Gestational Diabetes

GDM: Full Recommendations and Supporting Evidence (2016)

Recommendations

Below you will find a list of recommendations in the Gestational Diabetes Mellitus (GDM) Evidence-Based Nutrition Practice Guideline organized by nutrition care process and topic.

The project started with a review of the 2008 recommendations. Selected recommendations were reviewed. The Summary of Changes 2016 provides an overview of the recommendation revisions and updates.

To see the Recommendation Summary, just click on the Recommendation title. Also view the Executive Summary of Recommendations or print the 2016 guideline in PDF (Link to be Added) format.

Screening and Referral

GDM: Referral to an RDN

Nutrition Assessment

GDM: Nutrition Assessment

GDM: Assessment of Food/Nutrition-related History
GDM: Assessment of Anthropometric Measurement
GDM: Assessment of Biochemical Data, Medical Tests, and Procedures
GDM: Assessment of Nutrition-Focused Physical Findings and Client History

Nutrition Diagnosis

None.

Nutrition Intervention

GDM: Medical Nutrition Therapy

GDM: Medical Nutrition Therapy (MNT)
GDM: Frequency and Duration of MNT

GDM: Calories

GDM: Macronutrients

GDM: Macronutrient Requirements
GDM: Carbohydrate Prescription
GDM: Carbohydrate and Post Prandial Breakfast Glycemia

GDM: Vitamins and Minerals

GDM: Dietary Vitamin and Mineral Intake
GDM: Vitamin and Mineral Supplementation

GDM: Meal and Snack Distribution

GDM: High-Intensity Sweeteners

GDM: Alcohol

GDM: Physical Activity

Nutrition Monitoring and Evaluation

GDM: Nutrition Monitoring and Evaluation

Recommendation(s)

GDM: Referral to an RDN 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section.

- Recommendation(s)

GDM: Referral to an RDN

Pregnant women who are diagnosed with gestational diabetes mellitus (GDM), should be referred to a registered dietitian nutritionist (RDN) for medical nutrition therapy (MNT). Individualized MNT is important in helping pregnant women with GDM achieve and maintain normal glycemic levels and appropriate weight gain, while meeting essential nutrients for pregnancy to promote positive maternal and fetal outcomes.

Rating: Strong

Imperative

- Risks/Harms of Implementing This Recommendation

There are no potential risks or harms associated with the application of this recommendation.

- Conditions of Application

It is preferable to refer to an RDN who has experience in working with patients who have diabetes or who specialize in diabetes during pregnancy.

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andeal.org
management [e.g., Certified Diabetes Educator (CDE)].

**Potential Costs Associated with Application**

Costs of MNT sessions and reimbursement vary. However, MNT is essential for improved outcomes.

**Recommendation Narrative**

The recommendation GDM: Referral of Women with GDM to an RDN is based on the American Diabetes Association’s Standards of Medical Care in Diabetes 2016 and the Endocrine Society’s Diabetes and Pregnancy: An Endocrine Society Clinical Practice Guideline (Blumer et al, 2013) provide support for the recommendation as follows:

- **In Management of Diabetes in Pregnancy, the ADA makes the following clinical practice recommendation for women with gestational diabetes mellitus (GDM) (ADA 2016):**
  - Lifestyle changes are an essential component of management of GDM and may suffice for treatment for many women.
  - Medications should be added if needed to achieve glycemic targets. Rating: Level of Evidence: A**
  - Summary of support for the recommendation:
    - “After diagnosis, treatment starts with MNT, physical activity and weight management depending on pregestational weight and glucose monitoring aiming for the targets recommended by the Fifth International Workshop-Conference on GDM.”
    - “GDM is characterized by increased risk of macrosomia and birth complications and an increased risk of maternal diabetes after pregnancy. Although there is some heterogeneity, many randomized controlled trials suggest that the risk of GDM may be reduced by diet, exercise, and lifestyle counseling.”

The Endocrine Society’s Diabetes and Pregnancy: An Endocrine Society Clinical Practice Guideline (Blumer et al, 2013) make the following recommendations:

- **Management of Elevated Blood Glucose**
  - “2.3b. We recommend that the initial treatment of gestational diabetes should consist of MNT and daily moderate exercise for 30 minutes or more. Rating: 1/+++ (Strong recommendation / Moderate quality evidence)”
  - Summary of support for the recommendation:
    - “Lifestyle therapy for GDM results in a lower incidence of reduced birth weight, large-for-gestational-age births, and preclampsia. Both aerobic exercise and non-weight-bearing exercise have been shown to lower blood glucose levels in women with gestational diabetes.**
    - Nutrition therapy and weight gain targets for women with overt or gestational diabetes
      - “4.1. ‘We recommend medical nutrition therapy for all pregnant women with overt or gestational diabetes to help achieve and maintain desired glycemic control while providing essential nutrient requirements. Rating: 1/++ (Strong recommendation / Low quality evidence)”
    - Summary of support for the recommendation:
      - “Although nutrition intervention for overt diabetes and GDM is a fundamental treatment modality, there is a paucity of evidence-based data on this topic. Nevertheless, nutrition therapy has been shown to improve glycemic control for people living with overt diabetes and for women with GDM.”

**Recommendation Strength Rationale**

This topic was not included in the EAL systematic review. The Academy of Nutrition and Dietetics and the GDM Expert group concur with the American Diabetes Association’s Standards of Medical Care in Diabetes 2016 recommendation rating for “Management of Diabetes in Pregnancy (GDM)” and The Endocrine Society’s Diabetes and Pregnancy: An Endocrine Society Clinical Practice Guideline recommendation ratings for “Nutrition Therapy and Weight Gain Targets for Women with Overt or GDM” and Management of Elevated Blood Glucose.

**Minority Opinions**

None.

**Supporting Evidence**

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

**References**

**References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process**


**Gestational Diabetes**

**Gestational Diabetes (GDM) Guideline (2016)**

**Recommendations Summary**

**GDM: Nutrition Assessment 2016**

*Click here* to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the **Supporting Evidence Section** below.

**Recommendation(s)**

**GDM: Assessment of Food/Nutrition-related History of Women with GDM**

The registered dietitian nutritionist (RDN) should assess the food and nutrition-related history of women with gestational diabetes mellitus (GDM) including, but not limited to:

- Food, beverage and nutrient intake including:
  - Calorie intake
  - Types and amount of carbohydrate (including fiber), fat, protein; with special attention to high calorie, low-nutrient dense foods such as desserts, candy, sugar-sweetened beverages
  - Serving sizes
  - Meal and snack patterns, including frequency and duration
  - Recent changes
  - Preferences, avoidance, intolerances, allergies including:
    - In relationship to gastrointestinal discomforts (e.g., nausea, vomiting, heartburn, constipation, ptyalism)
    - Reaction to or changes in food tastes/smells related to pregnancy
  - Cultural and religious considerations.

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andeal.org
• Appetite and changes in appetite
• Eating environment and meals eaten away from home
• Diet history and behavior: previous diets and diet adherence, disordered eating
• Factors affecting access to food: Psychosocial/economic issues (e.g., social support) impacting nutrition therapy
• Method of food preparation, food safety
• Pharmacologic therapy (including insulin or oral glucose-lowering agent)
• Substance use: Alcohol, tobacco, caffeine, recreational drugs
• Use of dietary supplements, prenatal vitamins, over-the-counter medications, complementary and/or herbal
• Knowledge, beliefs or attitudes: Motivation, readiness to change, self-efficacy; willingness and ability to make lifestyle changes
• Physical activity and function: Exercise patterns, functionality for activities of daily living, sleep patterns.

Assessment of these factors is needed to effectively determine nutrition diagnoses and formulate a nutrition care plan. Inability to achieve optimal nutrient intake may contribute to poor outcomes.

**Rating: Consensus**

**Imperative**

**GDM: Assessment of Anthropometric Measurement of Women with GDM**

The RDN should assess the following anthropometric measurements in women with GDM, including but not limited to:

- Height, current weight, pre-pregnancy weight and body mass index (BMI)
- Weight changes during pregnancy.

Assessment of these factors is needed to effectively determine nutrition diagnoses and formulate a nutrition care plan.

**Rating: Consensus**

**Imperative**

**GDM: Assessment of Biochemical Data, Medical Tests, and Procedures of Women with GDM**

The RDN should evaluate available data of women with GDM and recommend as indicated: Biochemical data, medical tests and procedures including, but not limited to:

- Glycemic tests: Glucose challenge test (GCT), oral glucose tolerance test (OGTT), glycosylated hemoglobin (A1C), fasting glucose, random glucose.
- Use of self-monitoring blood glucose (SMBG) meters and urinary ketones, if recommended
- Gestational and fetal testing (e.g., ultrasounds, biophysical profile, non-stress testing)
- Nutritional anemia profile (e.g., hemoglobin, hematocrit, folate, B12, iron)
- Vitamin D and other micronutrient levels, as appropriate
- Thyroid function
- Kidney function.

Assessment of these factors is needed to effectively determine nutrition diagnoses and formulate a nutrition care plan.

**Rating: Consensus**

**Imperative**

**GDM: Assessment of Nutrition-Focused Physical Findings and Client History of Women with GDM**

The RDN should evaluate available data regarding the client history and nutrition-focused physical findings of women with GDM including, but not limited to:

**Patient/Family/Client Medical/Health history**

- Age
- Single or multiple fetuses
- Weeks of gestation; estimated date of delivery (EDD); method of delivery
- Previous obstetric history including GDM
- Risk factors for developing GDM of diabetes, including family history of diabetes
- General health; vital signs
- Pertinent medical and dental history including other diseases, conditions and illnesses
- Gastrointestinal discomforts: Nausea, vomiting, diarrhea, constipation, heartburn and ptyalism
- Health literacy and numeracy
- Education and occupation
- Social history: Psychological/socioeconomic factors (e.g., social support).

Assessment of these factors is needed to effectively determine nutrition diagnoses and formulate a nutrition care plan.

**Rating: Consensus**

**Imperative**

- **Risks/Harms of Implementing This Recommendation**
  
  There are no potential risks or harms associated with the application of these recommendations.

