The ocular phenotype of stiff-skin syndrome

Abstract

**Purpose**  Stiff skin syndrome (SSS; MIM#184900) is a rare autosomal dominantly inherited Mendelian disorder characterised by thickened and stone-hard indurations of the skin, mild hypertrichosis, and limitation of joint mobility with flexion contractures. It is autosomal dominant with high penetrance and results from mutations in the fibrillin 1 (FBNI; MIM*134797) gene. Here we present the associated ocular phenotype in a two generation nonconsanguineous Northern Irish family.

**Methods**  The affected patients underwent complete ophthalmic and orthoptic assessment and genetic testing.

**Results**  All three patients had ophthalmoplegia of varying degrees. Direct sequencing of the FBNI gene detected a heterozygous pathogenic mutation (c.4710G>C; p.Trp1570Cys) in all affected patients.

**Conclusions**  This is the first report of ophthalmoplegia in association with SSS.

Eye advance online publication, 16 October 2015; doi:10.1038/eye.2015.183

Introduction

Stiff skin syndrome (SSS; MIM#184900) is a rare Mendelian autosomal dominant disorder characterised by thickened, stone-hard indurations of the skin, mild hypertrichosis, and limitation of joint mobility with flexion contractures. It was first reported in 1971. It is caused by pathogenic mutations in theArg–Gly–Asp (RGD) sequence-encoding domain of fibrillin 1 (FBNI) gene that mediates intergrin binding. No previous descriptions of the ocular phenotype in SSS have been published and herein we report the ocular findings of an affected family.

Case reports

Three patients (a mother and her two daughters) were diagnosed with SSS at the Northern Ireland Regional Genetics Service, Belfast. There was an unaffected son from the sibship. Diagnosis was confirmed at the molecular level using direct sequencing; all affected family members had a heterozygous FBNI pathogenic gene mutation (c.4710G>C; p.Trp1570Cys). Ophthalmic and orthoptic examinations were subsequently completed.

**Case 1: Proband (54-year-old female)**

This patient wore glasses from the age of 5 years and had been diagnosed with right amblyopia, which was treated with occlusion therapy. Aged 51 years, she was diagnosed with bilateral posterior subcapsular cataracts and had uncomplicated phacoemulsification and intraocular lens implantation. The left eye was unexpectedly myopic post-operatively and was corrected by a LASIK (Laser Assisted in situ Keratomileusis) procedure.

She had been diagnosed with flexion contractures of fingers and elbows at 7 years of age and was found to have thickened skin on her hands and arms. By the age of 11 years the patient noticed stiffening/clawing of her hands, which by the age of 15 years was affecting her employment as a seamstress. By the age of 47 years predominately the distal and proximal interphalangeal joints of the hands were affected with evidence of cutaneous nodules. There has been some evidence of mild nail dysplasia. She has had respiratory symptoms and in particular shortness of breath from age of 33 years and has been under the care of a respiratory physician.

Her best-corrected visual acuity was 6/18 in both eyes. The left cornea showed evidence of previous LASIK. Both eyes showed well-centred stable posterior chamber IOLs. The posterior pole was normal in both eyes. Cover testing revealed orthophoria for distance and 10 prism dioptries of exophoria at near. The extraocular movements of both eyes were abnormal with restriction of elevation, depression, abduction, and adduction (Figure 1).
**Case 2 (23-year-old female)**

This patient was the first daughter of the proband and reported difficulty looking up and to the left. She tended to move her head instead of her eyes when looking to the side. She had worn glasses from the age of 8 years. She was noted to have fissured palms at birth, generalised stiff skin, flexion contractions at both elbows, tight hard skin on hands and feet. She toe walked and had Achilles tendon lengthening procedures at ages 8 and 14.

Her best-corrected visual acuity was 6/6 right eye and 6/5 left eye and N5 in both eyes. Her anterior and posterior chamber examination was normal. Cover testing revealed 1 prism dioptre of exophoria for distance with 10 prism dioptres of esophoria for near. Extraocular movements were restricted in upgaze and in lateral gaze (Figure 2).

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**Case 3 (22-year-old female)**

This patient who was the second daughter of the proband reported difficulty looking up and to the sides. She had worn glasses from 4 years of age. She had generalised stiff-skin flexion contractions of both elbows, a history of toe walking for which bilateral Achilles tendon release procedures were completed at age 7 years.

Her best-corrected visual acuity was 6/6 right eye and 6/9 left eye. Anterior segment examination was normal. The crystalline lens in the right eye was normal however in the left eye showed few refractile dot-like opacities in the nucleus. There was no phacodonesis or subluxation. Posterior segment examination was normal. Cover testing revealed 1 prism dioptre of exophoria for distance with 12 prism dioptres of esophoria for near. Extraocular movements were restricted in all positions of gaze (Figure 3).

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**Figure 1**  Patient 1 eight positions of gaze.

**Figure 2**  Patient 2 eight positions of gaze.
Discussion

SSS usually presents with stony hard thick skin that limits joint mobility congenitally or before the age of 6 years.3 The hard skin is bound firmly to the underlying tissues usually over the entire body, with the most common areas being the buttocks, thighs and shoulder areas that limits joint mobility and cause flexion contractures. Also cutaneous nodules that predominantly affect DIP joints and diffuse entrapment neuropathy are associated. Rarely SSS can be associated with scoliosis, restrictive pulmonary changes (as observed in the proband), focal lipodystrophy, hypertrichosis, muscle weakness, growth retardation, and/or relative short stature. The disease is felt to be slowly progressive. It is a rare autosomal dominantly inherited mendelian disorder with <40 reported cases in the literature. Fibrillin is essential in the mechanical strength of many tissues and contains binding site for integrin receptors which induce intracellular adaptations in response to extracellular changes. Mutations in FBN1 result in SSS and Marfan syndrome.2 The pathogenic mutations identified in SSS are in the Arg–Gly–Asp (RGD) sequence-encoding domain of fibrillin-1 that mediate integrin binding, resulting in a fibrotic phenotype.4 There is a wide phenotype variation described in the literature. The milder cases may represent a localised condition due to mosaic forms of SSS. Treatment for SSS is usually supportive with physiotherapy.

The ocular phenotype in this family with SSS consisted of a mechanical restriction of extraocular movement consistent with fascial shortening within the orbit. This is most likely due to the same pathophysiology as the limb contractions and dermatological findings; giant collagen fibril formation without inflammation in the extraocular fascia network.5,6 Ophthalmoplegia has not been documented in previous case reports of SSS.7–13

Summary

What was known before

- SSS is a rare autosomal dominant Mendelian disorder with high penetrance.
- It is caused by pathogenic mutations in the Arg–Gly–Asp sequence-encoding domain of the Fibrillin 1 (FBN1) gene.

What this study adds

- Restrictive ophthalmoplegia is part of the SSS ocular phenotype and ophthalmic assessment is recommended.
- Patients with this restrictive ophthalmoplegia should undergo systemic evaluation for SSS.

Conflict of interest

The authors declare no conflict of interest.

References


