Marfan’s syndrome

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Marfan’s syndrome is a systemic disorder of connective tissue caused by mutations in the extracellular matrix protein fibrillin 1. Cardinal manifestations include proximal aortic aneurysm, dislocation of the ocular lens, and long-bone overgrowth. Important advances have been made in the diagnosis and medical and surgical care of affected individuals, yet substantial morbidity and premature mortality remain associated with this disorder. Progress has been made with genetically defined mouse models to elucidate the pathogenetic sequence that is initiated by fibrillin-1 deficiency. The new understanding is that many aspects of the disease are caused by altered regulation of transforming growth factor β (TGFβ), a family of cytokines that affect cellular performance, highlighting the potential therapeutic application of TGFβ antagonists. Insights derived from studying this mendelian disorder are anticipated to have relevance for more common and non-syndromic presentations of selected aspects of the Marfan phenotype.

Marfan’s syndrome is a systemic disorder of connective tissue, first described more than 100 years ago by a Parisian professor of paediatrics, Antoine-Bernard Marfan, who reported the association of long slender digits and other skeletal abnormalities in a 5-year-old girl, Gabrielle.1 Today, the disorder that bears his name has benefited from decades of clinical and molecular investigation. This review will highlight the clinical manifestations, molecular pathogenesis, standard treatment, and emerging therapeutic options for this multisystem disorder.

Epidemiology

The incidence of classic Marfan’s syndrome is about 2–3 per 10000 individuals, although this estimate depends on complete recognition of all affected and genetically predisposed individuals. Gray reported the incidence in Scotland as one in 9802 livebirths.2 Various factors could have contributed to an underestimate of disease prevalence. First, the phenotype becomes more apparent with increasing age in most families. Second, many of the outward manifestations are common in the general population and many physicians miss the diagnostic significance of these findings. Third, while the disorder segregates as a dominant trait in families, about 25% of cases are sporadic due to de-novo mutations; a family history of Marfan’s syndrome is not always present as an obvious risk factor. Fourth, although it is known that mutations in the FBN1 gene, which encodes the matrix protein fibrillin 1, are the predominant (if not the sole) cause of classic Marfan’s syndrome, there is no rapid and efficient molecular diagnostic test.1,4

The disease occurs worldwide, with no predilection for either sex. Tall stature with dolichostenomelia (long-bone overgrowth) leads to increased incidence in certain athletes, including basketball and volleyball players, sometimes prompting a recommendation for screening by echocardiography.5 Kinoshita and co-workers6 screened 415 basketball and volleyball athletes in high school, college, or professional sports. Of these, four (1%) had aortic enlargement, with a diameter greater than 4.6 cm at the aortic root, and two were diagnosed with Marfan’s syndrome. Aortic enlargement was diagnosed independent of body surface area in this analysis. Although outdated criteria for the clinical diagnosis of the disorder were used, these data seem to indicate an increased incidence in this select population.

Because of the high incidence of aortic root aneurysm with associated risk of life-threatening aortic dissection, lifespan is often shortened. Before the successful use of surgical aortic root replacement, death from aortic dissection was far more common than it is today. A report in the early 1970s on life expectancy and cause of death in Marfan’s syndrome describes that life expectancy for affected individuals was about two-thirds that of unaffected individuals, with life-table mortality curves deviating in infancy between controls and affected individuals.5 Cause of death was cardiovascular (aortic dissection, congestive heart failure, or cardiac valve disease) in over 90% of cases. However, a more recent assessment of outcome in Marfan’s syndrome describes a nearly normal life expectancy, indicating improvement in the recognition and treatment of these conditions.7

Clinical manifestations

Marfan’s syndrome is a multisystem disorder with manifestations typically involving the cardiovascular, skeletal, and ocular systems. A consensus opinion regarding diagnostic criteria was outlined at the International Nosology of Heritable Disorders of Connective Tissue Meeting in Berlin in 1986.8 However, the recognition that many individuals diagnosed with the disorder do not carry the FBN1 mutation identified in more severely affected family members led to revised criteria several years later that were more specific and stringent (panel). These criteria placed greater emphasis

Search strategy and selection criteria

The material covered in this review is based on extensive review of published works, including journals and textbook articles, as well as our ongoing investigation in this area. For PubMed searches we used the search term “Marfan”. We did not restrict our search by language, and, when possible, references within the past 5 years were chosen.
on the diagnostic use of skeletal findings and a requirement that a positive family history of the disorder could only be used as a major criterion for diagnosis of a proband if at least one family member independently satisfied diagnostic criteria on the basis of physical manifestations alone.9

### Panel: Diagnostic criteria for Marfan’s syndrome

#### Index case
- If the family/genetic history is not contributory, major criteria in two or more different organ systems and involvement of a third organ system.
- If a mutation known to cause Marfan’s syndrome in others is identified, one major criterion in an organ system and involvement of a second organ system.

#### Relative of an index case who independently meets these strict diagnostic criteria
- Presence of a major criterion in the family history, one major criterion in an organ system, and involvement of a second organ system.

