

# Flex Therapist CEUs

## Muscular Dystrophy

### 1. Mutations in which gene cause Duchenne and Becker muscular dystrophy?

- A. Dystrophin
  - B. Sarcoglycans
  - C. Calpain
  - D. Dysferlin
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### 2. Which of the following is not as common in those with DMD?

- A. ADHD
  - B. Dyslexia
  - C. Executive Function Disorders
  - D. Addison's Disease
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### 3. What is the primary cause of death in individuals with Duchenne Muscular Dystrophy?

- A. Infectious diseases
  - B. Neurological Decline
  - C. Respiratory or Cardiac Complications
  - D. Renal Failure
- 

### 4. At what age do most individuals with Duchenne Muscular Dystrophy (DMD) typically lose the ability to walk independently?

- A. Before Age 8
  - B. Between 8-12
  - C. Age 15
  - D. After Age 18
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### 5. Out of these options, how can Duchenne Muscular Dystrophy (DMD) be inherited?

- A. From Carrier Fathers To Their Sons, Who Receive The Mutated X Chromosome.
  - B. From Carrier Mothers To Their Daughters, Who Have A 50% Chance Of Inheriting The Carrier Status.
  - C. From Fathers With DMD To Their Sons, Who Inherit The Mutated X Chromosome.
  - D. Duchenne Muscular Dystrophy (DMD) is typically inherited in an autosomal dominant manner.
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**6. What is a characteristic clinical finding in children with Duchenne Muscular Dystrophy (DMD) when rising from the floor?**

- A. Normal Stand
  - B. Gower's Sign
  - C. Wheelchair Assist
  - D. Need Lifting
- 

**7. What impact does progressive muscle weakness have on respiratory function in Duchenne Muscular Dystrophy?**

- A. Improved Respiratory Efficiency
  - B. Causes No Changes
  - C. Respiratory Weakness, Insufficiency and Failure
  - D. Increases Lung Volume and Effective Cough
- 

**8. Which type of mutations are commonly associated with Becker Muscular Dystrophy in the DMD gene?**

- A. Missense Mutations, Nonsense Mutations, Small Insertions, Or Deletions
  - B. Large Chromosomal Rearrangements
  - C. Duplication Of Entire Genes
  - D. Inversions Of DNA Segments Within The Gene
- 

**9. What symptoms are characteristic of Becker Muscular Dystrophy?**

- A. Progressive Muscle Weakness and Gait Abnormalities
  - B. Sudden Onset of Muscle Rigidity and Joint Stiffness
  - C. Rapid Development of Neurological Deficits
  - D. Immediate Loss of Sensory Functions and Reflexes
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**10. Which proteins are commonly affected by mutations in LGMD?**

- A. Proteins that Control Blood Sugar
  - B. Proteins for Muscle Stability
  - C. Proteins Involved in Brain Function and Neural Connectivity
  - D. Proteins that Regular Kidney Function
- 

**11. At what age do symptoms typically begin to manifest in individuals with Limb-Girdle Muscular Dystrophy?**

- A. One to Five
- B. Twenty to Thirty
- C. Forty-Fifty

D. 60 and older

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**12. What is a common clinical manifestation of LGMD?**

- A. Rapid Cognitive Decline
  - B. Progressive Muscle Weakness Around the Shoulders
  - C. Severe Neurological Symptoms
  - D. Rapid Vision Loss
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**13. What symptom is the earliest sign of FSHD?**

- A. Difficulty With Fine Motor Skills
  - B. Weakness In The Muscles Around The Scapulae
  - C. Respiratory Insufficiency During Sleep
  - D. Cardiac Complications Such As Arrhythmias
- 

**14. How does FSHD primarily affect muscle weakness?**

- A. Symmetrically Affecting Both Sides Of The Body Equally
  - B. Randomly Affecting Various Muscles Without Specific Pattern
  - C. Asymmetrically, With One Side Of The Body More Affected Than The Other
  - D. Mainly Affecting Lower Limbs Before Progressing To Upper Limbs
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**15. Which complication is associated with FSHD?**

