M2 - Endocrine, Winter 2008

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Abnormalities of Growth & Development

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OBJECTIVES

To understand the

- determinants of normal growth
- common variations in normal growth
- diagnostic approach to a child with abnormal growth
- principles of management of a child with abnormal growth
Topics **NOT** covered in today’s discussion

- Sexual differentiation
- Ambiguous genitalia and disorders of sexual differentiation
- Pubertal development
- Disorders of pubertal development - delayed / precocious
- Physiology of hormone secretion / action
Determinants of Normal Growth

Normal growth is the aggregate of hormonal, environmental, nutritional, and genetic factors.

Hormonal Factors

- **Thyroid** - essential for normal growth
  - hypothyroidism is a common cause of severe growth delay
- **Sex steroids** - bone maturation is dependent on estrogen
  - testosterone can enhance GH secretion
- **Glucocorticoids** - potent inhibitor of growth
GH/IGF-1 Axis

Image of GH/IGF-1 Axis removed
### Determinants of Normal Growth

<table>
<thead>
<tr>
<th>Hormone</th>
<th>Growth Rate $\text{cms/yr}$</th>
<th>Adult Height</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex Steroids $\uparrow$</td>
<td>Increase</td>
<td>Diminished</td>
</tr>
<tr>
<td>Sex Steroids $\downarrow$</td>
<td>Normal</td>
<td>Increased</td>
</tr>
<tr>
<td>Thyroxine $\uparrow$</td>
<td>Normal/± incr</td>
<td>Normal</td>
</tr>
<tr>
<td>Thyroxine $\downarrow$</td>
<td>Decreased</td>
<td>Diminished</td>
</tr>
<tr>
<td>GH $\uparrow$</td>
<td>Increase</td>
<td>Increased</td>
</tr>
<tr>
<td>GH $\downarrow$</td>
<td>Decrease</td>
<td>Diminished</td>
</tr>
<tr>
<td>Cortisol $\uparrow$</td>
<td>Decrease</td>
<td>Diminished</td>
</tr>
<tr>
<td>Cortisol $\downarrow$</td>
<td>Normal</td>
<td>Normal</td>
</tr>
</tbody>
</table>
Normal Growth

Anthropometric parameters

- Weight
- Measurement of height - Stadiometer
  - less than 2 yrs of age - length (supine)
  - greater than 2 yrs of age - height (erect)
- Head circumference
- Span
- Upper segment / lower segment ratio
Normal Growth

Anthropometric parameters

Upper / Lower Segment Ratio

Lower segment: symphysis pubis to floor
Upper segment: Ht (-) lower segment

Image of fetal and post-natal growth chart removed

<table>
<thead>
<tr>
<th>2m</th>
<th>5m</th>
<th>Birth</th>
<th>2yr</th>
<th>6yr</th>
<th>12yr</th>
<th>25yr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fetal</td>
<td>Post-natal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Girls 2-18 yrs

Source: Undetermined
Normal Growth

Growth Velocity

- measured in cms/yr
- should be measured over at least a 6-12 month period
- more the # of height points used to calculate GV – more reliable is the interpretation
- assessment of pubertal status is critical for interpretation of GV
- Normal GV – is a strong argument AGAINST a significant hormonal abnormality
Girls 2-18 yrs

Source: JM Tanner, et al.
Boys 2-18 yrs

Growth rate (cms/yr)

Age (yrs)

Source: JM Tanner, et al.
Normal Growth

- Chronological age
- Dental age
- Bone Age (skeletal maturation)

Beyond the neonatal age - X-ray of L wrist

Comparison with published standards (Greulich & Pyle)

Usefulness - prediction of final height

- Age of onset of puberty closely linked to bone age
- Corroborates diagnosis, but is never diagnostic

Caveats - imprecise / ethnic variability
SHORT STATURE

- Definition
- Classification
- Etiology
- Evaluation / Diagnostic Approach
- Treatment
Short Stature
height < 3rd percentile

Growth Retardation
growth velocity < 3rd percentile
Short Stature
height < 3rd percentile
Growth Retardation
growth velocity < 3rd percentile

