Endocrine Disrupters as Obesogens

Is the environment making us fat?

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The Worldwide Obesity Epidemic

- 60 million people in the US are clinically obese
  - >30% above “ideal” body weight

- Obesity accounts for 8% of healthcare costs in Western countries
  - $75 billion annually in US (2005)

- Obesity is associated with “metabolic syndrome” -> type 2 diabetes and cardiovascular disease
  - Central (abdominal obesity)
  - Atherogenic dyslipidemia (high triglycerides, high LDL, low HDL)
  - Hypertension
  - Insulin resistance
  - Prothrombotic state
  - Pro-inflammatory state (elevated CRP)
How does obesity occur?

- Prevailing wisdom - “couch potato syndrome”
  - Positive energy balance, i.e., too much food, too little exercise

- Other factors?
  - Stress (elevated glucocorticoids)
  - Inadequate sleep (stress?)
  - “Thrifty” genes which evolved to make the most of scarce calories
  - Viruses, SNPs

- What about role of prenatal nutrition or in utero experience?
  - Maternal smoking decreases birth weight and increases obesity

- What about the role of industrial chemicals in rise of obesity?
  - Baillie-Hamilton (2002) postulated a role for chemical toxins
  - Obesity epidemic roughly correlates with a marked increase in the use of chemicals (plastics, pesticides, etc.)

- Many chemicals have effects on the endocrine system
Hormonal control of weight

- Hormonal control of appetite and metabolism
  - Leptin, resistin adiponectin ghrelin are key players
    - Leptin, adiponectin, resistin - adipocytes
    - Ghrelin - stomach
  - Thyroid hormone/receptor
    - Sets basal metabolic rate

- Hormonal control of fat cell development and lipid balance
  - Regulated through nuclear hormone receptors RXR, PPARγ
  - PPARγ - master regulator of fat cell development
    - increased fat cell differentiation
    - Increased fat storage in existing cells
    - Increased insulin sensitivity

Nuclear Receptors - A Large Family of Ligand Modulated Transcription Factors

- Bind to specific DNA targets - Hormone Response Elements
  - Most are activators
  - Some constitutive
  - Few inactivate
- Ligands are small lipophilic molecules that freely enter cells
  - Diffuse from source & penetrate to a target
- Respond to low levels of hormone
  - Parts per billion levels
  - Regulation of levels is important
  - Can be disrupted by environmental contaminants
Endocrine Disrupting Chemicals (EDCs)

- *Endocrine disrupter* - a compound that mimics or blocks the action of endocrine hormones, either directly or indirectly

- Often persistent pollutants or dietary components

- Disturb development, physiology and homeostasis

- Frequently act through nuclear hormone receptors
  - Environmental estrogens
  - Anti-androgens
  - Anti-thyroid

- Are disturbances in endocrine signaling pathways involved in adipogenesis and obesity?
The Obesogen Hypothesis

- **Obesogens** - chemicals that inappropriately stimulate adipogenesis and fat storage, exist and contribute to obesity epidemic

- Pre- and postnatal exposure to environmental estrogens (ER) increases weight
  - DES, genistein

- Thiazolidinedione anti-diabetic drugs (PPARγ)
  - Increase fat storage and fat cell size at all ages in humans
  - Reduce insulin resistance in muscle but increases obesity which exacerbates diabetes

- Several compounds cause adipocyte differentiation in vitro (PPARγ)
  - Organotins, phthalates, BPA, PFOA, alkylphenols,

- Urinary phthalates correlate with waist diameter and insulin resistance in humans
Endocrine disruption by organotins

- Tributyltin -> causes imposex in molluscs
  - Imposition of male sex characteristics on female mollusks
- Impairs shell development in bivalve mollusks
- Sex reverses fishes (genetically female flounder and zebrafish -> males)
How do organotins cause endocrine disruption?

• How do organotins cause sex-reversal in mollusks and fishes?
  - Direct inhibition of aromatase (CYP19) activity (µM concentrations)
  - Inhibition of testosterone storage

• Transcriptional effects on aromatase expression
  - CYP19 in human ovarian granulosa cells is sensitive to inhibition by TBT, RXR- and PPARγ-specific ligands

• TBT alters sex determination - typically female -> male
  - TBT effects are seen at nM doses and below
  - Sex determination requires sex steroids at critical times
  - Sex steroids act through nuclear receptors

• Hypothesis - TBT alters the activity of one or more nuclear receptors, thereby causing endocrine disruption
  - Test nuclear receptors for activation or inhibition by organotins
  - Expect effect on steroid receptor activity
What is the effect of TBT on hormone receptors?

• TBT did not affect activity of sex steroid receptors at all

• TBT binds to and activates RXR-PPARγ with high affinity
  - TBT exposure causes RXR-PPARγ target genes to be expressed
  - Activation of RXR-PPARγ converts susceptible cultured cells into fat cells

• Prenatal TBT exposure predisposes animals to become obese
  - Exposed animals were about 15% heavier despite normal diet and exercise
  - Metabolism has been altered
How does TBT exposure cause weight gain?

- Changes in the hormonal control of appetite and satiety?
- Altered ability of adipocytes to process and store lipids?
- Increased number of adipocytes or pre-adipocytes?

Mesenchymal stem cells (MSCs) (now called multipotent stromal cells) precursors to many lineages including bone, cartilage, and adipose.
  - MSCs differentiate into adipocytes following rosiglitazone exposure
  - MSCs may (or may not) home to adipose depots after induction

TBT induces cultured MSCs to become adipocytes
  - Effects of TBT on human obesity likely to be via MSCs
Obesogens - Just the Tip of the Iceberg?

- Tributyl Tin
- Estradiol
- Genistein
- Organophosphate pesticides
- Phthalates
- DES
- Nicotine
- PCBs?
- PFOA
- Bisphenol A
- PBDEs?

What don’t we know yet?
- Body burdens in population
- Molecular targets of action beyond RXR-PPARγ
- Critical windows of exposure
- How does prenatal exposure alter adult phenotype?
- Is the prenatal reprogramming epigenetic?
Take Home Messages

- Diet and exercise are insufficient to explain obesity epidemic
- Obesogens inappropriately stimulate adipogenesis and fat storage
  - Environmental contaminants
    - TBT, environmental estrogens (BPA, DEHP), PFOA
  - Prescription drugs
    - Thiazolidinedione anti-diabetic drugs (Actos, Avandia)
    - Atypical antipsychotics (olanzapine)
    - Anti-depressants (tricyclics, SSRIs)
- Prenatal obesogen exposure reprograms the metabolism of exposed animals, predisposing them to obesity.
  - Likely to be epigenetic
- Obesogen exposure targets multiple cellular pathways, some of which involve nuclear receptors.