IUGR and Macrosomic Phenotypes – How They Develop and How They Change Over Time

Presented by
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In accordance with the Accreditation Council for Continuing Medical Education Standards, parallel documents from other accrediting bodies, and Annenberg Center for Health Sciences policy, the following disclosure has been made:

William W. Hay, Jr, MD
One time Consultant: Baxter–IV nutrition
Normal human fetuses are supposed to grow (regardless of the growth curve) at an **average** rate of ~17 g/kg/day from 28-40 weeks, with symmetrical growth of head and length.

**Average 50th %ile birth weights by gestational age**

From 6 sources adapted from Naeye R, Dixon J. *Pediatr Res.* 1978;12:987-991 (including “mile high” Denver, CO Lubchenco curve);

**Fenton** TR, Kim JH. *BMC Pediatr.* 2003;13:59;

17 g/kg/day is the average weight gain of the normal human fetus.

More energy in the maternal diet, fatter fetus, especially the LGA fetus, at risk for later obesity.

Not enough energy in the maternal diet, thinner fetus, at risk for later fat gain.

Both extremes of fetal growth (SGA/IUGR and LGA/Macrosomia) are associated with similar adult phenotypes of obesity, insulin resistance, diabetes, and cardiovascular disease. HOW?? And are these the results of similar or different mechanisms?
Under Nutrition

IUGR / SGA fetuses and neonates and consequences of trying to feed them more.
Glucose Stimulated Insulin Secretion (GSIS) is reduced near term) in IUGR fetal sheep, due to

Pancreatic Growth Failure—with later insulin deficiency (Beta Cell/Islet proliferation and Insulin Secretion—both reduced in the IUGR fetus)

Human IUGR Fetuses: Reduced glucose-stimulated insulin secretion

Nicolini et al, 1990

and β-cell mitosis

Limesand et al. 2006
Adverse effects of reduced oxygen

Hypoxia ➔ Increased Catecholamines

Decreased insulin secretion
Decreased insulin action

Hypoxia ➔ Increased Cortisol

Increased gluconeogenesis
Increased protein breakdown

Fetal insulin concentrations increase with blocking of catecholamines, proof that they suppress insulin secretion. Exaggerated in IUGR fetuses! After birth, good care might backfire!

Sean Limesand’s studies

IUGR islets also have increased **Fractional Islet Insulin Secretion**! (in culture, without or with catecholamines)

Positive adaptation to “chronic” reductions in fetal glucose, increasing **Glucose and Insulin Sensitivity**—with risk of later life development of increased fat storage and obesity.

These changes maintain glucose utilization and insulin sensitivity—ie, GUR/kg is normal at less than normal [Glu] and [Ins].

**Table:**

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<thead>
<tr>
<th></th>
<th>Control</th>
<th>IUGR</th>
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<tbody>
<tr>
<td>GUR</td>
<td>5 mg/kg/min = 5 mg/kg/min</td>
<td></td>
</tr>
<tr>
<td>[G]</td>
<td>20 mg/dL</td>
<td>10 mg/dL</td>
</tr>
<tr>
<td>[I]</td>
<td>12 μU/mL</td>
<td>5 μU/mL</td>
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Fetuses with chronic IUGR show reduced P-ERK-1,2, mTOR, and eIF4E, plus an increase in total/phosphorylated binding protein, 4EBP-1-P, indicating decreased capacity for synthesis of amino acids into protein and IGF-1 stimulated cell proliferation. A result? A potential cause? Or both?

Courtesy of Jed Friedman, PhD, University of Colorado School of Medicine, Aurora, CO.
Decreased fetal muscle mass in Placental Insufficiency IUGR model

Chronically restricted fetal AA supply, energy, insulin, and IGFs in IUGR fetuses (and preterm infants) decreases muscle growth (mass and myofiber area).

Mechanisms?
Chronic placental insufficiency induced IUGR leads to hepatic insulin resistance and development of persistent glucose production.

Hepatic insulin resistance in PI-IUGR fetal sheep. (A) In vivo glucose production rates and (B) liver PCK1 mRNA are increased in PI-IUGR fetuses (basal) and not suppressed in insulin-clamp (n=6-8 each).