- A. Kidney Failure
  - B. Increased Risk Of Infectious Diseases
  - C. Respiratory Muscle Weakness Leading To Breathing Difficulties
  - D. Neurodegenerative Disorders Such As Alzheimer's Disease
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**16. How does myotonic dystrophy (DM) typically affect life expectancy in type 1 (DM1) compared to type 2 (DM2)?**

- A. Both DM1 And DM2 Typically Do Not Affect Lifespan C) D)
  - B. DM1 Is Typically More Severe Than DM2, With A Significantly Shortened Lifespan In Many Cases
  - C. DM2 Has A Greater Impact On Lifespan Than DM1
  - D. DM1 And DM2 Increase Lifespan Due To Genetic Factors
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**17. Which symptom Is not typically associated With Myotonic Dystrophy?**

- A. Progressive Muscle Weakness
- B. Prolonged Muscle Contractions Known As Myotonia
- C. Rapid Improvement In Muscle Strength Over Time

## D. Difficulty With Swallowing And Speech Problems

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**18. What is a common symptom observed in infants with congenital muscular dystrophy (CMD)?**

- A. High Muscle Tone
  - B. Hypotonia
  - C. Rapid Cognitive Development
  - D. Increased Resistance to Passive Movement
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**19. What are the primary complications that may contribute to morbidity and mortality in congenital muscular dystrophy (CMD)?**

- A. Neurological And Cognitive Impairments
  - B. Respiratory Or Cardiac Issues Stemming From Muscle Weakness
  - C. Enhanced Metabolic Function
  - D. Increased Physical Strength And Mobility
- 

**20. Which type of muscular dystrophy is estimated to affect approximately 1 in every 3,500 to 5,000 male births worldwide?**

- A. Becker MD
  - B. Duchenne MD
  - C. Myotonic MD
  - D. LGMD
- 

**21. What is the primary cause of muscle fiber damage in Muscular Dystrophy?**

- A. Dysfunctional Proteins Compromise The Muscle Cell Membrane, Causing Calcium Influx
  - B. Excessive Myostatin Production Inhibits Muscle Growth
  - C. Reduced Inflammatory Cytokine Production
  - D. Uncontrolled Growth Due To Satellite Cell Proliferation
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**22. What is the purpose of measuring creatine kinase (CK) levels in the diagnosis of muscular dystrophy (MD)?**

- A. To Assess The Degree Of Liver Damage
  - B. To Evaluate The Extent Of Muscle Damage
  - C. To Determine The Level Of Glucose In The Blood
  - D. To Check The Efficiency Of Kidney Function
- 

**23. Which diagnostic test is used to identify specific protein deficiencies in muscular dystrophy, such as dystrophin in Duchenne Muscular Dystrophy (DMD)?**

- A. Electromyographic Testing

- B. Genetic Testing
  - C. Immunohistochemistry
  - D. Pulmonary Function Tests
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**24. Which condition is characterized by the degeneration of motor neurons, leading to muscle weakness, atrophy, and spasticity, and affects both upper and lower motor neurons?**

- A. Myasthenia Gravis
  - B. Amyotrophic Lateral Sclerosis (ALS)
  - C. Lambert-Eaton Myasthenic Syndrome
  - D. Spinal Muscular Atrophy (SMA)
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**25. What is the characteristic sign used to detect muscle weakness in Duchenne Muscular Dystrophy during clinical evaluation?**

- A. Trendelenburg
  - B. Babinski
  - C. Gowers'
  - D. Hoffman's
- 

**26. During what stage of Duchenne Muscular Dystrophy do children typically begin to use mobility aids such as braces?**

- A. Initial Symptoms Stage
  - B. Late Childhood
  - C. Pre-teen Years
  - D. Teenage Years
- 

