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Journal Title: BMJ Open Diabetes Research & Care
Volume: Volume 8, Number 1
Publisher: BMJ Publishing Group | 2020-01-01
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1136/bmjdr-2019-001062
Permanent URL: https://pid.emory.edu/ark:/25593/vnpbm

Final published version: http://dx.doi.org/10.1136/bmjdr-2019-001062

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Accessed November 9, 2022 6:43 PM EST
Long-term changes in carbohydrate tolerance, insulin secretion and action in African-American patients with obesity and history of hyperglycemic crises

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ABSTRACT

Introduction Many African-Americans (AA) with obesity with newly diagnosed diabetes presenting with diabetic ketoacidosis (DKA) or severe hyperglycemia (SH) discontinue insulin therapy and achieve near-normoglycemia remission (hemoglobin A1c (HbA1c) <7%, fasting blood glucose (FBG) <130 mg/dL) and able to be managed on oral antidiabetic agents (OAD) during follow-up. Using combined data from two randomized controlled trials, we assessed long-term carbohydrate tolerance and changes in insulin sensitivity and insulin secretion.

Research design and methods Seventy-five participants with DKA (n=33) and SH (n=42) underwent 2-hour 75g oral glucose tolerance test (OGTT) after insulin discontinuation and every 6 months until hyperglycemia relapse (FBG ≥130 mg/dL, HbA1c >7% or two random BG ≥180 mg/dL) while treated with OAD (metformin, sitagliptin or pioglitazone) or placebo. Glucose tolerance status was defined as per the American Diabetes Association. Sensitivity index (S) was calculated by oral minimal incretin model, insulin secretion as the incremental area under the curve of insulin (IncreAUCi) and disposition index (DI) as S = IncreASC.

Results During remission, OGTT showed normal glucose tolerance (NGT) (n=9 (12%), prediabetes (n=34 (45%)) and diabetes (n=32 (43%)). DI and S were higher in patients with NGT versus prediabetes versus diabetes (p<0.001), while IncreAUC was not significantly different among NGT, prediabetes and diabetes (p=0.14). Achieving NGT status did not prolong near-normoglycemia remission. OAD treatment significantly prolonged hyperglycemia relapse-free survival (log-rank p=0.0012) compared with placebo and was associated with lower hyperglycemia relapse (HR: 0.45, 95% CI: (0.21 to 0.96), p=0.04).

Conclusions In AA patients with obesity with history of DKA and SH, near-normoglycemia remission is associated with improved insulin secretion and action with half of patients achieving NGT or prediabetes, and only half having diabetes on OGTT. NGT and prediabetes on OGTT were not associated with prolonged hyperglycemia relapse-free survival.

Trial registration number NCT01099618, NCT00426413.

INTRODUCTION

More than half of newly diagnosed African-Americans with obesity presenting with unprovoked diabetic ketoacidosis (DKA) exhibit clinical, metabolic, immune and genetic features of type 2 diabetes during follow-up. At presentation, these patients have severe pancreatic beta (β)-cell dysfunction and increased insulin resistance.

Significance of this study

What is already known about this subject?
► Approximately 50% of African-Americans who present with new-onset diabetic ketoacidosis (DKA) or severe hyperglycemia at new onset of diabetes achieve near-normoglycemia remission after intensive insulin treatment.
► The period of near-normoglycemia remission is variable with some patients needing oral antidiabetic agents while some patients do not require any oral antidiabetic agents.

What are the new findings?
► The patients who achieve near-normoglycemia remission have a heterogeneous phenotype.
► Despite presenting with DKA or severe hyperglycemia, oral glucose tolerance testing (OGTT) at near-normoglycemia remission shows that ~50% of patients do not have diabetes while 12% have a normal OGTT.
► Normal glucose tolerance is characterized by higher insulin sensitivity compared with prediabetes or diabetes at remission.
► Initiation of oral antidiabetic agents is important to sustain near-normoglycemia remission.

