Genotype-phenotype analysis of F-helix mutations at the kinase domain of TGFBR2, including a type 2 Marfan syndrome familial study

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Purpose: Transforming growth factor beta receptor II (TGFBR2) gene mutations are associated with Marfan syndrome; however, the relationship between the mutations and clinical phenotypes are not clear.

Methods: Genomic DNA from peripheral blood leukocytes of a Chinese proband with Marfan syndrome, five of the proband’s relatives, and 100 unrelated Chinese control subjects were isolated and screened for fibrillin-1 (FBN1) and TGFBR2 gene mutations by direct sequencing, and a genotype-phenotype study was performed following a review of the literature on TGFBR2 mutations in the search area. Also, the structure of TGFBR2 protein before and after gene mutation was analyzed.

Results: The results identified a novel missense TGFBR2 mutation p.V453E (c.1358T>A) in the proband and two relatives that was located in the F-helix in the kinase domain of TGFBR2. No such genetic change was observed in the unrelated controls. No FBN1 mutation was detected in any of the subjects. Genotype-phenotype analyses indicated that F-helix mutations are related to type 2 Marfan syndrome and Loeys-Dietz syndrome, and these mutations can lead to severe cardiovascular (93.8%) and skeletal (81.3%) lesions and minor ocular lesions (25%). Losartan treatment can slow-down the progression of aortic lesions.

Conclusions: The findings extend the mutation spectrum of Marfan syndrome, and that mutations at the F-helix in the kinase domain of TGFBR2 may be associated with the development of severe cardiovascular and skeletal lesions and minor ocular lesions. These findings have implications for genetic testing, diagnosis, and treatment in individuals with transforming growth factor beta (TGF-β) signaling-related disorders.

Marfan syndrome (MFS) is an autosomal dominant extracellular connective tissue disorder that is clinically diagnosed according to the Ghent criteria [1]. Classic MFS presents as abnormal features in skeletal, ocular, and cardiovascular systems. The skeletal abnormalities of affected individuals typically include tall stature, long slender limbs (dolichostenomelia), scoliosis, arachnodactyly, and pectus excavatum or carinatum. About 80% of MFS sufferers have ectopia lentis, which is almost always bilateral. Progressive dilatation of the aorta is the main cardiovascular irregularity of MFS individuals, and acute aortic dissection originating from dilatation of the ascending aorta is the leading cause of premature death in untreated MFS individuals [2]. Other manifestations, including spontaneous pneumothorax, apical blebs, striae atrophicae, and lumbosacral dural ectasia, can also be observed in MFS patients [3,4].

MFS is most commonly caused by fibrillin-1 (FBN1) gene mutations at chromosome 15q21.1, with the mutation types being point mutations, insertions, large and small deletions, and splice mutations. The mutations are spread throughout almost the entire FBN1 gene without obvious predilection for any given region. Recent studies show that genetic heterogeneity exists in MFS individuals, and transforming growth factor beta receptor II (TGFBR2) gene mutations have been identified in a subset of patients with MFS – which was termed as type 2 Marfan syndrome (MFS2) [5]. Mutations in TGFBR2 have also recently been identified in patients with Loeys-Dietz syndrome, and familial thoracic aortic aneurysms [6,7].

Transforming growth factor beta superfamily signaling is of great importance in the maintenance of the extracellular matrix and tissue homeostasis [8]. One important member of this gene superfamily is TGFBR2, which encodes for the trans-membrane kinase receptor involved in signal transduction of the transforming growth factor beta (TGF-β) family of ligands [9]. This gene is composed of seven exons encoding 567 amino acids to form an NH₂-terminal ligand binding domain, a trans-membrane region, and a constitutively active COOH-terminal serine/threonine kinase.
domain [10]. These structures are critical for TGF-β signaling, which plays a key role in normal extracellular connective tissue functions. Several TGFBR2 mutations have been reported in association with vascular abnormalities [5, 11-13], with most being missense substitutions or nonsense mutations in the penultimate or final exons and predicted to disrupt the kinase domain. At present no strict genotype-phenotype correlations between TGFBR2 mutations and MFS2 have been established, and great clinical variability of cardiovascular, skeletal, and ocular system manifestations exists among TGFBR2 mutation carriers.

