

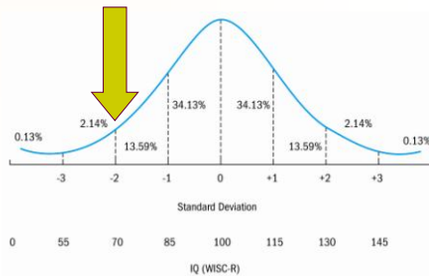
Intellectual Disability



DSM-IV Diagnostic Criteria

- A: Significantly sub-average intellectual functioning (an IQ of approximately 70 or below on an individually administered IQ test).
 - ≤ 2 SD of population mean
- B: Concurrent deficits or impairments in present adaptive functioning in at least two of the following areas: communication, self-care, home living, social/interpersonal skills, use of community resources, self-direction, functional academic skills, work, leisure, health, and safety.
- C: Onset is before age 18 years.

Intelligence



Levels of Intellectual Disability

Level	IQ Range	% of MR Pop.	Social	Communication	Sensorimotor	Academic Potential	Vocational/Living
Mild	50-55 to 70	85%	Develops in preschool years	Develops normally into preschool years	Minimal deficits	6 th grade	Usually achieves adult skills for self-support May need assistance; often successful
Moderate	35-40 to 50-55	10%	Benefits from training	Develops normally in early childhood		2 nd grade	Attend to personal care with support Supervised Community Living Perform unskilled/semi-skilled work
Severe	20-25 to 35-40	3-4%		May learn to talk by school age		Limited benefits of preacademic training	Minimal self-care Simple tasks with supervision as adults Community living with family or group home
Profound	Below 20-25	1-2%	Limited	Improvements with training	Impairments in childhood	Limited	Simple, supervised tasks Requires structure and constant supervision

DSM-IV Diagnostic Criteria

(Continued)

- Mild Mental Retardation: IQ level 50-55 to approximately 70.
- Moderate Mental Retardation: IQ level 35-40 to approximately 50-55.
- Severe Mental Retardation: IQ level 20-25 to approximately 35-40.
- Profound Mental Retardation: IQ level below 20-25.
- Mental Retardation, Severity Unspecified: There is a strong presumption of mental retardation, but the person's intelligence is untestable by standard tests.

Implications of Diagnostic Criteria

- Deficits in both intellectual functioning and adaptive behavior must be present.
- MR is not diagnosed when an individual is adequately meeting the demands of his/her environment.
- Assessment must focus on descriptions of present behavior.
- Individually administered intelligence tests are needed.
- The diagnosis is tied to the individual's age level.
- MR diagnosis does not rule out the presence of other disorders.

Adaptive Behavior

- “Adaptive behavior is defined as the effectiveness or degree with which individuals meet the standards of personal independence and social responsibility expected for age and cultural group” (Grossman, 1983, p.1).
- Assessment of adaptive behavior stresses an individual’s typical performance.
 - Actual behavior, and not abilities or constructs believed to underlie behavior, is important.
- Adaptive behavior varies as a function of age.
 - Increasing demand for meeting the demands of the environment.
- Cultural expectations will also be important, especially when evaluating social functioning.

Etiology of MR

- Etiological factors may be primarily biological, primarily psychosocial, or some combination of both.
- For 30-40 percent of individuals with MR seen in clinical settings, no clear etiology can be determined.
- Major predisposing factors:
 - Heredity (5%).
 - Early alterations of embryonic development (30%).
 - Pregnancy and perinatal problems (10%).
 - General medical conditions acquired in childhood (5%).
 - Environmental influences and other mental disorders (15-20%).

Helpfulness of Determining Etiology

- Family members often have a desire to understand why a child has cognitive and adaptive deficits.
- If a genetic basis is identified, there may be a need for other family members to pursue genetic counseling.
- With a clear etiology, clinicians may be able to provide information on long-term course and type of supports individual will need.
- There may be a clear treatment implication for certain etiologies.
- Determining the etiological basis allows individuals to be placed in more homogeneous groupings.

Etiology of MR

- Familial Group: This group is composed of individuals who fall in the mild range of MR.
 - Primarily in the lower portion of the normal distribution.
 - Likely the result of normal polygenic variation, but can also result from pathological factors interfering with brain functioning or the combined effect of below average heredity and below-average environment.
 - Generally don’t come to the attention of the professional community as adults.

Etiology of MR

(Continued)

- Organic Group (25-50%): This group is primarily composed of individuals in the more severely retarded range.
 - May be associated with a genetic component linked to single gene effects, chromosomal abnormalities, or brain damage.
 - Demonstrate a severe lag in behavioral development.
 - Fail to reach normal motor and language developmental milestones.

Epidemiology

1-3% of the general population
 More severe cases noticed earlier
 Childhood peak time for identification
 More prevalent in males
 Mild cases more prevalent in low SES group
 For mild MR, early intervention and training can result in a child no longer meeting the criteria for diagnosis

Factors Associated with MR

- Fetal Alcohol Syndrome.
- Phenylketonuria (PKU).
- Chromosomal Anomalies.
 - Down's Syndrome.
 - Klinefelter's Syndrome (XXY).
 - Fragile X.
- Birth factors and growth factors (e.g., inadequate prenatal care).
- Social-environmental factors (e.g., psychosocial disadvantage).