- **Conditions of Application**
  
  - RDNs should be appropriately trained to conduct a nutrition-focused physical exam
  - If necessary data are not available, the RDN should use professional judgment to request or obtain addition data
  - Women who have complicating conditions such as renal disease or eating disorders may require more indepth or specialized nutrition assessments [American Diabetes Association (ADA), 2016]
  - RDNs should be alert to psychosocial stressors, such as family and household strain, verbal or physical abuse, exposure to discrimination, food insecurity, unemployment, low resources, major or catastrophic life events and anxiety about the current pregnancy. Such stressors may indicate need for further screening and referral to a mental health professional for early treatment to prevent adverse pregnancy outcomes (Kaiser and Campbell, 2014).

- **Potential Costs Associated with Application**

  Accessibility and costs of additional testing should be considered.

- **Recommendation Narrative**

  The purpose of the nutrition assessment is to identify nutrition-related problems, their causes and their significance. Relevant data is verified and interpreted by the RDN through an ongoing, non-linear and dynamic process of collecting data and continual analysis of the patient or client's status, compared to specified criteria (eNCPT, 2016). Nutritional assessment encompasses changes in anthropometric, biochemical and clinical indicators throughout the course of pregnancy (Kaiser and Campbell, 2014). Data are obtained from the patient or client through interview, observation and measurements or may come from the medical record or other health care providers. Nutrition assessment findings are then documented in nutrition diagnosis statements and nutrition intervention goal setting (eNCPT, 2016).

  Nutrition assessment is organized under five domains (categories). These are: Food/Nutrition-Related History; Anthropometric Measurements; Biochemical Data, Medical Tests, and Procedures; Nutrition-Focused Physical Findings; Client History (eNCPT, 2016). The last two are combined in the narrative below.

- **Food or Nutrition-Related History**

  Food and nutrition-related history pertinent to diabetes and pregnancy include:
Dietary history includes a thorough review of usual food intake, pattern of intake (timing, meals and snacks) and previous history of diet adherence (ADA, 2013).

Educational knowledge, such as nutrition and meal-planning skills, barriers to dietary compliance, such as lack of family support, daily schedule or economic issues, etc. (ADA, 2013).

Macronutrient (especially carbohydrate and fiber) and micronutrient dietary intake (ADA, 2013).

Vitamin and mineral supplement use (prenatal and non-prenatal) or use of natural remedies, such as herbs or alternative therapies (ADA, 2013).

Food allergies or intolerances (ADA, 2013).

Use of alcohol, tobacco, caffeine or other substances [Joslin Diabetes Center & Joslin Clinic (Joslin), 2011].

Medications: Prescription (diabetes-related, non-diabetes-related) over the counter medications (ADA, 2013).

Screening for other nutrition risks (e.g., eating disorders, pica, adolescence, low literacy, low income, psychosocial issues) (Shields and Tsay, 2015).

Exercise pattern: Type, frequency, duration [American College of Obstetricians and Gynecologists (ACOG), 2015; Joslin, 2011].

Finally, language, cultural background, ethnic or religious beliefs should be taken into consideration (Shields and Tsay, 2015).

Anthropometric Measurements

Anthropometric measurements pertinent to diabetes and pregnancy include:

- Height, weight, weight history, pre-pregnancy weight and BMI should be assessed at the initial visit and weight should be tracked at each visit to determine if the gestational weight gain (GWG) is appropriate (within range), based on IOM revised guidelines for weight gain during pregnancy (IOM, 2009) (Shields and Tsay, 2015; Joslin, 2011; Kaiser and Campbell, 2014).

- While the total amount of weight gained in normal-term pregnancies varies in women (IOM, 2009), the IOM recommends that women achieve GWG within the range identified for their pre-pregnant BMI for singleton or multiple pregnancies, as appropriate (Kaiser and Campbell, 2014; IOM, 2009).

- Inappropriate weight gain (excess or inadequate weight gain) may require further assessment of food and kcal intake and adjustment in the nutrition prescription (ADA, 2016).

Biochemical Data, Medical Tests and Procedures

Clinical data related to medical tests pertinent to diabetes and pregnancy include:

- Glycemic tests, including GCT, OGTT, A1c, fasting and random glucose and fasting, pre-prandial, and post-prandial self-monitoring of blood glucose are recommended (ADA, 2016) to determine glycemic control throughout pregnancy (Shields and Tsay, 2015; ACOG, 2013).

- A nutritional anemia profile (hemoglobin/hematocrit, folate, B12, iron) and Vitamin D and other micronutrient screening, as needed help determine if the woman may benefit from additional counseling targeting specific nutrients (Shields and Tsay, 2015). For example, serum ferritin may be useful to identify pregnant women who would benefit from additional counseling about iron-rich foods and supplements (Kaiser and Campbell, 2014) or vitamin D screening may be considered for those who may be at risk of deficiency, such as lack of sun exposure, vegan or northern latitude (Kaiser and Campbell, 2014).

- Kidney function tests (creatinine clearance) (Shields and Tsay, 2015).

- Thyroid function (Shields and Tsay, 2015).

Nutrient-Focused Physical Findings and Client History

Nutrition-focused physical findings and client history related to diabetes and pregnancy include:

- Pertinent medical history (diseases, conditions, illnesses), previous obstetrical history

- Age, number of fetuses, weeks of gestation, EDD

- History of GDM duration of diabetes hypoglycemia, diabetes complications, family history (Shields and Tsay, 2015)

- GI discomforts, such as nausea, vomiting, etc. that may interfere with the ability to consume adequate nutrients

- Vital signs, such as blood pressure (Shields and Tsay, 2015) and general health

- Social history, living situation, health literacy, attitudes toward health including current diabetes knowledge (Shields and Tsay, 2015) and numeracy that may affect learning ability or needs and ability to implement dietary strategies or to make appropriate food choices

- Educational background and occupation, including financial and employment status (Shields and Tsay, 2015) may affect meal timing and schedule and healthy food purchasing ability.

Recommendation Strength Rationale

Consensus: This topic was not included in the EAL systematic review. The recommendations are based on consensus publications.

Minority Opinions

None.

Supporting Evidence

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

References

None.

References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process


Gestational Diabetes
- Gestational Diabetes (GDM) Guideline (2016)
Recommendations Summary

GDM: Medical Nutrition Therapy 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

- **Recommendation(s)**
  - **GDM: Medical Nutrition Therapy (MNT)**
    
    The registered dietitian nutritionist (RDN) should provide medical nutrition therapy (MNT) that includes an individual nutrition prescription and nutrition counseling for all women diagnosed with gestational diabetes mellitus (GDM). Research indicates that MNT provided by an RDN (or international equivalent) as part of a comprehensive nutrition intervention that includes individualization of MNT is effective in improving blood glucose control and neonatal and maternal outcomes in women with GDM. Improved outcomes included lower birth weight and a reduction in the following: Incidence of macrosomia (LGA), need for insulin therapy, hypertensive disorders of pregnancy and maternal hospitalizations, neonatal intensive care unit (NICU) admissions and neonatal deaths, premature births and rate of shoulder dystocia, bone fracture and nerve palsy.

  **Rating: Strong**
  **Imperative**

- **GDM: Frequency and Duration of MNT**
  
  The RDN should provide regular and frequent MNT visits to women with GDM to optimize outcomes. Visits should include an initial 60 to 90 minute MNT visit, followed by a second MNT visit (30 to 45 minutes) within one week, and a third MNT visit (15 to 45 minutes) within two to three weeks. Additional MNT visits should be scheduled every two to three weeks or as needed for the duration of the pregnancy. MNT assists the woman with GDM in meeting her blood glucose and weight gain targets, contribute to a well-balanced food intake and promote fetal and maternal well-being.

  **Rating: Consensus**
  **Imperative**

- **Risks/Harms of Implementing This Recommendation**
  
  There are no potential risks or harms associated with the application of these recommendations.

- **Conditions of Application**
  
  For the recommendation GDM: Frequency and Duration of MNT, barriers to attendance may include financial constraints, scheduling conflicts, inability to take time off work or school, lack of child care and lack of transportation.

- **Potential Costs Associated with Application**
  
  Costs of MNT sessions and reimbursement vary. However, MNT sessions are essential for improved outcomes.

- **Recommendation Narrative**
  
  **GDM: MNT**

  A total of five studies were included in the evidence supporting the recommendation:

  - Two positive quality randomized controlled trials (RCTs) (Landon et al, 2009; Reader et al, 2006)
  - One neutral quality RCT (Crowther et al, 2005)
  - One neutral quality prospective cohort study (Maher et al, 2013)
  - One neutral quality non-randomized controlled trial (Perichart-Perrera et al, 2009).

  **Evidence Summary**

  - Five studies evaluated the effectiveness of MNT intervention, provided by an RDN (or international equivalent) (specifically, dietitian, registered dietitian or nutritionist) on GDM-related outcomes. Four studies (three RCTs, one non-randomized controlled trial) compared MNT intervention to standard or usual care in women with GDM. One prospective cohort study (Maher et al, 2013) evaluated early MNT during the subjects' first trimester (mean, 10.2 weeks gestation), as part of multi-disciplinary intervention in women with a history of insulin-requiring GDM. All studies found that the MNT intervention improved fetal/neonatal and maternal outcomes in women with GDM (Crowther et al, 2005; Landon et al 2009; Perichart-Perrera et al, 2009; Reader et al; 2006) or with women with a history of insulin-requiring GDM (Maher et al, 2013).
  
  - The studies described MNT, provided by dietitians in a number of ways, including nutrition education or counseling (with nutrition assessment), diet therapy and dietary advice. All studies included individualization of MNT as part of a comprehensive intervention that included at least two of the following: education on diabetes, instructions in self-monitoring
of blood glucose (SMBG), regular follow up and monitoring with the physician, blood glucose (BG) monitoring and lifestyle (e.g., physical activity) counseling and advice. Insulin therapy was initiated, as required. Although the studies did not describe the actual number of MNT encounters during the intervention, all studies reported more than one MNT visit and one study (Reader et al, 2006) described a minimum of three MNT visits in the intervention. No studies described the frequency of MNT visits.

