INCLUSION BODY MYOSITIS and INCLUSION BODY MYOPATHY.

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Inclusion Body Myositis and Inclusion Body Myopathy are classified as types of muscular dystrophy.

Muscular dystrophy:
- Idiopathic inflammatory myositis (IIM) disorders:
  - Dermatomyositis (DM)
  - Polymyositis (PM)
  - Inclusion body myositis and inclusion body myopathies.
  - Idiopathic means the cause is unknown.

There are similarities between PM & IBM (and also big differences) but DM seems to be quite different.

People often confuse MD with a different disorder; MS: multiple sclerosis, a disorder affecting the nerves.
Sporadic inclusion body myositis (sIBM).

- Called a myositis to emphasize the inflammation of muscle that characterizes it. Sporadic means it just shows up here and there in people (it’s not inherited).
- This is the most common form, if you have been diagnosed with "IBM," this is likely what you have.
- sIBM is a progressive disorder of skeletal muscle cells: as more and more cells die off, the muscles shrink and become progressively weaker.
- sIBM is an age-related disease, as we get older, it gets more and more common, primarily after age 50.
- Symptoms emerge slowly, over months or years.
- The following diagram outlines the weakness seen.
Pattern of weakness seen in sIBM.

- Although there is a common pattern of weakness, it is important to note that to some degree, everyone is affected in slightly different ways, to different degrees & at different rates (Askanas & Engel, 2006).

sIBM symptoms.

- The quadriceps muscles in the front of the thighs are often affected first (and are often used for biopsy). This weakness is felt in climbing stairs, in getting up from chairs, etc. It leads to frequent falls.

- “Toe drop” (the leg muscles don’t raise the toe high enough in taking a step) commonly causes tripping.

- Usually, early and severe weakness of the muscles in the arm occurs and loss of wrist strength, finger dexterity and weak grip strength (making a fist) are common early symptoms or prominent symptoms.

- sIBM does not affect heart muscle or smooth muscle (the bowels).
More sIBM features.

- sIBM is not considered a fatal disorder (barring complications, sIBM will generally not kill you).

- sIBM is a relatively rare disorder, its incidence is about 15 per 1,000,000, overall, rising to over 50 per 1,000,000 in people over 50 (Needham, 2008).

- There is currently no effective treatment available and progression continues on to major or "total disability," usually within 10 to 15 years of symptom onset.
More sIBM features.

- sIBM is slightly more common in men.
- In from 40 to 85% of cases, progressive weakness in the muscles used in swallowing develops. If this occurs, the patient should be carefully evaluated (Oh et al, 2008).
- Problems swallowing (dysphagia) often cause choking and food to go into the lungs, resulting in a type of pneumonia which is often fatal in older people.
- sIBM may affect the muscles used in respiration leading to low volume and paradoxical breathing. Respiratory function should be checked in IBM cases.
Causes of sIBM.

- Currently, no cause has been identified. This makes developing a treatment more difficult.

- sIBM has two major aspects, one having to do with the immune system attacking and killing muscle cells (autoimmune aspect), the other, a deterioration of the proteins in muscle cells (degenerative aspect). The two aspects appear to occur in parallel in muscle cells.

- It is not clear which aspect comes first, if one causes the other, or if some other factor causes both aspects.
Causes of sIBM.

- One theory is that an undiscovered \textit{virus} triggers sIBM, setting in motion an ongoing immune response which attacks and kills the muscle cells.
- IBM was named for the observation that “inclusions” (inclusion bodies = clumps) and strands of abnormal proteins form in the muscle cells affected by sIBM.
- Based upon this, another theory is that \textit{abnormal proteins} form in the muscle cells causing sIBM and triggering the immune response that is seen.
- It is likely that a combination of features will be discovered as causing sIBM, involving both immune and degenerative aspects, environmental and genetic factors and their interaction with each other.
Genetics of sIBM.

- sIBM is **not** an inherited disorder – that is, the disorder is not passed on to the children of patients with sIBM.

- sIBM has been **linked** with a group of genes related to the immune system that are commonly seen in Caucasians from Northern European ancestry.

- This group of genes is called the **Major Histocompatibility Complex (MHC)** and is found on chromosome #6.