Courtesy Stephanie Wesolowski, PhD, University of Colorado, Denver, CO.
Mixed Hyper- and Hypo-glycemia in IUGR infants.
More complicated than what you thought!
(a clinical problem explained by basic science research)

1. Hyperglycemia
   • Reduced pancreatic β-cell number and insulin production.
   • Hypoxia and catecholamine suppression of insulin secretion and insulin action.
   • Hepatic insulin resistance (hypoxia and high catecholamines) and increased glucose production (increased cortisol).

2. Hypoglycemia
   • Greater head/brain to body/liver ratio, thus greater body weight-specific glucose utilization rate.
   • Increased peripheral tissue glucose uptake capacity, from increased or at least maintained glucose transporters.
   • Increased fractional insulin secretion (particularly after reduction in catecholamine concentrations) and greater susceptibility to metabolic stimulation of insulin secretion.
Other mixed outcomes

Catch-up Growth in Exclusively Breast Fed SGA Infants

- Growth occurs in sequence
- Normalization of lean mass occurs by 4 months of age
- Normalization of bone mineral content occurs by 12 months
- Fat mass still reduced at 12 months of age

Catch-up Growth in Formula Fed SGA Infants

- Early catch up of fat mass, and then production of obesity
- Associated with elevated levels of IGF-1 and low levels of adiponectin (adipocyte-secreted hormone that enhances insulin sensitivity and glucose metabolism), risk factors for later development of obesity and insulin resistance.

“Normal” fetuses and preterm infants of the same gestational age need more protein to grow. The key is the right amount at the right time!

Between 24 and 30 weeks, amino acid requirements = 3.6-4.8 g/day.

Between 32 and 37 weeks, fractional growth rate decreases, as does the protein requirement for growth, to 2.5-3.5 g/kg/day.

At term, protein requirements decrease to those of the normal breast fed infant, or 1.5-2.0 g/kg/day.

Caution—Long-term adaptation to energy and protein deficiency in IUGR fetuses.

Reduced rate of growth, but return to normal cellular energy/oxidative metabolism.

Higher protein diets in chronically IUGR infants might not work as well.
Promotion of faster weight gain in infants born SGA: Is there an adverse effect on later blood pressure?

- Infants e37 weeks gestation from 5 hospitals in the UK
- Birth weight <10 percentile
- Randomized: standard infant formula (n=147; n=83 age 6-8 years) or nutrient enriched formula until 9 months of age (n=152; n=70 age 6-8)

<table>
<thead>
<tr>
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<th>Standard (n=83)</th>
<th>Nutrient Enriched (n=70)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diastolic</td>
<td>61.3</td>
<td>64.8</td>
<td>0.01</td>
</tr>
<tr>
<td>MAP</td>
<td>76.9</td>
<td>79.9</td>
<td>0.03</td>
</tr>
<tr>
<td>Systolic</td>
<td>100.5</td>
<td>102.5</td>
<td>0.2</td>
</tr>
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Adjusted for age, sex, socioeconomic status, z score for weight and height

- Faster early weight gain programmed higher later blood pressure in both groups of SGA infants.

Given the growth restriction phenotype, growth restriction appears to be a “set up”.

Adaptations to under nutrition, when presented with different later life conditions (eg, high calorie diet, limited exercise), lead to several specific adult disorders—

- Decreased pancreatic growth, development, and insulin secretion
  Type 2 Diabetes
- Increased insulin and glucose sensitivity for glucose uptake
  Obesity
- Decreased amino acid synthesis into protein and for cell growth
  Short stature, poor neurodevelopment and cognitive capacity
- Development of hepatic insulin resistance and glucose production
  Hyperglycemia, Type 2 diabetes
- Increased development of later obesity
  Inflammation, cardiovascular disease (hypertension, stroke, MIs)
Later life metabolic and nutritional problems of the IUGR infant

Developmental programming of excess fat production, aggravated by fat/high glycemic index diets to develop—

more obesity
insulin resistance
glucose intolerance
type 2 diabetes
systemic inflammation
cardiovascular disease

Metabolic Syndrome
Over Nutrition

LGA-MACROSOMIC INFANTS

and consequences of trying to feed them more.
First off, they don’t look like they need more food.
And not just IDMs—many infants are fatter and fatter these days as maternal obesity grows and grows.