- MNT, as part of a comprehensive intervention improved blood glucose control (Perichart-Perrera et al, 2009; Reader et al, 2006) and improved the following adverse outcomes:
  - Maternal outcomes:
    - Fewer hypertensive disorders of pregnancy and pre-eclampsia (Landon et al, 2009; Perichart-Perrera et al, 2009)
    - Fewer maternal hospitalizations (Perichart-Perrera et al, 2009)
    - Fewer premature births (Perichart-Perrera et al, 2009)
    - Reduced need for insulin therapy (Reader et al, 2006).
    - Fewer caesarian deliveries (Landon et al, 2009)
  - Neonatal outcomes:
    - Fewer neonatal deaths (Crowther et al, 2005; Perichart-Perrera et al, 2009)
    - Fewer NICU admissions (Perichart-Perrera et al, 2009)
    - Lower birth weight (Crowther et al, 2005; Perichart-Perrera et al, 2009) and reduced neonatal fat mass (Landon et al, 2009)
    - Fewer LGA (Crowther et al, 2005; Landon et al, 2009) and lower prevalence of macrosomia (Crowther et al, 2005; Landon et al, 2009; Maher et al, 2013; Perichart-Perrera et al, 2009)
    - Reduced rate of shoulder dystocia (Landon et al, 2009; Crowther et al, 2005), bone fracture, and nerve palsy (Crowther et al, 2005).


**GDM: Frequency and Duration of MNT**

- No evidence was identified to evaluate the optimal frequency and duration of MNT visits by an RDN (or international equivalent) to improve fetal and maternal outcomes. However, the following guidance from Joslin Diabetes Center & Joslin Clinic (2011) provides support for the recommendation:
  - A minimum of three encounters with a Certified Diabetes Educator (CDE) (RDN, RN) for assessment and meal plan modification (and SMBG instruction, if RDN is adequately trained) are recommended as follows.
    - Visit One (60 to 90 minutes) individual or group visit
    - Visit Two (30 to 45 minutes) one week after initial visit
    - Visit Three (15 to 45 minutes) in one to three weeks.
    - Additional visits every two to three weeks and as needed until delivery.

**Recommendation Strength Rationale**

- Conclusion statement supporting the recommendation GDM: MNT is Grade II
- Consensus: The recommendation GDM: Frequency and Duration of MNT is based on consensus publications. This topic was included in the EAL systematic review. However, no evidence was found to answer the research question.

**Minority Opinions**

None.

**Supporting Evidence**

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

**In women with GDM, what is the effectiveness of MNT intervention, provided by an RDN on fetal/neonatal and maternal outcomes?**

- References
GDM: Calories 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section, below.

**Recommendation(s)**

**GDM: Calorie Prescription**

For women with gestational diabetes mellitus (GDM), the registered dietitian nutritionist (RDN) should individualize the calorie prescription based on a thorough nutrition assessment with guidance from relevant references [Dietary Reference Intakes (DRI), Institute of Medicine (IOM)] and encourage adequate caloric intake to promote fetal/neonatal and maternal health, achieve glycemic goals, and promote appropriate gestational weight gain (GWG). No definitive research suggests there is a specific optimal caloric intake for women with GDM or if calorie needs are different than pregnant women without GDM. Limited research in women with GDM whose pre-pregnancy weights ranged from normal to obese showed no significant differences in most fetal/neonatal and maternal outcomes with various reported caloric intakes. In a study of obese women only, GWG slowed after women with GDM reportedly consumed 30% below their caloric requirements, without adverse effects.

**Rating: Fair**

**Imperative**

- **Risks/Harms of Implementing This Recommendation**

  There are no potential risks or harms associated with the application of this recommendation.

- **Conditions of Application**

  - Refer to the GDM: Nutrition Assessment and GDM: Nutrition Monitoring & Evaluation recommendations for factors to consider in determining and adjusting an individualized calorie prescription, such as pre-pregnancy weight and BMI, total and rate of GWG, single vs. multiple fetuses, physical activity level, etc.
  - Individual caloric requirements during pregnancy vary among women and are dependent on several factors, including pre-pregnancy BMI and fat mass, and changes in physical activity (Ho et al, 2005).

- **Potential Costs Associated with Application**

  Costs may include expenses related to medical nutrition therapy (MNT) visits from an RDN and higher food costs, if a caloric increase is needed (e.g., types and amounts of food).

**Recommendation Narrative**

**GDM: Calorie Prescription**

A total of three studies were included in the evidence analysis supporting the recommendation:

- One positive quality prospective cohort study (Ho et al, 2005)
- One positive quality randomized controlled trial (RCT) (Rae et al, 2000)
- One neutral quality prospective cohort study (Romon et al, 2001).

**Evidence Summary**

Three international studies evaluated the impact of calorie intake on fetal/neonatal and maternal outcomes (glycemic control, maternal weight gain, fetal growth/birth weight and adverse outcomes), in women with GDM.

- One positive quality prospective cohort study by Ho et al, 2005, evaluated a caloric prescription of 30kcal/kg body weight (BW) in 62 women with a non-obese pre-pregnancy weight (BMI range: 22.4±3.2 to 23.1±4.2kg/m²).
  - Caloric intake in the women who was categorized into three tertiles (calculated kcal/kg values): 1, 86±3kcal/kg (lowest tertile), 1, 69±2kcal/kg (middle tertile) and 1, 38±1kcal/kg BW (highest tertile), noting the women had a tendency to over-restrict their calorie intake.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.4±7.9, 22±8.3, 22±9.5 pounds, respectively.
  - There was no significant (NS) difference between calorie intake tertiles and either maternal GWG or infant birth weight.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.2±7.9, 22±8.3, 22±9.5 pounds, respectively.
- One neutral quality prospective cohort study (Romon et al, 2001).
  - One positive quality prospective cohort study by (Ho et al, 2005) evaluated a calorie prescription of 30kcal/kg BW (highest tertile), 1, 69±2kcal/kg BW (middle tertile) and 1, 38±1kcal/kg BW (lowest tertile), noting the women had a tendency to over-restrict their calorie intake.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.4±7.9, 22±8.3, 22±9.5 pounds, respectively.
  - There was no significant (NS) difference between calorie intake tertiles and either maternal GWG or infant birth weight.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.2±7.9, 22±8.3, 22±9.5 pounds, respectively.
  - There was no significant (NS) difference between calorie intake tertiles and either maternal GWG or infant birth weight.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.4±7.9, 22±8.3, 22±9.5 pounds, respectively.
  - There was no significant (NS) difference between calorie intake tertiles and either maternal GWG or infant birth weight.
  - Women in the highest tertile had significantly higher post-dinner glucose concentration after controlling for pre-pregnancy weight and height. Total GWG was 20.4±7.9, 22±8.3, 22±9.5 pounds, respectively.

References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process


- Gestational Diabetes
- Gestational Diabetes (GDM) Guideline (2016)

**Quick Links**

**Recommendations Summary**

GDM: Calories 2016


- Reference(s)
  - References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process

Australia) * [1, 560kcal intervention vs. 1, 630kcal control; NS], despite no prescriptive calorie restriction in the control group.

- The mean rate of GWG slowed once dietary intervention started and 54.1% of the intervention vs. 40.7% of the control groups failed to gain or lost weight (NS). Women in both groups lost 1.68kg [SE 0.33, range 7.5-0.00 and SE 0.32, range 5.0-0.00 intervention vs. control, respectively].
- Total GWG was 25.4 pounds in the intervention and 21.3 pounds in the control group (NS).
- There were no differences between the groups in maternal outcomes (anemia, pre-existing hypertension, pre-eclampsia, premature rupture of membranes (PROM), threatened preterm labor, percent requiring insulin, ketonuria (34.5 vs. 38.5% nondetectable ketones), serum beta-hydroxybuterate, BG control, HbA1c, GWG) or neonatal outcomes (rates of delivery mode, labor induction, fetal distress birth trauma, GA and incidence of macrosomia).
- Intervention group infants had greater average abdominal skinfold thickness, but mean total skinfold measurements were similar between the two groups.
- Control group infants had greater incidences of polycythemia and shoulder dystocia.
- More research is needed to elucidate the effect of caloric consumption (kcals/kg pre-pregnancy BW), independent of other factors, on fetal/neonatal and maternal outcomes.

*The kcal intake per kg of body weight for the Romon et al, 2001, and Rae et al, 2000, studies could not be calculated for the groups.

**Recommendation Strength Rationale**

- Conclusion statement supporting GDM: Calorie Prescription is Grade III.
- Results of the studies were confounded by use of reported vs. actual caloric intakes (possible underreporting), tendency of the women to over-restrict caloric intake vs. prescribed, inconsistent stratification by pre-pregnancy BMI, and pre-pregnancy weights not described, making comparison and synthesis of the research challenging.

**Minority Opinions**

None.

**Supporting Evidence**

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what is the effect of caloric consumption on fetal/neonatal and maternal outcomes?

**References**


**References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process**


**Gestational Diabetes**

**Gestational Diabetes (GDM) Guideline (2016)**

**Quick Links**

**Recommendations Summary**

**GDM: Macronutrients 2016**

*Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.*

**Recommendation(s)**

**GDM: Macronutrient Requirements**

In women with gestational diabetes mellitus (GDM), the registered dietitian nutritionist (RDN) should provide adequate amounts of macronutrients to support pregnancy, based on nutrition assessment, with guidance from the Dietary Reference Intakes (DRI). The DRI for all pregnant women, including those with GDM, recommends a minimum of 175g carbohydrate (CHO), a minimum of 71g protein (or 1.1g per kg per day protein) and 28g fiber.
GDM: Carbohydrate Prescription

The RDN should individualize both the amount and type of CHO for women with GDM based on nutrition assessment, treatment goals, blood glucose response and patient needs. Limited evidence does not confirm an ideal amount (grams or percent of total calories) of CHO for all women with GDM, but suggests an interaction between the amount and type of CHO. Several studies showed positive effects on glycemic control and neonatal/fetal and maternal outcomes in women with GDM, when evaluating varying amounts and types of CHO:

- Low glycemic index (GI) (less than 55) or medium GI (55 to 69) diets, containing a range of 36.7% to more than 60% CHO
- Dietary Approaches to Stop Hypertension (DASH) diets (greater than 65% CHO)

However, when two studies evaluated the amount of CHO alone (without specifying the type of CHO) mixed results were found:

- A CHO prescription of 202g CHO per day was more effective at reducing post-prandial blood glucose (PPBG), compared to >270g CHO per day.
- A 3% incidence of large-for-gestational-age (LGA) infants was found with CHO intake of less than 211g per day, but no LGA when greater than 211g per day.

