

UC Irvine

UC Irvine Previously Published Works

Title

Clinical and molecular studies in a unique family with autosomal dominant limb-girdle muscular dystrophy and Paget disease of bone

Permalink

<https://escholarship.org/uc/item/9pr3c16h>

Journal

Genetics in Medicine, 2(4)

ISSN

1098-3600

Authors

Kimonis, Virginia E
Kovach, Margaret J
Waggoner, Brook
[et al.](#)

Publication Date

2000-07-01

DOI

10.1097/00125817-200007000-00006

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at <https://creativecommons.org/licenses/by/4.0/>

Peer reviewed



Published in final edited form as:

Genet Med. 2000 ; 2(4): 232–241. doi:10.1097/00125817-200007000-00006.

Clinical and molecular studies in a unique family with autosomal dominant limb-girdle muscular dystrophy and Paget disease of bone

Virginia E. Kimonis, MD, MRCP¹, Margaret J. Kovach, PhD¹, Brook Waggoner, BS¹, Suzanne Leal, PhD², Ambar Salam, BS², Lisa Rimer, MS³, Katherine Davis¹, Romesh Khardori, MD⁴, and David Gelber, MD⁵

¹Division of Genetics and Metabolism. Department of Pediatrics, Southern Illinois University-School of Medicine.P.O. Box 19658, Springfield, IL 62791-9658,

²Laboratory of Statistical Genetics, Rockefeller University, New York, New York,

³Children's Hospital, Dayton, Ohio; Departments of Internal Medicine Southern Illinois University-School of Medicine, Springfield, Illinois

⁴Internal Medicine and Southern Illinois University-School of Medicine, Springfield, Illinois

⁵Neurology, Southern Illinois University-School of Medicine, Springfield, Illinois

Abstract

Purpose: To characterize the clinical features and perform linkage analysis of candidate loci in a large Illinois family with autosomal dominant limb-girdle muscular dystrophy (LGMD) and Paget disease of bone (PDB).

Methods: The family includes 11 affected individuals (8 M, 3 F). Clinical, biochemical and radiologic evaluations were performed to delineate clinical features of the disorder. Linkage analysis with polymorphic markers was performed for previously identified LGMD, PDB and cardiomyopathy loci.

Results: Onset of PDB is early, at a mean age of 35 y, with classic distribution involving the spine, pelvis, and skull. Muscle weakness and atrophy is progressive with mildly elevated to normal creatine phosphokinase levels. Muscle biopsy in the oldest male revealed vacuolated fibers, however, in others revealed nonspecific myopathy. Affected individuals die from progressive muscle weakness, and respiratory and cardiac failure in their 40s-60s. Linkage analysis excluded autosomal dominant and recessive LGMD, PDB, and cardiomyopathy loci.

Conclusion: Autosomal dominant LGMD associated with PDB is an unusual disorder. Linkage analysis indicates a unique locus in this family.

Keywords

Limb-girdle muscular dystrophy; Paget disease of bone; autosomal dominant; vacuolar myopathy

We describe a family with autosomal dominant limb-girdle muscular dystrophy (LGMD) with the associated finding of early onset Paget disease of bone (PDB) and cardiomyopathy in a few individuals. Three similar associations of neuromuscular disease and Paget disease

of bone have been previously reported. Familial PDB in association with dystrophia myotonica was reported in one family by Caughey et al.¹ (MIM 16,0900²). McBride³ found an association of PDB and a limb girdle type of muscular dystrophy in four of six siblings and Tucker et al.⁴ identified a family with autosomal dominant amyotrophic lateral sclerosis (ALS, MIM 10,5400²) and PDB.

Characterized by progressive proximal muscle dystrophy and weakness particularly of the shoulder and pelvic girdle with relative sparing of the other muscle groups, limb-girdle muscular dystrophies are a clinically and genetically heterogeneous group of neuromuscular diseases. At least 4 autosomal dominant and 8 autosomal recessive forms of LGMD have been reported. LGMDIA, located on chromosome 5q22-q31, is caused by a mutation in myotilin, a sarcomeric structural protein.⁵⁻⁷ LGMDIC, caused by mutations in the muscle specific protein, caveolin-3, is located on chromosome 3p25.⁸⁻⁹ The genes for LGMDIB (1q11-q21),¹⁰ LGMDID (7q),¹¹ and Edstrom myopathy, a related autosomal dominant myopathy that maps to 2q24-q31,¹² are yet to be identified.

The genes for eight of the recessive forms of LGMD have been identified, the majority of which encode for components of the dystrophin-sarcoglycan complex (Table 1). None of these loci have been implicated in autosomal dominant neuromuscular or bone disease.

Autosomal dominant limb-girdle muscular dystrophy is associated with cardiomyopathy in LGMDIB and also in another family with dilated cardiomyopathy with conduction defect (CDCD3) located on chromosome 6q23.¹³ The loci for cardiomyopathy were therefore considered candidates for the disease in our family.

Paget disease of bone is characterized by an increase in bone remodeling and hypertrophy leading to abnormal bone structure, bone pain, and deformity.¹⁴ McKusick¹⁵ reviewed 37 families with familial Paget disease of bone (PDB). Only one family exhibited male-to-male transmission suggesting the rarity of autosomal dominant families. Subsequently, two genetic loci have been linked to autosomal dominant PDB. The first to be identified, PDB1, was obtained with the HLA locus on chromosome 6p.¹⁶ PDB2 was assigned to chromosome 18p21-22¹⁷ in a large Northern Irish family with PDB, and recently mutations in the gene TNFRSF11A were found in association.¹⁸ TNFRSF11A encodes for a receptor activator of nuclear factor- κ B (RANK), which is essential for osteoclast formation. In addition to PDB1 and PDB2, there is evidence for further genetic heterogeneity in Paget disease of bone.¹⁹

Linkage analysis with genetic markers for LGMD, cardiomyopathy and PDB excluded all dominant and recessive forms of LGMD as well as 6 loci for dominant dilated cardiomyopathy and 2 loci reported for dominant Paget disease of the bone (Table 2). These studies indicate that a unique locus exists for autosomal dominant LGMD in association with Paget disease of bone.

MATERIALS AND METHODS

Clinical studies

Clinical studies were approved by the Springfield Committee for Research Involving Human Subjects, and consent was obtained from each subject prior to participation. Several members of this six-generation family (Fig. 1) were examined for the presentation of LGMD and PDB through clinical, bio-chemical, and radiologic studies. Diagnosis for LGMD was based on physical examination for proximal muscle weakness and laboratory test results of creatine phosphokinase (CPK) levels in the blood. Skeletal surveys including the skull, spine, hips, long bones, and hands and feet were performed to evaluate Paget disease in family members. X-rays were read by a radiologist without prior knowledge of the individual's affected status. Pyridinoline/deoxypyridinoline collagen cross-linking metabolites in urine samples and quantitation of bone-specific alkaline phosphatase and osteocalcin levels in blood were performed by Quest Diagnostics (San Juan Capistrano, CA) and Emory University (Atlanta, GA), respectively.