**27. At what stage of Becker Muscular Dystrophy might individuals begin using mobility aids like canes or braces to assist with walking and balance?**

- A. Early Stage
  - B. Intermediate Stage
  - C. Advanced Stage
  - D. Late Stage
- 

**28. What is a common cardiovascular complication associated with certain types of Muscular Dystrophy (MD) such as Duchenne and Becker?**

- A. Hypertension
  - B. Cardiomyopathy
  - C. Aortic Dissection
  - D. Deep Vein Thrombosis
-

**29. What is the primary goal of pharmacologic treatments in managing Duchenne Muscular Dystrophy (DMD)?**

- A. To Completely Cure The Disease
  - B. To Increase Muscle Enzyme Levels
  - C. To Slow Muscle Degeneration And Improve Quality Of Life
  - D. To Decrease Cardiac Output
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**30. Which diagnostic tool is routinely used to monitor heart function in patients with Duchenne Muscular Dystrophy?**

- A. Spirometry
  - B. Electromyography
  - C. Gastrointestinal Endoscopy
  - D. Echocardiogram
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**31. In managing Becker Muscular Dystrophy (BMD), what role do ACE inhibitors play?**

- A. They Help Manage Cardiac Issues
  - B. They Help Increase Muscle Mass
  - C. They Are Used To Treat Neuropathic Pain
  - D. They Directly Correct Genetic Mutations
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**32. What is the primary purpose of using non-invasive ventilation like BiPAP in Muscular Dystrophy care?**

- A. To Enhance Physical Mobility
  - B. To Support Breathing as Respiratory Muscles Weaken
  - C. To Increase Heart Rate
  - D. To Monitor Brain Activity
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**33. Which intervention is specifically used to manage scoliosis in patients with Muscular Dystrophy?**

- A. Surgical Intervention
  - B. Corticosteroid Therapy
  - C. Psychological Counseling
  - D. Genetic Therapy
- 

**34. What is the primary purpose of conducting a range of motion (ROM) assessment in patients with muscular dystrophy (MD)?**

- A. To Evaluate Physical Strength and Endurance
- B. To Determine Cardiovascular Fitness

- C. To Measure Cognitive Function
  - D. To Assess Joint Integrity and Muscular Function
- 

**35. Which test is commonly used to evaluate endurance and functional mobility in patients with muscular dystrophy?**

- A. Gait Analysis
  - B. 6MWT
  - C. MMT
  - D. Dynamometer Testing
- 

**36. During a neurological examination for muscular dystrophy, what gait pattern is characterized by a dropping of the hip on the side opposite to the lifted leg due to gluteal weakness?**

- A. Trendelenburg Gait
  - B. Toe Walking
  - C. Steppage Gait
  - D. Waddling Gait
- 

**37. What type of equipment might physical therapists evaluate for its appropriateness and effectiveness in later stages of muscular dystrophy?**

- A. Manual and Power Wheelchairs
  - B. Stationary Bike
  - C. Cane
  - D. Rollator Walker
- 

**38. What is emphasized during the early stages of physical therapy for patients with muscular dystrophy?**

- A. High-Intensity Interval Training
  - B. Immediate Transition To Advanced Aerobic Exercises
  - C. Delaying The Progression Of Muscle Atrophy And Maintaining Neuromuscular Function
  - D. Rapid Improvement In Functional Mobility
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**39. In the context of muscular dystrophy, why are isometric exercises recommended over high-load resistance activities?**

- A. They Are Less Likely To Exacerbate Muscle Breakdown
  - B. More Effective in Building Muscle Mass
  - C. They Can Completely Reverse Muscle Weakness
  - D. They Increase the Speed of Disease Progression
-

**40. What is a primary goal of physical therapy in the management of muscular dystrophy (MD)?**

- A. To Completely Reverse Muscle Weakness
  - B. To Prevent Complications Such As Contractures And Respiratory Distress
  - C. To Ensure Complete Recovery From The Disease
  - D. To Increase Muscle Mass
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