Article

GDF6 Knockdown in a Family with Multiple Synostosis Syndrome and Speech Impairment

Raymond A. Clarke 1,* , Zhiming Fang 1 , Dedee Murrell 2 , Tabrez Sherriff 2 and Valsamma Eapen 1

1 Ingham Institute, School of Psychiatry, University of NSW, Sydney, NSW 2170, Australia; z.fang@alumini.unsw.edu.au (Z.F.); v.eapen@unsw.edu.au (V.E.)
2 School of Medicine, St George Hospital, University of NSW, Sydney, NSW 2217, Australia; d.murrell@unsw.edu.au (D.M.); tabrez.sherriff@gmail.com (T.S.)

* Correspondence: r.clarke@unsw.edu.au

Abstract: Multiple synostoses syndrome type 4 (SYNS4; MIM 617898) is an autosomal dominant disorder characterized by carpal-tarsal coalition and otosclerosis-associated hearing loss. SYNS4 has been associated with GDF6 gain-of-function mutations. Here we report a five-generation SYNS4 family with a reduction in GDF6 expression resulting from a chromosomal breakpoint 3′ of GDF6. A 30-year medical history of the family indicated bilateral carpal-tarsal coalition in ~50% of affected family members and acquired otosclerosis-associated hearing loss in females only, whereas vertebral fusion was present in all affected family members, most of whom were speech impaired. All vertebral fusions were acquired postnatally in progressive fashion from a very early age. Thinning across the 2nd cervical vertebral interspace (C2-3) in the proband during infancy progressed to block fusion across C2-7 and T3-7 later in life. Carpal-tarsal coalition and pisiform expansion were bilaterally symmetrical within, but varied greatly between, affected family members. This is the first report of SYNS4 in a family with reduced GDF6 expression indicating a prenatal role for GDF6 in regulating development of the joints of the carpals and tarsals, the pisiform, ears, larynx, mouth and face and an overlapping postnatal role in suppression of aberrant ossification and synostosis of the joints of the inner ear (otosclerosis), larynx and vertebrae. RNAseq gene expression analysis indicated >10 fold knockdown of NOMO3, RBMXL1 and NEIL2 in both primary fibroblast cultures and fresh white blood cells. Together these results provide greater insight into the role of GDF6 in skeletal joint development.

Keywords: multiple synostosis syndrome; vertebral fusion; GDF6; SYNS4; Klippel-Feil; progressive ossification; pisiform; skeletal morphology

1. Introduction

Bone morphogenetic proteins (BMPs) regulate skeletal development, bone morphology and joint formation. Joints form when skeletal elements segment and develop articulations that provide flexibility, strength and versatility through specialised functions and appendages [1]. Much of what we now know regarding the molecular basis of skeletal joint development derives from our understanding of genetic changes affecting BMP13 and BMP14, more commonly referred to as growth and differentiation factor 6 (GDF6) and 5 (GDF5), respectively [1–5].

GDF6 joint phenotypes overlap GDF5 joint phenotypes, including carpal, tarsal and vertebral fusion [6–8]. GDF5 gain-of-function mutations increase the downstream signalling of GDF5 causing proximal symphalangism (SYM) and multiple synostoses syndrome type 2 (SYNS2; MIM 610017) which is characterised by the fusion of the carpals, tarsals and vertebrae [6–8]. By comparison, GDF5 loss-of-function mutations cause brachydactyly (BDA and BDC) [8,9]. GDF6 gain-of-function mutations cause multiple synostoses syndrome type 4 (SYNS4) [3–5] which is characterised by synostoses of the carpals and tarsals and otosclerosis-associated conductive hearing loss but not vertebral fusion [3–5].
Leber congenital amaurosis-17 (LCA17, MIM 615360) and isolated cases of microphthalmia (MCOP4, MIM 613094) have also been reported in association with GDF6 loss-of-function mutations [10,11]. In contrast, increased levels of GDF6 expression, which derive from the recurrent duplication of the GDF6 gene in Leri’s pleonostosis are characterised by vertebral fusion, reduced joint flexibility and mobility and flexion contractures of the interphalangeal joints [12].

In this study we report a SYNS4 family with classical carpal-tarsal coalition and acquired otosclerosis-related conductive hearing loss [3–5]. All affected family members presented with variable degrees of vertebral fusion and most were speech impaired.