Rating: Fair

GDM: Carbohydrate and Post Prandial Breakfast Glycemia

The RDN should individualize both the amount and type of CHO at breakfast based on nutrition assessment, treatment goals, blood glucose response and patient needs. If the woman with GDM continues to experience elevated PPBG after breakfast, the RDN may further modify the amount or the type of CHO at breakfast to achieve blood glucose targets. Limited evidence examining the impact of CHO on PPBG after response and patient needs. If the woman with GDM continues to experience elevated PPBG after breakfast, the RDN may further modify the amount or the type of CHO at breakfast to achieve blood glucose targets.

- In women with GDM who followed low or medium glycemic index (GI) diets containing 42-60% total CHO (GI for breakfast meal <55; CHO range 15g to 60g or more) met PPBG targets after breakfast.
- One study evaluating a 45% CHO diet overall (without specifying the type of CHO), found improved PPBG after breakfast, compared to one that contained 60% CHO.
- No studies evaluated the effect of only restricting individual foods (e.g., fruit or milk) at breakfast, although one study showed improved PPBG when fruit bread and milk were eaten in a low GI breakfast vs. a high GI breakfast with CHOs from other sources.

Rating: Consensus

Risks/Harms of Implementing This Recommendation

There were no potential risks or harms associated with the application of these recommendations.

Conditions of Application

For the recommendation GDM: Carbohydrate and Post Prandial Breakfast Glycemia, the RDN should use clinical judgment in individualizing the breakfast meal. Although no evidence was found to support a specific or range of carbohydrate (CHO) distribution at meals or snacks, customary practice suggests limiting the amount and type of CHOs at breakfast (Joslin Diabetes Center & Joslin Clinic, 2013). The CHO intake is reassessed at subsequent visits for possible adjustment according to the blood glucose records (Shields & Tsay, 2015). See the recommendation GDM: Distribution of Meals and Snacks

Potential Costs Associated with Application

- Costs may include expenses associated with receiving care from an RDN
- There may be extra time needed or increased food costs associated with improving the quality of the diet as recommended.

Recommendation Narrative

A total of 12 studies were included in the evidence analysis supporting these recommendations:

- Six positive quality randomized controlled trials (RCTs) (Asemi et al, 2014; Grant et al, 2011; Moreno-Castillo et al, 2013; Louie et al, 2011; Moses et al, 2009; Pennach-Perera et al, 2012)
- Two positive quality randomized crossover trial (Hernandez et al, 2014; Louie et al, 2013)
- Three neutral quality RCTs (Afagh et al, 2013; Asemi et al, 2013 (a, b); Cypryk et al, 2007)
- One neutral quality retrospective cohort (Ronen et al, 2001).

Impact of the amount of CHO consumed on fetal/neonatal and maternal outcomes (independent of dietary patterns based on the Dietary Approaches to Stop Hypertension (DASH) diet and Glycemic Index)

Conclusion: Limited evidence was found to demonstrate the impact of the amount of CHO consumption on neonatal or fetal and maternal outcomes in women with gestational diabetes mellitus (GDM). In one study, women prescribed a minimum of 1, 800kcal per day found reductions in post-prandial blood glucose (PPBG) levels at all three meals with 202g per day CHO, while those prescribed >270g CHO showed reductions in PPBG at two meals only. Another study of women with an average intake of 1, 852±343kcal per day found zero incidence of LGA when CHO intake was >211g per day, but a 23% incidence in women consuming less CHO. No other differences in fetal and maternal outcomes were found in these studies.

Overview

Three studies evaluated the impact of the amount of CHO consumed (independent of dietary patterns including DASH and glycemic index) on fetal or neonatal and maternal outcomes (glycemic control, maternal weight gain, fetal growth or birth weight, and adverse outcomes), in women with GDM:

- A neutral quality RCT (Cypryk et al, 2007) assigned 30 women with GDM to consume an 1, 800kcal diet with either low-CHO (LC) content (45% CHO (202g), 25% PRO, 30% FAT) or high-CHO (HC) content (>60% CHO (>270g), 25% PRO, 15% FAT) for two weeks. Actual intake was not specified. There was a significant reduction in PPBG concentrations for all meals in the LC group, while the HC group realized significant BG reductions after lunch and dinner only. No significant (NS) differences were observed between groups for ketonuria or obstetric outcomes.
- A positive quality RCT (Moreno-Castillo, 2013) assigned 150 women with GDM to either a LC diet (40% CHO) or a HC diet (55% CHO). There was no difference between groups regarding the need for insulin therapy. Women in the HC group gained more weight than women in the LC group, but the difference disappeared when time to follow-up was included in the statistical analysis. There was no difference in the women in the LC group. There were no other significant differences between groups for other maternal or neonatal outcomes.
- A neutral quality prospective cohort study (Ronen et al, 2001) evaluated actual intake in 80 women with an elevated pre-pregnancy BMI. Women consumed an average of 1, 852±343kcal per day [43.4% CHO (202±43g), 18.5% PRO, 37.9% FAT], Mild ketonuria was present in 45% of the women during week 1, but decreased to 16%. No relationship between the reported calorie intake and infant birth weight was found. However, an inverse relationship between CHO intake and infant birthweight was found. The 32 women who consumed >211g CHO per day and proportionately lower fat had no LGA infants, whereas in women consuming less CHO, 11 women (or 23%) had LGA infants.

Conclusion: Dietary patterns based on the DASH diet, (which contained higher amounts of CHO and dietary fiber, and less sucrose, total fat, dietary cholesterol and less sodium) when compared to a control diet, were effective in improving both fetal and maternal outcomes in women with GDM who did not require insulin at the time of diagnosis. Improvements were found in glucose tolerance,
glycosylated hemoglobin levels, insulin resistance, less need for insulin, lipid profile, systolic blood pressure and biomarkers of oxidative stress. There was also a lower incidence of Cesarean-section deliveries. Infant birthweights, head circumferences, ponderal indices and the incidence of macrosomia were lower in infants whose mothers consumed the DASH diet.

**Overview**

- **Two RCTs** by Asemi et al [2013 (a) and 2014] evaluated the impact of dietary patterns based on the DASH eating plan on fetal/neonatal and maternal outcomes (glycemic control, maternal weight gain, fetal growth/birth weight and adverse outcomes), in Iranian women with GDM who did not require insulin* at time of diagnosis. A third paper by Asemi et al [2013 (b)] reported additional outcomes for the same subjects of the Asemi et al [2013 (a)]. The study design was a randomized controlled trial in which the intervention diet was a standard prescribed diet for the duration of pregnancy. A fourth study by Asemi et al [2013 (c)] compared two low to medium GI (LGI-MGI) diets with inverse macronutrient percentages (GI for carbohydrates ≤55) in women with GDM. In women with GDM, four of five studies reported a significant improvement in glycemic control and had similar neonatal or fetal and maternal outcomes, in women with GDM in most studies. In a few studies, an LGI diet (36-47% CHO) was found to significantly reduce the need for insulin therapy and prevent excessive maternal weight gain. One study comparing an LGI-MGI diet (40% CHO, 45% fat) to an LGI-MGI diet (60% CHO, 25% fat) found that both diets achieved glycemia within target levels.

**Impact of dietary patterns based on the glycemic index on fetal/neonatal and maternal outcomes**

- **Conclusion:** A low (LGI) or medium GI (MGI) dietary pattern resulted in glycemic control and had similar neonatal or fetal and maternal outcomes in women with GDM in most studies. In a few studies, an LGI diet (36-47% CHO) was found to significantly reduce the need for insulin therapy and prevent excessive maternal weight gain. One study comparing an LGI-MGI diet (40% CHO, 45% fat) to an LGI-MGI diet (60% CHO, 25% fat) found that both diets achieved glycemia within target levels.

**Overview**

- **Six studies** evaluated the impact of dietary patterns based on the glycemic index (GI) on fetal or neonatal and maternal outcomes (glycemic control, maternal weight gain, fetal growth or birth weight, and adverse outcomes), in women with GDM.
- **A low (LGI) or medium GI (MGI) dietary pattern** resulted in glycemic control and had similar neonatal or fetal and maternal outcomes in women with GDM in most studies. In a few studies, an LGI diet (36-47% CHO) was found to significantly reduce the need for insulin therapy and prevent excessive maternal weight gain. One study comparing an LGI-MGI diet (40% CHO, 45% fat) to an LGI-MGI diet (60% CHO, 25% fat) found that both diets achieved glycemia within target levels.

- **GI parameters for the purpose of study comparison** are as follows: Low GI=0-55; Medium GI=56-69; and High GI=70 or greater (Augustin et al, 2015).

- **A no-study review** compared LGI to high GI diets. However, three studies compared an LGI diet to other LGI diets with variations in CHO or fiber. Louie et al, 2011 compared an LGI diet (GI 47±1) to a conventional high fiber (HF) diet (GI 53±1) and Perichart-Perrera et al, 2012 compared an LGI (GI 47±6.9) to an all CHO diet (GI 48±6.4). A third study by Afaghi et al, 2013 compared an LGI diet with added wheat fiber (GI ≤55) to a conventional high fiber (same composition). Two studies compared LGI diets to MGI diets. Grant et al, 2011 compared an LGI diet (GI 49±0.8) to an MGI diet (GI 58±0.5) and Moses et al, 2009 compared an LGI diet (GI 48) to a conventional HF diet (GI 56). The macronutrients as a percentage of total calories in the comparison groups. One study (Hernandez et al, 2014) compared two low to medium GI (LGI-MGI) diets with inverse macronutrient percentages (GI for breakfast meals was <35.7). A macronutrient and GI comparison of the studies and outcome results are below.

- **In women with GDM, four of five studies reported** (Abati et al, 2011; Grant; Hernandez et al, 2014). Perichart-Perrera and two of five studies showed the need for insulin therapy was significantly reduced (Afaghi, Moses), when following either an LGI diet. Afaghi et al, 2013 demonstrated improved glycemic control and reduced need for insulin in women who followed an LGI diet at each meal with added wheat fiber to those who didn’t. Moses et al, 2009 reported that fewer women on an LGI diet required insulin to achieve BG control, compared to a conventional HF diet (GI 53±1). Three other studies did not find differences in insulin use among women with GDM. (Hernandez et al, 2011; Louie et al, 2011). Perichart-Perrera et al, 2012 found that an LGI diet was significantly more effective than a control glycemic diet, as a diet including all types of CHO with a similar GI. Grant et al, 2011 found that women who consumed an LGI diet, compared to an MGI diet had more PPBG values within target range, but other BG values were NS. However, neither study found significant between-group differences for insulin initiation. In a study by Louie et al, 2011 (N=52) found significantly fewer women in the DASH group required c-sections and insulin injections after the intervention compared to those in the control group. Significantly fewer infants were macrosomic in the DASH group and had significantly lower birthweight compared to those controlled to the control mothers. The DASH diet outcomes remained significant even after controlling for pre-pregnancy body mass index (BMI <30kg/m² and >30kg/m²), baseline maternal FGF and maternal age. NS differences were found between groups in Apgar scores, percentage of fetal/neonatal polyhydraminos or gestational age in weeks.

