Molecular etiology of arthrogryposis in multiple families of mostly Turkish origin

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BACKGROUND. Arthrogryposis, defined as congenital joint contractures in 2 or more body areas, is a clinical sign rather than a specific disease diagnosis. To date, more than 400 different disorders have been described that present with arthrogryposis, and variants of more than 220 genes have been associated with these disorders; however, the underlying molecular etiology remains unknown in the considerable majority of these cases.

METHODS. We performed whole exome sequencing (WES) of 52 patients with clinical presentation of arthrogryposis from 48 different families.

RESULTS. Affected individuals from 17 families (35.4%) had variants in known arthrogryposis-associated genes, including homozygous variants of cholinergic γ nicotinic receptor (CHRNG, 6 subjects) and endothelin converting enzyme-like 1 (ECEL1, 4 subjects). Deleterious variants in candidate arthrogryposis-causing genes (fibrillin 3 [FBN3], myosin IXA [MYO9A], and pleckstrin and Sec7 domain containing 3 [PSEN3]) were identified in 3 families (6.2%). Moreover, in 8 families with a homozygous mutation in an arthrogryposis-associated gene, we identified a second locus with either a homozygous or compound heterozygous variant in a candidate gene (myosin binding protein C, fast type [MYBPC2] and vacuolar protein sorting 8 [VPS8], 2 families, 4.2%) or in another disease-associated genes (6 families, 12.5%), indicating a potential mutational burden contributing to disease expression.

CONCLUSION. In 58.3% of families, the arthrogryposis manifestation could be explained by a molecular diagnosis; however, the molecular etiology in subjects from 20 families remained unsolved by WES. Only 5 of these 20 unrelated subjects had a clinical presentation consistent with amyoplasia; a phenotype not thought to be of genetic origin. Our results indicate that increased use of genome-wide technologies will provide opportunities to better understand genetic models for diseases and molecular mechanisms of genetically heterogeneous disorders, such as arthrogryposis.

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Conflict of interest: J.R. Lupski has stock ownership in 23andMe, is a paid consultant for Regeneron Pharmaceuticals, has stock options in Lasergen Inc., and is a coinventor on multiple US and European patents related to molecular diagnostics for inherited neuromuscular, eye diseases, and bacterial genomic fingerprinting. The Department of Molecular and Human Genetics at Baylor College of Medicine derives revenue from the chromosomal microarray analysis (CMA) and clinical exome sequencing offered in the Baylor Miraca Genetics Laboratory (BMGL, http://www.bmgl.com/BMGL/Default.aspx). R.A. Gibbs is interim Chief Scientific Officer of BMGL, and J.R. Lupski is on the Scientific Advisory Board of BMGL.

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Introduction

Arthrogryposis, also known as arthrogryposis multiplex congenita, is clinically defined as congenital joint contractures or movement restriction in multiple body areas. It should be distinguished from isolated congenital contractures that affect a single joint such as congenital clubfoot deformity or dislocated hip. Arthrogryposis is a symptom rather than a specific diagnosis and occurs in between 1/3,000–1/5,000 live births (1). To date, more
than 400 disorders have been described that include arthrogryposis as an endophenotype (2).

In general, arthrogryposis occurs by virtue of a secondary effect of decreased fetal joint mobility (i.e., fetal akinesia). Disorders leading to fetal akinesia can result from abnormalities of the CNS, neuromuscular system, skeletal system, and connective and cartilage tissue disturbances. Maternal diseases or environmental factors such as intrauterine space limitations, maternal exposures to drugs or chemicals, compromise of blood supply to the fetus, and metabolic disturbances may also lead to fetal akinesia (2–4).

Distal arthrogryposis (DA) syndromes describe a heterogeneous subgroup of arthrogryposis characterized by multiple congenital contractures that mainly involve the distal parts of the upper and lower limbs without a primary neuromuscular disease (5, 6). According to different classifications, up to 19 clinical subtypes of DA were described (2, 5, 7). In most cases (~50%), mutations in genes encoding contractile proteins of skeletal muscles were shown to cause the DA phenotype, which consists of nonprogressive congenital contractures of the joints located at distal limbs (8–12). Genes determined to contribute to the molecular etiology of DA include TNNI2, MYH3, MYH8, PIEZO2, TNNT2, TNNT3, and TPM2. Recently, homozygous ECEL1 mutations, most representing loss-of-function alleles and resulting in absence of the encoded protein, were identified in patients with DA type 5D (OMIM 615065). In this group of DA, patients presented a distinctive clinical presentation with severe camptodactyly of the hands and wrists; milder camptodactyly of toes; limited knee flexion; talus and/or varus deformity of the ankles; ptosis and distinctive facial features of round-shaped, mask-like faces; high-arched eyebrows; bulbous noses; and micrognathia (13, 14).