This study describes the genetic and clinical features of a Chinese family, the proband of which displays the MFS2 phenotype genetically caused by mutation at the TGFBR2 locus. A literature review was also undertaken with the aim of exploring the relationship between mutations at the region in this study and clinical phenotypes.

METHODS

Study subjects: This study was conducted in accordance with the Declaration of Helsinki and approved by the Fuwai Cardiovascular Center Ethics Committee.

An MFS patient and family and 100 unrelated controls participated in this study. All subjects were of Han Chinese origin. Written informed consent was obtained from all subjects (or their parents if the subject was less than 18 years of age).

The pedigree of the enrolled family is presented in Figure 1. The proband of the enrolled family was a 12-year-old female (III: 2, in Figure 1), who presented for examination of her cardiovascular system because previous X-ray results indicated aortic root dilation. Other family members (II: 1, II: 2, III: 1, III: 2, III: 3 in Figure 1) of the proband were recruited and underwent multidisciplinary clinical and noninvasive instrumental studies. Our assessments covered all the diagnostic criteria for MFS [5]. A physical examination and echocardiography were performed on the unrelated control subjects to exclude MFS and other congenital abnormalities. The aortic root diameter was evaluated by echocardiography.

Genetic analysis: Genomic DNA was isolated from peripheral venous blood leukocytes of all subjects using a commercially available kit (Qiagen Inc., Germantown, MD). Following amplification by polymerase chain reaction (35 cycles), mutation screening (which allowed for the scanning of 65 FBN1 exons and flanking regions, including the splice sites up to the branching regions [primers used for bidirectional sequencing are presented in Appendix 1]) was performed using an ABI3730xl Sequencer (Applied Biosystems, Foster City, CA). The primers used for bidirectional sequencing of TGFBR2 are presented in Table 1. TGFBR2 mutation screening of all seven encoding exons was performed after finding that all the exon-intron boundaries of FBN1 were negative. Mutation numbering refers to the TGFBR2 cDNA GenBank reference sequence: NM_003242.5, with the A of the ATG translation initiation codon as nucleotide +1.

In silico prediction of the effects caused by TGFBR2 mutations: The human TGFBR2 reference sequence accession NP_003233.4 was used to perform in silico prediction, and the SWISS-MODEL tool was used [14,15]. Mutagenesis and visualization were performed to determine the possible effects of the mutation on the structure of their corresponding modules during in silico 3D modeling analysis.

Genotype-phenotype investigation: A literature search of TGFBR2 mutations and related clinical manifestations was performed and mutations were classified according to their locations in the gene and functional domains. Correlations between mutations at the region where the mutation in this study was found and their clinical features were analyzed.

Statistical analysis: Statistical analyses were performed using SPSS software version 15.0 (SPSS Inc., Chicago, IL). Phenotypes detected which involved separate organ systems were compared with the Pearson χ² test; or using a Fisher’s exact test for small samples. All the reported p values are two-sided, and a p value of <0.05 was considered significant.

RESULTS

Clinical features of the patients: The clinical manifestations of all family members are summarized in Table 2. The
The proband was confirmed as an MFS patient [5], and a familial history suggestive of MFS was reported. A chest computed tomography of the proband revealed that she had an ascending aortic aneurysm, and echocardiography showed an aortic root diameter of 50 mm. The patient underwent a ROSS procedure (graft replacement of the ascending aorta) to treat her aortic aneurysm. A characteristic habitus with tall stature, long slender limbs (dolichostenomelia) was observed. She was 168 cm tall at 12 years of age, and her arm span to height ratio was 1.15. Pectus carinatum, scoliosis (25°), dolichostenomelia, arachnodactyly, and positive wrist and thumb signs were also noted; the findings indicated that her skeletal system was involved. Striae atrophicae was also found.

