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Number

8

Subject

Neurodegenerative disorders / part 3

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LECTURE 8

Neurodegenerative disorders / part 3

Topics to be covered:

- 1. Spinocerebellar ataxia
- 2. Frederick ataxia
- 3. Ataxia telangiectasia
- 4. Amyotrophic lateral sclerosis
- 5. Metabolic and toxic effects on the CNS

Spinocerebellar ataxia

- Ataxia; Greek = lack of order
- = the loss of full control of bodily movements





Spinocerebellar ataxia

- Heterogeneous group of diseases
- Caused by trinucleotide repeat expansion mutations.

Examples of diseases caused by trinucleotide repeat mutations

		range	
Disease	repeats	normal	mutant
*Huntingon Disease	(CAG)n	11 - 34	36 - 120
*Spinocerebellar ataxia I	(CAG)n	6-40	40 - 82
'Machado-Joseph ataxia	(CAG)n	13 - 40	68-69
Friedrich's ataxia	(GAA)n	10 - 21	200 - 900
Myotonic dystrophy	(CTG)n	5 - 30	50 - 2000
Fragile X syndrome	(CGG)n	6 - 50	60-1000

* The (CAG)n is within an exon and produces polyglutamines in the protein product

• Group of diseases that differ in the mutation type, inheritance pattern, age of onset and clinical symptoms.

- Affects cerebellar cortex, spinal cord, other brain regions and peripheral nerves
- SO: symptoms are related to the area/s affected and vary from one person to another.

Clinical symptoms include

- Cerebellar ataxia
- sensory ataxia
- Spasticity
- Sensorimotor peripheral neuropathy

Cerebellar ataxia

• Cerebellar ataxia presents with symptoms of an inability to coordinate balance, gait, extremity and eye movements.

Symptoms of Ataxia

- Gait/Posture abnormalities Difficulty maintaining normal upright posture, balance, coordinated walking, and running. Unsteady gait, staggering, tripping, falling, unsteadiness on stairs or maintaining balance on moving platforms, such as escalators or boats.
- Fine motor incoordination Difficulty with handwriting, cutting food, opening jars, buttoning clothes, sewing, typing, playing an instrument or a sport.
- **Speech and swallowing difficulties** Speech: slurred, slow, indistinct, abnormal in rhythm. Swallowing: difficulty swallowing or choking (especially with liquids).
- Visual abnormalities Blurred vision or double vision. Reading: difficulty moving from word to word. Problems following moving objects or shifting gaze from one object to another.
- Increased fatigue Patients with ataxia due to cerebellar atrophy often experience unexpected fatigue when performing normal activities. The impaired regulation of coordinated movements may lead to increased fatigue because of the need to expend more effort to perform activities that are no longer coordinated. Patients with ataxia often report needing to "concentrate on" their movements.

Symptoms of ataxia

 Cognitive and Mood Problems - In addition to motor dysfunction, patients with cerebellar degeneration may have cognitive and emotional difficulties. The cerebellum plays a role in some forms of thinking. Patients with cerebellar atrophy may have impaired recall of newly learned information or difficulty with "executive functions" such as making plans and keeping thoughts in proper sequence. Personality and mood disorders, such as increased irritability, anxiety, and depression, are more common in persons with cerebellar degeneration than in control subjects.

Sensory ataxia

- It occurs if there is spinal cord or peripheral nerves' involvement
- Sensory ataxia is distinguished from cerebellar ataxia by the presence of near-normal coordination when the movement is visually observed by the patient, but marked worsening of coordination when the eyes are shut, indicating a positive Romberg's sign.
- Patients complain of loss of balance in the dark, typically when closing their eyes in the shower or removing clothing over the head.

Romberg's sign



Friedreich ataxia

- Is a type of spinocerebellar ataxia
- Autosomal recessive
- Manifests in the first decade of life
- Gait ataxia and hand clumsiness and dysarthria
- Associated with high incidence of cardiac disease and Diabetes mellitus

Genetic mutation in Friedreich ataxia

- Due to GAA repeat expansion.. Coding for frataxin; a protein that regulates cellular iron level
- The repeat expansion causes transcriptional silencing .. This results in decreased frataxin level

Friedreich ataxia

- Frataxin... important in iron levels especially in the mitochondria
- The repeat mutation causes transcription silencing resulting in decreased frataxin..
- This causes mitochondrial dysfunction and increased oxidative damage

- Frataxin consists of two α helices and seven β strands .
- It assists iron-sulfer protein synthesis in the electron transport chain to ultimately generate adenosine triphosphate (ATP)
- It also regulates iron transfer in the mitochondria in order provide a proper amount of reactive oxygen species (ROS) to maintain normal processes
- Without frataxin, 1. the energy in the mitochondria fails and 2. excess iron causes extra ROS to be created, leading to further cell damage.

Ataxia Telangiectasia

- Autosomal recessive disorder.
- Starts in early childhood
- Progressive and patients die in the second decade
- Symptoms: ataxia, dyskinesia, telangiectasias in conjunctiva and skin, immunodeficiency
- Complications: Infections are common and Lymphoid neoplasms may occur

genetics

- Mutation in ATM gene = ataxia mutated gene
- ATM encodes a kinase important for regulating cellular response to double stranded DNA breaks
- Also ATM protein facilitates apoptosis, maintains telomeres and has other actions.

morphology

- Loss of Purkinjie cells in the cerebellum
- Also there is degeneration of dorsal columns, spinocerebellar tracts, anterior horn cells and peripheral neuropathy.
- There is telangiectasia in the CNS

Amyotrophic lateral sclerosis

 Amyotrophic lateral sclerosis (ALS), also known as motor neuron disease (MND), is a specific disease that causes the death of neurons which control voluntary muscles. [Some also use the term "motor neuron disease" for a group of conditions of which ALS is the most common. ALS is characterized by stiff muscles, muscle twitching, and gradually worsening weakness due to muscles decreasing in size. This results in difficulty in speaking, swallowing, and eventually breathing.