2. Materials and Methods

Clinical Investigation: Radiology of the wrists, feet and spine were performed using routine practices [13]. We conducted a long-term review of radiographs and reports compiled over 30 years. We performed a detailed review of radiographs of the hands, feet and spine from four affected family members IV-5, IV-9, IV-10 and IV-12 (Figure 1 and Table 1). Other members of this family have been described previously with what was misdiagnosed as congenital vertebral fusion (Klippel-Feil anomaly; MIM 118100) [13]. Hearing tests and endoscopic examinations of the larynx and skeletal radiographs were performed at clinics across Australia. There were no reports of vision impairment or other ocular anomalies in this family.

![Figure 1. SYNS4 family pedigree. ■ Multiple synostoses, vertebral fusion, speech impairment; chromosomal breakpoint 3' of GDF6; * Decreased GDF6 expression confirmed; Proband arrowed; □ Unaffected.](image URL)

Primary cell culture: Primary fibroblasts acquired from skin biopsies were cultured in DMEM supplemented with 10% fetal calf serum (FCS). Cells were centrifuged at 1000 rpm for 5 min, resuspended and washed in PBS before RNA isolation and comparative rtPCR analysis as described elsewhere [14].

RNA isolation: Total RNA was extracted from fresh white blood cells and fibroblast cultures using Trizol according to the manufacturer’s instructions (Thermo Fisher Scientific, Sydney, Australia). Purelink RNA mini kit (12183025) spin columns(#12183025 Thermo Fisher Scientific, Sydney, Australia) were used for RNA purification. RNA was resuspended in DEPC-treated water and quality tested using a fragment Analyser (Agilent, Mulgrave, Australia) followed by the addition of 1 uL of RNASEOUT ((#10777019 Thermo Fisher Scientific, Sydney, Australia)).
Table 1. Clinical details of selected family members.

| Family Member | Sex | Ages | Anomalies |
|---------------|-----|------|-----------|
| Proband (IV-12) | F   | 0–27 | Carpal Tarsal Coalition, Pisiform elongated, No hearing impairment or congenital vertebral fusion, Postnatal vertebral fusion, speech impaired, short tongue and microstomia. |
| Brother (IV-10) | M   | 7–26 | Carpal Tarsal Coalition, Pisiform elongated, No hearing impairment or congenital vertebral fusion, Postnatal vertebral fusion, No speech impairment, short tongue and microstomia. |
| Brother (IV-9)  | M   | 12–19| Carpal Tarsal Coalition, Pisiform elongated, No hearing impairment, Postnatal vertebral fusion, Severe speech impairment, short tongue and microstomia. |
| Cousin (IV-5)   | M   | 17–50| Carpal Tarsal Coalition, Pisiform not tested, No hearing impairment, Progressive vertebral fusion, Severe speech impairment, short tongue and microstomia. |

Comparative rtPCR: First-strand cDNA synthesis was performed using the SuperScript™ III First-Strand synthesis rtPCR Kit (Invitrogen Cat# 11752-050, Thermo Fisher Scientific, Sydney, Australia) according to manufacturers’ instructions: 10 µL of 2 x RT Reaction Mix, 2 µL RT Enzyme Mix and 50 pg of purified RNA were made up to 20 µL with DEPC-treated water and incubated at 25 °C for 10 min and again at 42 °C for 50 min. Reactions were terminated at 85 °C for 5 min, then chilled on ice for 5 min followed by a short spin in the microfuge. Then 1 µL (2 U) of E. coli RNase H was added and incubated at 37 °C for 20 min.

PCR master mix was prepared from a common stock reaction mix. Volumes for a 25 µL reaction were as follows: 12.5 µL of Platinum®® SYBR®® Green qPCR SuperMix-UDG (#11733-046 Thermo Fisher Scientific, Sydney, Australia.), 1 µL each of 10 µM primer stocks specific for the genes of interest (Table 2), 2.5 µL of cDNA and DEPC-treated water to 25 µL. Reactions were incubated at 50 °C for 2 min and an initial denaturation step of 94 °C for 2 min. PCR was performed for 40 cycles with denaturation at 94 °C for 15 sec, annealing at 55 °C for 10 sec and extension at 72 °C for 20 sec. Comparative PCR profiles were independently normalised against the expression of two house-keeping genes (GAPDH and 18sRNA) to remove any non-biological variation.