The proband’s mother had a characteristic habitus with tall stature and long slender limbs (arm span to height ratio was 1.09); dolichostenomelia, arachnodactyly, and wrist and thumb signs were noted; and dilatation of the ascending aorta (aortic root diameter was 45 mm) and aortic regurgitation were observed.

No ocular lesions were found in either the proband or her mother.

Further findings positive for MFS in other family members are presented in Table 2. No positive findings were detected in the control subjects.

Losartan (25 mg per day) was used as previously reported [16] in the proband after surgery to prevent aortic enlargement and associated cardiovascular pathologic changes, and no recurrent aortic lesion was found after losartan treatment. Losartan (50 mg per day) was also used in the other individuals (II: 1 and II: 2) who had TGFBR2 mutations. No obvious progression of aortic root dilation was detected after losartan therapy.

Mutation analysis: A novel missense mutation (p.V453E (c.1358T>A) in exon 5) was identified in TGFBR2 in the proband (III: 2), the proband’s mother (II: 2), and the proband’s uncle (II: 1; see Figure 2). This alteration was not detected in the 100 unrelated chromosomes from ethnically matched Han Chinese controls.

The mutation identified affects a highly conserved nucleotide and is predicted as damaging protein production or affecting protein function. The mutation located in the F-helix in the kinase domain of TGFBR2, which was formed by the 440–457 residues encoded by the 1318–1370 base pairs. No FBN1 mutation was found in any of the subjects of the study.

Protein structure changes due to mutations: The missense mutation involves a highly conserved valine residue that is

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### Table 1. Primers used for the amplification of TGFBR2.

| Exon | Forward primer (5'→3') | Reverse primer (5'→3') | Melting temperature (°C) | Product length (bp) |
|------|------------------------|------------------------|-------------------------|---------------------|
| Exon1 | AGCTGTTGGCGGAGGAGTTCCTCGTTT | GCGGCTTCAAGATAACCAACTTCTCAAC | 58 | 765 |
| Exon2 | TTGACAAAAAGCAAATGGCTACTC | GGAAGGGGAATGGGAACAG | 58 | 539 |
| Exon3 | AAAACAGAAAGAGTAAAGAAAGCATAGG | TGATGAGAAAGGCCCCAACAACCTT | 58 | 574 |
| Exon4 | AGCAGGGGATGACGCAACAGA | GAAGGATTTGAAGTGGAGAGGAA | 58 | 1226 |
| Exon5 | TTTTATGCTTCTTTCAGGGGTTTTC | CCAATAGTCTGGGATGGTTTGA | 58 | 507 |
| Exon6 | AAAACCTAAGCTCCGTGAC | TTAACAGGCCATAGAAC | 58 | 545 |
| Exon7 | GTTTGAGGTGTAGTGTGTA | CAATGTCAAGGGCATAGATT | 58 | 695 |

Sequences are given in the 5’→3’ direction. TGFBR2: transforming growth factor beta receptor II gene.