* The authors noted that in previous studies conducted in Iran, insulin therapy was often recommended upon diagnosis of GDM, to reduce complications.

**The authors noted c-section rates in Iran among women with GDM are very high (almost 90%).**
between groups. The risk of pre-eclampsia, intrauterine and neonatal death was the same for both groups.

- Low GI vs. Medium GI Diets
  - Grant et al, 2011 (N=38) found NS differences in fasting serum glucose, mean post-prandial SMBG, HbA1C, fasting insulin, lipids or CRP between a group of women with GDM consuming an LGI diet (GI 49±0.8; GL 98.2±5.1) and a control group consuming a significantly higher GI (GI 58±0.5; GI215±8.8). The LGI diet group had significantly fewer fasting serum glucose measurements below target, and over twice as many on target, than the control group. Fewer PPBG measurements were above target in the LGI group. There was NS difference between groups in the number of women started on insulin, and insulin treatment was associated with significantly greater maternal weight gain. There were NS differences between groups in infant birth weight, LGA infants and SGA infants.
  - Alonso et al, 2009 (N=63) found that women with GDM who consumed an LGI diet (36.7±1% CHO; GI 48.3) needed insulin therapy significantly less of the children (68±1% CHO; GL 56.8). Of women in the HF group who met criteria for insulin therapy, nearly half were able to avoid insulin use after switching to the LGI diet. There were NS differences between groups for maternal weight gain, induction of delivery or type of delivery. NS differences were found between groups for fetal outcomes including gestational age at delivery, fetal or birth centile, or ponderal index.

- Low-Medium GI vs. Low-Medium GI Diets
  - Hernandez et al, 2014 (N=16) found no between-diet differences for fasting or preprandial glycemia in a group of women with GDM consuming either a low CHO/high fat diet (LC/CONV) (40% CHO, 45% fat; low to medium GI; breakfast GI 35.7) or a higher complex CHO diet (CHO>56% fat; low to medium GI; breakfast 34.8). When considered as a mean across three meals, 1- and 2-hour PPBG and daytime mean glycemia were modestly higher on the CHOICE diet. However, both diets produced results well within current treatment targets for daytime, nocturnal, post-prandial and mean BG levels. Free fatty acid (FFA) levels were significantly lower for the CHOICE diet.

Impact of the type or amount of CHO consumed on post-prandial breakfast glycemia

Conclusion: Limited evidence was found to demonstrate the impact of the type or amount of carbohydrate (CHO) consumption on post-prandial breakfast glycemia in women with gestational diabetes mellitus (GDM). Three studies that evaluated glycemic index (GI) reported that lower GI diets containing 42-60% total CHO (fat for breakfast meal <55; CHO range 15g to 60g or more) improved glycemic control after breakfast. One study that did not consider the glycemic index showed that lower CHO (45% vs. 60% of kcal) improved post-prandial blood glucose after breakfast. No studies evaluated the effect of only restricting individual foods (e.g., fruit or milk) at breakfast although one study showed improved blood sugars when fruit bread and milk were eaten in a low GI breakfast over a high GI breakfast with carbohydrates from other sources.

Overview

- Five studies evaluated the impact of the type or amount of CHO on breakfast PPBG values in women with GDM. Three studies evaluated dietary patterns based on the GI with varying amounts of CHO (Grant et al, 2011; Hernandez et al, 2014; Perichart-Perrera et al, 2012) and one study (Loi et al, 2013) evaluated an LGI high GI breakfast meal alone. One study evaluated the amount of CHO as a percentage of calories (Cypryk et al, 2007). GI parameters for the purpose of study comparison of the GI studies are as follows: Low GI=55; Medium GI=56-69; and High GI=70 or greater (Augustin et al, 2015). Three of these studies (Hernandez 2014, Perichart-Perrera 2012, and Louie 2013) specifically tested a controlled breakfast meal.

A brief description of the comparison diets and specific glycemic outcomes and results for each study are as follows:

- Cypryk et al, 2007 (N=39) compared a low-CHO diet (45% CHO, 30%, 15% fat) and a high-CHO diet (>60% CHO, 15% fat) and found a significant reduction in PPBG concentrations for all meals, including breakfast in women consuming a low-CHO diet (45% CHO, 30%, 15% fat) compared to women consuming a high-CHO diet (>60% CHO, 15% fat). The high-CHO group showed a larger increase in BG only at two hours after the meal. The low-CHO group consumed significantly fewer all CHO at breakfast, 1- and 2-hour PPBG and nocturnal PPBG and daytime mean glucose. However, post-intervention, the low-CHO group consumed significantly more dietary fiber than controls (30±1.6 and 23.1±0.9, respectively; P<0.001) and had a significantly lower GI than control (GI 49±0.8 vs. 58±0.5; P<0.001).

- Grant et al, 2011 (N=38) compared an LGI diet (% CHO NR; GI 49±0.8) to an MGI diet (% CHO NR; GI 58±0.5) and found a significant relationship between post-prandial SMBG after breakfast and BMI baseline (P<0.001) in two study groups of women (control and low GI) diagnosed with gestational hyperglycemia. During the study, NS differences were observed between the changes in the low GI fasting, -0.48±0.11 mmol/L; postprandial, -0.54±0.19 mmol/L; SMBG, -0.45±0.19 mmol/L; and control fasting, -0.44±0.21 mmol/L, groups, although the fasting and (P<0.001) and mean postprandial (P<0.001) fell significantly. Also, the control group demonstrated a strong positive relationship between pre-pregnancy BMI and SMBG after breakfast (R=0.75, P<0.001); although the low-GI group did not demonstrate this correlation, although significant (R=0.01, P=0.02). However, post-intervention, the LGI group consumed significantly more dietary fiber than controls (30±1.6 and 23.1±0.9, respectively; P<0.001) and had a significantly lower GI than control (49±0.8 vs. 58±0.5; P<0.001).

The result was the LGI group had a significantly lower GL than control post-intervention (98.2±5.1 vs. 125±8.8; P=0.014).

- Hernandez et al, 2014 (N=16) compared two low medium GI (LGI-MGI) diets (40% CHO, 45% fat; GI 36.7 vs. 35.7) vs. (80% CHO; 25% fat; GI NR; breakfast 34.8) and found a significant relationship between post-prandial SMBG in women on a higher complex CHO/lower fat diet (60% CHO, 25% fat; low to medium GI; breakfast GI 34.8), compared to a lower CHO/high-fat diet (40% CHO, 45% fat; low to medium GI; breakfast GI 35.7). However, PPBG after breakfast was within current treatment targets 1 hour (all, P<0.02) and 2 hours, 107±3mg/dL, P<0.01; 2 hours, 106±3 vs. 97±5 mg/dL, P=0.001). The breakfast meal contained 25% of total kcal and reflected the overall macronutrient percentage for each diet.

- Louie et al, 2013 (N=10), crossover study design provided two groups of 10 subjects each, in a study focused on only a controlled breakfast meal compared a low-GI breakfast (GI 45, GL 21) that included fruit bread and milk, with a high-GI breakfast (GI 82, GL 31). Louie et al found significantly lower peak BG levels (6.7±0.3mmol/L vs. 8.6±0.4mmol/L; P=0.001) in women consuming a low GI breakfast (GI 45) compared to those consuming a high-GI breakfast (GI 82; GL 36; 42.7g CHO*). The area under the glucose curve (IAUGC glucose) was significantly lower for the LGI group (212.7±22.9 vs. 340.8±23.4; P=0.001). There was large inter-subject variability in the timing of the peak BG level between the test meals, with peaks occurring at 45-75 minutes for the LGI meal and between 30-60 minutes for the high GI meal.

- Perichart-Perrera et al, 2012 (N=107; N=52 GDM; N=55 T2D) compared an LGI diet (46.6±9.1% CHO; GI 47.2±6.9) to an all types of CHO diet (45.8±8.3% CHO; GI 48.6±8.4) and found that an LGI diet was equally effective in improving glycemic control, as a diet including all types of CHO. Breakfast was limited to 15-30g CHO in both diets. At the end of the study, there was a statistically significant increase in the number of women in the LGI group who met glycemic targets after lunch, pre-dinner and post-dinner (P<0.05); and in all the types of CHO diet group, the increase was only at the post-lunch time (P<0.03). There was NS difference between groups or within groups for two-hour post-prandial breakfast glucose throughout the study.

*calculated value.

Recommendation Strength Rationale

The recommendation GDM: Macronutrient Requirements is based on consensus publications. The topic was not included in the EAL systematic review.

Three conclusion statements supporting the recommendation GDM: Carbohydrate Prescription are grade III

- Interpretation of results for the topic on impact of dietary patterns based on the DASH diet on fetal or neonatal and maternal outcomes is limited due to inability to compare diets across studies and none of the studies included a comparison to the high GI (HGI) diet.
- Interpretation of results for the topic on impact of dietary patterns based on the DASH diet on fetal or neonatal and maternal outcomes is limited due to subjects consuming relatively the same diet (despite differences in prescribed GI), limited population.
- Interpretation of results for the topic on impact of the amount of CHO consumed (independent of dietary patterns including DASH and GI/GL) on fetal and maternal outcomes and the topic of impact of dietary patterns based on the DASH diet, maternal and maternal outcomes is limited because results of these studies vary widely among studies.

Conclusion statement supporting the recommendation GDM: Carbohydrate and Post Prandial Breakfaet Glycemia is grade III

- Interpretation of results for the topic on impact of the type or amount of CHO consumed on post-prandial breakfast

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andeal.org
The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

In women with GDM, what impact does the amount or type of carbohydrate consumed have on post-prandial breakfast glycemia?