Table 2. PCR Primers.

| Primer Direction | Primer Sequence |
|------------------|----------------|
| GDF6—forward     | CCTGTTGCTTGTGTGGTTC|
| GDF6—reverse     | GCTGTCCATTCTCTTTGCG|
| 18s rRNA—forward | GTAACCCGTTGAACCCCATT |
| 18s rRNA—reverse | CCATCCAATCGGTAGTAGCG |
| GAPDH—forward    | CCACCATGGGCAATCCTATGGGA |
| GAPDH—reverse    | TCTAGACGGCCAGTGACAGTCCAGCCC |
| Stealth control—sense | CAAGAACACGCCAGAGCAGCGCGUCA |
| Stealth control—antisense | UGACGGCCUCUUCUCGCGUGUUCUG |

mRNA sequencing by Illumina HiSeq /Novaseq: Total RNA of each sample was extracted using TRIzol Reagent and Purelink RNA mini kit columns. Total RNA of each sample was quantified and qualified by an Agilent 2100/2200 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA), NanoDrop (Thermo Fisher Scientific, Sydney, Australia). 1 µg total RNA was used for library preparation. Next generation sequencing library preparations were constructed according to the manufacturer’s protocol by Genewiz China.
Differential expression analysis used the DESeq2 Bioconductor package, a model based on the negative binomial distribution. The estimates of dispersion and logarithmic fold changes incorporate data-driven prior distributions, Padj of genes were set <0.05 to detect differentially expressed genes. For expression analysis transcripts in Fasta format were converted from known gff annotation file and indexed properly. Then, with this file as a reference gene file, HTSeq (v0.6.1) estimated gene and isoform expression levels from the pair-end clean data. For GO and KEGG enrichment analysis GOSeq (v1.34.1) was used to identify Gene Ontology (GO) terms that annotate a list of enriched genes with a significant padj < 0.05. TopGO was used to plot DAG. Differentially expressed genes are presented in order of fold change relative to unaffected control.

3. Results
3.1. Radiological Review
This is the first long-term review of a SYNS4 family. We review a 30-year history of radiological films and reports for the affected family (Figure 1) where ~50% of the affected displayed carpal and tarsal coalition, all affected family members displayed expansion of the pisiform and six of the seven affected females tested presented with otosclerosis-associated conductive hearing loss that was absent from affected males. All affected family members presented with variable degrees of vertebral fusion and most were speech impaired.

3.1.1. Female Proband (IV-12)
The female proband presented with bilateral carpal and tarsal coalition. In the feet there was bilateral fusion between the navicular and cuboid. In the hands there was bilateral fusion between the triquetrum and lunate and between the hamate and capitate and partial fusion between the hamate and pisiform (Figure 2). Bilateral elongation of the pisiforms was symmetrically similar in the proband (Figure 3a) and all other affected family members tested, but varied greatly in degree between affected family members (Figure 3b). Pisiform elongation associated with restricted wrist rotation/supination and grasping and capacity to write. Prenatal ultrasound of the proband provided no evidence of vertebral fusion (Figure 4a). Spinal radiographs at age 10 weeks for the proband confirmed the absence of congenital vertebral fusion in the cervical spine (Figure 4b). At age 12 months the proband had developed fusion of the C2-3 apophyseal joints and anterior regions of the spinous processes (Figure 4c). At age 13 there was complete fusion of the C2-3 vertebral bodies, fusion of the anterior edges of C3-4 and C4-5 vertebral bodies, progressive ossification of the anterior edges between C5-6 vertebral bodies, partial fusion of the apophyseal joints and spinous processes at C6-7 (Figure 4d) and partial fusion of T3-7 vertebral bodies. At age 27 spinal fusion of the vertebrae had progressed to form block fusion across C2-7 (Figure 4e) and a continuous set of partial fusions of the anterior edges of the vertebral bodies between T3-T7 (Figure 4f). The bilaterally elongated pisiforms of the proband did not increase in length or size over time nor did the fusion of carpals or tarsals appear to progress. Pisiform elongation was associated with restricted flexion and supination movement of the wrist. The proband presented with stiffness in the Achilles tendon with associated toe walking. She was the only female (age 27) tested that was negative for hearing loss and she was dysphonic from birth and displayed severe speech impairment which was associated with malformation of laryngeal cartilages in her speech-impaired father. Short tongue and microstomia was evident in the proband and most other affected family members in association with overcrowding of the teeth that required teeth removal.
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**Figure 2.** Radiographs of proband’s wrist highlighting: (a) Fusion of lunate and triquetrum (large white arrow), nearly complete fusion of hamate and pisiform (small white arrow) and elongation of the pisiform (black arrow). (b) Elongated pisiform (large white arrow) and fusion of hamate and pisiform (small white arrow).