#### Organ systems

**Skeletal**

At least four of the following constitutes a major criterion in the skeletal system:

- Pectus carinatum
- Pectus excavatum, needing surgery
- Reduced upper-segment to lower-segment ratio or arm span to height ratio > 1.05
- Wrist and thumb signs
- Scoliosis of > 20° or spondylolisthesis
- Reduced extension at the elbows (< 170°)
- Medial displacement of the medial malleolus, causing pes planus
- Protrusio acetabulae of any degree (ascertained on radiographs)

**Minor criteria**

- Pectus excavatum of moderate severity
- Joint hypermobility
- Highly arched palate with crowding of teeth
- Facial appearance (dolichocephaly, malar hypoplasia, enophthalmos, retrognathia, down-slanting palpebral fissures)

For involvement of the skeletal system, at least two features contributing to major criteria, or one feature from the list contributing to the major criterion and two of the minor criteria must be present.

#### Cardiovascular system

**Major criteria** (either of the following)

- Dilatation of the ascending aorta, with or without aortic regurgitation, and involving at least the sinuses of Valsalva
- Dissection of the ascending aorta

**Minor criteria**

- Mitral valve prolapse with or without mitral valve regurgitation
- Dilatation of the main pulmonary artery, in the absence of valvular or peripheral pulmoni stenosis or any other obvious cause, younger than age 40 years
- Calcification of the mitral annulus younger than age 40 years
- Dilatation or dissection of the descending thoracic or abdominal aorta younger than age 50 years

For involvement of the cardiovascular system, only one of the minor criteria must be present.

#### Pulmonary system

**Major criteria**

- None

**Minor criteria**

- Spontaneous pneumothorax
- Apical blebs

For involvement of the pulmonary system, only one of the minor criteria must be present.

#### Skin and integument

**Major criteria**

- None

**Minor criteria**

- Striae atrophicae (stretch marks) without marked weight gain, pregnancy, or repetitive stress
- Recurrent or incisional herniae

For involvement of the skin and integument, only one of the minor criteria must be present.

#### Dura

**Major criterion**

- Lumbosacral dural ectasia

**Minor criteria**

- None

#### Family/genetic history

**Major criteria** (any one of the following)

- Having a parent, child, or sibling who meets these diagnostic criteria independently
- Presence of a mutation in FBN1, which is known to cause Marfan’s syndrome
- Presence of a haplotype around FBN1, inherited by descent, known to be associated with unequivocally diagnosed Marfan’s syndrome in the family

**Minor criteria**

- None
**Skeletal system**

Disproportionate overgrowth of the long bones is often the most striking and immediately evident manifestation. Anterior chest deformity is caused by overgrowth of the ribs, pushing the sternum anteriorly (pectus carinatum) or posteriorly (pectus excavatum). Overgrowth of arms and legs can lead to an arm span greater than 1.05 times the height or a reduced upper to lower segment ratio (in the absence of severe scoliosis). Arachnodactyly (overgrowth of the fingers) is generally a subjective finding. The combination of long fingers and loose joints leads to the characteristic Walker-Murdoch or wrist sign: full overlap of the distal phalanges of the thumb and fifth finger when wrapped around the contralateral wrist. The Steinberg or thumb sign is present when the distal phalanx of the thumb fully extends beyond the ulnar border of the hand when folded across the palm. Although commonly present, thoracolumbar scoliosis must be sufficiently severe (≥20°) to contribute to the skeletal criteria for the syndrome. Protrusio acetabuli, which is generally asymptomatic in young adults, is best identified with radiographic imaging. Pes planus (flat feet) are frequently present, and vary from mild and asymptomatic to severe deformity wherein medial displacement of the medial malleolus results in collapse of the arch and often reactive hip and knee disturbances. Curiously, a subset of individuals with the disorder present with an exaggerated arch, pes cavus. Although joint laxity or hypermobility is frequently identified, joints can be normal or even develop contractures. Reduced extension of the elbows is common and can contribute to the designation of major involvement of the skeleton. Contracture of the fingers (camptodactyly) is also commonly observed, especially in children with severe and rapidly progressive Marfan’s syndrome. Several craniofacial manifestations are frequently present, but are not specific enough to the disorder for inclusion in the major criteria. These include a long narrow skull (dolicocephaly), a high-arched palate, tooth crowding, retroglossal (recessed lower mandible) or micrognathia (small chin), malar flattening, and downward-slaingting palpebral fissures.

**Ocular system**

A major criterion is present with ectopia lentis (dislocation of the ocular lens) of any degree, although, once again, this condition is not unique to Marfan’s syndrome. Ectopia lentis has been shown to occur in around 60% of patients with the disorder. When identified, it should prompt further assessment for Marfan’s syndrome, although homocystinuria, Weill-Marchesani, and familial ectopia lentis are also associated with this condition. Other manifestations of the ocular system include early and severe myopia, flat cornea, increased axial length of the globe, hypoplastic iris, and ciliary muscle hypoplasia, causing decreased miosis. Individuals with Marfan’s syndrome can also have retinal detachment and a predisposition for early cataracts or glaucoma.