Cytogenetic analysis

High-resolution karyotype analysis at 800–850 band resolution performed on case 9 was normal excluding a deletion or other rearrangements as the cause of this unusual disorder (Genzyme Genetics, Santa Fe, NM, and Washington University, St. Louis, MO).

DNA marker analysis

Peripheral venous blood was obtained by standard veni-puncture. Genomic DNA was extracted by means of the PUREGENE DNA Isolation Kit (Gentra Systems; Minneapolis, MN). In cases where blood samples could not be obtained, buccal scraping were collected and processed using standard methods. Genotyping was performed to exclude/include linkage to known chromosome locations of the autosomal dominant and recessive forms of LGMD, dominant dilated cardiomyopathy, and PDB using microsatellite markers (Research Genetics, Inc., Huntsville, AL).

Linkage analysis

Linkage analysis was carried out on 23 members of the family (9 affected, 13 unaffected, and 1 unaffected spouse) with molecular markers for established loci of LGMD, Edstrom myopathy, PDB and cardiomyopathy (Table 2). The amyotrophic lateral sclerosis locus on 21q was also excluded in view of its association with PDB.⁴ The power to detect linkage with the available pedigree was evaluated by simulating a marker locus with 5 alleles of equal frequency (heterozygosity = 0.8) completely linked to the disease locus ($\theta = 0.0$) utilizing the SLINK program.^{20–21} The analysis was carried out under an autosomal dominant model. All individuals included in the analysis were above 20 years of age. Three age dependent liability classes were used to generate the pedigrees and to analyze them. It was assumed that there were no phenocopies and ratio of being wild type to being a disease allele carrier is as follows: 20–29 years old (1.11:1); 30–34 years old; (5:1), and 35 years and over (50:1). Two-point linkage analysis was carried out using the MLINK program of the FASTLINK computer package.²² The marker-allele frequencies were estimated from the

data by means of both observed and reconstructed genotypes of founders within the pedigrees. Linkage was considered established to a locus if a positive LOD score of 3 was obtained and was considered excluded if a LOD score of -2 was obtained.

RESULTS

Clinical studies

Several members of this 6-generation family have been examined. The family includes several affected individuals; however, many are deceased prematurely from cardiomyopathy or other complications related to LGMD/PDB. The disorder in this family begins insidiously in the late 20s–30s in many individuals with back and hip pain, in addition to weakness and atrophy of the proximal arm and leg muscles. Nerve conduction studies were normal in most individuals. EMG shows muscle denervation. Bone X-rays of the spine, pelvis, shoulder, and skull indicate coarse trabeculation, cortical thickening, and spotty sclerosis consistent with the diagnosis of PDB. The disorder progresses to wheelchair confinement, quadriparesis, bedfastness, respiratory failure, cardiac failure, and death in the 40s–60s. Two individuals have had limited benefit from intravenous pamidronate several years ago. Two individuals received calcitonin with limited benefit, and one female is receiving alendronate with clinical improvement.

The following individuals were identified as affected from previous medical evaluations and/or clinical examination performed. Clinical data of the these affected individuals are summarized in Table 3, and laboratory data including alkaline phosphatase, osteocalcin and CPK levels are summarized in Table 4.

Case 1, the proband, a 48 y old female, developed back pain at the age of 34 y. X-rays of the spine and hip, and bone scan, revealed increased uptake in the L2 lumbar spine and the left ileum, suspicious for malignancy. A bone biopsy, however, revealed Paget disease and no evidence of malignancy. At age 34 y, she also noted proximal muscle weakness causing difficulty raising her arms. She had previously noticed tiredness and pain in her legs. At age 37 y, she had difficulty walking up the stairs, which she climbed by pulling herself up with her arms. She was diagnosed with LGMD at the age of 44 y. On physical examination, she had weakness of the proximal muscles of the arms and legs. There was a lumbar lordosis and her gait was wide based. The reflexes, sensation, and muscle tone in her limbs was normal. Bone specific alkaline phosphatase levels were markedly elevated.

Case 2 is a 47 y old male who admitted to a history of weakness of arms and legs and muscle pain for the previous 5 years. An examination at age 47 y revealed minimal weakness of the proximal muscles. A subsequent muscle biopsy revealed variation in fiber size with fiber splitting, fiber nesting, and atrophic and hypertrophic fibers. The muscle was described as having mild myopathic changes. In addition, there was focal endomysial fibrosis, but no necrosis, regeneration, or degeneration of muscle fibers, features consistent with muscular dystrophy. An EMG at 45 y showed moderate fibrillation potentials, increased muscle fiber irritability, and small amplitude motor potentials, indicating a primary muscle disorder. Nerve conduction velocities (NCV) of the left upper and lower extremities were normal. All laboratory tests were within normal range and x-rays did not

show evidence of Paget disease. He had a myocardial infarct at age 42 y and subsequently was diagnosed with cardiomyopathy.

Case 3, a 49 y old female, has a 10 y history of back pain and occasional pain of the knees and calves. She complained of “Charlie horses” in the bottom of her feet, shoulders, and neck. Radiologic evaluation revealed sclerosis and lucencies seen in the right parietal bone with cortical thickening, which was also seen in the calvarium, mandible, mastoids, and sphenoid. Lumbar spine revealed Paget disease of L2 and L3, with coarse trabeculation of the T3 and T9 vertebral bodies. A bone scan revealed abnormal intense uptake in the calvarium, manubrium, left scapula, lower dorsal vertebral body, right pubic bone, and left distal femur consistent with Paget disease. At the age of 44 y, she received intravenous bisphosphonate for her PDB, which gave her temporary relief from the pain. She has also had an 8 y history of constant aching of the pectoral, shoulder, and thigh muscles. She is unable to raise her arms up and has difficulty climbing stairs. Physical evaluation reveals the typical weakness of the iliopsoas, quadriceps, and mild atrophy of the forearms bilaterally. She has a widebased gait. At 48 y, a muscle biopsy containing atrophic fibers revealed muscle with no pathologic diagnosis. An EMG showed small amplitude motor units in the biceps and vastus lateralis consistent with a myopathic process. She is currently on disability for her muscular dystrophy.

Case 4 is a 57 y old male currently on disability from PDB and LGMD. He is unable to raise his arms and walks slowly with the assistance of a cane. He has moderately severe pain in his thoracic and lumbar spine. He is a smoker and has respiratory difficulties. Skeletal X-rays and bone scan reveals active PDB in the skull, shoulder, hip, and lumbar and vertebral spine. An echocardiogram was normal. Physical examination reveals tenderness of the spine and hip in addition to features of LGMD. Treatment includes bisphosphonates and calcitonin. An EMG showed small amplitude motor potentials in the leg muscles suggestive of a primary muscle disorder and focal abnormalities of upper extremity nerves. The NCV were normal.

Case 5, the 33-year-old son of case 4, has had back and right hip pain for approximately 18 months. X-ray studies suggested PDB in the hip, skull, and spine. Examination revealed an individual who walked slowly with a limp because of back pain. He had moderately severe tenderness of the back over the thoracic and lumbar spine. Bone specific alkaline phosphatase and osteocalcin levels were markedly elevated. He is being treated with analgesics and calcitonin. He did not have evidence of muscular dystrophy clinically, and his CPK was normal.