### Table 2. Clinical manifestation of the proband and her family members.

| Parameters | III:2 | II:2 | II:1 | I:1 |
|------------|-------|------|------|-----|
| Age        | 12    | 35   | 39   | Death at 43 |
| Sex        | Female | Female | Male | Male |
| Cardiovascular system | Aortic root dilation (aortic root diameter=50 mm); Aortic regurgitation; Mitral valve regurgitation; Dilatation of the main pulmonary artery. | Aortic root dilation (aortic root diameter=45 mm); Aortic regurgitation. | Aortic root dilation (aortic root diameter=48 mm) | Aortic root dilation; Dissection of the ascending aorta |
| Skeletal system | Arm span to height ratio=1.15; Wrist and thumb signs; Scoliosis of=25°; Pectus excavatum; Dolichostenomelia; Arachnodactyly. | Arm span to height ratio=1.09; Dolichostenomelia; Arachnodactyly; Wrist and thumb signs. | Arm span to height ratio=1.05; Dolichostenomelia; Arachnodactyly; Joint hypermobility. | ? |
| Ocular system | Nil | Nil | Nil | ? |
| Pulmonary system | Nil | Nil | Nil | ? |
| Skin and integument | Striae atrophicae | Nil | Nil | ? |
| Gicht nosology | Fulfilled | Not fulfilled | Not fulfilled | ? |
| TGFBR2 mutation | c.1358T>A, p.V453E | c.1358T>A, p.V453E | c.1358T>A, p.V453E | ? |

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located in the F-helix in the kinase domain of TGFBR2. The F-helix in the kinase domain plays an important role in maintaining the function of TGFBR2. The mutant residue is predicted to induce steric clashes with its surroundings. In silico modeling analysis showed that four additional hydrogen bonds were formed; and the conformation changed after the valine residue was replaced by glutamic acid (Figure 3E,F). This structural alteration may perturb the crucial F-helix structural components of the protein kinase domain. Previously reported mutations at the F-helix can alter the protein kinase domain of TGFBR2 to impair the TGFBR2 pathway (Figure 3C,D), and can therefore lead to MFS or MFS-like phenotypes [17].

Genotype-phenotype relationships: The results of the literature review of the F-helix of the kinase domain of TGFBR2 are shown in Table 3. Nine kinds of mutations (all of which are point mutations, except for one deletion change) were found in this domain, as reported in 16 studies [5,12, 13,18-25]. Almost all mutation carriers (15/16, 93.8%) displayed major cardiovascular system abnormalities. Four cases (4/16, 25%) were reported as showing minor ocular lesions, and no ectopia lentis was reported. Skeletal system abnormalities were reported in 81.3% (13/16) of cases, and 31.3% (5/16) had major skeletal system lesions. The cardiovascular and skeletal system had significantly higher involvement rates when compared to the ocular system (p<0.01 of both compares), but no significant difference in involvement rates was found between the cardiovascular system and the skeletal system (93.8% versus 81.3%, p=0.60).

MFS2 was reported in 37.5% (6/16 cases) of all the patients, but only 12.5% (2/16 cases) fulfilled the Ghent nosology criteria. The remaining 62.5% are Loeys-Dietz Syndrome individuals. The occurrence rates of MFS2 and Loeys-Dietz Syndrome caused by mutations in this region were comparable (p=0.29). Mutations in this region tend to attack the carriers before adulthood, and a significant higher illness detection rate was found in patients <16 years old (p=0.012).

DISCUSSION
Marfan syndrome is an autosomal dominant connective tissue disorder involving multiple organ systems. Patients with
Table 3. Phenotypes caused by genetic changes at F-helix in the kinase domain of TGFBR2.