**Cardiovascular system**

Manifestations of Marfan’s syndrome in the cardiovascular system are conveniently divided into those affecting the heart and those affecting the vasculature. Within the heart, the atrioventricular valves are most often affected. Thickening of the atrioventricular valves is common and often associated with prolapse of either the mitral or tricuspid valves or both atrioventricular valves. Variable degrees of regurgitation may be present. In children with early onset and severe Marfan’s syndrome, insufficiency of the mitral valve can lead to congestive heart failure, pulmonary hypertension, and death in infancy; this insufficiency represents the leading cause of morbidity and mortality in young children with the disorder. A survey of 166 patients with apparent Marfan’s syndrome (mean age 11.9 years), identified more than half of them to have auscultatory or echocardiographic evidence of mitral valve dysfunction, most commonly prolapse. In this same analysis, more than 25% of these individuals had progression of mitral valve prolapse to mitral regurgitation by adulthood, with twice as many women as men progressing in their mitral valve dysfunction. Calcification of the mitral annulus in individuals younger than age 40 years constitutes a minor criterion in the cardiovascular system.

Aortic valve dysfunction is generally a late occurrence, attributed to stretching of the aortic annulus by an expanding root aneurysm. Both the aortic and atrioventricular valves seem to be more prone to calcification in individuals with Marfan’s syndrome. Ventricular dysrhythmia has been described among a small cohort of children with the disorder. In association with mitral valve dysfunction, supraventricular arrhythmia, such as atrial fibrillation, may be seen. Several groups have described increased prevalence of prolonged QT interval on electrocardiographic surveys of patients with Marfan’s syndrome.

Dilated cardiomyopathy, beyond that explained by aortic or mitral valve regurgitation, seems to occur with increased prevalence in patients with Marfan’s syndrome, perhaps implicating a role of the extracellular matrix protein fibrillin 1 in the cardiac ventricles. However, the occurrence rate seems to be low, and the occurrence of mild to moderate ventricular systolic dysfunction is often attributed to mitral or aortic insufficiency, or to the use of β adrenergic receptor blockade. Meijboom and colleagues reported their findings in 234 patients with the disorder, without significant valve disease or aortic root surgery. They identified 17 of these people with left ventricular enlargement, but none had fractional shortening less than 25%, with an average age among the cohort of 29 years (SD 11).

Aortic aneurysm and dissection remain the most life-threatening manifestations of Marfan’s syndrome. This finding is age dependent, prompting life-long monitoring by echocardiography or other imaging modalities. Dilatation at the sinuses of Valsalva can begin in utero in...
severe cases, although some unequivocally affected individuals never reach an aortic size that needs surgical intervention. By contrast with atherosclerotic aneurysms and some other forms of ascending aortic aneurysms, dilatation is generally greatest at (and often restricted to) the aortic root. Normal aortic dimensions vary with both age and body size; proper interpretation of aortic dimensions needs comparison to age-dependent nomograms (figure). The two most important determinants of risk of dissection of the aorta are the maximal dimension and family history of dissection. Surgical repair of the aorta is recommended when its greatest diameter reaches about 50 mm in adults. Early intervention is considered in the presence of a family history of early dissection. There are no definitive methods to guide the timing of surgery in childhood. The observation that dissection is extremely rare in this age-group irrespective of aortic size has prompted many centres to adopt the adult criterion of 50 mm. Early surgery is often undertaken given a rapid rate of growth (>1 cm in a year) or the emergence of significant aortic regurgitation. Most patients with acute aortic dissection have classic symptoms, including severe chest pain, often radiating along the path of dissection. This path almost invariably begins at the aortic root (type A) and dissection can remain isolated (type II) or propagate along the length of the descending aorta (type I). Acute onset congestive heart failure typically indicates severe aortic valve insufficiency, complicating the aortic dissection. Depending on the involvement of the carotid arteries, some patients can have neurological sequelae due to cerebrovascular injury. Involvement of the coronary arteries can lead to myocardial infarction or sudden cardiac death. The mechanism of death usually includes rupture into the pericardial sac with subsequent pericardial tamponade. Chronic aortic dissection and intimal tears usually present more insidiously, often without chest pain.

Dilatation of the main pulmonary artery, in the absence of valvular or peripheral pulmonic stenosis, or any other obvious cause, at age younger than 40 years, constitutes a minor criterion in the cardiovascular system. Dilatation or dissection of the descending thoracic or abdominal aorta at age younger than 50 years is also regarded as a minor criterion in the cardiovascular system.

Pulmonary system

Several factors can result in pulmonary disease in patients with Marfan’s syndrome. Pectus excavatum or progressive scoliosis can contribute to a restrictive pattern of lung disease. Widening of the distal airspaces with or without discrete bullae or (often apical) blebs can predispose to spontaneous pneumothorax, which is found in 4–15% of patients with the disorder. During assessment of pulmonary volumes and function, one must recognise that long-bone overgrowth affecting the lower extremities can lead to reduction in the normalised forced vital capacity and total lung capacity. However, if normalised to thoracic size or sitting height, pulmonary function testing is often normal in patients with the disorder.

Skin and integument

By contrast with several other connective tissue disorders, such as the Ehlers-Danlos syndromes, patients with Marfan’s syndrome typically have normal skin texture and elasticity. The most common manifestation in the skin is striae atrophicae, which occurs in about
two-thirds of patients.26 By contrast with striae in those without a connective tissue disorder, stretch marks tend to occur in the absence of obesity, rapid gain in muscle mass, or pregnancy, and at sites not associated with increased skin distension (eg, the anterior shoulder area and the lower back). Another common manifestation is inguinal hernia, either occurring at birth or acquired in adolescence. There is an increased risk of surgical and recurrent hernias in the Marfan population.