Case 6 was evaluated after his death at age 50 y. Diagnosis of LGMD, PDB, and cardiomyopathy was at 38 y. He had a prior history of diabetes mellitus, stroke, obesity, and coronary artery disease for which he had a balloon angioplasty. Medical records indicate that his alkaline phosphatase levels were consistently elevated. Physical examination revealed left-sided hemiparesis from the stroke, shoulder and pelvic muscle atrophy. He had not received any prior treatment for Paget disease of bone. He had been bedridden for several months prior to death from cardiac failure.

Case 7, a 45 y female, was recently diagnosed with PDB because of back and shoulder pain. MRI of the back showed Paget disease in the L5 vertebral body. Total alkaline phosphatase levels were elevated. On examination, she had proximal muscle weakness of the arms and hips and wasting of the small muscles of the hands consistent with LGMD. She is currently receiving treatment with Fosamax (alendronate sodium), cyclobenzaprine, and analgesics.

Case 8, age 46 y, was diagnosed with LGMD at age 40 y when he noticed muscle weakness and atrophy of bilateral upper arms. He has weakness in his legs and upper arms and has difficulty getting up from a squatting position. Muscle biopsy reveals atrophic fibers and end stage myopathy. EMG shows mild spontaneous activity and small amplitude motor potentials consistent with a primary muscle disorder. Nerve conduction studies were normal. CPK levels were highly elevated and alkaline phosphatase levels were normal. X-rays did not reveal definite evidence of PDB. His father had a history of Paget disease, and died at 73 y due to respiratory compromise related to LGMD.

Case 9 first noticed muscular weakness at age 50 y. The disease progressed to cause severe weakness of the muscles, making it difficult for him to get out of a chair. He was wheelchair bound for the last year and required supplemental O₂. Additionally, he had peripheral neuropathy causing loss of sensation in his extremities. Deep tendon reflexes were absent in the leg. He reported no disability from PDB, which was discovered at 37 y, after a spine x-ray was performed to investigate a fall from a ladder. A bone scan revealed intense activity in the right clavicle, L3 vertebra, right hemipelvis, sacrum, and left acetabulum. Laboratory results showed his total alkaline phosphatase was greatly elevated. His CPK was borderline elevated. His echocardiogram was grossly normal. A series of muscle biopsies in case 9 shows a progression of the disorder. At 40 y, there were minimal and nonspecific changes with occasional atrophic fibers favoring a “lower motor neuron disease” rather than muscular dystrophy. Aside from these minor variations, the biopsy was reported as within normal limits. At 55 y, degenerating and regenerating fibers, along with fiber size variation was noted. Small vacuolated fibers were also seen scattered throughout the muscle (Fig. 2). A final biopsy at 63 y revealed severe, chronic myopathy with vacuolar changes, including rimmed vacuoles, and atrophic fibers in large groups of whole fascicles. NCV studies suggested mild carpal tunnel syndrome. EMG showed mild myopathic changes. He died at age 65 y from respiratory failure.

Cases 10 and 11 were not examined. Medical information was obtained from medical records.

Case 10 died at the age of 59 y after being bedridden for the preceding three years. Symptoms of the LGMD began with progressive weakness of both his upper and lower extremities at 51 y. He had difficulty walking up the stairs and experienced aching and weakness of his trunk muscles and lower extremities, which affected the thighs more than the lower legs. He demonstrated winging of the scapulae bilaterally. Alkaline phosphatase levels were elevated, while CPK levels were within the normal range. A muscle biopsy was performed and adipose tissue was seen between some of the muscle bundles with no specific findings of intrinsic muscle disease.

Case 11 died at age 61 y of congestive heart failure and PDB. He had a history of diabetes mellitus, pulmonary obstructive disease, chronic bronchitis and bronchial asthma, and Paget disease involving the lumbar spine, pelvis, and proximal portions of both femurs. His alkaline phosphatase levels were greatly elevated. He demonstrated winging of the scapulae and weakness in the arms. He also experienced intermittent hyper-tension and mild coronary insufficiency. His EKG showed slight ST depression in V5 and V6 and his alkaline phosphatase levels were greatly elevated.

Laboratory evaluation

Paget disease of bone—All individuals showing radiologic evidence of PDB had corresponding elevated alkaline phosphatase levels (Table 4). The mean value of total alkaline phosphatase in affected individuals was 747.7 IU/liter (normal range 30–130 U/liter) with a range of 88–1724 IU/liter. Cases 2 and 8 were not clinically diagnosed with PDB and had normal alkaline phosphatase levels. Osteocalcin was elevated only in the youngest individual, 33 y, to a value of 54.6 ng/mL (normal range 8–52 ng/mL). All other individuals had normal levels with a mean of 20.0 ng/mL. Additionally, urine pyridinium cross link studies were obtained in all individuals, but results were not helpful in distinguishing affected from unaffected individuals (data not shown).

Limb-girdle muscular dystrophy—All individuals diagnosed with LGMD were characterized by severe muscle weakness particularly in the arms, legs, shoulder, and pelvic girdle. Laboratory evaluations of the CPK were not found helpful in screening for LGMD in the majority of individuals. Only 2/11 individuals had elevations of CPK (normal, 60–222 IU/liter).

Muscle biopsy—Six out of eleven individuals had muscle biopsies. Histology in 5 individuals showed nonspecific myopathic changes (Table 4). A series of muscle biopsies in case 9 shows a progression of the disorder. At age 40 y, the muscle was almost normal. At 55 y, generating and regenerating fibers were present with fiber size variation and small vacuolated fibers (Fig. 2). At age 63 y, he had severe, chronic myopathy with vacuolar changes, rimmed vacuoles, and atrophic fibers in large groups of whole fascicles.

Electromyographic (EMG) studies—EMG was performed on 5 affected individuals. Overall, the studies revealed spontaneous fibrillation, small polyphasic motor units and mild membrane irritability consistent with a myopathic process as seen in LGMD. One individual (case 9) had additional neuropathic changes clinically and on EMG.

Linkage analysis—Linkage analysis was carried out on 23 members of the family (9 affected, 13 unaffected, 1 spouse) with molecular markers for established loci of autosomal dominant LGMD and auto-somal dominant PDB. Linkage to the autosomal recessive forms of LGMD as well as 6 loci for dominant cardiomyopathy was also examined for this pedigree. Sufficiently negative LOD scores were obtained for all markers evaluated (Table 2), thus excluding all the candidate genes from linkage.

DISCUSSION

Limb-girdle muscular dystrophy associated with Paget disease is an unusual disorder. Paget disease of bone affects approximately 2–3% of the population > 50 y, and autosomal dominant LGMD is extremely rare; therefore, the co-occurrence of both disorders by chance in 8/11 individuals in our family is unlikely.²³

Three reports of families with Paget disease of bone and neuromuscular disease have been described previously (Table 5). McBride³ reported a family of 6 siblings: three brothers and one sister were diagnosed with widespread Paget disease of bone and muscular dystrophy, one sister had localized Paget disease in one lumbar vertebra, and one brother was unaffected. Possible mild phenotype in the mother was reported so the inheritance pattern was difficult to deduce.