Etiology

Normal Variant  Pathological
**Short Stature**

**Normal Variant**

**Familial / Genetic**
- Final height appropriate for parental height
- Normal size at birth
- GV may be ↓ in 0-3 yrs of age
- BA = CA

**Constitutional Delay of Growth & Puberty**
- "Late Bloomer"
- Family history
- Normal size at birth
- Normal GV
- Delayed puberty
- BA < CA

\[ \text{BA} = \text{bone age} \]
\[ \text{CA} = \text{chronological age} \]
14 yr old boy
h/o “shortest in his class”
h/o “always a small boy”
h/o father did not “grow” till he entered college

Prepubertal
GV = 5.0 cm/yr
Normal BUN / ESR
Normal Free T4 & TSH
Low-normal IGF-1
Normal IGFBP3 (for Tanner stage)
Bone age = 11.5 yrs

MRI
Normal

Testosterone 50 mg / q 4wks x 3 doses

Source: Undetermined
Short Stature
height < 3rd percentile
Growth Retardation
growth velocity < 3rd percentile

Definitions
- Proportionate
- Disproportionate

Etiology
- Normal Variant
- Pathological
  - Proportionate
  - Disproportionate
Hypothyroidism
GH deficiency
Cushing’s syndrome

Malabsorption
Inflammatory bowel disease
Celiac disease

Chronic renal failure
Renal tubular acidosis

Cardiac
Pulmonary
Liver
Infection

Psychosocial Dwarfism
Emotional Deprivation Syndrome
4 yr old boy
Voracious appetite / drinks urine - toilet bowl
Withdrawn / flat affect
No dysmorphic features
Chaotic home situation - abusive father

Ht age = 1 yr old

All lab tests normal
4 yr old boy
No dysmorphic features
Chaotic home situation - abusive father

Emotional deprivation syndrome
Psychosocial dwarfism

Admitted to hospital for observation

Source: Undetermined
6 yr old girl
GV = 3.0 cm/yr
No dysmorphic features
Chaotic home situation – parent incarcerated – shuttled through couple of foster homes

Adopted by a family
Stable home environment

All lab tests normal

Emotional deprivation syndrome
Psychosocial dwarfism

Source: Undetermined
SHORT STATURE

Skeletal Abnormalities
- Dysplasia
- Achondroplasia
- Rickets
- Vertebral anomalies

Dysmorphic Syndromes
- Turner
- Down
- Russell-Silver
- Prader-Willi
- Pseudo-hypoparathyroidism

Pathological Disproportionate
SHORT STATURE

Evaluation
Clinical History

Prenatal
- maternal infection, alcohol

Pattern of growth
- birth wt and length

Family History
- onset of puberty

Nutrition

Systemic Disease

Drugs
- steroids

Neurological
- headache, vision, enuresis

Psychosocial
First sign of puberty on PE:
♀ breast dev / ♂ incr in testicular volume

Anthropometric
ht, wt, head circ., arm span, U/L ratio

Nutritional state

Tanner Staging for Pubertal Development

Dysmorphic Features

Neurological exam

Thyroid Gland
Target Height (in cms)

girl = \frac{[father's \ ht + mother's \ ht] - 13}{2}

boy = \frac{[father's \ ht + mother's \ ht] + 13}{2}

normal range is ± 8 cms
Key Parameter - Growth Velocity

Normal GV
- Familial
- Constitutional

Impaired GV
- Malnutrition
- Chronic systemic illness
- IUGR
- Psychosocial
- Chromosomal abnormalities
- Endocrine
- Malabsorption
- Bone dysplasias
SHORT STATURE

Screening Tests

CBC, ESR, BUN
FT_{4}, TSH
IGF-1, IGFBP3
Tissue Transglutaminase ab

KARYOTYPE
- in girls to exclude TURNER
- dysmorphic features

Evaluation Laboratory Tests

RADIOLOGICAL
- bone age
- skeletal survey
SHORT STATURE

Growth Hormone Deficiency (GHD)
Neonatal - normal size / hypoglycemia / jaundice / micropenis / midline defect
Decreased growth velocity
Delayed dentition / mid-facial hypoplasia
Increase in adiposity
*SHORT STATURE*  