**Dural ectasia**

Widening of the dural sac or root sleeves constitutes ectasia of the dura, which is present in 63–92% of people with the disorder.27–29 Although dural ectasia can result in lumbar back pain, it is often asymptomatic and should be assessed by lumbosacral imaging with CT or MRI. When present, it constitutes a major criterion for the clinical diagnosis of Marfan’s syndrome, although the specificity and predictive value of this finding are not known. Incidence of dural ectasia in other heritable disorders of connective tissue has not been rigorously assessed, although the results of one trial indicate that 24% of patients with Ehlers-Danlos syndrome had dural ectasia.30

**Family/genetic history**

By contrast with the consensus criteria for Marfan’s syndrome established in 1986, the revised criteria emphasise the need for a first-degree relative to independently meet criteria before including family history as a major criterion for the diagnosis of the disorder. The presence of an \( FBN1 \) mutation or the \( FBN1 \) locus, inherited by descent and unequivocally associated with a relative diagnosed with Marfan’s syndrome in the family, constitutes a major criterion in the diagnosis of the disorder.

**Differential diagnosis**

Several disorders are included in the differential diagnosis of Marfan’s syndrome on the basis of similar skeletal, cardiac, or ophthalmological manifestations (table). Many individuals referred for possible Marfan’s syndrome are shown to have evidence of a systemic disorder of the connective tissue, including long limbs, deformity of the thoracic cage, striae atrophicae, mitral valve prolapse, and mild and non-progressive dilatation of the aortic root, but do not meet diagnostic criteria for the disorder. This constellation of features, not fulfilling the diagnostic requirements of the disorder, is referred to by the acronym MASS phenotype, emphasising the mitral, aortic, skin, and skeletal manifestations.31,32 MASS phenotype can segregate in large pedigrees and remain stable over time. This diagnosis is most challenging in the context of an isolated and young individual. In this setting, careful follow-up is needed to distinguish MASS phenotype from emerging Marfan’s syndrome, especially in children. Other fibrillinopathies, such as familial mitral valve prolapse syndrome and familial ectopia lentis, also include subclinical manifestations and can be due to mutations in the gene encoding fibrillin 1.31,32

Also included in the differential diagnosis of the disorder is homocystinuria caused by a deficiency of cystathionine \( \beta \) synthase. Patients with homocystinuria often have tall stature, long-bone overgrowth, and ectopia lentis, but do not typically have aortic enlargement or dissection. By contrast with Marfan’s syndrome, the inheritance of homocystinuria is autosomal recessive, and affected individuals often have mental retardation, a predisposition to thromboembolism, and a high incidence of coronary artery atherosclerosis. Observation of severely raised concentrations of plasma homocystine is an efficient mechanism to distinguish homocystinuria from Marfan’s syndrome.

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Familial thoracic aortic aneurysm syndrome segregates as a dominant trait and can show vascular disease identical to that seen in Marfan’s syndrome, including aortic root aneurysm and dissection. These individuals generally do not show any of the systemic manifestations of Marfan’s syndrome. Other families show the association between bicommissural aortic valve and ascending aortic aneurysm, which can also segregate as a dominant trait. Here, maximum dilatation often occurs further up in the ascending aorta, beyond the sinotubular junction. There is emerging evidence that the bicommissural aortic valve and aneurysm both represent primary manifestations of a single gene defect, and that family members of a proband can show aneurysm without the accompanying valve abnormality. Once again, affected individuals do not show systemic features of a connective tissue disorder. Unlike Marfan’s syndrome, families with both thoracic aortic aneurysm syndrome and with bicommissural aortic valve and ascending aortic aneurysm show reduced penetrance and variable onset of aortic dilatation. Although genetic loci have been described for thoracic aortic aneurysm syndrome, no specific genes (or molecular tests) have been described for these disorders, mandating ongoing follow-up of all at-risk family members. The management principles that have been generated for Marfan’s syndrome have proven effective for these other forms of familial aortic aneurysm.

A clear exception to this rule applies to an aortic aneurysm syndrome that associates some systemic features of Marfan’s syndrome with other features that are quite unique. As in Marfan’s syndrome, patients with Loeys-Dietz aortic aneurysm syndrome show malar hypoplasia, arched palate, retroglossia, pectus deformity, scoliosis, joint laxity, dural ectasia, and aortic root aneurysms and dissection. Although their fingers tend to be long, overgrowth of the long bones can be subtle and is often absent. These patients do not show ectopia lentis. Unique features include a high frequency of hypertelorism, broad or bifid uvula, arterial tortuosity, and aneurysms with dissection throughout the arterial tree. The aneurysms often dissect at sizes not associated with risk in Marfan’s syndrome, and frequently lead to death in young childhood. Other less consistent features include blue sclerae, translucent skin, easy bruising, craniosynostosis, cleft palate, Chiari type I malformation of the brain, learning disability, congenital heart disease (patent ductus arteriosus, atrial septal defect, bicommissural aortic valve), and clubfoot deformity.