In women with GDM, what impact does the amount or type of carbohydrate consumed (independent of dietary patterns based on the DASH diet and glycemic index) have on fetal/neonatal and maternal outcomes?

In women with GDM, what impact do dietary patterns based on the DASH diet have on fetal/neonatal and maternal outcomes?
Imperative
Rating: Consensus

GDM: Vitamin and Mineral Supplementation

The RDQ should consider recom mendating dietary supplementation within the Dietary Reference Intakes (DRI) for pregnancy with a prenatal multi-vitamin/mineral or specific vitamin or mineral supplement(s) to address inade quate dietary vitamin and mineral intake (e.g., iron, folate, calcium, vitamin D, choline and iodine) or documented micronutrient deficiency. Dietary supplements may be indicated in pregnant women at high risk for inadequate micronutrient intake, such as food insecurity; alcohol, tobacco or other substance dependency; anemia; strict vegetarian (vegan) diet; or poor eating habits.

Rating: Consensus

Conditions of Application

- Some individuals may not tolerate vitamin or mineral supplementation
- In general, pregnant women should seek medical consultation before or while taking a non-prescribed over-the-counter (OTC) micronutrient supplement that exceeds the Tolerable Upper Levels (UL) for a particular vitamin or mineral (Kaiser & Campbell, 2014) or if taking herbal supplements.

Risks/Harms of Implementing This Recommendation

- Consideration should be given to the total intake of micronutrients from all sources in the diet, such as fortified foods and beverages (e.g., calcium-fortified juice; grains enriched with iron, folate, and other B vitamins) and prescribed or non-prescribed vitamin and mineral supplements
- The RDQ should use professional judgment when assessing nutrition status and determining the need for vitamin and mineral supplementation for those at high risk of nutrient deficiencies, including history of malabsorptive disorders (bariatric surgery), multi-fetal pregnancy, omission of food groups and eating disorders (Kaiser and Campbell, 2014; Shields and Tsay, 2015).

Potential Costs Associated with Application

There is an increased cost for vitamin and mineral supplements.

Recommendation Narrative

- The micronutrient needs of pregnant women with GDM are the same as for those without diabetes. Consuming sufficient calories to support recommended weight gain and eating a variety of foods to meet nutrient needs are beneficial for pregnant women (Marrs and Boyar, 2009). As long as good food choices are made, the higher intake of calories, coupled with the increased absorption and efficiency of nutrient utilization that occurs in pregnancy are generally adequate to meet the needs for most nutrients, when good food choices are consistently made.

- The micronutrient needs of pregnant women with GDM: Vitamin and Mineral Supplementation are the same as for those without diabetes. Consuming sufficient calories to support recommended weight gain and eating a variety of foods to meet nutrient needs are beneficial for pregnant women (Marrs and Boyar, 2009). As long as good food choices are made, the higher intake of calories, coupled with the increased absorption and efficiency of nutrient utilization that occurs in pregnancy are generally adequate to meet the needs for most nutrients, including history of malabsorptive disorders (bariatric surgery), multi-fetal pregnancy, omission of food groups and eating disorders (Kaiser and Campbell, 2014; Shields and Tsay, 2015).

- Consideration should be given to the total intake of micronutrients from all sources in the diet, such as fortified foods and beverages (e.g., calcium-fortified juice; grains enriched with iron, folate, calcium, vitamin D, choline and iodine) or documented micronutrient deficiency. Dietary supplements may be indicated in pregnant women at high risk for inadequate micronutrient intake, such as food insecurity; alcohol, tobacco or other substance dependency; anemia; strict vegetarian (vegan) diet; or poor eating habits.

- In general, pregnant women should seek medical consultation before or while taking a non-prescribed over-the-counter (OTC) micronutrient supplement that exceeds the Tolerable Upper Levels (UL) for a particular vitamin or mineral (Kaiser & Campbell, 2014) or if taking herbal supplements.

- The RDQ should use professional judgment when assessing nutrition status and determining the need for vitamin and mineral supplementation for those at high risk of nutrient deficiencies, including history of malabsorptive disorders (bariatric surgery), multi-fetal pregnancy, omission of food groups and eating disorders (Kaiser and Campbell, 2014; Shields and Tsay, 2015).

- Consideration should be given to the total intake of micronutrients from all sources in the diet, such as fortified foods and beverages (e.g., calcium-fortified juice; grains enriched with iron, folate, calcium, vitamin D, choline and iodine) or documented micronutrient deficiency. Dietary supplements may be indicated in pregnant women at high risk for inadequate micronutrient intake, such as food insecurity; alcohol, tobacco or other substance dependency; anemia; strict vegetarian (vegan) diet; or poor eating habits.

- The overall prevalence of iron deficiency is nearly 18% in pregnant women in the United States, with anemia at 5% of pregnant women. Rates of iron deficiency increase across trimesters from 6.9% to 14.3% to 28.4% (Siu, 2015)

- The risk for iron deficiency is increased if there is iron deficiency anemia during the first two trimesters of pregnancy. Iron supplementation is needed to meet the increased maternal and fetal demand for iron throughout pregnancy (Procter & Campbell, 2014)

- The Recommended Dietary Allowance (RDA) for iron is 27 mg/day (Institute of Medicine (IOM) 1997-2011). Iron supplementation (30mg per day) is generally recommended starting at the first prenatal visit. However, while routine iron supplementation during pregnancy may improve maternal hematologic status and reduce the incidence of iron deficiency and iron deficiency anemia (IDA) in the short term, no clear or consistent evidence was found that prenatal iron supplementation has a beneficial clinical impact on maternal or infant health (Siu, 2015). Prenatal vitamins generally contain 30mg of iron. Women who have iron deficiency anemia during pregnancy should be prescribed 60mg to 120mg of elemental iron per day (Shields and Tsay, 2015).

- Folic acid is recognized for preventing neural tube defects and is important before (pre-conception) and during pregnancy (Procter and Campbell, 2014). Prenatal women should consume 600mcg of dietary folate equivalents (DFE) daily from all food sources (IOM, 1997-2011; Procter and Campbell, 2014). The DFE is equal to 1.0mcg food folate or 0.6mcg of folic acid from fortified food or as a supplement consumed with food (or 0.5mcg if supplement is taken on an empty stomach) (IOM, 1997-2011). Most prenatal multi-vitamin and mineral supplements contain 600mcg of folic acid and can assist the woman in meeting folic acid requirements of pregnancy (Shields and Tsay, 2015).

- Calcium: The RDA for calcium is 1,000mg per day for women ages 19 to 50 years. The DRI for calcium in pregnancy is the same as for women of the same age who are not pregnant, due to increased efficiency in calcium absorption and maternal bone calcium mobilization during pregnancy (Procter and Campbell, 2014). Women with calcium intakes under 500mg per day may need supplementation (Procter and Campbell, 2014).

- Vitamin D: Vitamin D’s function during pregnancy for both mother and fetus is not fully defined at this time and vitamin D supplementation during pregnancy remains controversial, although ongoing research suggests higher levels of supplementation appear to be safe and effective. The RDA for vitamin D is 600 IU (15mcg) per day (IOM 1997-2011) to meet the needs of most North American adults, including pregnant women (Procter and Campbell, 2014).

- Choline: Because of its high rate of transport from mother to fetus, choline is considered an essential nutrient during pregnancy. A deficiency can interfere with normal fetal brain development (Procter and Campbell, 2014). The adequate intake (AI) for choline is 450mg (IOM, 1997-2011). Most pregnant women do not consume the AI for choline, despite its presence in many foods (Procter and Campbell, 2014).

- Iodine: Iodine is required for normal brain development and growth. Iodine requirements increase during pregnancy. Iodine deficiency is a growing concern and recent nutrition surveys suggest a subset of pregnant women may have mild to moderately inadequate intake of iodine. The IOM recommends an iodine intake of 150mcg per day before conception and 220mcg per day for pregnant women (Procter and Campbell, 2014).

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andeal.org
GDM: Meal and Snack Distribution 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

**Recommendation(s)**

**GDM: Meal and Snack Distribution**

In women with gestational diabetes mellitus (GDM), the registered dietitian nutritionist (RDN) should distribute the total calories and carbohydrate (CHO) into smaller meals and multiple snacks per day. The distribution should be individualized, based on blood glucose levels, physical activity and medication, if any (e.g., insulin) and adjusted as needed. Three meals and two or more snacks helps to distribute CHO intake and reduce post-prandial blood glucose fluctuations.

**Rating:** Consensus

**Imperative**

**Risks/Harms of Implementing This Recommendation**

There are no risks or harms associated with the application of this recommendation.

**Conditions of Application**

- The RDN should consider the following when individualizing the distribution of meals and snacks: Usual food intake, food preferences, pharmacotherapy, blood glucose levels (hypo- or hyperglycemia), activity level, sleep pattern, treatment goals, work schedule (e.g., shift work, night schedule), food insecurity and access, and cultural/religious beliefs and practices (e.g., fasting), etc.
- The RDN should encourage regular and timely consumption of meals and snacks and avoid fasting beyond 10 to 12 hours (e.g., skipping the evening snack at bedtime or skipping breakfast) to promote blood glucose control (Buchanan et al, 1990; Metger et al, 1982; Mills et al, 1998)
- Pregnant women who opt to fast beyond 12 hours, due to cultural or religious reasons (e.g., Ramadan), should be medically evaluated prior to the fasting period for intensive management, self-management instruction, appropriate diet and insulin adjustment, if prescribed (Al-Arouj et al, 2010).

**Potential Costs Associated with Application**

Costs may include expenses related to medical nutrition therapy (MNT) visits from an RDN, blood glucose monitoring and associated medical follow-up.

**Recommendation Narrative**

No evidence was found to evaluate the impact of meal and snack distribution for women with gestational diabetes mellitus (GDM). However, the following provide for the consensus recommendation:

- Six to eight small meals and snacks are suggested to decrease post-prandial hyperglycemia in women with GDM. The distribution of calories should be individualized and based on usual intake, preferences and medication regimen (Joslin Diabetes Center & Joslin Clinic (Joslin), 2011)
- Three meals and two to three snacks is suggested to distribute glucose intake and reduce fluctuations in post-prandial glucose in women with GDM [American College of Obstetricians and Gynecologists (ACOG), 2013]
- Three meals and several snacks is recommended to prevent hypoglycemia, particularly for women with GDM who are taking glitazibure or multiple insulin injections. The food distribution should be individualized to tolerance and preference (Shields and Tsay, 2015)
- In pregnancy, decreased insulin sensitivity occurs as hormonal production increases (Shields and Tsay, 2015). Although no evidence was found to support a specific or range of CHO distribution at meals or snacks, customary practice suggests limiting the amount and type of CHOs at breakfast (Joslin, 2011). The CHO intake is reassessed at subsequent visits for possible adjustment according to the blood glucose records (Shields and Tsay, 2015). See the recommendation GDM:...
Macronutrients (GDM: Carbohydrate and Post Prandial Breakfast Glycemia).