**Figure 3.** Radiographs of wrists of: (a) Proband—display bilateral symmetry of elongation of the pisiforms. (b) Proband’s father—displays bilateral symmetry of elongation of the pisiforms which are different in length and morphology from those of the proband. Both father and daughter had restricted wrist rotation/supination and grasping and writing capacity.
The proband's oldest brother (IV-9) presented with postnatal fusion in the cervical spine which progressed in a cranio-caudal direction from C2-3. At age 7 this boy presented with fusion of the apophyseal joints and spinous processes of the 2nd and 3rd cervical (C2-3) vertebrae (Figure 4g) which by age 14 had progressed to complete fusion of the C2-3 vertebral bodies and thinning of the vertebral interspace at C3-4 (Figure 4h). At age 17 ossification of intervertebral disc spaces had progressed to where there was thinning of the C3-7 vertebral interspaces and fusion of the apophyseal joints at C2-6 on the right and C6-7 on the left (Figure 4i) which by age 26 had progressed to complete block fusion of C2-7 (Figure 4j). At age 26 he presented with vertebral fusion in the upper thoracic spine where there had only been thinning of these interspaces 9 years earlier (Figure 4k). At age 26 there was partial anterior fusion of the vertebral bodies at T4-T6 (Figure 4i) comparable to the thoracic fusion profile in the spine of his affected father III-6 (Figure 4m). Aged 7 years, he presented with very mild dysphonia, habitual toe walking, severely restricted flexion and supination movement in the hands, microtia, low set ears, short tongue, microstomia and overcrowding of the teeth that required teeth removal. Mobility of other joints appeared normal and there was no conductive hearing loss evident in this man or any other affected male member of the family.

3.1.3. Brother (IV-9)

The proband’s oldest brother (IV-9) presented with postnatal fusion in the cervical spine which progressed in a cranio-caudal direction from C2-3. A solitary C2-3 fusion identified at age 12 (Figure 4n) had progressed 7 years later to include a C4-5 fusion...
and thinning of the C6-7 interspace (Figure 4o). There was no evidence of conductive hearing loss in this brother or any other affected male members of the family. He suffered severe dysphonia from birth associated with severe speech impairment, microstomia and overcrowding of the teeth.

3.1.4. Male Cousin (IV-5)

The proband’s male cousin (IV-5) presented with progressive postnatal fusion of the cervical spine. Fusion of the C2-3, C4-5 and C6-7 cervical vertebrae at age 17 (Figure 4p) had progressed to C2-7 block fusion at age 24 (Figure 4q). At age 50 he experienced severe stiffness of the neck, shoulders and back. There was no evidence of conductive hearing loss. Severe speech impairment was evident from an early age. Short tongue and microstomia was associated with overcrowding of the teeth that required removal of eight teeth from him and all three of his affected sisters (IV-6, -7 and -8).

3.1.5. Familial Skeletal Anomalies

Near 50% of affected family members tested presented with variable degrees of carpal and tarsal coalition. Carpal and tarsal fusions were bilaterally similar in extent and morphology within affected family members but varied between affected family members. Likewise, expansion of the pisiform was bilaterally similar in extent and appearance within affected family members but varied between affected family members. A total of six out of the seven affected female family members tested for hearing loss were diagnosed with otosclerosis and/or unilateral conductive hearing loss, including the proband’s grandmother (II-4) age 45, aunty (III-3) age 40, aunty (III-5) age 36, female cousin (IV-6) age 14, female cousin (IV-8) age 20 and niece (V-1) age 5, with evidence of deterioration with age. The only female tested that was negative for hearing loss was the proband, age 27 years (Figure 1).