There is significant overlap between Loeys-Dietz aortic aneurysm syndrome and the Shprintzen-Goldberg syndrome that includes craniosynostosis, hypertelorism, arched palate, learning disability, bone overgrowth, pectus deformity, and scoliosis. Vascular disease is described as a rare manifestation of Shprintzen-Goldberg syndrome. Furlong and colleagues described one patient with Marfanoid habitus, craniosynostosis, hypertelorism, normal intelligence, and aortic root aneurysm with dissection. Relative to other aortic diseases, features unique to Loeys-Dietz aortic aneurysm syndrome include cleft palate, bifid uvula, arterial tortuosity, and diffuse aneurysms and dissections. In view of the aggressive behaviour of vascular disease, the distinction of Loeys-Dietz aortic aneurysm syndrome is essential to individualise management.

**Molecular genetics and pathophysiology**

Both skin and aorta from patients with Marfan’s syndrome show decreased elastin content and fragmentation of elastic fibres. However, phenotypic manifestations in tissues without elastin, such as ciliary zonules and bone, helped to exclude the elastin gene as the primary site of mutations causing the disorder. Additionally, linkage analysis mapped the Marfan’s syndrome locus to 15q21.1, distant from the chromosomal locus of elastin at 15q21.2. It encodes a 350 kDa glycoprotein, which is highly conserved among different species. Mutations occur within the 47 tandemly repeated epidermal growth factor-like domains, many disrupting one of the six predictably spaced cysteine residues that interact via disulfide linkage to determine domain folding or residues that affect calcium binding to fibrillin 1. Such perturbations lead to enhanced cleavage and proteolytic degradation.

Several findings suggested that dominance in Marfan’s syndrome is caused by adverse activity of the mutant protein on the deposition, stability, or function of the normal protein encoded by the normal copy of the FBN1 gene—a so-called dominant-negative effect. First, this was a well-established framework for other connective tissue disorders, such as type I collagen defects, causing osteogenesis imperfecta. Second, fibrillin-1 monomers aggregate to form complex structures called microfibrils, providing an opportunity for interference. Third, tissues from affected patients often have a dramatic paucity of fibrillin-1, far less than 50% that would be predicted from simple loss of contribution of one allele. Fourth, some patients with mutations associated with low production of mutant protein have a mild, subdiagnostic variant of the disorder. However, this association between abundance of mutant transcript and severity of disease was refuted by the report of patients with classic severe disease and very low levels of mutant transcripts containing a premature termination codon.

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Evidence suggests that half-normal production of normal protein (haploinsufficiency), rather than the production of mutant protein, could be critical to reach the threshold loss of fibrillin-1 function needed for clinical expression of the disorder. First, transgenic overexpression of mutant fibrillin 1, in the context of two normal *Fbn1* alleles, was shown to be insufficient to cause the vascular changes of Marfan’s syndrome seen in mice heterozygous for a comparable missense mutation (C1039G). Second, mice heterozygous for a null *Fbn1* allele showed similar alterations in aortic wall architecture as C1039G/+ mice (Ramirez F, Hospital for Special Surgery, New York, USA, personal communication). Finally, the transgenic addition of a wild-type allele to C1039G heterozygous mice rescued the aortic phenotype. Thus, a normal complement of fibrillin 1 might be needed to initiate productive microfibrillar assembly. Although the haploinsufficiency-imposed reduction in microfibrillar abundance may or may not be sufficient in isolation to lead to classic Marfan’s syndrome, it seems critical to the context within which a dominant negative effect can achieve clinical significance. These data suggest that boosting fibrillin-1 expression could be a productive therapeutic strategy and implicate genetic modifiers in the modulation of disease severity. For example, Hutchinson and colleagues have described a family with Marfan’s syndrome in whom phenotypic severity correlated inversely with the expression level of fibrillin 1 by the allele without a mutation.

Murine models of Marfan’s syndrome have provided the opportunity to assess some of the earliest pathogenetic abnormalities resulting in aneurysm and dissection. They have shown that fibrillin-1 microfibrils are not needed to assemble an elastic fibre, as previously inferred. This model had suggested that infants with the disorder are born without assembled elastic fibres—in essence an obligate structural predisposition for tissue failure later in life. This notion boded poorly for the development of productive treatment strategies. Rather, microfibrils are needed to maintain elastic fibres during postnatal life. In the absence of proper connections between elastic fibres and vascular smooth-muscle cells mediated by fibrillin 1, the cells adopt an abnormal synthetic repertoire that includes matrix-degrading enzymes, such as matrix metalloproteinases 2 and 9. Subsequent events include elastic fibre calcification, vascular wall inflammation, intimal hyperplasia due to migration and proliferation of vascular smooth-muscle cells, and structural collapse of the vessel wall. Similar vascular pathology has been identified in large muscular arteries from seven patients diagnosed with the disorder.