How might these results change the focus of research or clinical practice?
► Further studies need to be performed regarding the long-term treatment of patient who have normal glucose tolerance.
Furthermore, our group and others have shown that African-American patients with obesity who present with severe hyperglycemia without ketoacidosis have similar impaired β-cell dysfunction and insulin resistance. In ~70% of patients, aggressive management with insulin in patients with obesity presenting with DKA and severe hyperglycemia results in significant improvement in pancreatic β-cell function and insulin sensitivity, sufficient to allow discontinuation of insulin and maintain near-normoglycemia remission for several months to years with lifestyle modification alone or with oral antidiabetic agents (OAD).

The duration of near-normoglycemia remission is variable. We performed two studies that showed that initiation of metformin, sitagliptin or pioglitazone after insulin discontinuation prolongs remission. In addition, the patients who achieve remission represent a heterogeneous population. In a study by Banerji et al, oral glucose tolerance test (OGTT) performed at time of near-normoglycemia remission showed that 16.7% of patients achieved normalization of the OGTT while 33.3% had impaired glucose tolerance and 50% remained with diabetes. In the Diabetes Prevention Program (DPP), people who achieved normalization of the OGTT at least once during the study period showed a 56% lower incidence of diabetes compared with people who stayed with prediabetes irrespective of being randomized to physical activity, placebo or metformin. In the DPP study, normalization of the OGTT was associated with higher insulin sensitivity and secretion. In this study, we aimed to examine the differences in insulin secretion and action based on OGTT status at near-normoglycemia remission and whether OGTT status influenced long-term hyperglycemia relapse-free survival independent of OAD use. Since each study had a small number of participants with a normalized OGTT, inclusion/exclusion criteria were similar in both studies and performed at the same center, we combined the OGTT results from both studies to assess the effect of OGTT status on hyperglycemia relapse-free survival.

### RESEARCH DESIGN AND METHODS

#### Study participants

This study was a post hoc analysis of data from two randomized controlled studies conducted at Grady Memorial Hospital in Atlanta, Georgia, USA. Briefly, participants were of African-American ancestry, aged 18–65 years, overweight or obese (body mass index (BMI) ≥28 kg/m²), presenting with unprovoked DKA or severe hyperglycemia at diagnosis of diabetes. Participants with contraindications to any of the medications used in the individual randomized controlled studies (metformin, sitagliptin or pioglitazone), history of pancreatitis, moderate or severe heart failure, significant anemia, pregnant or unable to consent were excluded. DKA was defined as blood glucose (BG) ≥250 mg/dL (13.9 mmol/L), pH <7.30, bicarbonate levels <18 mmol/L and positive ketonemia defined as beta-hydroxybutyrate ≥3 mmol/L. Severe hyperglycemia was defined as BG >100 mg/dL (5.6 mmol/L) and 2-hour BG <140 mg/dL (7.8 mmol/L). Prediabetes was defined as FBG between 100 and 125 mg/dL (5.5–6.9 mmol/L).

#### Measured outcomes and calculations

Glucose levels from the OGTT at month 0 was used to define the different glucose tolerance groups as recommended by the American Diabetes Association. Normal glucose tolerance was defined as FBG <100 mg/dL (5.6 mmol/L) and 2-hour BG <140 mg/dL (7.8 mmol/L). Prediabetes was defined as FBG between 100 and 125 mg/dL (5.5–6.9 mmol/L) or 2-hour BG between 140 and 199 mg/dL (7.8–11.1 mmol/L). Diabetes was defined as

#### Study protocol

After acute resolution of DKA/hyperglycemia, all participants were treated intensively with subcutaneous insulin to a target fasting BG (FBG) and premeal BG between 70 and 130 mg/dL (3.9–7.2 mmol/L). Insulin was decreased up to 12 weeks until near-normoglycemia remission was achieved. Near-normoglycemia remission was defined as FBG <130 mg/dL (7.2 mmol/L) and random BG <180 mg/dL (10 mmol/L), at least a week off insulin therapy and with a hemoglobin A1c (HbA1c) <7% (53 mmol/L). HbA1c and BG levels were checked a week after insulin discontinuation. After discontinuing insulin, participants were randomized to an oral antidiabetic agent (OAD) or placebo. In one study, participants were randomized to three groups: sitagliptin 100 mg daily, metformin 1000 mg daily or placebo. In the second study, participants were randomized to two groups: pioglitazone 30 mg daily or placebo. Prior to randomization, all participants received a 75 g OGTT. After randomization (month 0), study participants were followed at the Grady Diabetes Clinic every 4 weeks until 3 months and then every 3 months until 27 months or until they had a hyperglycemic relapse while on oral medication.