Syndrome	Prev.	Cause	Cognitive	Physical	Behavioral
Down	1:1,000	Trisomy 21 (95% of cases)	Moderate to severe MR Delayed speech Expressive language not affected Deficits in verbal STM & auditory processing Intact VS abilities Dementia in adulthood	Upward slant and folds at corners of eyes Flat facial features Fissured tongue Broad hands and feet Poor muscle tone Heart defects	Relatively good social skills and mild manner Noncompliance, stubbornness, inattention, overactivity Depression developing in adult.
Fragile X	1:4,000 (M) 1:8,000 (F)	200+ DNA nucleotide repeats (CGG) → FMR-1 gene not expressed	Moderate MR Lang. skills plateau @ age 4 Cog. & adaptive behavior slow by age 5 and plateau by late childhood Weaknesses: VS, sequential processing, motor coord., math, and executive func. Strengths: Verbal STM & LTM	Boys: Velvetlike skin, double-jointed thumbs Adol.: Long faces, large ears, oversized testicles	Inattention, hyperactivity, stereotyped movements, anxiety, social avoidance, poor peer interactions Co-occurring autism assoc. w/ > dev. delay
Williams	1:20,000	Small deletions of several genes on Chromosome 7	Deficient depth perception Deficient VS STM & perf. IQ Inability to perceive gross diff. in spatial orientation and copy simple stick figures Strengths: facial expression recognition, verbal STM, verbal IQ, grammar, sophisticated vocab.	"Elflike" features (small lower jaw, prominent cheeks) Growth deficiency Often an "aged" appearance in late adult/early adulthood Cardiac & kidney problems Sound hypersensitivity	Anxiety, fears and phobias, inattention, hyperactivity, indiscriminate and overly friendly social interaction, poor social judgment
Frazer-Wills	1:15,000	Deletion of genes in certain area of paternal chrom. 15 (70% or both from mother (30%))	Borderline to moderate MR Deficient verbal IQ, high VS abilities (paternal deletion) Poor VS abilities, fewer facial characteristics, severe depress. & soc. impairments (maternal)	Hypotonia Flat face w/ almond-shaped eyes and prominent forehead Small hands and feet Short, underdeveloped gonads Obesity	Hyperphagia (excessive eating) & food hoarding Obsessions and compulsions Aggression, anxiety, impulsivity, & tantrums

Genetic Syndromes

Table 11-8

Down Syndrome

Most common single disorder
Caused by **Trisomy 21**
Higher risk with maternal age
Alzheimer's
Moderate to severe MR
Delayed speech, verbal short-term memory and auditory processing deficits

Fragile X

Most common inherited form
Fractured X chromosome
More common in boys - they have more severe forms
Long faces, prominent jaws, large ears (males)
Visual-spatial, sequential processing, motor coordination and executive function deficits
Social impairments

Fragile X Syndrome

X-linked recessive, carrier mother

Unaffected father
Carrier mother

Unaffected son
Carrier daughter
Affected son

CGG repeat: 5-45 (normal), 100-1000 (expanded)

U.S. National Library of Medicine

Genetic Syndromes

Williams Syndrome

Rare
Deletions on Chromosome 7
Cardiac and kidney problems, sound perception weaknesses
Mild to moderate MR
General knowledge & **visual spatial deficits**
Relative strengths in language
Elfin appearance

Williams Syndrome

Supernarrow bridge
Upturned nostrils
Blue eyes
Long upper lip
Small mouth
Small chin

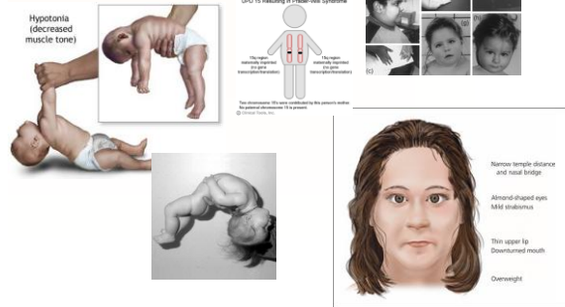
Normal nasal bridge
Low nasal bridge

Model Drawing
Williams Syndrome
Down Syndrome

Prader-Willi Syndrome

- 70% of cases result from paternal deletion chromosome 15
- IQ ranges from borderline to moderate impairment
- Hyperphagia and food hoarding
- Other compulsions, skin picking
- Strengths and weakness may vary depending on cause

Prader-Willi Syndrome



Family

- Adjustment is a lifelong process
- May be related to the severity of the impairment
- Stressors
 - Diagnostic
 - Medical
 - Financial/employment
 - Social
 - Marital
 - Parental distress
- Coping
 - Ethnic differences
 - Beliefs/parenting style
 - Skills
 - Support
 - Parental IQ
 - Siblings
 - Stressors
 - Rewards
- Access to resources is key