In addition to the obvious structural role of fibrillin-1-rich microfibrils, it is becoming increasingly clear that these microfibrils also have a critical role in regulation of cytokines, molecules that affect tissue development and homoeostasis via regulation of cellular performance—proliferation, migration, synthetic repertoire, and death. Fibrillin 1 shares a high degree of homology with the latent transforming growth factor β (TGFβ) binding proteins. The TGFβ cytokines are secreted as large latent complexes, consisting of TGFβ, latency-associated peptide, and one of three latent TGFβ-binding proteins. On secretion, the large latent complex is sequestered by the extracellular matrix. TGFβ signalling requires release of the mature cytokine, interaction with cell-surface receptors, and initiation of a downstream signalling cascade. The homology between the fibrillins and latent TGFβ-binding proteins prompted the hypothesis that extracellular microfibrils might participate in the regulation of TGFβ activation. This proposal was advanced through study of distal airspace enlargement in fibrillin-1-deficient mice. Similar pulmonary changes occur in Marfan’s syndrome and have been attributed to the effects of physiological stress acting on a structurally predisposed tissue—ie, classic destructive emphysema. Subsequent analyses in fibrillin-1-deficient mice showed that airspace enlargement manifested a primary failure of distal alveolar septation during lung morphogenesis in the absence of lung inflammation or destruction. These animals showed a substantial increase in both free TGFβ and TGFβ-signalling at a developmental stage relevant to alveolar septation. Furthermore, administration of TGFβ-neutralising antibody was sufficient to rescue septation in fibrillin 1-deficient mice.

It seemed plausible that this pathogenetic mechanism was relevant to other manifestations of Marfan’s syndrome—especially those that were difficult to reconcile on the basis of pathogenetic models that singularly invoke structural failure of the tissues. Likely candidates included bone overgrowth, myxomatous changes of the mitral valve, craniofacial abnormalities, and muscle or fat hypoplasia. Increased local activity of TGFβ has recently been shown to be responsible for the myxomatous cardiac valve disease in fibrillin-1-deficient mice. As demonstrated in the lungs, atrioventricular valve lengthening, thickening, and dysfunction is prevented when TGFβ-neutralising antibody is given to mouse models of Marfan’s syndrome. Increased TGFβ activity has also been recorded in other tissues, including the vasculature and the dura.

Further evidence for the broad relevance of dysregulated TGFβ signalling came with the identification of the genetic defect in a subset of patients previously characterised as having classic Marfan’s syndrome and all individuals with Loeys-Dietz aortic aneurysm syndrome. First, Mizuguchi and colleagues described that mutations in the gene encoding the type II TGFβ receptor (*TGFBR2*) exactly recapitulate the classic Marfan phenotype. While the presentation of these individuals is atypical for Marfan’s syndrome (eg, absence of lens dislocation, normal height, lack of dolichostenomelia, high rate of non-penetrance), the researchers reported heterozygous loss-of-function mutations in patients with many features of the disorder. It was difficult, however, to reconcile the inferred mechanism of decreased TGFβ
signalling with the increase seen in mouse models of Marfan’s syndrome.\(^{44-46}\) Loey and colleagues\(^{47}\) went on to show that all patients with Loey-Dietz aortic aneurysm syndrome were heterozygous for loss-of-function mutations in either of the genes encoding the type I or type II TGFβ receptor (TGFBR1 or TGFBR2). Furthermore, heterozygosity for these mutations did not impair responsiveness to TGFβ in cultured cells from patients and led paradoxically to enhanced signalling in the aortic wall, as seen in both mice and people deficient for fibrillin 1.

Clinical genetic testing

The role of genetic testing in establishing a diagnosis remains limited. To date, over 500 mutations resulting in the disorder have been catalogued in an international database.\(^{68}\) Over 90% are private mutations unique to an individual or family. Even within families where all affected individuals share the same mutation, phenotypic variation is prominent. Thus, it is difficult to derive significant genotype–phenotype correlations.\(^{69,70}\) Due to the large size of \(FBN1\) (65 exons), the cost of routine sequence analysis of all exons could be prohibitive. By contrast, haplotype segregation analysis may be done more easily, and could provide the desired information regarding who in the family has inherited the predisposing copy of the \(FBN1\) gene.\(^{70-72}\) However, around 25% of patients with the disorder have a de-novo mutation, limiting the feasibility of haplotype segregation analysis.

As previously mentioned, many other conditions can be caused by mutations in \(FBN1\), including MASS phenotype, Shprintzen-Goldberg syndrome, mitral valve prolapse syndrome, familial ectopia lentis, isolated Marfanoid habitus, and Weill Marchesani syndrome.\(^{44,47,72-74}\) However, mutations in \(ADAMTS10\) have also been described as causing Weill Marchesani syndrome, and many patients with Shprintzen-Goldberg syndrome have not been shown to have a mutation in \(FBN1\), thus exhibiting locus heterogeneity for these disorders.\(^{75}\) Phenotype or severity are often difficult or impossible to predict simply on the basis of the nature or location of a mutation in \(FBN1\). Furthermore, best estimates suggest that about 10% of mutations that cause classic Marfan’s syndrome are missed by conventional screening methods.\(^{7}\) In essence, gene testing lacks both sensitivity and specificity. Under these circumstances, mutational or linkage analysis seems to be best used to identify who in a family has inherited the predisposition for the phenotype seen in that family. This notion extends to the use of molecular methods for prenatal diagnosis.\(^{75}\) At the present time, diagnosis is largely based on clinical assessment.

Management

The diagnosis of Marfan’s syndrome requires a multidisciplinary assessment that generally includes a geneticist for general assessment and anthropomorphic measurements, an ophthalmologist for a slit-lamp examination, and a cardiologist for cardiovascular imaging.