- Recommendation Strength Rationale

Consensus: This topic was included in the EAL systematic review. However, no evidence was found to answer the research question. The recommendation is based on consensus publications.

- Minority Opinions

 None.

- Supporting Evidence

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

- References

- References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process

- Gestational Diabetes

- Gestational Diabetes (GDM) Guideline (2016)

Recommendations Summary

GDM: High-Intensity Sweeteners 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

- Recommendation(s)

GDM: Use of High-Intensity Sweeteners

In pregnant women with gestational diabetes mellitus (GDM), who choose to consume high-intensity sweeteners, the registered dietitian nutritionist (RDN) should educate the woman to select only those approved or generally recognized as safe (GRAS) by the US Food and Drug Administration (FDA) and to limit her intake to the acceptable daily intake (ADI), established by the FDA. The FDA has concluded the safety of six high-intensity sweeteners [saccharin, aspartame, acesulfame potassium (Ace-K), sucralose, neotame and advantame] when consumed within the ADI by the general population, including pregnant women. Steviol glycosides and Luo Han Guo (monk fruit) extracts are also GRAS when consumed within the ADI.

Rating: Consensus

Conditional

- Risks/Harms of Implementing This Recommendation

In a 1985 review of saccharin, the American Medical Association suggested pregnant women should consider avoiding saccharin, due to limited epidemiological studies in pregnant women and children (Council on Scientific Affairs, 1985). To date, more than 30 human studies have found that saccharin is safe for human consumption (FDA, 2015). Saccharin is approved for use as a non-nutritive, high-intensity sweetener by the FDA (FDA, 2015).

- Conditions of Application

- This recommendation applies to pregnant women with GDM who are considering use of high-intensity sweeteners as a replacement for higher calorie sweeteners (sucrose, fructose, honey, etc.)

- This recommendation applies only to high-intensity sweeteners within ADI levels (amount considered safe for consumption for each day over a person’s lifetime) approved by the FDA and those that are GRAS in the United States

- Individuals with phenylketonuria (PKU), a rare hereditary disease, have difficulty metabolizing phenylalanine which is a component of aspartame. Those with PKU should control their intake of phenylalanine from all sources, including aspartame (FDA, 2015).

- Individuals should not add high-intensity sweeteners to their diet with the intention of improving blood sugar levels. Consumption of high-intensity sweeteners has not been found to have a significant impact on glycemic control in non-pregnant adults with Type 1 and Type 2 diabetes (AND, 2015).

- Potential Costs Associated with Application

There are no obvious costs associated with the application of this recommendation.

- Recommendation Narrative

High-intensity sweeteners are commonly used as sugar substitutes or sugar alternatives because they are much sweeter than sugar, but contribute little to no carbohydrates or calories. High-intensity sweeteners are regulated by the FDA as a food additive, unless they are GRAS. Use of a GRAS substance does not require premarket approval by the FDA. For a substance to be determined GRAS there must be common knowledge within the expert scientific community regarding safety, and reasonable certainty that the substance is not harmful within intended use (Federal Register, 2016).

- Six high-intensity sweeteners are FDA-approved as food additives in the United States: Saccharin, aspartame, acesulfame potassium (Ace-K), sucralose, neotame and advantame. These high-intensity sweeteners met safety standards for consumption by the general population, including pregnant women under certain conditions of use, as identified by the FDA (FDA, 2015).

- An ADI level, which is the amount considered safe for consumption for each day over a person’s lifetime, was set for each of
the high-intensity sweeteners. GRAS notices have been submitted to the FDA for steviol glycosides and Luo Han Guo extracts (FDA, 2015).

- The position of the Academy of Nutrition and Dietetics is that use of nutritive sweeteners is acceptable during pregnancy, but there is very little evidence to support the use of nonnutritive sweeteners in GDM. However, FDA-approved nonnutritive sweeteners are acceptable, with the exception of aspartame in women with PKU (Fitch & Keim, 2012; Kaiser and Campbell, 2015). Based on FDA guidance, the American Diabetes Association (ADA, 2007) and Sweet Success (Shields and Tsay, 2015) also state that use of high-intensity sweeteners within the ADI are considered safe for consumption in the pregnant population.

- **Recommendation Strength Rationale**

  Consensus: This topic was not included in the EAL systematic review. The recommendation is based on consensus publications.

  - **Minority Opinions**

    None.

- **Supporting Evidence**

  The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

- **References**

- **References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process**


- **Kaiser LL, Campbell CG; Academy Positions Committee Workgroup. Practice paper of the Academy of Nutrition and Dietetics abstract: nutrition and lifestyle for a healthy pregnancy outcome. J Acad Nutr Diet. 2014 Sep;114(9): 1, 447. PMID: 25699300.**


- **Gestational Diabetes**

- **Gestational Diabetes (GDM) Guideline (2016)**

**Recommendations Summary**

GDM: Alcohol 2016

[Click here](http://www.andeal.org/template.cfm?template=guide_summary&key=4332) to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

- **Recommendation(s)**

  **GDM: Alcohol Intake**

  The registered dietitian nutritionist (RDN) should reinforce abstinence from alcohol during pregnancy for women with gestational diabetes mellitus (GDM). The safest choice for all pregnant women is to abstain from alcohol to eliminate the risk for alcohol-related birth defects such as behavioral or neurological defects, growth deficiencies, facial abnormalities and impaired intellectual development.

  **Rating: Consensus**

  **Impervasive**

  - **Risks/Harms of Implementing This Recommendation**

    There are no potential risks or harms associated with the application of this recommendation.

  **Conditions of Application**

  - **Alcohol use during pregnancy is typically addressed by their health care provider and other health professionals during the first obstetric visit and reinforced throughout pregnancy**

    - **Women who are unwilling or unable to refrain from alcohol consumption during pregnancy (e.g., heavy drinker, alcohol dependency, binge drinker) should be referred for supportive services, such as counseling and possible treatment (O’Leary and Bower, 2012).**

  **Potential Costs Associated with Application**

  There are no obvious costs that may be associated with the application of this recommendation.

  **Recommendation Narrative**

  - **Alcohol exposure during pregnancy has been linked to birth-related defects in the fetus including fetal alcohol spectrum disorders and low birth weight [Centers for Disease Control & Prevention (CDC), 2016].**

    - Evidence-based scientific information is limited given the nature of any research involving the alcohol intake of pregnant women. However, fetal alcohol spectrum disorders are the leading cause and the most commonly identifiable cause of preventable developmental delays and intellectual disabilities (CDC, 2016).

    - While there is a lack of consensus among healthcare providers regarding the amount of alcohol that would be considered detrimental to the fetus, the Academy of Nutrition and Dietetics (The Academy) and several health organizations, including American Academy of Pediatrics (Williams et al, 2015), American College of Obstetricians and Gynecologists (2012), March of Preemies (2015), and the American Foundation for Suicide Prevention (2012), recommend that pregnant women avoid alcohol consumption.

    - The Academy of Nutrition and Dietetics (AND) and the Academy of Nutrition and Dietetics (AND) states that use of nutritive sweeteners is acceptable during pregnancy, but there is very little evidence to support the use of nonnutritive sweeteners in GDM. However, FDA-approved nonnutritive sweeteners are acceptable, with the exception of aspartame in women with PKU (Fitch & Keim, 2012; Kaiser and Campbell, 2015). Based on FDA guidance, the American Diabetes Association (ADA, 2007) and Sweet Success (Shields and Tsay, 2015) also state that use of high-intensity sweeteners within the ADI are considered safe for consumption in the pregnant population.

    - The registered dietetic nutritionist (RDN) should reinforce abstinence from alcohol to eliminate the risk for alcohol-related birth defects such as behavioral or neurological defects, growth deficiencies, facial abnormalities and impaired intellectual development.

    - The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).
Dimes (2015), CDC (2016) have advised pregnant women to abstain from alcohol.

- The position of The Academy on alcohol use during pregnancy is as follows: "Alcohol should not be consumed by pregnant women or those who may become pregnant. Drinking alcohol during pregnancy, especially in early pregnancy, may result in behavioral or neurological defects in the offspring and affect a child’s future intelligence. No safe level of alcohol consumption during pregnancy has been established (Procter & Campbell, 2014)."

**Recommendation Strength Rationale**

Consensus: This topic was not included in the EAL systematic review. The recommendation is based on consensus publications.

**Minority Opinions**

None.

**Supporting Evidence**

The recommendations were created from the evidence analysis on the following questions. To see more detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

**References**

- References not graded in Academy of Nutrition and Dietetics Evidence Analysis Process.

**References**


**Gestational Diabetes**

**Gestational Diabetes (GDM) Guideline (2016).**

### Recommendations Summary

**GDM: Physical Activity 2016**

*Click here* to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

**Recommendation(s)**

**GDM: Physical Activity**

Unless contraindicated, the registered dietitian nutritionist (RDN) should encourage women with gestational diabetes mellitus (GDM) to engage in a goal to achieve daily moderate exercise of 30 minutes or more per day. In addition to a healthy diet, exercise can help improve blood glucose control and achieve weight gain recommendations. Both aerobic exercise and non–weight-bearing exercise (e.g., stretching, swimming, yoga, etc.) have been shown to lower blood glucose levels in women with GDM. Lifestyle therapy for GDM results in lower birth weight and a lower incidence of large-for-gestational-age births and pre-eclampsia.

**Rating:** Strong Conditional

**Risks/Harms of Implementing This Recommendation**

- High-intensity or prolonged exercise in excess of 45 minutes can lead to hypoglycemia [American College of Obstetricians and Gynecologists (ACOG), 2015]
- Pregnant women engaging in physical activity should be advised to ensure adequate caloric intake and to remain well hydrated (ACOG, 2015)
- Contact sports (ice hockey, boxing, soccer, basketball), activities with a high risk of falling (sking surfing, off-road cycling, gymnastics, horseback riding), scuba diving, sky diving, and hot yoga or hot pilates should be avoided (ACOG, 2015).