Treatment of MR

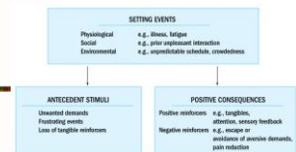
Prevention

- Prenatal care and diet
 - e.g., Phenylketonuria (PKU)
- Education on the impact of toxins
- Early intervention programs
- Educational services



Behavioral Intervention

- Discrete trial learning
- Naturalistic or incidental learning
- Operant conditioning to build adaptive skills
- Positive behavioral support
 - Functional assessment vs. functional analysis
- Research supports use of behavioral interventions for increasing prosocial and adaptive behaviors, and reducing maladaptive behaviors, e.g., self-injurious behavior (SIB)



Functional Analysis

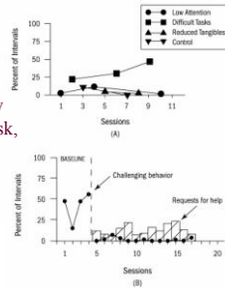
Matt, 5 y.o. boy w/ moderate MR

Observations during 4 conditions:

low teacher attention during a task, low access to a preferred object during a task, a more difficult task, and a control condition

Observations determined Matt engaged in SIB most during difficult tasks

Matt trained to ask for help with difficult tasks



Intervention

Pharmacological

- Frequently prescribed
- Usage increases when child exhibits behavioral problems
- Stimulant medications
- Antipsychotic medications
 - Overused
 - Lack of research

Psychotherapy

- Talk therapies not widely employed or researched
- Modifications necessary

Comorbid Diagnoses

- Between 20% and 35% of non-institutionalized individuals with MR have a comorbid psychiatric diagnosis or behavioral disorder.
- The rates of psychiatric and behavioral diagnoses are four to five times those of individuals without MR.
- Rates increase with age and cognitive impairment.

Relevant Legislation

Individuals with Disabilities Education Act (IDEA)

- In 1975, Congress passed Public Law 94-142 (the Education for All Handicapped Children Act).
- This law was updated in 1990, and its name was changed to IDEA.
- IDEA was reauthorized and amended in 1997, becoming IDEA '97.

IDEA '97 Principles of Interest

- Children with disabilities must receive a **Free Appropriate Public Education** that provides special education and related services.
- Children with disabilities should be placed in the **Least Restrictive Environment** to the maximum extent possible.
 - "Mainstreaming".
 - This decision is made by the IEP team after evaluation.
 - School systems must maintain a full continuum of alternative placements to meet the needs of children with disabilities.

IDEA '97 Principles of Interest

Continued

- Each child being considered for special education must receive a **full, individualized, and appropriate evaluation**.
 - *Schools are responsible* for finding, identifying, and evaluating these children.
 - Assessment measures must be *nondiscriminatory*.
 - *No single procedure* must be used as the sole criteria for making determinations of eligibility.
 - Standardized tests must be *validated* for the specific purpose in which they are used and must be administered by *trained* personnel.

IDEA '97 Principles of Interest

Continued

- Procedural safeguards are provided so that:
 - The rights of children with disabilities are protected.
 - Children and their parents are provided with the information they need to make informed decisions about the educational opportunities available.
 - Procedures are in place to resolve disagreements between the parents and the school district.

IDEA '97 Principles of Interest

Continued

- State- or district-wide group tests administered to children without disabilities should also be administered to children with disabilities.
 - Accommodations can be made.
 - Alternative assessments can be provided.

IDEA '97 Principles of Interest

Continued

- The Individualized Education Plan (IEP) and the IEP team.
 - It is determined that children are eligible for services when they have a disability and that disability affects their educational performance adversely.
 - The IEP spells out the needs of the child and how the agency will meet these needs.
 - The IEP must be reviewed at least once per year, and the child must be reevaluated at least once every three years.

IDEA Part C: Infants and Toddlers with Disabilities

- In 1986, P.L. 99-457 amended P.L. 94-142 and authorized early intervention programs for infants and toddlers with disabilities and *extended the rights of P.L. 94-142 to children with disabilities from ages 3 through 5 years*.
 - These rights can be extended to 2-year olds.
- In identified cases, an Individualized Family Service Plan is developed with parental consent.

Section 504 of the Rehabilitation Act of 1973

- It was designed to protect individuals with disabilities from discrimination in any setting receiving funds from the federal government.
- Reasonable accommodations must be made for children and adults with disabilities.

Americans with Disabilities Act (ADA)

- Also known as P.L. 101-336.
 - It provides protection from discrimination for individuals with disabilities in all settings, regardless of whether federal funding is involved.
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Potential Quiz Questions (Chapter 12a)

- The primary symptoms of autism will fall within three broad areas...name these.
 - Girls are affected with autism more often than boys. True or false?
 - List 3 categories of secondary symptoms of autism.
 - List 3 well-established facts regarding treatment of autism according to Schreibman (2000)
 - List 3 hypothesized causes of autism.
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