Caughey et al.¹ reported three brothers with dystrophia myotonica, two of whom had PDB. Individuals had typical features of dystrophia myotonica including stiff muscles, inability to relax the grip, myotonia of the tongue and hand muscles, hypertrophy of the biceps and forearms, muscle size disproportionate muscle strength, and weakness of the abdominal and sternomastoid muscles. One of the brothers exhibiting Paget of the vertebrae and pelvis had a neoplastic process compressing the spinal cord. The parents were unaffected, possibly suggesting autosomal recessive inheritance.

Tucker et al.⁴ described a family with lower motor neuron degeneration and “Paget-like bone disease.” In the nine individuals pelvic and shoulder muscle weakness and atrophy started at 35–40 y, with death occurring before the age of 60 y. Of the nine individuals, seven were known to exhibit signs of Paget disease. The reported features were very similar to our family; however, there was no instance of male-to-male transmission in the family possibly suggesting an X-linked mode of inheritance. An isolated case report of 53 y old male with PDB and lower motor neuron was reported by Varelas and Kapati²⁴ who estimated that the combined incidence of PDB and ALS ranges between 0.012 and 0.108/100,000.

Cardiac failure and cardiomyopathy occurs in the latter stages of the disease in some individuals in our family. Two of these individuals are heavy smokers and have coronary artery disease; therefore, the significance of cardiomyopathy is uncertain in them. Messina et al.¹³ reported a four-generation family with conduction defect (CDCD3) and muscle weakness, which they mapped to 6q23. Van der Kooi et al.¹⁰ identified a family demonstrating LGMDIB with related cardiac abnormalities in 62.5% of the affected individuals and sudden cardiac death at approximately 50 y. Paget disease of bone has been known to lead to high output congestive heart failure due to the increased vascularity of abnormal bone.²⁵ Linkage analysis was undertaken for candidate loci for cardiomyopathy; however, they were excluded.

Pharmacological treatment of PDB has included calcitonin, a peptide hormone that inhibits osteoclast activity. Bisphosphonates, are more successful in the management of Paget disease, induce marked and prolonged inhibition of bone resorption through a direct effect on mature osteoclasts. Disruption of cellular metabolism and induction of osteoclast apoptosis are the most likely mechanisms.²⁶ Etidronate was the first bisphosphonate to be

used in the management of Paget disease.²⁷ Introduction of additional bisphosphonates include pamidronate, tiludronate, alendronate, and risedronate, respectively, with each increasing in potency and efficacy.

Therapy in the muscular dystrophies has been less successful and has included coenzyme Q,²⁸ steroids,²⁹ myoblast transfer³⁰ and gene therapy clinical trials in the autosomal recessive types of LGMD.^{31–32}

In conclusion, the results from our linkage analysis confirm that autosomal dominant LGMD with associated PDB is genetically distinct. Discovery of new genes involved in dominant LGMD should provide insight into the pathogenesis of muscle disease as they have done in the autosomal recessive types.³³ It is possible that a biochemical overlap exists between the biological processes of bone and muscle maintenance and regeneration that could potentially be influenced by a single gene mutation. Elucidation of the gene responsible for LGMD associated with PDB may help unravel the biochemical nature of this unique disorder, eventually resulting in the development of novel treatment protocols including gene therapy.

Acknowledgments

This research has been funded by a grant from the Central Research Committee and Excellence in Academic Medicine Award from Southern Illinois University and the Muscular Dystrophy Association.

We thank the members of the family for their participation in our study; Dr. Gregory Gregg, Radiology, St. Johns Hospital, for interpretation of x-rays; Dr. Jhaveri, Dr. Onsi Kamel, and Rod Angeli, Pathology Department, St. John's Hospital, for assistance with laboratory results; Dr. Stephen Jennison and Claire Call, Prairie Heart Institute; Shawnia Forrester, Division of Genetics, Southern Illinois University; and Katerina Kimonis from Springfield, IL; Dr. Marzia Pasquale, Emory University, Atlanta, GA; Dr. Michael Whyte, Metabolic Research Unit, Shriners Hospital for Children, St. Louis, MO; and Beverly McElroy, RN, NINDS, National Institutes of Health, Bethesda, MD, for their contributions.

References

1. Caughey JE, Gwynne JF, Jefferson NR. Dystrophia myotonica associated with familial Paget's disease (osteitis deformans) with sarcomata. *J Bone Joint Surg Br* 1957;39: 316–325. [PubMed: 13438974]
2. MIM (Online Mendelian inheritance in man) Baltimore: Johns Hopkins University, Center for Medical Genetics 2000 Available at: <http://www3.ncbi.nlm.nih.gov/omim/>
3. McBride TI, Paget's disease and muscular dystrophy: report of an unusual association in one family. *Scott Med J* 1966;11:238–243. [PubMed: 5953945]
4. Tucker WS, Jr, Hubbard WH, Stryker TD, Morgan SW, Evans OB, Freemon F, Theil GB. A new familial disorder of combined lower motor neuron degeneration and skeletal isorganization. *Trans Assoc Am Phys* 1982;95:126–134. [PubMed: 7182974]
5. Hauser MA, Salmikangas P, Horrigan SK, Viles KD, Torian UM, Taivainen L, et al. Positional cloning of the gene responsible for limb-girdle muscular dystrophy 1A. *Am J Hum Genet* 1999;65:574A.
6. Speer MC, Yamaoka LH, Gilchrist JH, Gaskell CP, Stajich JM, Vance JM, Kazantsev A, Lastra AA, Haynes CS, Beckman JS, et al. Confirmation of genetic heterogeneity in limb-girdle muscular dystrophy: linkage of an autosomal dominant form to chromosome 5q. *Am J Hum Genet* 1992;50:1211–1217. [PubMed: 1598902]
7. Bartoloni L, Horrigan SK, Viles KD, Gilchrist JM, Stajich JM, Vance JM, Yamaoko LH, Pericak-Vance MA, Westbrook CA, Speer MC. Use of a CEPH meiotic breakpoint panel to refine the locus of limb-girdle muscular dystrophy type 1A (LGMD1A) to a 2-Mb interval on 5q31. *Genomics* 1998;54:250–255. [PubMed: 9828127]