All randomized participants underwent OGTTs at randomization (0 months), 3, 9, 15, 21 and 27 months or until hyperglycemia relapse (defined as an FBG >130 mg/dL (7.2 mmol/L), random blood glucose >180 mg/dL (10 mmol/L) for two consecutive days or HbA1c ≥7% (53 mmol/L)).

#### Medications

Sitagliptin was provided by Merck & Co and pioglitazone by Takeda Pharmaceuticals. Metformin was purchased from the research pharmacy at Grady Memorial Hospital. The pharmaceutical companies were not involved in study design, subject recruitment, data analyses or writing of the manuscript.

#### Modified oral glucose tolerance test

After an 8–10 hours overnight fast, a 75 g OGTT was performed in participants between 08:00 and 10:00 hours. Glucose and insulin levels were checked at 0, 15, 30, 60, 90 and 120 min.
Table 1  Baseline characteristics of patients presenting with hyperglycemic crises at near-normoglycemia remission according to OGTT

<table>
<thead>
<tr>
<th></th>
<th>NGT (n=9)</th>
<th>Prediabetes (n=34)</th>
<th>Diabetes (n=32)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>45±13</td>
<td>45±10</td>
<td>49±10</td>
<td>0.45</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>39.6±16.8</td>
<td>34.9±10.3</td>
<td>36.6±8.8</td>
<td>0.76</td>
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<tr>
<td>Sex, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1 (11)</td>
<td>15 (44)</td>
<td>12 (37.5)</td>
<td>0.22</td>
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<tr>
<td>Male</td>
<td>8 (89)</td>
<td>19 (56)</td>
<td>20 (62.5)</td>
<td></td>
</tr>
<tr>
<td>Diagnosis, n (%)</td>
<td></td>
<td></td>
<td></td>
<td>0.95</td>
</tr>
<tr>
<td>DKA</td>
<td>4 (44)</td>
<td>14 (41)</td>
<td>15 (47)</td>
<td></td>
</tr>
<tr>
<td>Severe hyperglycemia</td>
<td>5 (56)</td>
<td>20 (59)</td>
<td>17 (53)</td>
<td></td>
</tr>
<tr>
<td>HbA1c, %</td>
<td>13.0±1.8</td>
<td>12.7±2.2</td>
<td>13.0±2.4</td>
<td>0.82</td>
</tr>
<tr>
<td>Family history of type 2 diabetes, n (%)</td>
<td>6 (67)</td>
<td>30 (88)</td>
<td>27 (84)</td>
<td>&gt;0.99</td>
</tr>
<tr>
<td>Insulin dose, units/kg/day</td>
<td>0.46±0.17</td>
<td>0.57±0.21</td>
<td>0.67±0.30</td>
<td>0.13</td>
</tr>
<tr>
<td>Duration of insulin treatment (weeks)</td>
<td>&gt;0.99</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤8</td>
<td>3 (33)</td>
<td>13 (38)</td>
<td>13 (41)</td>
<td></td>
</tr>
<tr>
<td>&gt;8</td>
<td>6 (67)</td>
<td>21 (62)</td>
<td>19 (59)</td>
<td></td>
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<tr>
<td>OAD, n (%)</td>
<td>7 (78)</td>
<td>27 (82)</td>
<td>16 (50)</td>
<td>0.07</td>
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<tr>
<td>Metformin</td>
<td>2 (22)</td>
<td>7 (21)</td>
<td>7 (22)</td>
<td></td>
</tr>
<tr>
<td>Sitagliptin</td>
<td>1 (11)</td>
<td>10 (30)</td>
<td>5 (16)</td>
<td></td>
</tr>
<tr>
<td>Pioglitazone</td>
<td>4 (44)</td>
<td>10 (30)</td>
<td>4 (13)</td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>2 (22)</td>
<td>6 (18)</td>
<td>16 (50)</td>
<td></td>
</tr>
</tbody>
</table>