All four of the affected family members reviewed here in detail (Table 1) presented with progressive postnatal acquisition of vertebral fusions (Figure 4). Ultrasound before birth for two affected family members (IV-12 and V-3) confirmed the absence of congenital vertebral fusion which later developed postnatally through progressive ossification. Vertebral fusion, carpal and tarsal coalition, bilateral pisiform elongation and vocal impairment were all variable in extent between affected family members. Bilateral deviation of the proximal phalanges of toes 2–5 was evident in two affected female members of the family (IV-8, see Figure 4r) and (III-2, see Figure 4s). The former female’s mother (III-5) developed painful age-related bilateral spurs on the patella at age 65 (Figure 4t) and age-related conductive hearing impairment age 40. Other skeletal anomalies included Perthes of the hip in one teenage male in association with a minimal fusion bridge between the C2-3 vertebrae and no obvious speech impairment (V-3). There was no evidence of restriction of the elbows or the shoulders in affected family members. Most affected family members presented with varying degrees of speech impairment from a young age in association with malformation of laryngeal cartilages including flattening of the anterior commissure and shortening of the vocal cords [15]. Surgical intervention in one affected family member reported that the vocal cords were shorter, failed to meet in the midline, were of a different complexion/composition and did not vibrate normally and that other vocal ligaments were ossified [15]. Short tongue and microstomia with overcrowding of the teeth were common. In more severely speech-affected members of the family there were deficits in verbal fluency and significant difficulty in generating words beginning with a certain letter [15]. Stature and intelligence appeared within the normal limits for all affected family members. Speech and hearing impairment affected learning; pisiform expansion restricted wrist rotation supination and grasping and capacity to write; narrowing of the oesophageal and laryngeal canals restricted swallowing and anaesthetic intubation; Achilles tendon stiffness was associated with toe walking; and arthritis progressed with age in many of the affected family members.
3.2. Gene Expression Analyses

3.2.1. Comparative rtPCR Gene Expression Analysis

RNA isolated from fresh white blood cells derived from two severely affected family members (III-6 and IV-10) and five unaffected control individuals were analysed for changes in GDF6 expression using comparative rtPCR expression analysis (Figure 5). GDF6 expression levels were reduced in both affected family members when compared to the mean expression level for the five unaffected control individuals (Figure 5).

![Mean Percentage Control Value](image)

**Figure 5.** Comparative rtPCR analysis of GDF6 expression in SYNS4 family. GDF6 expression levels in fresh white blood cells expressed as the mean percentage change for two affected male members of the family compared with the mean for five age- and gender-matched unaffected controls independently normalised against the expression of two housekeeping reference genes GAPDH and 18sRNA.

3.2.2. RNA Sequencing Gene Expression Analysis (RNAseq)

RNA isolated from a severely affected family member with reduced GDF6 expression (III-6) was used for RNAseq gene expression analysis. RNAseq analysis identified 68 and 61 genes with >10 fold differential gene expression in primary fibroblast cultures and fresh white blood cells, respectively (p < 0.05) (Table 3). Of these, 3 genes (NOMO3, RBMXL1 and NEIL2) exhibited >10 fold knockdown in both primary fibroblasts and white blood cells from the affected family member compared with an age, gender and racially matched unaffected control individual (Table 3).
### Table 3. Differentially expressed genes in RNA sequence analysis.