| Reference | Nucleotide | Protein | Diagnosis | CS | OS | SS | S | LDS symptoms | Age (years) | Sex | Ghent nosology |
|-----------|------------|---------|-----------|----|----|----|---|---------------|-------------|-----|----------------|
| [12]      | c.1318G>A  | p.E440K | LDS       | M  |  - | m  |   |  Involve      |  9          | Male| Not fulfilled  |
| [22]      | c.1322C>T  | p.S441F | MFS2      | M  |  - | m  |   | Not involve   | 14         | Male| Fulfilled      |
| [19]      | c.1324T>C  | p.F442L | LDS       | m  |  - | m  |   |  Involve      |  36         | Female| Not fulfilled  |
| [21]      | c.1336G>A  | p.D446H | LDS       | m  |  - | m  |   |  Involve      |  1          | Male| Not fulfilled  |
| [23]      | c.1336G>C  | p.D446H | LDS       | m  |  - | m  |   |  Involve      |  7          | Female| Not fulfilled  |
| [13]      | c.1336G>A  | p.D446H | LDS       | m  |  - | m  |   |  Involve      |  1          | Male| Not fulfilled  |
| [18]      | c.1336G>C  | p.D446N | MFS2      | M  |  - | m  |   | Not involve   |  4          | Female| Not fulfilled  |
| [23]      | c.1336G>A  | p.D446N | LDS       | M  |  - | m  |   |  Involve      | 26         | Female| Not fulfilled  |
| [25]      | c.1342T>A  | p.Y448N | LDS       | ?  |  - | ?  |   |  Involve      |  ?          | ?   | Not fulfilled  |
| [5]       | c.1346C>T  | p.S449F | MFS2      | M  |  - | M  |   | Not involve   | 12         | Female| Not fulfilled  |
| [25]      | c.1346C>T  | p.S449F | LDS       | M  |  - | M  |   |  Involve      | 13         | Female| Not fulfilled  |
| [23]      | c.1351_1356del | p.A451_L452del | LDS       | M  |  - | M  |   |  Involve      | 1          | Male| Not fulfilled  |
| This study| c.1358T>A  | p.V453E | MFS2      | M  |  - | M  |   | Not involve   | 12         | Female| Fulfilled      |
| [24]      | c.1370T>A  | p.M457K | MFS2      | M  |  - | m  |   | Not involve   | 9          | Female| Not fulfilled  |
| [20]      | c.1370T>A  | p.M457K | LDS       | M  |  - | M  |   |  Involve      | 9          | Female| Not fulfilled  |
| [23]      | c.1351_1356del | p.A451_L452del | LDS       | M  |  - | M  |   |  Involve      | 1          | Male| Not fulfilled  |

M: major criterion fulfilled, m: minor criterion fulfilled, -: organ system not involved, ?: not mentioned. CS: Cardiac system, OS: Ocular system, SS: Skeletal system, S: Skin and integument. LDS: Loeys Dietz Syndrome, MFS: Marfan Syndrome. Nucleotide numbering reflects cDNA numbering with +1 corresponding to the A of the ATG translation initiation codon in the reference sequences (TGFBR2: NM_003242.5). The initiation codon is codon 1.
young-onset cardiovascular lesions failing to meet the Ghent criteria have been classified as Marfan-like connective tissue disorders. After the first report by Boileau et al. [26] in 1993, a Marfan-like phenotype that was not linked to the FBN1 mutation was subsequently designated Marfan syndrome type 2 (MFS2). Diagnosis of MFS2 consists of severe cardiovascular findings, which can include the sudden death of affected people at a young age owing to a thoracic aortic dissection, and typical MFS skeletal features, but ocular findings were rare [27]. Genetic analysis of MFS2 patients by Mizuguchi et al. [5] identified a de novo chromosomal rearrangement involving chromosome 3p24.1 that can disrupt TGFBR2 function to result in protein truncation due to abnormal splicing. Further studies demonstrate that the MFS2 phenotype is caused by mutations in TGFBR2.

TGF-β family cytokines form a complex signaling pathway and control several cellular processes, including cell proliferation and differentiation, apoptosis, and extracellular matrix formation [28]. Incorrect TGF-β signaling may lead to a decreased and disordered incorporation of fibrillin into the connective tissue matrix, and this abnormality in turn results in human MFS phenotypes [29]. Two types of trans-membrane receptors, type I (TbRI) and type II (TbRII), transduce the signals of TGF-β into the cell. TbRI and TbRII are encoded by TGFBR1 and TGFBR2, respectively [30,31]. The identification of TGFBR2 mutations in patients with MFS2 provided the first direct link between a human connective tissue disorder and abnormal TGF-β signaling. TGFBR2 mutations may cause a loss in function of TbRII and lead to MFS2, Loeys-Dietz Syndrome [32] and Type 2 Thoracic aortic aneurysms and dissections.