After establishing a clinical diagnosis of Marfan’s syndrome, routine monitoring of aortic growth is essential to decrease the risk of aortic dissection. Yearly assessment by transthoracic echocardiography allows serial measurements of the proximal aorta, including the size of the aortic root at greatest dimension, the sinotubular junction, and the ascending aorta. Both the absolute size of the aorta in greatest diameter and the rate of growth of the aorta are monitored to establish the appropriate time to intervene surgically. At times, technical difficulties limit views of the proximal aorta, especially in patients with significant pectus deformity of the sternum. In such cases, CT angiography or MRI can provide accurate images of the proximal aorta, and should be used on at least a yearly basis. More frequent imaging is indicated if the aortic size is approaching a threshold that would initiate surgery or if rapid growth is observed.

While ectopia lentis is a major criterion in establishing a clinical diagnosis of Marfan’s syndrome, these patients are also at increased risk of retinal detachment, glaucoma, and cataracts. Myopia is reported in most patients, and may predispose to amblyopia in early childhood.\(^{76}\) Thus, patients with the disorder should undergo comprehensive yearly assessments by an ophthalmologist, ideally with training or expertise in this disorder. Lens dislocation can generally be managed with eyeglasses or contact lenses; occasionally, surgical aphakia is needed to achieve adequate vision.\(^{77}\) This can be followed by artificial lens implantation once growth is complete.

Severe orthopaedic issues will need involvement of a skilled orthopaedist. Bracing is generally inadequate to manage severe and progressive scoliosis, which often requires surgical stabilisation.\(^{78}\) This situation must be monitored closely during growth. Pectus deformity is largely a cosmetic issue, although often significant enough to warrant surgery.\(^{79}\) Although many patients with pectus deformity have restrictive lung disease, some studies have shown little or no effect on lung capacity after repair.\(^{80,81}\) Others have suggested that improvement in symptoms could be due to increased cardiac function.\(^{82}\) A recent report of 45 patients who underwent minimally invasive surgical intervention (Nuss procedure) and bar removal for pectus excavatum described a significant increase in forced vital capacity after surgery, with improvement most prominent in patients aged older than 11 years.\(^{83}\) We have never observed a case where pectus excavatum was proven to critically impair cardiovascular function. Good cosmetic results for pectus excavatum can generally be achieved with the minimally invasive Nuss procedure.\(^{84}\) If done too early, however, continued rib growth will lead to recurrent deformity.\(^{85}\)

Medical treatment to prevent or delay aortic aneurysm

β-adrenergic receptor blockade to delay or prevent aortic aneurysm and dissection is currently regarded as the standard of care for patients with the disorder. This strategy was first proposed in 1971 by Halpern and
colleagues. The rationale for this treatment strategy is primarily to decrease proximal aortic shear stress, or change in pressure over time (dP/dT). β blockers are probably beneficial both through negative inotropic and negative chronotropic effects. Although small, most published studies have shown benefit of treatment with β blockers in Marfan’s syndrome, including in children. The only randomised trial assessing the effect of β blockade in patients with the disorder was published in 1994. Herein they describe 70 patients with the syndrome, and treated 32 with propranolol. Using an open-label protocol, treated patients received a dose of propranolol targeted for a heart rate below 100 beats per minute during exercise or resulting in a 30% rise in the systolic time interval (corrected for the heart rate). Patients were analysed for aortic growth, with change in body size accounted for in an aortic ratio by dividing the measured aortic diameter by the diameter predicted from the patient’s height, weight, and age. Fewer patients treated with propranolol reached a primary clinical endpoint of aortic regurgitation, aortic dissection, cardiovascular surgery, congestive heart failure, and death (five in treatment group, nine in control group). Furthermore, the normalised rate of aortic dilatation was lower in the propranolol group than in the control group (0·023 vs 0·084 peryear, p<0·001).

A recent trial compared 58 patients treated with β blocker (propranolol or atenolol) with those treated with an ACE-inhibitor, enalapril. Patients were not randomised and the dose of β blocker was not titrated to physiological response. Aortic change was not reported with the dose of β blocker needed to achieve a therapeutic response is highly variable and is not predicted by bodyweight or age. Failure to titrate the dose of medication represents a major limitation of this study.

Around 10–20% of patients with Marfan’s syndrome are intolerant to β blockers. Reasons include asthma, depression, and fatigue. Among such individuals, a trial of verapamil may be justified on the basis of a report showing that such treatment can slow the rate of aortic growth, although this trial was non-randomised and uncontrolled. Among the trials that have assessed use of pharmacological therapy for aortic aneurysm in Marfan’s syndrome, it is important to point out that aortic growth is not stopped or reversed, but typically is slowed in response to treatment. Thus, in the setting of aortic enlargement, despite treatment with pharmacological agents to slow the rate of aortic growth, one should remain vigilant for further aortic enlargement with at least yearly measures of aortic dimensions.

Surgery for aortic aneurysm

The rate of acute aortic dissection is directly proportional to the maximum diameter of the aorta. Elective surgery to repair the aortic root is recommended when the maximum aortic diameter reaches 5 cm. Additional considerations include the rate of aortic growth and family history of aortic dissection at a size less than 5 cm. Earlier surgical intervention is recommended for individuals with an increase in aortic diameter exceeding 1 cm per year.