Courtesy Patrick Catalano, MD, The MetroHealth System, Cleveland, OH
Excessive maternal weight gain during pregnancy, directly related to their dietary fat and high glycemic index food intake, directly correlates with fetal macrosomia.

Odds Ratio for High Birth Weight (> 4000 g).
(mean and 95% confidence intervals)

Early and marked return to obesity in childhood in overfed children who were macrosomic infants of diabetic mothers.

And the obesity that is produced and regenerated in macrosomic IDM offspring leads to Type 2 Diabetes.

And not just in IDMs
Prevalence of overweight at 2, 5, and 8 years of age in children in relation to birth size (non-IDMs)
Prevalence of **METABOLIC SYNDROME** at any age among children grouped according to birth weight and maternal diabetes.

Metabolic and nutritional problems of the macrosomic infant

Already programmed with excess fat from maternal obesity and high fat/high glycemic index diet to further develop obesity and its consequences—

- more obesity
- insulin resistance
- glucose intolerance
- diabetes
- systemic inflammation
- cardiovascular disease

Metabolic Syndrome
What could we do to prevent such rapid gains in weight and the inevitable consequences?
Breastfeeding, considered dichotomously (yes or no), and the Odds Ratio for Later Obesity

- Obesity in the US affects 35%, of adults (~17% of youth aged 2-19).
- Women that were OW/OB prior to conception carried 60% of all US pregnancies (2013).
- Children born to obese mothers have increased likelihood of childhood obesity.
- **Exclusive breastfeeding is protective against elevated obesity risk.**

![Figure 6. Breastfeeding, considered dichotomously (yes or no), and the OR for later obesity.](image-url)

Reproduced with permission from *Pediatrics*, Vol. 114, Pages 1146-1173, Copyright © 2004 by the AAP.
But—milk has to be “healthy” too. Possible that the milk n-6/n-3 ratio is more important than maternal BMI and fat gain in producing offspring fat mass?

Each 1-unit increase n-6/n-3 = 0.6–1.4 g fat mass/day.

The benefit might be very long lasting—
Adult mice who received a LOW n-6/n-3 fatty acid ratio diet as neonates had better glucose tolerance post 4-weeks of High Fat/High Sucrose diet: Eating well might be beneficial!

High neonatal n-6/n-3 fatty acid ratio diet—more glucose intolerant
Low neonatal n-6/n-3 fatty acid ratio diet—more glucose tolerant

Unpublished data courtesy Michael Rudolph, PhD, University of Colorado School of Medicine, Denver, CO. 2015
The benefits of Calorie Restriction (*percentage increase in lifespan over controls*) in relation to the extent of restriction (*percentage decrease in intake relative to controls*); studies of both *rats* and *mice*.

The effect of Calorie Restriction expressed relative to the same effect when initiated at weaning as a function of the proportion of expected lifespan that had elapsed when restriction was initiated. Several studies of rodents (mice and rats).

The earlier you start restricting calories, the longer you will live.

Modeled benefits of Calorie Restriction (percentage extension of remaining life) in relation to the proportion of expected life remaining when restriction was initiated. Even when expressed relative to the remaining lifespan, the benefits decline as age of onset increases.

If you’re over 60, LIVE IT UP!! as there is little further increase in life span in response to calorie restriction.

Although maybe **YOU** should be careful of too much dessert.

Courtesy Richard Johnston, MD, PhD.
Summary

1. Growth patterns, from small to large, are a continuum, and represent more or less nutrition.

2. Nutrition does matter, and the mechanisms—programming via epigenetic phenomena in the context of developmental plasticity—are clearly well established and fundamental.

3. Over feeding energy for any infant (preterm, IUGR, SGA, LGA, IDM) has only produced more fat gain, which persists, contributing measurably to obesity and the metabolic syndrome in later life.

4. Individual fatty acids, particularly the n-3 long chain PUFAs, may be more important than the amount of lipid fed to infants.

5. While human milk appears to be best, mother has to eat well to optimize it. Healthy mothers produce healthy offspring!

6. Establish healthy eating early, continue it for life.
University of Colorado School of Medicine

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• Kendra Hendrickson, RD (UCH)
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