Data are expressed as mean±SD unless otherwise noted. OAD, NGT (fasting glucose <100 mg/dL and 2-hour glucose <140 mg/dL), prediabetes (fasting glucose 100–125 mg/dL or 2-hour glucose 140–199 mg/dL), diabetes (fasting glucose ≥126 mg/dL or 2-hour glucose ≥200 mg/dL).

BMI, body mass index; DKA, diabetic ketoacidosis; HbA1c, hemoglobin A1c; NGT, normal fasting glucose/normal glucose tolerance; OAD, oral antidiabetic agent; OGTT, oral glucose tolerance test.

FBG ≥126 mg/dL (7.0 mmol/L) or 2-hour BG ≥200 mg/dL (11.1 mmol/L).

Insulin sensitivity (S_i) and β-cell function were calculated using OGTT-derived measures. We had previously validated our 120 min 6-point oral minimal model S_i, which was similar to the 120 min 7-point minimal model. Therefore, we calculated S_i using glucose and insulin levels from the 6-point OGTT using the oral minimal model. Incremental area under the curve of insulin levels (IncreAUC_i) using the trapezoidal method was used to calculate insulin secretion. Disposition index was calculated as the product of S_i and IncreAUC_i.

Analytic techniques

Assays for glucose and insulin were performed at the Endocrinology/Lipoprotein Laboratory at the University of Tennessee Health Sciences Center. HbA1c and GAD-65 antibody tests were measured at the central laboratory at Grady Memorial Hospital.

Statistical analyses

The main outcome was to assess differences among the different OGTT categories in disposition index, S_i and IncreAUC_i. Since both studies had similar inclusion/exclusion criteria and conducted at the same site, data were combined for both studies. In each of the individual randomized studies, there were no differences between the DKA and severe hyperglycemia groups in hyperglycemia relapse-free survival. Therefore, the DKA and severe hyperglycemia groups were also combined for this analysis. The metformin, sitagliptin and pioglitazone groups had small numbers, therefore all three medication groups were combined. Hyperglycemia relapse-free survival was compared between the different glucose tolerance groups using Kaplan-Meier curves and log-rank tests. We performed univariate and multivariate Cox proportional hazards analysis to evaluate the associations between multiple factors and hyperglycemia relapse-free remission. Continuous variables were compared using Kruskal-Wallis tests for three group comparisons and Wilcoxon rank-sum tests for two group comparisons. Continuous variables are expressed as mean±SD. Discrete variables were compared using χ² tests and expressed as per cent. All analyses were performed using SAS V.9.4 (Cary, North Carolina, USA).

RESULTS

Seventy-five participants with DKA or severe hyperglycemia underwent an OGTT after near-normoglycemia...
remission within 1 week after stopping insulin therapy. Of them, 12% (n=9) had normal glucose tolerance, 45% (n=34) with prediabetes and 43% (n=32) with diabetes. There were no differences in baseline demographics between the normal glucose tolerance, prediabetes and diabetes groups (table 1). Even though not statistically significant, a higher percentage of participants with normal glucose tolerance and prediabetes were randomized to the OAD compared with the placebo group (table 1). Treatment allocation to OAD and placebo is shown in table 2. There was one participant who received an OGTT at remission but did not get randomized to an OAD. Therefore, 50 participants were randomized to OAD and 24 were randomized to placebo after near-normoglycemia remission (table 2). Of these, 32% received metformin and sitagliptin while 36% received pioglitazone. There were no differences in the age, BMI, family history of type 2 diabetes, hemoglobin A1c or insulin doses (table 2). There were significantly more participants with severe hyperglycemia in the placebo group compared with the OAD group (table 2).