8. Minetti C, Sotgia F, Bruno C, Scanzini P, Broda P, Bado M, Masetti E, Mazzocco M, Egeo A, Dondti MA, Volonte D, Galbiati F, Cordone G, Bricarelli FD, Lisanti MP, Zara F. Mutations in the Caveolin-3 gene cause autosomal dominant limb-girdle muscular dystrophy. *Nat Genet* 1998;18:365–368. [PubMed: 9537420]
9. Song KS, Scherer PE, Tang Z, Okamoto T, Li S, Chafel M, Chu C, Kohty DS, Lisanti MP. Expression of caveolin-3 in skeletal, cardiac, and smooth muscle cells: caveolin-3 is a component of the sarcolemma and co-fractionates with dystrophin and dystrophin-associated glycoproteins. *J Biol Chem* 1996;271:15160–15165. [PubMed: 8663016]
10. Van der Kooi AJ, van Meegen M, Ledderhof TM, McNally EM, de Visser M, Bolhuis PA. Genetic localization of a newly recognized autosomal dominant limb-girdle muscular dystrophy with cardiac involvement (LGMD1B) to chromosome 1q11–21. *Am J Hum Genet* 1997;60:891–895. [PubMed: 9106535]
11. Speer MC, Vance JM, Grubber JM, Lennon Graham F, Stajich JM, Viles KD, Rogala A, McMichael R, Chutkow J, Goldsmith C, Tim RW, Pericak-Vance MA. Identification of a new autosomal dominant limb-girdle muscular dystrophy loca on chromosome 7. *Am J Hum Genet* 1999;64:556–562. [PubMed: 9973293]
12. Nicolao P, Xiang F, Gunnarsson LG, Giometto B, Edstrom L, Anvret M, Zhang Z. Autosomal dominant myopathy with proximal weakness and early respiratory muscle involvement maps to chromosome 2q. *Am J Hum Genet* 1999;64:788–792. [PubMed: 10053013]
13. Messina DN, Speer MC, Pericak-Vance MA, McNally EM. Linkage of familial dilated cardiomyopathy with conduction defect and muscular dystrophy to chromosome 6q23. *Am J Hum Genet* 1997;61:909–917. [PubMed: 9382102]
14. Delmas PD, Meunier PJ. The management of Paget disease of bone. *N Engl J Med* 1997;336:558–566. [PubMed: 9023094]
15. McKusik VA, Paget's disease of the bone In: Heritable disorders of connective tissue, 3rd Ed. St. Louis: C.V. Mosby 1960:718–723.
16. Fotino M, Haymivits A, Falk CT. Evidence for linkage between HLA and Paget's disease. *Transplant Proc* 1977;9: 1876–1868.
17. Cody JD, Singer FR. Genetic linkage of Paget's disease of the bone to chromosome 18q. *Am J Hum Genet* 1997;61:1117–1122. [PubMed: 9345096]
18. Hughes AE, Ralston SH, Marken J, Bell C, MacPherson H, Wallace RGH, van Hul W, Whyte MP, Nakatsuka K, Hovy L, Anderson DM. Mutations In TNFRSF11A, affecting the signal peptide of RANK, cause familial expansile osteolysis *Nat Genet* 2000;24:45–48. [PubMed: 10615125]
19. van Hul W, Paget's disease from a genetic perspective. *Bone*, 1999;24(Suppl 5):29–30. [PubMed: 9916781]
20. Ott J Computer-simulation methods In human linkage analysis. *Proc Natl Acad Sci U S A* 1989;86:4175–4178.
21. Weeks DE, Ott J, Lathrop GM. Slink: a general simulation program for linkage analysis. *Am J Hum Genet* 1994;47(Suppl):A204.
22. Cottingham RW, Jr, Idury RM, Schaffer AA. Faster sequential genetic linkage computations. *Am J Hum Genet* 1993;53:252–263. [PubMed: 8317490]
23. Schmorl G. Ueber Ostitis deformans Paget. *Virchows Arch Pathol Anat Physiol Klin Med* 1932;238:694–751.
24. Varelas PN, Bertoni TE, Kapaki E, Papagerogiou CT. Paget's disease of bone and motor neuron disease. *Muscle Nerve* 1997;5:630.
25. Hultgren HN. Osteitis deformans (Paget's disease) and calcific disease of the heart valves. *Am J Cardiol* 1998;81:1461–1464. [PubMed: 9645898]
26. Hughes DE, Wright KR, Uy HL. Bisphosphonates promote apoptosis In murine osteoclasts In vitro and in vivo. *J Bone Res* 1995;10:1478–1487.
27. Smith R, Russell RG, Bishop M, Woods C, Bishop M. Paget's disease of bone-experience with diphosphonate in treatment. *Q J Med* 1973;42:235–238. [PubMed: 4206518]
28. Folkers K, Wolaniuk J, Simonsen R, Morishita M. Biochemical rationale and the cardiac response of patients with muscle disease to therapy with coenzyme Q. *Proc Natl Acad Sci U S A* 1985;82:901–904. [PubMed: 3856239]

29. Dubrovsky AL, Angelini C, Bonifati DM, Pegoraro E, Mesa L. Steroids in muscular dystrophy: where do we stand? *Neuromuscul Disord* 1998;8:380–384. [PubMed: 9713854]
30. Mendell JR, Kissel JT, Amato AA, King W, Signore L, Prior TW, Sahenk Z, Benson S, McAndrew PE, Rice R. Myoblast transfer in the treatment of Duchenne muscular dystrophy. *N Engl Med* 1995;333:832–838.
31. Karpati G, Pari G, Molnar MJ. Molecular therapy for genetic muscle disease—status 1999. *Clin Genet* 1999;55:1–8. [PubMed: 10066023]
32. Muscular Dystrophy Association. New York. New York 2000 <http://www.mdausa.org/research>.
33. Bushby K Making sense of the limb-girdle muscular dystrophies. *Brain* 1999;122: 1403–1420. [PubMed: 10430828]
34. Beckmann JS, Richard I, Hillaire D, Broux O, Antignac C, Bois E, Cann H, Cottingham RW, Jr, Feingold N, Feingold J, et al. A gene for limb-girdle muscular dystrophy maps to chromosome 15 by linkage. *C R Acad Sci III* 1991;312:141–148. [PubMed: 1901754]
35. Richard I, Broux O, Allamand V, Fougousse F, Chiannikulchai N, Bourg N, Brenguier L, Devaud C, Pasturand P, Roudent C, et al. Mutations in the proteolytic enzyme calpain 3 cause limb-girdle muscular dystrophy type 2A. *Cell* 1995;81:27–40. [PubMed: 7720071]
36. Bashir R, Strachan T, Keers S, Stephenson A, Mahineh I, Marconi G, Nashef L, Bushby KM. A gene for autosomal recessive limb-girdle muscular dystrophy maps to chromosome 2p. *Hum Mol Genet* 1994;3:455–457. [PubMed: 8012357]
37. Noguchi S, McNally EM, Ben Othmane K, Hagiwara Y, Mizuno Y, Yoshida M, Yamamoto H, Bonnemant CG, Gussoni E, Denton OH, et al. Mutations in the dystrophin-associated protein gamma-sarcoglycan in chromosome 13 muscular dystrophy. *Science* 1995;270:819–821. [PubMed: 7481775]
38. Roberds SL, Leturcq F, Allamand V, Piccolo F, Jeanpierre M, Anderson RD, Lim LE, Lee JC, Tome FM, Romeo NB, et al. Missense mutations in the adhalin gene linked to autosomal recessive muscular dystrophy. *Cell* 1994;78:625–633. [PubMed: 8069911]
39. Lim LE, Duclos F, Broux O, Bourg N, Sunada Y, Allamand V, Meyer J, Richard I, Moomaw C, Slaughter C, et al. Beta-sarcoglycan: characterization and role in limb-girdle muscular dystrophy linked to 4q12. *Nat Genet* 1995;11:257–265.
40. Nigro V, Moreira ES, Piluso G, Vainzof M, Belsito A, Politano L, Puca A, Passos-Bueno MR, Zatz M. Autosomal recessive limb-girdle muscular dystrophy, LGMD2F, is caused by a mutation in the delta sarcoglycan gene. *Nat Genet* 1996;14: 195–198. [PubMed: 8841194]
41. Passos-Bueno MR, Moreira ES, Vainzof M, Marie SK, Zatz M. Linkage analysis in autosomal recessive limb-girdle muscular dystrophy (ARLGMD) maps a sixth form to 5q33–34 (LGMD2F) and indicates that there is at least one more subtype of ARLGMD. *Hum Mol Genet* 1996;5:815–820. [PubMed: 8776597]
42. Moreira ES, Passos-Bueno MR, Wiltshire TJ, Faulkner G, Nilforoushan A, Vainzof M, Suzuki OT, Valle G, Reeves R, Zatz M, Passos-Bueno MK, Jenne DE. Limb-girdle muscular dystrophy Type 2G is caused by mutations in the gene encoding the sarcomeric protein telethonin. *Nat Genet* 2000;24:163–166. [PubMed: 10655062]
43. Moreira ES, Vainzof M, Marie SK, Sertie AL, Zatz M, Passos-Bueno MR. The seventh form of autosomal recessive limb-girdle muscular dystrophy is mapped to 17q11–12. *Am J Hum Genet* 1997;61:151–159. [PubMed: 9245996]
44. Weiler T, Greenberg CR, Zelinski T, Nysten E, Coghlan G, Crumley MJ, Fuliwara TM, Morgan K, Wrogemann K. A gene for autosomal recessive limb-girdle muscular dystrophy in Manitoba Hutterites maps to chromosome region 9q31–q33: evidence for another limb-girdle muscular dystrophy locus. *Am J Hum Genet* 1998;63: 140–147. [PubMed: 9634523]