- **Genetic predisposition:** It is likely that some **combination** of genes interact with each other and with environmental variables to **increase the likelihood** of sIBM developing in a given individual.
Genetic predispositions in sIBM?

- There is a subset of these MHC genes called the \textit{8.1 ancestral haplotype}. A further subset of this group called the Human Leukocyte Antigen (\textit{HLA}) genes have been linked to patients with sIBM (and fIBM).

- Chemicals controlled by HLA genes play an important role in regulating immune responses to certain environmental factors.

- People with certain groups of HLA genes are \textit{predisposed to develop autoimmune disorders}.

- sIBM is associated with HLA-DR3 genes in about 70\% of patients (Badrising, 2004).
Diagnosis.

- Muscle disorders are generally difficult to diagnose.

- sIBM is often initially misdiagnosed as another muscle disorder called polymyositis.

- Usually a specialist (neurologist or rheumatologist) is required to diagnose the disorder.

- Most patients say that diagnosis took a long time and was a frustrating process.

- Many say that medications were suggested or tried (generally, no medication is indicated for IBM).

- These methods and tests are usually used in diagnosis:
Diagnosis of sIBM.

- **Clinical signs** are the symptoms the doctor can see when you are examined. The doctor looks for the clinical signs of sIBM; a *characteristic pattern* of muscle weakness.

- A **blood test** is done for creatine kinase (CK) (also known as "phosphocreatine kinase," or CPK), an enzyme in the blood showing muscle damage. CK is at most about 10X normal, although this may vary.

- **Electromyography (EMG):** Electrical studies of the muscle done with a computer (an electromyograph) will display abnormalities in these disorders.

- **Muscle Biopsy:** Surgical removal of a piece of muscle and its subsequent study in the laboratory.
Familial Inclusion Body Myositis (fIBM)

- fIBM occurs in two or more siblings in a family in the same generation (but it is not passed on to children).
- The symptoms and features of fIBM are very similar to those seen in the sporadic form of IB myositis.
- fIBM is also linked with similar genes as sIBM, raising the possibility that fIBM and sIBM share similar inherited predispositions.
- The familial occurrence of such a rare disorder highlights the importance of genetic predisposition in the causation of sIBM (Needham 2007).
Sporadic versus hereditary forms.

- Sporadic inclusion body myositis (sIBM) features inflammation in the muscle and abnormal proteins. It is not directly linked to a genetic mutation (not inherited).

- Hereditary inclusion body myopathy (hIBM):
  - "Myopathy" is used here as this type does not show muscle inflammation (myo: muscle, pathy: disease).
  - Hereditary is used to indicate that it is inherited.

- The rest of this presentation will focus on the hereditary forms which are exceedingly rare (in some forms, only a few hundred known cases in the world).

- See slide 26 for a general conclusion.
Sporadic versus hereditary forms.

- Today, several different hereditary inclusion body myopathies, are recognized. These different forms can display different symptoms but all share similar underlying structural features in common.

- Various hereditary types follow either autosomal recessive or autosomal dominant patterns of inheritance.

- As these disorders were discovered, a complex series of names were assigned to them before it was recognized that some shared commonalities or were the same.
Various names for the recessive form.

- As the autosomal recessive form was initially described, several different names were given to it:
  - Hereditary Inclusion Body Myopathy (hIBM2).
  - Inclusion Body Myopathy 2 (IBM2);
  - Autosomal recessive Inclusion Body Myopathy (AR-hIBM);
  - Quadriceps-sparing Inclusion Body Myopathy (QSM);
  - Nonaka distal myopathy with rimmed vacuoles;
  - Distal myopathy with rimmed vacuoles (DMRV).
- This is the most common type of inherited IBM.
Various autosomal dominant forms.

- **Autosomal dominant forms:**
  - **IBM 3:** was called “AD myopathy with congenital joint contractures, ophthalmoplegia and rimmed vacuoles” (linked to mutations in a gene on chromosome 17).
  - **AD-IBM:** Autosomal dominant inclusion body myopathy.
  - **IBMPFD:** IBM associated with Paget’s disease and frontotemporal dementia (linked to a gene on chromosome 9, located at 9p13-p12).
Features of hIBM 2.