In the participants who had normal glucose tolerance at remission, disposition index was significantly higher than in participants who had prediabetes and diabetes, p<0.001 (figure 1A). This difference in disposition index was due to differences in insulin sensitivity (S) rather than insulin secretion (IncreAUCi). Participants with normal glucose tolerance had higher S compared with prediabetes or diabetes group, p<0.001 (figure 1B), while IncreAUCi did not differ (figure 1C). Disposition index,
Figure 2  Disposition index (A), insulin sensitivity (B) and incremental area under the curve (C) did not differ at time of remission in patients randomized to oral antidiabetic agents (black bars) compared with placebo (clear bars). Data are shown as mean±SEM. incremental AUCi, incremental area under the curve of insulin; OAD, oral antidiabetic agents.

Figure 3  Survival curve of hyperglycemia relapse-free survival after near-normoglycemia remission according to oral glucose tolerance test status. Hyperglycemia relapse-free survival did not differ among patients with normal glucose tolerance, prediabetes and diabetes (log-rank p=0.20).

Figure 4  Survival curve of hyperglycemia relapse-free survival after remission in patients with diabetic ketoacidosis and severe hyperglycemia. Hyperglycemia relapse-free survival in patients randomized to oral antidiabetic agents (OAD) was longer compared with patients randomized to placebo, log-rank p=0.0012.

Figure 3  Survival curve of hyperglycemia relapse-free survival after near-normoglycemia remission according to oral glucose tolerance test status. Hyperglycemia relapse-free survival did not differ among patients with normal glucose tolerance, prediabetes and diabetes (log-rank p=0.20).

DISCUSSION

Our study shows that African-American patients with obesity presenting with DKA and severe hyperglycemia who achieve near-normoglycemia represent a heterogeneous group. Despite presenting with DKA and/or severe hyperglycemia, 57% participants achieved a normal

S_i and IncreAUC_i calculated from OGTT at remission did not differ significantly in participants who were randomized to OAD compared with placebo (figure 2).

The median time to hyperglycemia relapse or end of follow-up was 392 days. Hyperglycemia relapse-free survival did not differ significantly according to the OGTT groups (figure 3); however, initiation of an OAD was associated with significantly increased hyperglycemia relapse-free survival (log-rank p=0.0012) (figure 4). There were no significant differences in prolongation of near-normoglycemia remission among participants receiving metformin, pioglitazone or sitagliptin. Univariate analysis showed that receiving an OAD was associated with lower risk of hyperglycemia relapse compared with receiving placebo (HR 0.33, 95% CI (0.16 to 0.67), p=0.002). Presentation with DKA was also associated with lower risk of hyperglycemia relapse compared with participants presenting with severe hyperglycemia (HR 0.39, 95% CI (0.17 to 0.87), p=0.02). In a multivariable Cox hazards model, after adjustment for OGTT status, initiation of OAD was still significantly associated with lower risk of hyperglycemia relapse (HR 0.37, 95% CI (0.18 to 0.77), p=0.008). In a multivariate model which adjusts for age, OAD use and the presentation with DKA versus severe hyperglycemia, OAD use was associated with lower risk of hyperglycemia relapse (HR 0.45, 95% CI (0.21 to 0.96), p=0.04) while presentation with DKA or severe hyperglycemia did not remain significant (p=0.07).
glucose tolerance or prediabetes at near-normoglycemia remission, with only 43% of participants having diabetes on OGTT. Normalization of the OGTT was associated with higher disposition index than prediabetes or diabetes, however, there was no association with duration of hyperglycemia relapse-free survival.

Our results are in agreement with the report by Banerji et al., who reported in black participants with obesity presenting with DKA and severe hyperglycemia, 17% had normalized OGTT and with a third of the participants with impaired glucose tolerance and another half of the participants with diabetes. In that study, attaining normal glucose tolerance status did not predict long-term relapse-free survival. Presentation with lower glucose level and female sex led to longer hyperglycemia relapse-free survival over an 8-year period.