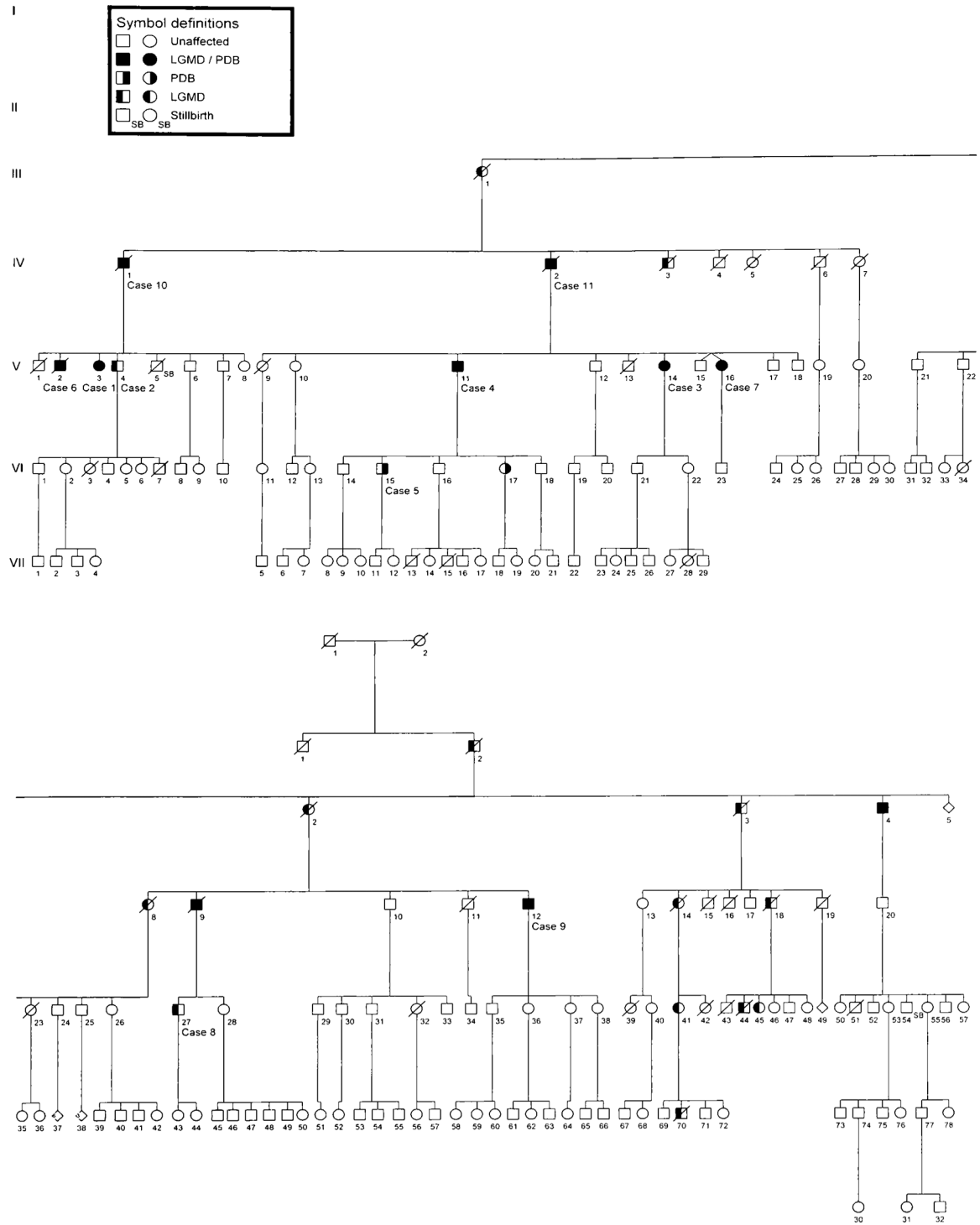


Fig. 1. Pedigree of family affected with autosomal dominant LGMD and PDB. Circles denote females, squares denote males, slashed symbols denote deceased individuals.

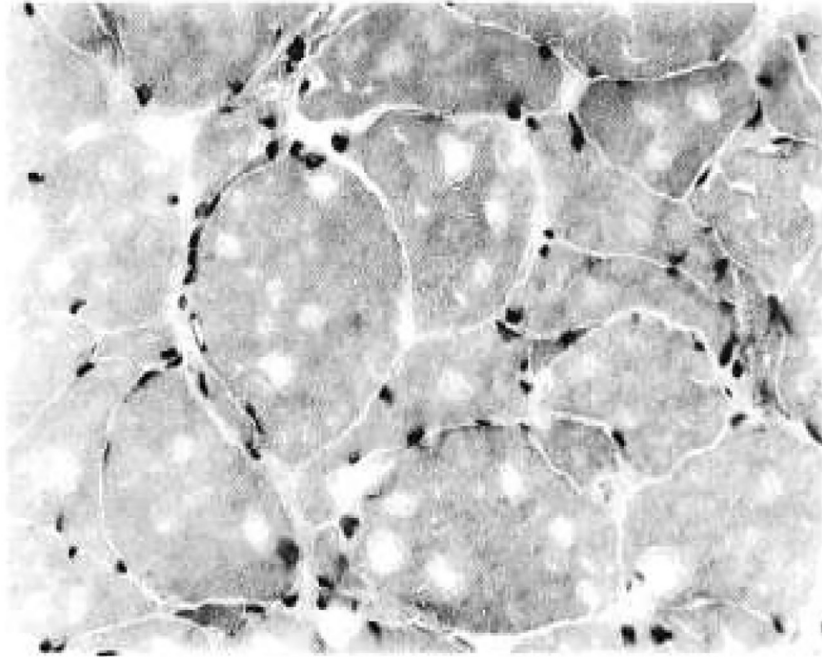


Fig. 2. High powered magnification (hematoxylin and eosin, $\times 400$) of muscle from the right deltoid of case 9 showed variation in fiber size with frequent angular fibers. Many of the fibers contained blue, rimmed vacuoles.