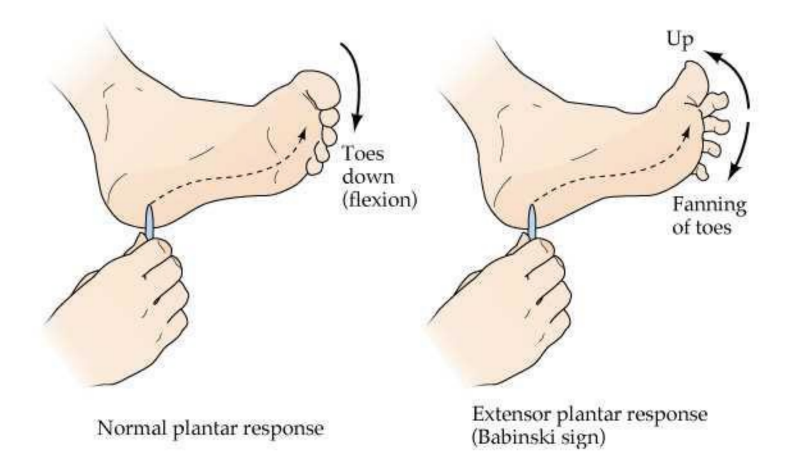
Amyotrophic lateral sclerosis (ALS)

• Results from death of **lower** motor neurons in the spinal cord and brain stem and of **upper** motor neurons in the motor cortex.

symptoms

- Loss of lower motor neurons results in denervation of muscle, muscle atrophy (amyotrophic) and weakness and fasciculation (a brief, spontaneous contraction affecting a small number of muscle fibers, often causing a flicker of movement under the skin).
- Loss of upper motor neurons results in paresis, hyperreflexia, , spasticity and Babinski sign,
- As a consequence of upper motor neuron loss there is degeneration of corticospinal tracts in lateral portion of spinal cord (lateral sclerosis)

Babinski sign





ALS

- Sensation NOT affected.
- Cognitive impairment occurs.
- Males slightly more than females
- Majority of cases are sporadic
- 5-10% inherited; autosomal dominant
- familial cases: earlier onset but disease progression similar



 Respiratory muscles affected later in the course of the disease resulting in recurrent pulmonary infections which is the usual cause of death

Genetic factors

- Several genetic mutations implicated in the familial cases
- 20% of cases: mutation in superoxide dismutase gene SOD 1 on chromosome 21
- This mutation causes abnormally folded SOD 1 protein
- The misfolded SOD 1 causes death of neurons

morphology

• Grossly: thin grey anterior roots of the spinal cord.

: motor cortex might become mildly atrophic especially in the severe cases

- Micro: reduction in the number of the anterior horn cells + reactive gliosis
- Similar micro features seen in motor cranial nerve nuclei except those supplying the extraocular muscle which are spared except in very longstanding survivors

Stephen Hawking

- Born 1942
- Diagnosed with ALS at the age of 21
- Expected to live for 2 years only
- He's 75 now
- Well known physicist and scientist.. Won Nobel prize
- Married twice and have 3 children.



Euthanasia = mercy killing

CARENOTKILING



Acquired metabolic and toxic disturbances

-Toxic and metabolic diseases are relatively common causes of neurologic illness

-the brain has a high metabolic demand which makes it vulnerable to nutritional diseases and changes in metabolic stat.

Acquired metabolic and toxic disturbances

- 1. nutritional diseases: thiamin and B12 deficiency
- 2. metabolic disorders: hypoglycemia, hyperglycemia, hepatic encephalopathy
- 3. toxins

Thiamine deficiency

- Thiamine def. causes Wernicke encephalopathy :
- 1. Confusion
- 2. Abnormal eye movements
- 3. Ataxia
- -Treatment: thiamine.. Things go back to normal
- -If thiamine def. untreated: irreversible memory disturbances: Korsakoff syndrome.

Wernicke-Korsakoff

Causes of thiamin def. :

- -Alcoholism
- Gastric disorders affecting thiamine absorption: tumors, chronic gastritis
- - Chronic vomiting

morphology

• Foci of hemorrhage and necrosis mainly in mammillary bodies

Vitamin B12 def

- Vit B 12 def causes subacute combined degeneration of the spinal cord
- Combined means that ascending and descending tracts affected
- Symptoms: Ataxia, lower extremity numbness and tingling
- Can progress to lower limb weakness
- Sometimes complete paraplegia can occur and this carries a worse prognosis.

Metabolic disorders

- 1. Hypoglycaemia: effect similar to global hypoxia...hippocampal neurones and Purkinje cells first affected.
- 2.Hyperglycaemia: ketoacidosis or hyperosmolar coma: confusion then coma

Note: hyperglycemia must be corrected gradually because rapid correction can produce severe cerebral edema.

• **3.Hepatic encephalopathy**: decreased consciousness and coma due to increased ammonia, inflammation and hyponatremia

Toxic disorders

- Alcohol
- Ionizing radiation
- Pesticides
- Carbon monoxide
- And many other toxic agents can affect the brain function