- Argov and Yarom (1984) first described the disorder in Jews of Persian (Iranian) origin.
- The onset of this disorder usually occurs after the age of 20 but before the middle of the fourth decade. Proximal and distal muscle weakness and wasting of the upper and lower limbs are progressive and result in severe incapacitation within 10 to 20 years.
- Sparing of the quadriceps muscles, even in advanced stages of the disorder, is a feature unique to this inherited form of inclusion body myopathy.
- hIBM 2 shares several features with sIBM, including “muscle holes” and protein inclusions (but with NO inflammation or quadriceps muscle involvement).
Mapping to a specific gene location.

- Eisenberg et al. (1999, 2001) traced the disorder to a gene on an autosome: chromosome #9 (9p12-p11).
- All patients of Middle Eastern descent were found to share a single homozygous missense mutation in the GNE gene. This gene produces GNE protein. (GNE stands for: UDP-N-acetylglucosamine 2-epimerase/N-acetylmannosamine kinase).
- GNE is an enzyme involved in the production of sialic acid: a chemical involved in the management of glycoproteins and glycolipids in the cell.
- To date, more than 40 different mutations in the GNE gene have been reported that lead to hIBM 2.
Who is affected?

- hIBM 2 is a very rare disorder, affecting about 500 people worldwide. The disorder predominantly affects Iranian Jews, who have a 5 to 10 percent chance of carrying the hIBM 2 (GNE) gene mutation.

- Argov et al. (2003) concluded that this mutation first appeared about 1,300 years ago and today, is not limited to those of Jewish descent.

- hIBM 2 has also been seen in some Japanese families (first described by Dr. Nonaka in 1981) as well as in culturally diverse families and in various parts of the world.
Autosomal recessive disorders.

- A **mutation** of the GNE gene on one of the # 9 pair of autosomal chromosomes from each parent is required to cause the disorder. People with only one abnormal gene in the pair will be carriers, but since the gene is *recessive* they do not exhibit the disorder.

- To develop symptoms, recessive traits require that both chromosomes in the pair (one received from mom and the other from dad) carry the mutation.

- A child inheriting 1 mutated gene will be a carrier, a child receiving 2 mutated genes will have the disorder.

- The following diagram outlines the statistical situation.
If both parents are carriers.

- **Expected outcomes:**
  - 1 child with 2 normal chromosomes (normal: 25%)
  - 2 children with 1 normal and 1 abnormal chromosome (carriers, no disorder: 50%)
  - 1 child with 2 abnormal chromosomes (shows the disorder: 25%).


- These are **average odds**, you could be lucky and have 4 normal children in a row, or you could be unlucky and have 4 children with the disorder.
Other scenarios.

- One parent is a carrier, the other normal:
  - 50% chance a child will be normal.
  - 50% chance a child will be a carrier.

- Both parents are afflicted with the disorder:
  - All children will be afflicted with the disorder

- One parent afflicted with the disorder, one carrier:
  - 50% afflicted with the disorder, 50% carriers

- One parent afflicted with the disorder, one normal:
  - All children will be carriers.
Mutations plus other factors?

- GNE mutations have been found in some people who never show symptoms, showing that other factors must play a role in the expression of the disorder.

- These factors also seem to be important in the clinical expression of hIBM 2 in all patients; although the gene defect is present from conception, symptoms may not appear until the 30s or 40s.

- Researchers are struggling to discover what these factors may be (Needham, 2007).

- Many genetic conditions show interactions with environmental factors that may speed or slow their expression (the development of symptoms).

Conclusion.

- Inclusion body myositis and myopathy are very complex disorders of the skeletal muscle.
  - It is not clear what causes the sporadic form.
  - The mutations behind the inherited forms are being described but possible interacting environmental factors remain to be discovered.

- These can be frustrating disorders to diagnose and all previous attempted treatments have failed.

- Although the slow loss of mobility and arm function is very frustrating, patients must be encouraged to adapt and “make the best of things.”
Some abbreviations used.

- Sporadic inclusion body myositis (sIBM).
- Familial Inclusion Body Myositis (fIBM)
- Hereditary inclusion body myopathy (hIBM).
- Polymyositis (PM)
- Muscular dystrophy (MD)
- myo = muscle; pathy = disease; itis = inflammation.
References.


