among patients with type 2 diabetes (−0.56% [95% CI, −0.72% to −0.41%]) than among patients with type 1 diabetes (−0.34% [95% CI, −0.43% to −0.25%]) (P value for interaction = .003). In addition, a sensitivity analysis revealed a dose-response association between CGM adherence (0, 1, or ≥2 claims for CGM transmitters) and changes in HbA1c level and hypoglycemia health care utilization.

These studies are important for several reasons. First, they confirm that CGM is a technology that can be effectively used by patients with type 2 diabetes to improve glycemic control. The trial by Martens et al8 recruited a diverse sample of patients who have disproportionately had barriers to fully accessing health care and health care–related technology and also have had disproportionately lower rates of adherence to diabetes treatment plans. Most patients in this RCT were non-White persons (53%), had less than a college degree education (55%), and did not have private insurance (58%). Exploratory analyses suggested that the reduction in HbA1c level did not differ across age groups, baseline diabetes control, education level, and diabetes numeracy, thus indicating a broad population benefit for CGM among patients with type 2 diabetes. The observational study from Karter et al9 demonstrated the benefits associated with CGM in usual care settings and found a greater improvement in diabetes control among patients with type 2 diabetes than those with type 1 diabetes.

Second, the clinical trial by Martens et al8 demonstrated the promise of using CGM in primary care settings, where most patients with type 2 diabetes receive their care. This trial, in which study clinicians met with trial participants during
in-person clinic visits followed by virtual visits, provides a model that could be replicated or modified in many primary care practices throughout the US. For example, having an initial consultation with an endocrinologist followed by telehealth visits with advanced practice nurses in an endocrinology practice could allow for download and interpretation of the CGM data in the specialty practice without requiring primary care practices to develop this expertise. A recent telehealth program that included remote monitoring of CGM demonstrated statistically significant reductions in HbA1c levels among 594 patients with type 2 diabetes.10 Project Extension for Community Health Outcomes (ECHO) successfully used remote learning as a venue for subspecialists to train primary care physicians to treat a range of conditions, including complex diabetes care,11 and could be an alternative strategy for integrating CGM usage into primary care practice.

Third, these studies suggest that patients with type 2 diabetes who use less intensive insulin regimens may have similarly robust glycemic benefit as those who require more intensive regimens. In both the clinical trial, in which the intervention group received basal insulin only, and the observational study, in which 97% of the type 2 diabetes CGM group was taking basal/bolus insulin, the difference in HbA1c reduction compared with the group that did not initiate CGM was −0.4%. This has significant implications for health policy. While patients in the RCT were taking basal insulin only and monitoring their blood glucose 3 or more times per week, the current American Diabetes Association grade A guidelines for CGM use include multiple daily injections of insulin (or an insulin pump) and Medicare guidelines require 3 or more daily injections of insulin (or an insulin pump) and self-monitoring of glucose 4 or more times daily.12,13 The RCT by Martens et al8 demonstrates that CGM is effective in patients with type 2 diabetes who are treated with less intensive insulin regimens and adds to the body of evidence that CGM is effective among patients with less intensive blood glucose monitoring.14 The Medicare criteria have created significant administrative barriers to CGM use even for patients who are currently eligible because of the substantial documentation requirements that are unfamiliar, time-consuming, or both to clinicians and their staff. These criteria also create access barriers for patients who could clinically benefit from CGM but are not currently eligible. It is time to revise the Medicare criteria for CGM to reflect the current scientific evidence and simultaneously mitigate disparities in CGM access and diabetes control.13,15

Fourth, the RCT results suggest that patient engagement (ie, improved insulin adherence, changes in diet, or increased physical activity in response to CGM readings) was the most likely source of improved glycemic control because there were no differences in the total amount of insulin between study groups or in the amount of medication adjustments by clinicians. Activated patients are a powerful part of achieving diabetes control.6 Patients in the clinical trial by Martens et al8 reported high rates of satisfaction with the CGM, including high mean “benefits” scores and low mean “hassle” scores, suggesting a willingness of this diverse patient population to engage with the technology. Access to diabetes-related technology, including CGM, has been restricted among marginalized populations. These studies add to the literature by demonstrating that persons from racial and ethnic minority populations, low-income persons, and those with low numeracy want to be engaged, and can successfully be engaged, in diabetes-related technology that enhances self-management and improves diabetes control.