**Conditions of Application**

- Healthcare provider consultation is warranted or required prior to beginning any exercise program (ACOG, 2015)
- This recommendation applies to women with GDM for whom physical activity during pregnancy is not contraindicated.
  - Absolute contraindications include, but are not limited to: Hemodynamically significant heart disease, restrictive lung disease, incompetent cervices or cerclage, multiple gestation at risk of premature labor, persistent second or third trimester bleeding, placenta previa after 26 weeks of gestation, premature labor during the current pregnancy, ruptured membranes, preeclampsia or pregnancy induced hypertension (HTN) and severe anemia, and women with relative (ACOG, 2015)
  - Relative contraindications include, but are not limited to: Anemia, unevaluated maternal cardiac arrhythmia, chronic bronchitis, poorly controlled type 1 diabetes, extreme morbid obesity, extreme underweight [body mass index (BMI) below 12kg/m²], history of extremely sedentary lifestyle, intrauterine growth restriction in current pregnancy, poorly controlled HTN, orthopedic limitations, poorly controlled seizure disorder, poorly controlled hyperthyroidism, heavy smoker (ACOG, 2015)
- Pregnant women with relative contraindications to physical activity may be able to incorporate physical activity with individualized recommendations provided by their health care provider (ACOG, 2013).
- Pregnant women who were sedentary or obese prior to their pregnancy should follow a gradual progression to physical activity goals (ACOG, 2015)
- Overweight or obese women should start with low-intensity, short periods of exercise and gradually increase as able (ACOG, 2015).
- Pregnant women who regularly engaged in physical activity before they were pregnant can continue to engage in high intensity aerobic activity in the absence of contraindications (ACOG, 2015)
- Safe activities within moderate intensity include: Walking, swimming, stationary cycling, low-impact aerobics, modified yoga, modified pilates, running or jogging, racquet sports or strength training (ACOG, 2015).
Potential Costs Associated with Application
There are no obvious costs that may be associated with the application of this recommendation.

Recommendation Narrative
The recommendation is based on The Endocrine Society’s Diabetes and Pregnancy: An Endocrine Society Clinical Practice Guideline (Blumer et al, 2013) as follows:

- **Management of Elevated Blood Glucose**
  - "2.3b. We recommend that the initial treatment of gestational diabetes should consist of medical nutrition therapy and daily moderate exercise for 30 minutes or more. Rating: 1/+++ (Strong recommendation / Moderate quality evidence)"
  - Summary of support for the recommendation:
    - "Lifestyle therapy for GDM results in a lower incidence of reduced birth weight, large-for-gestational-age births, and preclampsia. Both aerobic exercise and non–weight-bearing exercise have been shown to lower blood glucose levels in women with GDM."

Summary
Physical activity recommendations for pregnant women do not differ from recommendations for the general public (ACOG, 2015). Physical activity during pregnancy is considered safe and desirable for pregnant women in the absence of complications or contraindications (ACOG, 2015; Office of Disease Prevention & Health Promotion, 2008). After a thorough evaluation from the healthcare provider, initial treatment of GDM should include a working goal toward daily moderate exercise of 30 minutes or more (Blumer et al, 2013). Pregnant women should use perceived exertion as a method to monitor exercise intensity. Physical activity has been found to lower blood glucose levels and promote recommended weight status in women with GDM (ACOG, 2013, 2015; Blumer et al, 2013).

Recommendation Strength Rationale

Minority Opinions
None.

Supporting Evidence
The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

References


- Gestational Diabetes
- Gestational Diabetes (GDM) Guideline (2016)

Recommendations Summary

GDM: Nutrition Monitoring and Evaluation 2016

Click here to see the explanation of recommendation ratings (Strong, Fair, Weak, Consensus, Insufficient Evidence) and labels (Imperative or Conditional). To see more detail on the evidence from which the following recommendations were drawn, use the hyperlinks in the Supporting Evidence Section below.

- **Recommendation(s)**
  - GDM: Nutrition Monitoring and Evaluation

Following the nutrition intervention of women with gestational diabetes mellitus (GDM), to check progress, the registered dietitian nutritionist (RDN) should monitor and evaluate the following components at each visit and compare to desired individual outcomes relevant to the nutrition diagnosis and nutrition intervention. This may include, but is not limited to:

Food/Nutrition-Related History Outcomes

- Daily food intake in relation to post-meal glucose readings
- Food, beverage and nutrient intake including
  - Calorie intake; types and amount of carbohydrate (including fiber), fat, protein; with special attention to high calorie, low-nutrient dense foods such as desserts, candy, sugar-sweetened beverages
  - Serving sizes
  - Meal and snack patterns, including frequency and duration
  - Recent changes to food choices and/or eating patterns
  - Preferences, avoidance, intolerances, allergies including
    - In relationship to gastrointestinal discomforts (e.g., nausea, vomiting, heartburn, constipation, ptosisism)
    - Reaction to or changes in food tastes/smells related to pregnancy
    - Cultural and religious considerations.
  - Appetite and changes in appetite
  - Frequency and intake of meals and snacks; meals eaten away from home
  - Methods of food preparation; food safety
  - Recommendation to add pharmacologic therapy (oral and/or insulin therapy) to maintain nutrient intake and achieve glycemic targets
  - Pharmacologic therapy – dose of diabetes medications: Oral glucose-lowering agent and insulin.
  - Changes in substance use: alcohol, tobacco, caffeine, recreational drugs
  - Knowledge, beliefs or attitudes: Motivation, readiness to change, self-efficacy; willingness and ability to make lifestyle

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andeal.org
Anthropometric Measurements

- Weight changes compared to previous obstetric visit or medical nutrition therapy (MNT) visit.

Biochemical Data, Medical Tests, and Procedure Outcomes:

- Self-monitoring blood glucose (SMBG) records, including meter downloads
- Ketone testing records (if previously recommended because of weight loss or inadequate calorie intake)
- Updated fetal and maternal testing or lab values.

Nutrition monitoring and evaluation of these factors is needed to correctly/effectively diagnose nutrition problems that should be the focus of further nutrition interventions. Inability to achieve optimal nutrient intake may contribute to poor outcomes or initiation of or changes in pharmacologic therapy.

Rating: Consensus

Imperative

- **Risks/Harms of Implementing This Recommendation**
  There are no potential risks or harms associated with the application of this recommendation.

- **Conditions of Application**
  - If necessary data are not available, the RDN should use professional judgment to request or obtain addition data
  - Women who have complicating conditions such as renal disease or eating disorders may require more indepth or specialized nutrition assessments [American Diabetes Association (ADA), 2016]
  - RDNs should be alert to psychosocial stressors, such as family and household strain, verbal or physical abuse, exposure to discrimination, food insecurity, unemployment, low resources, major or catastrophic life events and anxiety about the current pregnancy. Such stressors may indicate need for further screening and referral to a mental health professional for early treatment to prevent adverse pregnancy outcomes (Kaiser and Campbell, 2014).

- **Potential Costs Associated with Application**
  Accessibility and costs of additional testing should be considered.

**Recommendation Narrative**

The purpose of nutrition M&E is to assess the effectiveness of nutrition intervention through monitoring, measuring, and evaluating changes in nutrition care indicators. The RDN determines the progress made for the nutrition intervention and whether the patient/client’s nutrition related goals or desired outcomes are being achieved (eNCPT, 2016).

Outcomes are measured by data collection of appropriate nutrition outcome indicator(s).

Nutrition M&E in GDM is organized under three domains (categories): Food/Nutrition-Related History Outcomes; Anthropometric Measurement Outcomes; and Biochemical Data, Medical Tests, and Procedure Outcomes (eNCPT, 2016).

**Food or Nutrition-Related History**

Food and nutrition-related history pertinent to diabetes and pregnancy include:

- Dietary history includes a thorough review of usual food intake, pattern of intake (timing, meals and snacks) and previous history of diet adherence (ADA, 2013)
- Educational knowledge, such as nutrition and meal planning skills, barriers to dietary compliance, such as lack of family support, daily schedule or economic issues, etc. (ADA, 2013)
- Macronutrient (especially CHO and fiber) and micronutrient food intake (ADA, 2013)
- Vitamin and mineral supplement use (prenatal and non-prenatal) or use of natural remedies, such as herbs or alternative therapies (ADA, 2013)
- Food allergies/intolerances (ADA, 2013)
- Use of alcohol, tobacco, caffeine, or other substances [Joslin Diabetes Center & Joslin Clinic (Joslin), 2011]
- Medications: prescription (diabetes-related, non-diabetes-related); over the counter medications (ADA, 2013)
- Exercise pattern — type, frequency, duration [American College of Obstetricians and Gynecologists (ACOG), 2015; Joslin, 2013]

**Anthropometric Measurements**

Anthropometric measurements pertinent to diabetes and pregnancy include:

- Weight should be tracked at each visit to determine if the gestational weight gain (GWG) is appropriate (within range), based on Institute of Medicine (IOM) revised guidelines for weight gain during pregnancy (IOM, 2009) (Shields and Tsay, 2015; Joslin, 2011; Kaiser and Campbell, 2014)
- Inappropriate weight gain (excess or inadequate weight gain) may require further assessment of food and calorie intake and adjustment in the nutrition prescription. (ADA, 2016).

**Biochemical Data, Medical Tests and Procedures**

Clinical data related to medical tests pertinent to diabetes and pregnancy include:

- Post-prandial SMBG is recommended (ADA, 2016) to determine glycemic control throughout pregnancy (Shields and Tsay, 2015; ACOG, 2013)

**Recommendation Strength Rationale**

Consensus: This topic was not included in the EAL systematic review. The recommendation is based on consensus publications.

**Minority Opinions**

None.

**Supporting Evidence**

The recommendations were created from the evidence analysis on the following questions. To see detail of the evidence analysis, click the blue hyperlinks below (recommendations rated consensus will not have supporting evidence linked).

**References**

- Institute of Medicine (IOM) and National Research Council. Weight Gain During Pregnancy: Reexamining the Guidelines.

© 2017 Academy of Nutrition and Dietetics (A.N.D.), Evidence Analysis Library. Printed on: 03/30/17 - from: http://www.andel.org