Table 1 Clinical and genetic heterogeneity of limb-girdle muscular dystrophy and Paget disease of bone

Type	Reference	Inheritance	Chromosomal locus	Protein Defect	Clinical Features
LGMD1A	5, 6	Dominant	5q22.3-q31.3	Myotilin	Onset 18–35 y, progressive muscle disease, mildly elevated CPK, tight Achilles tendon, distinctive dysarthric speech, anticipation
LGMD1B	10	Dominant	1q11–q21	Unknown	Onset muscle weakness <20 y in 50%, CPK normal, mild muscular dystrophy on EMG and muscle biopsy. Death from cardiomyopathy
LGMD1C	8, 9	Dominant	3p25	Caveolin 3	Calf hypertrophy, mild to moderate weakness, CPK is 4–25× normal, rapid progression in one family
LGMD1D	11	Dominant	7q	Unknown	No related cardiomyopathy, 1 family had early onset muscle weakness
Edstrom Myopathy	12	Dominant	2q24–q31	Unknown	Normal to mild elevation of CPK, myopathic changes on EMG, no necrosis or fibrosis on muscle biopsy
LGMD2A	34, 35	Recessive	15q15.1–q21.1	Calpain	Majority onset 8–15 y, calf hypertrophy, Achilles tendon contractures, respiratory complications severe, no cardiac involvement
LGMD2B	36	Recessive	2p13–p16	Dysferlin/DYSF	2 families. Onset in late teens, with slow progression, CPK is 25–56 × normal, EMG myopathy, muscle biopsy necrotizing myopathy
LGMD2C	37	Recessive	13q12	Gamma sarcoglycan	Onset before 5 y, calf shows muscle pseudohypertrophy, CPK elevated early, lumbar lordosis. Death typically before 20 y
LGMD2D	38	Recessive	17q21	Alpha sarcoglycan	Typical childhood onset, calf hypertrophy, Achilles tendon contractures, elevation in CPK, respiratory impairment
LGMD2E	39	Recessive	4q12	Beta sarcoglycan	Progressive weakness, elevated CPK, dystrophic changes in muscle biopsy
LGMD2F	40, 41	Recessive	5q33–q34	Delta sarcoglycan	Onset 2–7 y, confinement to a wheelchair at 11–16 y, death from 9–19 y, serum CPK elevated 10–50× normal
LGMD2G	42, 43	Recessive	17q11–12	Telethonin	Mean age onset 12.5 y, difficulty to walking on heels, absent tendon reflexes, foot drop, no cardiac disease, CPK 3–17× normal, muscle biopsy showed myopathy and 1 rimmed vacuole/fiber
LGMD2H	44	Recessive	9q31–q33	Unknown	Onset between 8–27 y, CPK elevated 2–30 × normal, facial muscles involved, usually ambulant until late in life
PDB1	16	Dominant	6p	Unknown	Classic distribution of Paget in 29 individuals in 3 families
PDB2	17, 18	Dominant	18q21–22	RANK	Alkaline phosphatase 25–2000 U/L, classic distribution of Paget disease in 8 affected individuals in an Irish family

Table 2
 TWO point LOD score analysis reflecting linkage between LGMD/PDB disease and known disease loci.*

Marker	MIM Number ²	Locus	Chromosome	0.00	0.01	0.02	0.03	0.04	0.05	0.10	0.20	0.30	0.40	Exclusion
D5S479	159000	LGMD1A	5q22.3-q31.1	-9.96	-4.29	-3.55	-3.09	-2.76	-2.49	-1.63	-0.77	-0.33	-0.09	7.05 cM
D5S500	159000	LGMD1A	5q22.3-q31.1	-6.47	-3.30	-2.68	-2.30	-2.03	-1.82	-1.14	-0.51	-0.23	-0.08	4.01 cM
D1S305	159001	LGMD1B/CDCD1	1q11-q21	-9.72	-4.29	-3.49	-2.99	-2.63	-2.35	-1.46	-0.62	-0.24	-0.06	6.03 cM
AFOA2	159001	LGMD1B/CDCD1	1q11-q21	-14.04	-7.11	-5.71	-4.88	-4.28	-3.82	-2.42	-1.11	-0.46	-0.12	12.24 cM
D3S1263	601253	LGMD1C/CDCD2	3p25	-11.05	-5.30	-4.39	-3.80	-3.36	-3.01	-1.88	-0.82	-0.33	-0.09	9.10 cM
D7S550	603511	LGMD1D	7q	-7.74	-4.28	-3.56	-3.12	-2.80	-2.54	-1.74	-0.91	-0.43	-0.15	7.05 cM
D2S138	603689	Edström's myopathy	2q24-q31	-6.75	-0.27	0.11	0.33	0.48	0.59	0.83	0.79	0.50	0.17	On marker
D2S1245	603689	Edström's myopathy	2q24-q31	-3.58	-0.63	-0.24	-0.01	0.15	0.26	0.53	0.51	0.26	0.06	On marker
D21S1255	105400	ALS1	21q22.1	-6.69	-3.04	-2.48	-2.10	-1.82	-1.59	-0.87	-0.28	-0.09	-0.04	3.00 cM
D15S778	253600	LGMD2A	15q15.1-q21.1	-2.70	-1.17	0.87	-0.68	-0.55	-0.44	-0.09	0.17	0.18	0.08	On marker
D2S2113	253601	LGMD2B	2p13-p16	-16.53	-5.12	-3.93	-3.23	-2.75	-2.38	-1.28	-0.39	-0.07	0.00	6.03 cM
D13S115	253700	LGMD2C	13q12	-6.55	-3.30	-2.71	-2.34	-2.06	-1.84	-1.13	-0.48	-0.18	-0.05	4.00 cM
ADL	600119	LGMD2D	17q21	-5.73	-2.74	-2.18	-1.80	-1.51	-1.28	-0.55	0.04	0.16	0.09	2.00 cM
D4S2690	604286	LGMD2E	4q12	-4.24	-1.05	-0.72	-0.51	-0.36	-0.25	0.09	0.28	0.23	0.08	On marker
D5S470	601287	LGMD2F	5q33-q34	-12.58	-4.49	-3.73	-3.26	-2.90	-2.62	-1.72	-0.83	-0.35	-0.10	8.07 cM
D17S1818	601954	LGMD2G	17q11-q12	-8.93	-3.42	-2.82	-2.48	-2.23	-2.04	-1.42	-0.70	-0.27	-0.06	5.02 cM
D17S250	601954	LGMD2G	17q11-q12	-10.97	-7.50	-6.23	-5.42	-4.82	-4.34	-2.80	-1.31	-0.57	-0.17	14.38 cM
D9S302	254110	LGMD2H	9q31-q33	-11.67	-4.37	-3.54	-3.02	-2.65	-2.36	-1.52	-0.85	-0.52	-0.23	6.03 cM
D6S105	167250	PDB1	6p21.2	-9.93	-4.31	-3.63	-3.19	-2.86	-2.59	-1.67	-0.72	-0.26	-0.05	7.05 cM
D18S60	602080	PDB2	18q21-q22	-4.76	-2.04	-1.71	-1.52	-1.37	-1.26	-0.89	-0.47	-0.21	-0.05	1.00 cM
D18S42	602080	PDB2	18q21-q22	-9.06	-4.72	-3.94	-3.44	-3.06	-2.76	-1.77	-0.79	-0.31	-0.07	8.07 cM
D9S152	600884	CMD1B	9q13-q22	-7.70	-1.53	-1.19	-0.99	-0.84	-0.72	-0.35	-0.03	0.06	0.05	On marker
D9S153	600884	CMD1B	9q13-q22	-7.88	-1.42	-1.07	-0.87	-0.72	-0.61	-0.28	-0.05	-0.01	-0.02	On marker
D10S201	601493	CMD1C	10q21-q23	-13.09	-4.98	-4.12	-3.57	-3.16	-2.83	-1.79	-0.79	-0.30	-0.06	8.07 cM
D1S414	601493	CMD1D	1q32	-5.31	-0.79	-0.25	0.05	0.24	0.38	0.70	0.71	0.47	0.19	On marker
D6S262	602067	CMD1F/CDCD3	6q23	-5.42	-1.67	-1.30	-1.05	-0.87	-0.71	-0.22	0.16	0.20	0.10	On marker
D6S1040	602067	CMD1F/CDCD3	6q23	-5.93	-3.06	-2.52	-2.16	-1.90	-1.69	-1.02	-0.42	-0.14	-0.03	3.00 cM