In summary, the studies by Karter et al3 and Martens et al8 provide additional evidence that patients with type 2 diabetes benefit from the use of CGM in terms of improved HbA1c level, time spent in the target blood glucose range, and reduced hypoglycemic episodes. The glycemic benefits may be primarily due to patient factors, such as insulin adherence and lifestyle modifications, and provide a powerful narrative that CGM may be a useful technology that helps control diabetes among multiple patient groups. Important policy changes in Medicare eligibility to CGM for type 2 diabetes and institutional changes that promote its use in primary care will go a long way to improving diabetes control and reducing complications, particularly among the populations most in need. The time has come to broaden access to CGM for patients with type 2 diabetes.

ARTICLE INFORMATION

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Conflict of Interest Disclosures: Dr Peek reported being a paid speaker for PRIME Education and giving a talk on continuous glucose monitoring. Dr Thomas reported no disclosures.

REFERENCES

Realizing the Potential of Maternal Influenza Vaccination

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Influenza viruses cause substantial morbidity and mortality in pregnant women and neonates worldwide annually, with higher incidence during pandemics.1,2 To reduce disease incidence in these vulnerable populations, the World Health Organization recommended in 2012 that countries prioritize pregnant women for influenza vaccination.2 Over 2 decades, compelling evidence has accumulated about the complex interplay among risks of influenza virus infection to the mother, the fetus, and the offspring, vs the benefits and safety of vaccination.1,4

Physiologic changes during pregnancy increase the risk for influenza complications in pregnant women. Severe complications after infection can occur throughout gestation, but risk is greatest during the third trimester. In an individual participant data meta-analysis including 27,699 participants from 9 studies, risk of hospitalization in pregnant women with influenza was substantially higher than in nonpregnant women (adjusted odds ratio, 6.8 [95% CI, 6.0-7.7]).3 Pregnant women with comorbidities, including underlying cardiac conditions, chronic respiratory diseases, and obesity, are at even greater risk of hospitalization from influenza.

Maternal infection and complications can also affect the pregnancy, possibly resulting in fetal demise or preterm birth.1,4 Fetal development might be adversely affected by maternal infection, with effects varying by gestation period.

In a retrospective cohort study of more than 27,000 women, self-reported symptoms of cold or flu with fever during pregnancy was associated with increased risk of several congenital abnormalities when compared with no symptoms of cold or flu, whereas no association was found among women with cold or flu without fever.7 The concern is that these adverse effects might be related to maternal fever (a common occurrence with influenza virus infection) during early gestation or maternal immune activation after infection resulting in systemic inflammation and imbalances in cytokines.8,9 During the first few months of life, neonates are at increased risk of severe influenza because of factors such as waning levels of maternally derived antibodies, undeveloped immune systems, lack of antibodies from prior infections, and ongoing alveolarization and changing lung physiology.10 In turn, adverse effects of infection on mother and offspring might have commensurate psychological, social, and economic implications for the family and society.

Maternal influenza vaccination can provide dual benefits, helping to prevent various complications in pregnant women and their fetuses. Maternal vaccination offers benefits to the infant through transplacental antibody transfer, which can protect infants during the high-risk first 6 months of life when they are not age-eligible for vaccination.2,4 These benefits of vaccination have been well documented, with studies showing immunogenicity in pregnant women, transplacental antibody transfer to the infant, and clinical efficacy and effectiveness against influenza-associated illness in mothers and infants. In a pooled analysis of 3 randomized trials involving 10,002 women, maternal influenza vaccination (with follow-up of infants up to 6 months) with an inactivated vaccine was associated with an efficacy of 50% in mothers (absolute rates of 18.0 vs 36.0 per 1000 person-years) and 35% in infants (absolute rates of 62.3 vs 95.4 per 1000 infant-years) against laboratory-confirmed influenza in Mali, Nepal, and South Africa.4 The clinical trials from Bangladesh and Nepal found that infants born to mothers vaccinated vs unvaccinated against influenza had higher mean birth weight and a reduced risk of low birth weight, but similar findings were not observed in South Africa and Mali.4

Against these documented benefits of vaccination, the potential risks of maternal vaccination require consideration. From the safety perspective, a systematic review and meta-analysis including 48 cohort studies, case-control studies, and randomized clinical trials did not identify any association between maternal influenza vaccination and adverse fetal outcomes including preterm birth, small for gestational age, congenital malformation, or fetal death.3 Concerns also