We found that normalization of OGTT was characterized by higher insulin sensitivity rather than secretion compared with prediabetes and diabetes at near-normoglycemia remission. These data differ from the DPP where participants with prediabetes who were able to normalize their glucose tolerance had higher insulin sensitivity and secretion. Data from Banerji et al. were similar to our study indicating no differences in insulin secretion among the three groups, but insulin sensitivity was significantly higher in the participants who attained a normal OGTT. Similar to our study, insulin secretion was determined by AUC with insulin levels from the OGTT time points. We likely did not see a difference in insulin secretion due to defects in insulin clearance in our participants with prediabetes and diabetes. Insulin clearance is directly correlated with insulin sensitivity and insulin clearance is decreased in insulin-resistant individuals with obesity. It is possible that insulin clearance is impaired in the more insulin-resistant participants who remained with prediabetes and diabetes leading to lack of differences in our measures of insulin secretion.

We investigated the effect of antidiabetic therapy on hyperglycemia remission after insulin discontinuation. Initiation of an OAD resulted in prolonged remission in our cohort. More importantly, initiation of an OAD was the only predictor for sustaining near-normoglycemia remission independent of having normalization of the OGTT, prediabetes or diabetes on OGTT. The effect of initiation of an OAD was significant even when adjusted for presentation with DKA or severe hyperglycemia. This is consistent with published studies from our group and others where initiation of an oral agent prolonged remission after discontinuation of insulin therapy in patients with a history of DKA or severe hyperglycemia.

There are several limitations to our study. Since this is a post hoc analysis of two randomized controlled studies and participants were already started on an OAD, we were not able to assess the natural course of glucose control based on the different OGTT groups at remission. Metformin, sitagliptin and pioglitazone have differing mechanisms of actions, which could have affected hyperglycemia relapse. However, we did not have adequate numbers to discern the differences in insulin sensitivity and secretion in response to metformin, sitagliptin or pioglitazone. Since one study had two and the other study had three randomization groups, when data were combined, the study was biased toward more participants randomized to the placebo group. We also did not have sample to measure C-peptide levels to calculate insulin clearance. Nevertheless, this study shows important metabolic differences in patients who undergo remission and emphasized the importance of starting OAD after remission from insulin.

CONCLUSIONS

In summary, we showed that African-Americans with obesity who present with DKA and severe hyperglycemia with near-normoglycemia remission represent a heterogeneous group of patients. Despite presenting with DKA and severe hyperglycemia, more than half of patients have normal glucose tolerance or prediabetes, with less than half having diabetes on OGTT at the time of insulin remission. Normalization of OGTT was characterized by higher insulin sensitivity rather than insulin secretion. Although initiation of an OAD was the significant predictor of long-term near-normoglycemia remission, more studies are needed to explore the underlying pathophysiology of patients achieving a normal glucose tolerance. Future studies with larger number of patients are needed to characterize metabolic parameters along with the natural clinical course in African-Americans with obesity presenting with hyperglycemic crises.

Contributors PV conceived the study, wrote the first draft and edited the manuscript. DS performed the minimal model analysis. DDS and IIA conducted the study. LP performed data analysis. GEU conceived and performed the study. All authors critically edited the manuscript. PV is the guarantor of the work.

Funding PV is supported in part by K12HD085850 and K23DK113241-01A1. This work was funded by K08DK0830351 (DDS) and was supported by the National Center for Advancing Translational Sciences Award number (UL1TR002378) and 1P30DK111024-01 from the National Institutes of Health and National Research Resources. GEU is partly supported by research grants from the NIH/NATS UL1 TR002378 from the Clinical and Translational Science Award program, and 1P30DK111024-01 from NIH and National Center for Research Resources.

Competing interests PV has received consulting fees from Boehringer-Ingelheim and Merck. GEU has received unrestricted research support for inpatient studies (to Emory University) from Novo Nordisk, Dexcom and Sanofi. DDS has received fees for serving on the Speaker Bureau for Novo Nordisk, Merck and Sanofi. The other authors do not have any significant conflicts of interest.

Patient consent for publication Not required.

Ethics approval The Institutional Review Board at Emory University approved to report the combined results of the two randomized controlled studies (NCT01099618 and NCT004246413, www.ClinicalTrials.gov).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request from the PI. Please email at pvellan@emory.edu for any requests or questions.

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