**GH Deficiency (GHD) Causes**

- Tumor - craniopharyngioma
- Trauma - surgery / irradiation
- Idiopathic
- Congenital Aplasia / Hypoplasia / Septic-optic dypslasia
- Genetic Defects -
  - Isolated Growth Hormone Deficiency (IGHD)
  - PROP1 / POU1F1 (*Pit1*)
**Criteria for diagnosing GH deficiency**

- Clinical (NOT laboratory) diagnosis
  - GV < 2 SD
  - Low IGF-1 & IGFBP-3
  - Provocative GH Level < 7-10 ng/ml

**Corroborative evidence**

- Delayed BA
- Related pathology
**SHORT STATURE**

**GH Deficiency Diagnosis**

**Measurement of GH**

- Spontaneous pulsatility of GH precludes random measurement
- Provocative test after overnight fast
  - Insulin induced hypoglycemia is the “Gold standard”

**IGF-1 / IGFBP3**

- Altered by nutritional status
- Normal range related to age & pubertal status
Growth hormone deficiency
Turner syndrome
Renal disease, before transplant
Small for gestational age
Prader-Willi syndrome
Idiopathic short stature
SHORT STATURE

**Treatment**

GH Replacement Therapy

s/c injection
7 days/wk

Side Effects

- Secondary/tertiary hypothyroidism
- Worsening of scoliosis
- Slipped capital femoral epiphysis
- Pseudotumor cerebri

Monitor

GV, Free T₄, IGF-1, IGFBP3
8 1/2 yr old girl
h/o poor growth x 12-18 months
recent h/o vague headaches
school performance has recently deteriorated
recent episodes of enuresis

Prepubertal
GV = 1.5 cm/yr
Low Free T<sub>4</sub>, Normal TSH
Low IGF-1 & IGFBP3
Karyotype = 46 XX
Bone age = 6 yrs

MRI
craniopharyngioma

Source: Undetermined
8 1/2 yr old girl
h/o poor growth x 12-18 months
h/o vague abdominal discomfort

Prepubertal
$GV = 2.5 \text{ cm/yr}$
Normal Free $T_4$ & TSH
Low IGF-1
Normal IGFBP3
Karyotype = 46 XX
Bone age = 7.5 yrs

Decreased serum albumin, microcytic anemia
ESR - 30

Tissue transglutaminase antibodies +ve
Small Intestine Biopsy - CELIAC DISEASE

Source: Undetermined
5 yr old girl
GV = 3.0 cm/yr
subtle dysmorphic features - clinodactyly, webbing of neck ±, t carrying angle
GV = 3.0 cm/yr
Normal Free T₄ & TSH
Normal IGF-1
Normal IGFBP3
Bone age = 5.0 yrs
Karyotype = 45,X
TURNER SYNDROME
Source: Undetermined
Turner Syndrome
Described in 1938 by Dr. Henry Turner
Most common sex chromosomal abnormality in females -- X chromosome
Frequency 1:1500 to 1:2500 in live born infant girls
15% of spontaneous abortions = TS
Karyotype 45, X
Turner Syndrome

Clinical Features - Postnatal

- Growth Failure 80-100%
- Gonadal Dysgenesis 80-100%
- Inverted/ widespaced nipples 60%
- Nail dysplasia 60-80%
- High narrow palate 60-80%
- Cardiac malformation 40-60%
- Renal dysplasia 40-60%
- Low hairline/webbing 30-40%
- Pigmented nevi common
Lymphedema at birth is highly correlated with 45,X karyotype and congenital heart abn

BY: Johannes Nielsen, et al.
Growth velocity (and NOT height) is the key anthropometric parameter.

Normal growth velocity virtually excludes a pathological cause for short stature.

Always exclude Turner’s syndrome in a girl with short stature.

Diagnosis of a child with growth problems is made more on CLINICAL and less on laboratory criteria.