* CDCD, cardiomyopathy, dilated, conduction defect; CMD, cardiomyopathy, dilated.

Table 3

Clinical data of affected individuals*

Case	Age (y)	Sex	PDB	LGMD	AgeDx PDB(y)	AgeDx LGMD (y)	Treatment	Activity Level
1	48	F	+	+	34	44	None	Difficulty going up stairs and raising arms, back/leg pain, lumbar lordosis, wide based gait
2	47	M	-	+	-	45	None	Weakness in proximal arm and leg muscles
3	49	F	+	+	44	48	Previous bisphosphonates	Difficulty climbing stairs, lifting arms, extremity atrophy of extremities, lordotic posture, waddling gait
4	57	M	+	+	42	54	Calcitonin and bisphosphonates	Walks slowly with cane, severe pain and tenderness in spine and hip
5	33	M	+	-	31	-	Analgesics and calcitonin	Walks slowly with a limp, tenderness in back, pain in hip
6	50	M	+	+	38	38	None	Bedridden for several months, left side hemiparesis from stroke
7	47	F	+	+	45	45	Alendronate, cyclobenzaprine	Difficulty sitting, getting up from chair, back pain and weakness of the arms and hips
8	46	M	-	+	-	40	None	Unable to run, difficulty with stairs
9	64	M	+	+	37	40	None	Wheelchair bound, loss of sensation in extremities
10	59	M	+	+	UK	51	None	Bedridden 3 y prior to death, winging of the scapulae, weakness of trunk & lower extremities
11	61	M	+	+	61	UK	None	Difficulty walking, pain in spine and pelvis and femur
MEAN	50.9	3F/8M	9	10	41.5	45.6		

* + = present; - = absent; UK = unknown

Table 4

Laboratory data of affected individuals*

Case	Alkaline Phosphatase (30–130 IU/L)	Bone Specific Alkaline Phosphatase (39–151 IU/L)	Osteocalcin (8.0–52.0 ng/mL)	CPK (20.0–222.0 U/L)	Muscle Biopsy Findings	Echocardiogram Findings	X-ray Findings of Paget
1	1137	1133	34.4	60	NT	Normal	L1, L2, L3, and pelvis
2	93	NT	25.2	220	Fiber size variation, splitting & nesting	Cardiomyopathy	No Paget
3	354	351	19.1	98	Atrophic fibers	NT	L2, L3, and pubic bone
4	786	764	5	268	Myopathy	Normal	Spine, hip, and skull
5	1724	1724	54.6	90	NT	NT	Spine, hip, and skull
6	240	NT	NT	82	NT	Cardiomyopathy	NT
7	329	NT	NT	NT	NT	NT	L5
8	88	85	1	1145	End stage myopathy	Normal	Possible early Paget
9	1508	1488	1	220	Muscle atrophy, vacuolar myopathy	Aortic valve calcification, mild cardiomyopathy	Widespread
10	431	NT	NT	84	Adipose tissue between muscle bundles	NT	NT
11	1535	NT	NT	NT	NT	Normal	Lumbar spine and pelvis
Mean	747.7	924.2	20.0	251.9			

* NT, not tested; CPK, creatine phosphokinase.

Clinical, radiological and biochemical data in Paget and neuromuscular disease (NMD) reported in the literature

Table 5

Author	Case	Sex	PBD	NMD	Alkaline Phosphatase (4.5–13 KA units, 30–115 U/L)	CPK (0–100 U/L)	Paget Disease Involvement	Neuromuscular Disease
McBride et. al ²	1	M	+	+	161 KA units	Normal	Widespread	Onset at 33 y, proximal muscle weakness, muscle biopsy consistent with LGMD
	2	M	+	+	154 KA units	Normal	Lumbar spine, pelvis, upper femora, left tibia	Myocardial ischaemia. EMG showed myopathic pattern, weakness and wasting of shoulder girdle and pelvis, hands severely affected
	3	M	+	+	23 KA units	Normal	Lumbar spine	Deterioration of mental condition, wasting of upper limb muscles, thighs, and hands
	4	F	+	+	27 KA units	Normal	Widespread	Scoliosis, scapular winging, wasting of hand muscles, waddling gait
	5	F	+	-	14.8 KA units	Normal	Single lumbar vertebra	Diagnosed with schizophrenia, no muscle weakness or wasting
Caughey et. al ¹	1	M	+	+	68 KA units	NA	Pelvis, spine and skull	Myotonic dystrophy features; baldness at 17 y, stiff leg muscles, bilateral cataracts, myotonia of the tongue, weak abdominal muscles
	2	M	+	+	NA	NA	Spine and pelvis	Weakness of neck flexors and abdomen, myotonia of hands and tongue, frontal baldness, leukoderma of face and limbs
	3	M	-	+	NA	NA	No evidence of Paget	Myotonia of the hands, hypertrophy of the thigh and calf muscles, muscle size disproportionate to strength
Tucker et. al ³	1	M	+	+	222 U/L	125 U/L	Pelvic, L5, femoral heads, skull	Amyotrophic lateral sclerosis, moderate atrophy of limb girdle muscles, winging of scapulae, waddling gait, absent tendon reflexes, onset at 40 y

NA = not available