In the present study we directly sequenced the entire coding region of FBN1 and TGFBR2, and found a previously unreported TGFBR2 mutation, p.V453E (c.1358T>A). V453 is a highly conserved residue located in the F-helix in the kinase domain of TbRII. The F-helix functions as a scaffold to correctly position catalytic, substrate-binding, and substrate residues [33]. The D-helix in the protein kinase domain is either close to or makes direct contact with the substrate. Mutations at the F-helix will perturb F-helix-D-helix communication and impair TGFBR2 signaling. In this study, the V453E mutation replaced the hydrophobic valine with a hydrophilic glutamic acid to form more hydrogen bonds, which altered the conformation of the F-helix (Figure 3E,F). This alteration destabilized the F-helix, which in turn perturbs the protein structure and function of TGFBR2. Therefore, the newly identified p.V453E missense mutation could seriously disturb the TGF-β signaling pathway, the consequence of which is MFS2 phenotypes in affected individuals. These findings aid in understanding that MFS can be caused not only by mutations in FBN1 but also by mutations in TGFBR2.

More research is needed to better understand the relationship between TGFBR2 mutations and MFS2 phenotypes. Great clinical heterogeneity was observed among TGFBR2 mutations carriers [34]. In the present study, we found that F-helix mutation carriers had a very high major cardiovascular system involvement rate (93.8%). TGF-β signaling drives aneurysm progression in MFS. Habashi et al. [35] and Holm et al. [36] found that an angiotensin II receptor blocker, losartan, can inhibit TGF-β-mediated activation of extracellular signal–regulated kinase. Therefore, losartan can be widely used to treat patients with MFS. In our study, no obvious progression of aortic lesion was observed in all of the affected individuals after losartan treatment.

The proband with a TGFBR2 mutation fulfilled the current diagnostic criteria of MFS2 [22], indicating an association between the genotype of V453E alterations and MFS2. Nine kinds of mutations at the F-helix of TGFBR2 have been reported, with residue sites 446 and 449 having higher mutation frequencies. Patients with mutations at this region are more prone to severe cardiovascular events and skeletal lesions (Table 3). Because mutations at the F-helix can seriously impair the TGF-β signaling pathway, individuals who have mutations at this region display a higher ratio (p=0.012) of developing severe clinical manifestations before adulthood.

Previous studies indicate that ectopia lentis is rare in MFS2 individuals [5,18], and a study performed in Chinese patients with Marfan-related disorders also found no ocular involvement [32]. In the Hilhorst-Hofstee et al. [37] study, they thought that ectopia lentis is caused by the lower production of fibrillin-1 and not by perturbation of the TGF-β signaling. Consistent with these results, no ocular involvement was observed in patients of our study who had TGFBR2 mutations, and no ectopia lentis was found in patients carrying a mutation at the F-helix. Only four F-helix mutation carriers (4/16, 25%) were reported as showing minor ocular lesions. One MFS2 patients had myopia, and in the three patients with Loeys-Dietz Syndrome, two blue sclerae and two strabismus were reported (Table 3).

In conclusion, this study reports a novel TGFBR2 mutation in a Chinese family with a MFS2 proband. Mutations at the F-helix in the kinase domain of TGFBR2 are prone to developing severe cardiovascular phenotypes and skeletal irregularities. Losartan can slow-down the progression of aortic lesion. Mutations in this region are also related to young-onset clinical phenotypes. Minor ocular abnormalities can be found in a low percentage of carriers with mutation at the F-helix. These findings have implications for genetic testing, diagnosis, and treatment in individuals with MFS2, and other TGF-β-signaling-related disorders.

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Appendix 1

Primers used for the amplification of FBN1. To access the data, click or select the words “Appendix 1.” This will initiate the download of a compressed (pdf) archive that contains the file.