Composite surgical replacement of the aortic root and valve was pioneered by Bentall and De Bono in 1968. Modifications of this technique have evolved with various methods of coronary reimplantation. Gott and colleagues reported outcomes for 675 patients with the disorder who underwent aortic root replacement surgery at ten experienced surgical centres (seven in North America and three in Europe). Mortality for elective surgery was 1·5% compared with 2·6% for urgent surgery (taking place within 7 days after surgical consultation). Mortality was 11·7% among patients who underwent emergency surgery within 24 h after surgical consultation.

Due to risks of thromboembolism and the lifetime requirement of warfarin anticoagulation in the setting of a mechanical prosthetic aortic valve, recent surgical efforts, pioneered by David and colleagues, have attempted to maintain the native aortic valve among eligible patients with the disorder. To date, no randomised clinical trials of valve replacement versus valve-sparing aortic root surgery have been undertaken, and long-term data on the outcomes with valve-sparing surgery are not yet available. Nevertheless, the short-term data are encouraging, with an extremely low rate of operative mortality (equivalent to that seen with composite graft repair). The valve-sparing procedure has seen a tremendous evolution, largely focused on preserving function of the native aortic valve. Initial attempts to resuspend the valve into the Dacron graft were complicated by damage to the leaflets on maximum excursion. A remodelling procedure that involved sewing the graft to the top of the aortic annulus showed an unacceptable rate of valve splaying with subsequent dysfunction and reoperation. Most recently, a modified resuspension procedure is being employed with crafting of artificial sinuses above the valve. This procedure has shown excellent short-term results and is now the preferred treatment in all eligible patients who present for surgical intervention. The valve-sparing approach is especially attractive for young women who anticipate pregnancy.

Lifestyle modifications

Due primarily to the risk of acute aortic dissection, patients should be counselled not to engage in contact sports, competitive athletics, or isometric exercise. Sudden death in the setting of unrecognised Marfan’s syndrome...
among several high profile athletes has helped to emphasise the importance of early recognition and activity limitations for this disorder. Most patients, however, should be encouraged to remain active with aerobic activities performed in moderation. This will promote skeletal, cardiovascular, and psychosocial health in the long term. More detailed descriptions of specific activities appropriate for patients with the disorder are reviewed elsewhere.\textsuperscript{101,102}

The issue of pregnancy in Marfan’s syndrome draws attention to concerns about the risk of transmission of this disease. Genetic counselling should be undertaken and one should inform prospective parents of the 50% risk of offspring of an affected individual inheriting a genetic predisposition to the disorder. Before current standards for surgical repair of the aortic root, early experience indicated a high risk of aortic dissection during pregnancy.\textsuperscript{103} More recent analyses have indicated that the risk of aortic enlargement is related to the size of the aortic root before pregnancy, and in the setting of an aortic root size less than 4 cm, the risk of pregnancy is low.\textsuperscript{104,105} Finally, one should discuss the issue of anticoagulation and pregnancy with women who have Marfan’s syndrome. Systemic anticoagulation with warfarin during pregnancy in the setting of a mechanical prosthetic aortic valve is associated with increased risk of fetal demise and warfarin embryopathy.\textsuperscript{106} Low molecular weight heparin, given subcutaneously during pregnancy, seems safer than warfarin as it does not cross the placenta, although it is expensive, can result in heparin-induced thrombocytopenia, and might not prevent prosthetic valve thrombosis.\textsuperscript{107–109} Pregnancy in patients with the disorder should precede significant aortic enlargement or follow a valve-sparing procedure. Close monitoring by an obstetrician and a cardiologist remains mandatory.

**New developments**

The prevailing view had been that people with Marfan’s syndrome are born with a structural weakness of the tissues that imposes an obligate risk of tissue failure later in life. Strategies to strengthen the connective tissue throughout the body were difficult to conceive much less implement. The new realisation that Marfan’s syndrome manifests postnatally acquired tissue pathology, and that this may largely indicate a failed regulatory (as opposed to structural) role of the extracellular matrix, presents a far more optimistic outlook. With the possible exception of ectopia lentis, most manifestations of Marfan’s syndrome seem to implicate dysregulation of TGF\(\beta\) activity and signalling. Furthermore, emerging data from mouse models suggest that these phenotypes can be productively modified in the postnatal period through manipulation of TGF\(\beta\) activity, plausibly including the use of drugs that are already in development or in use for other indications (unpublished data). These treatments have the potential to not only slow or prevent progression of aortic root aneurysms, but also to attenuate the multisystem pathogenesis of disease.

**Conclusions**

Progress in the past century has led to an improved understanding of the cause, pathophysiology, and treatment of Marfan’s syndrome. As our knowledge of the consequences of fibrillin-1 deficiency develops, we anticipate that treatment will continue to advance, allowing improved length and quality of life for patients with the disorder. Indeed, Marfan’s syndrome represents a paradigm success that foreshadows future challenges and opportunities that will derive from the human genome project and increased sophistication in the elucidation of the genetic basis of disease. As we modify the natural history of disease, it will be important to observe and respond to the new reality of ageing with selected genetic disorders, including Marfan’s syndrome, which shows an already apparent predisposition for accelerated joint disease and a predisposition for primary involvement of the descending aorta in older individuals.

**Conflict of interest statement**

We declare that we have no conflict of interest.

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**References**


Seminar