| Gene        | Change in Fibroblast Cell Lines | Gene        | Change in White Blood Cells |
|-------------|---------------------------------|-------------|-----------------------------|
| NOMO3       | −15.3854                        | MDGA1       | −15.077                       |
| DNAH1       | −15.0439                        | AK4         | 11.0154                      |
| PRDM10      | −14.9457                        | GPR68       | 11.1429                      |
| IDO1        | −14.7973                        | CDK19       | 11.8143                      |
| SP140L      | −14.7586                        | NPR1A       | 11.8187                      |
| MYSM1       | −14.6305                        | ABHD16B     | 12.0527                      |
| RB10        | −14.5996                        | ZCCHC4      | 12.1243                      |
| PTGS2BLOC1S2| −14.4498                        | BBS5        | 12.1532                      |
| PTGS2BLOC1S2| −14.4498                        | UCP2        | 12.1681                      |
| TME260      | −14.1642                        | ATP6V0A1    | 12.3948                      |
| SNX13       | −14.0789                        | CEP8        | 12.4649                      |
| RCBT8       | −14.0487                        | NMNAT1      | 12.6268                      |
| CCP110      | −14.0126                        | SYNS4       | 12.6402                      |
| RBMXL1      | −13.7795                        | IPOC        | 12.8599                      |
| EPHX2       | −13.7398                        | HLA        | 13.0328                      |
| NF2         | −13.6138                        | UBL4A       | 13.0354                      |
| GZF1        | −13.3585                        | GOLPH3L     | 13.0845                      |
| CDA8        | −13.2922                        | MED19       | 13.1062                      |
| AMZ2        | −13.2741                        | GLI2        | 13.1362                      |
| POLR1B      | −13.2696                        | ZBED8       | 13.1382                      |
| TTI1        | −13.2425                        | SCML1       | 13.1582                      |
| WDR19       | −13.2212                        | MORC2       | 13.2509                      |
| ABCB7       | −13.1889                        | NEDD9       | 13.2561                      |
| SMc6        | −13.1802                        | ROR1        | 13.3067                      |
| DERa        | −13.104                         | ZC3H8       | 13.3068                      |
| ZNF33B      | −12.7709                        | CEP44       | 13.3209                      |
| POLR3B      | −12.6132                        | ZNF274      | 13.4586                      |
| NEIL2       | −11.8528                        | POGZ        | 13.617                       |
| BEGAIN      | −11.8236                        | TENT5A      | 13.6751                      |
| −11.801     | −11.801                         | TN1K        | 13.8448                      |
| −15.0439    | −15.0439                        | BTRC        | 13.8448                      |
| −14.1642    | −14.1642                        | INPP4A      | 13.9705                      |
| −14.0789    | −14.0789                        | PIK3R1      | 14.1731                      |
| −14.0487    | −14.0487                        | PIK3R1      | 14.3364                      |
| −14.0126    | −14.0126                        | ELAPOR2     | 14.3602                      |
| −13.7795    | −13.7795                        | NAV3        | 14.5402                      |
| −13.7398    | −13.7398                        | GALNT15     | 14.8302                      |
| −13.6138    | −13.6138                        | NUP155      | 14.9429                      |
| −13.3585    | −13.3585                        | ADGRB2      | 15.0595                      |
| −13.2922    | −13.2922                        | SLC4A3      | 15.5187                      |
| −13.2741    | −13.2741                        | MPDZ        | 15.8286                      |
| −13.2696    | −13.2696                        | MDGA1       | 16.3832                      |
| −13.2425    | −13.2425                        | ZNF50       | 16.5375                      |
| −13.2212    | −13.2212                        | ZNF58       | 16.7575                      |
| −13.1889    | −13.1889                        | ZNF530      | 16.9263                      |
| −13.1802    | −13.1802                        | NT5M        | 17.0263                      |
| −13.104     | −13.104                         | TENT5A      | 17.0263                      |
| −12.7709    | −12.7709                        | REEF5       | 17.0263                      |
| −12.6132    | −12.6132                        | TENT5A      | 17.0263                      |
| −11.8528    | −11.8528                        | TENT5A      | 17.0263                      |
| −11.8236    | −11.8236                        | TENT5A      | 17.0263                      |
| −11.801     | −11.801                         | TENT5A      | 17.0263                      |

* Genes downregulated in both fibroblast cell lines and fresh white blood cells.

4. Discussion

In this study we report the first SYNS4 family with reduced GDF6 expression. The three previously reported SYNS4 families had GDF6 gain-of-function mutations [3–5]. The family phenotype included the classical bilateral carpal and tarsal coalition and otosclerosis associated conductive hearing loss typical of SYNS4 [3–5]. In addition, there was progressive postnatal acquisition of vertebral fusions in the cervical and thoracic spine from an early age. The extent of the vertebral fusion was variable between affected family members; notwithstanding, all affected family members displayed some degree of fusion across the C2-3 vertebral interspace at the cranial end of the cervical spine. Most affected family members were also speech impaired in association with malformation of the laryngeal cartilages and what appeared to be progressive ossification of laryngeal ligaments and joints.
The decrease in GDF6 expression in this family was associated with a chromosomal breakpoint 3′ of the GDF6 gene [13]. GDF6 encodes a bone morphogenetic protein which functions in a dose-dependent fashion in its extracellular regulation of skeletal development [1,16]. In consequence, the varying degrees of joint ossification and skeletal deformation in this family likely reflect variations in the reduction of GDF6 dose at different skeletal sites in different affected family members, possibly due to position effects on GDF6 regulatory elements located near the chromosomal breakpoint. Indeed, a conserved long-range GDF6 pharyngeal specific enhancer (ECR5) that functions from the earliest stages of pharyngeal and otolaryngeal development has been located between GDF6 and the chromosomal breakpoint in this family [1,17].

Otosclerosis-associated conductive hearing loss represents one of the classical acquired characteristics of SYSN4 [3–5]. In this family, otosclerotic hearing loss presented at varying ages (5–40 years of age) in affected females only, but not in affected males. This finding would appear to indicate a gender effect on the progressive ossification of the ossicles in this family. In contrast, speech impairment in this family was associated with congenital malformation and aberrant postnatal ossification of the larynx which was more severe in males compared to females and further exacerbated during descent of the thyroid cartilage during male puberty. Notwithstanding, the one affected male member of the family without speech impairment (V-3) also displayed the least degree of vertebral fusion.

Gdf6 is expressed in discrete patterns within the developing joints of the mouse [1,2]. These same joints correspond precisely with those joints that were fused and malformed in the affected family including the joints of the carpals, tarsals, vertebrae, ear and larynx [1,2]. Moreover, knockout of Gdf6 in mice results in the fusion of the carpals and tarsals but not the vertebrae [1]. This study identifies the biological basis of vertebral fusion in this family as aberrant progressive postnatal ossification and synostosis of the spinal joints from an early age. This mechanism is comparable and consistent with the aberrant postnatal ossification within the inner ear causing otosclerosis in SYSN4 [3–5]. In contrast, the precise mechanism of the carpal and tarsal fusions in this family remains uncertain. Notwithstanding, the bilateral symmetry of the carpal, tarsal and pisiform fusions/malformations within affected family members appears more consistent with a prenatal error of development, possibly effecting aberrant and excessive bone condensation/ossification prenatally. This fusion scenario is supported by animal studies which indicate a role for Gdf6 in stimulating chondrogenesis at early stages of development [16] while in vitro studies indicate that GDF6 has a distinct inhibitory effect on ossification and mineralization at later-stage, differentiated chondrocytes or osteoblasts [18]. GDF6 is a bone morphogenetic protein that acts extracellularly as a morphogen during development. Albeit the pathway down stream of GDF6 in determining cell fate and function has not been fully elucidated [3–5]. For insight into this pathway we searched for changes in gene expression in a severely affected family member (III-6) with reduced GDF6 expression (Figure 5 and Table 3). RNAseq expression analysis identified >10 fold knockdown of three genes, NOMO3, RBMXL1 and NEIL2, in both primary fibroblast cultures and fresh white blood cells from this severely affected family member (Table 3) [19–22]. This limited but close correlation in differential gene expression between disparate cell types provides a high level of confidence with respect to ongoing pathway and gene therapy investigations, not only in these three genes but the other genes differentially expressed in this patient (Table 3). It is therefore likely that one or more of these down-regulated genes has a role in the GDF6 pathway to skeletal joint development and ossification. RBMXL1 is of particular interest as it strengthens DNA heterochromatin binding, impedes the activity of transcription factors, suppresses gene transcription and serves as a barrier to direct cell conversion. Knockdown of RBMXL1 increases gene transcription [19]. NOMO3 is another molecule of interest as it is a positive modulator of the morphogen Nodal which is down regulated >5 fold in patients with facial asymmetry and jaw malformations. Nodal is a transcription factor regulated by asymmetric cascades of morphogens. Nodal initiates the molecular pathway that induces
chirality and asymmetry in endoderm and mesoderm germ layers during late gastrulation and neurulation [20,23].

5. Conclusions
Prenatal ultrasound followed with long-term radiological evaluation of the skeleton helped to differentiate between congenital, acquired, postnatal and progressive skeletal ossification and associated skeletal malformations. This study indicates a role for GDF6 in the prenatal development of the joints of the hands and feet and larynx and in the progressive postnatal ossification of the joints of the inner ear, larynx and vertebral column. These findings further suggest that all GDF6-associated vertebral fusions may result from aberrant postnatal ossification, including those vertebral fusions previously assumed but never proven to be congenital errors of segmentation [10,13].

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Institutional Review Board Statement: The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the University of New South Wales Human Ethics Committee (Approval Number: H8853).

Informed Consent Statement: Informed consent was obtained from all subjects involved in the study.

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