Another phenotypically and genetically heterogeneous subgroup of arthrogryposis is multiple pterygium syndrome (MPS), which is characterized by multiple pterygia, scoliosis, and congenital contractures of the limbs. MPS has both lethal (OMIM 253290) and nonlethal (Escobar variant) (OMIM 265000) types. Although most of the MPS cases were reported with a pattern of autosomal-recessive inheritance (15), autosomal-dominant transmission was also observed in a few cases (16–18). While homozygous and compound heterozygous mutations of CHRNA7 (19, 20) were associated with both lethal and nonlethal types of MPS, mutations of CHRND (21) and CHRNA1 (21) are also responsible for lethal MPS. Recently, homozygous mutations in VRK1 (22) and autosomal-dominant inheritance with MYH3 mutations were reported in MPS patients (23).

Genetic causes of disorders with arthrogryposis include single gene mutations, chromosomal abnormalities, and mitochondrial defects (2, 24, 25). To date, variants in more than 220 genes with different modes of inheritance have been associated with arthrogryposis disorders (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI84457DS1), but in the considerable majority of these disorders, the underlying molecular etiology remains unknown. To better understand the genetics of arthrogryposis and identify potential novel molecular etiologies underlying this genetically heterogeneous group of diseases, we applied whole exome sequencing (WES) to a cohort of 52 patients from 48 families, including 22 with reported consanguinity yet with different clinical subtypes of arthrogryposis.

Results

The approach used for gene identification is outlined in Supplemental Figure 1. We first assessed WES data of all patients for rare variants in known arthrogryposis genes before investigating for potential novel genes. A list of known genes (Supplemental Table 1) was constructed from multiple online database resources (http://www.omim.org/, http://www.ncbi.nlm.nih.gov/pubmed) or abstracted from recently published articles (2, 3, 26–35). Deleterious variants in the known arthrogryposis-associated genes were identified in 17 of 48 (35.4%) unrelated families (Table 1). To the best of our knowledge, specific rare variants in known genes were not reported previously in 12 of 17 families, and 3 of those (CENPJ, GBE1, and IDS) were associated with traits suggesting phenotypic expansion (Table 1). In 3 families (6.2%), we identified deleterious variants in 3 novel genes, including FBNS, MYO9A, and PSD3. In 8 families (16.7%), we describe a second homozygous or compound heterozygous variant in a novel gene or in another known gene, in addition to the homozygous variant in a known gene (vacuolar protein sorting 8 [VPS8] novel) with POLR3A, MYBPC2 [novel] with GPR126, CHRN with ERC22, COL6A3 with BICD2, LIFR with PI4KA, LIFR with MYH14, MOI8B with MYH7B, and RIKP4 with LMNA (Table 2). Identified variants in these apparently solved cases with a molecular diagnosis obtained from WES and family pedigrees are provided in Supplemental Table 2 and Supplemental Figure 2, respectively.

While 58.3% of families manifesting arthrogryposis could be explained by a molecular diagnosis identified on WES, subjects from 20 families remained unsolved, in that no definitive molecular diagnosis could be concluded. Since amyoaplasia is the most common form of arthrogryposis that appears to be a sporadic nongenetic condition, we have carefully reexamined the clinical information of the patients in the 20 unsolved families. Five of these indeed match the clinical diagnosis of amyoaplasia. However, the remaining 15 probands had different types of arthrogryposis-related disorders, such as spinal muscular atrophy, DA, MPS, and other unspecified multiple joint contractures. There were no disease-causing variants in known genes in these patients, and we could not identify any variant in a novel gene with convincing supporting data. Thus, we concluded that 5/20 unsolved cases had a clinical presentation consistent with amyoaplasia, providing a possible explanation as to why no molecular diagnosis was achieved, at least in these cases.

We identified 1 homozygous stop-gain (in 2 families) and 2 homozygous missense mutations in ECEL1 in 4 unrelated consanguineous families, one of which is an Arab family (HOU2530). In 2 unrelated Turkish families (HOU1530 and HOU2432) with DA, we identified the same homozygous stop-gain mutation in ECEL1 (c.1147C>T; p.Gln383X). In addition, for 2 unrelated consanguineous families (HOU2460 and HOU2530), we detected 2 different homozygous missense ECEL1 mutations (c.2023G>A; p.Ala675Thr and c.1210C>T; p.Arg404Cys, respectively [Figure 1A]).

From a clinical perspective, patient BAB3934 from family HOU1530 had contractures of both hands and feet, limited knee flexion, and bilateral hip dislocation. Electromyelography (EMG) study revealed mild myogenic involvement. The subject had a flat upturned nose. Clinical findings of the patient BAB6500 from family HOU2460 revealed mild myogenic involvement. The subject had a flat upturned nose. Clinical findings of the patient BAB6500 from family HOU2530 included mild myogenic involvement. The subject had a flat upturned nose. Clinical findings of the patient BAB6500 from family HOU2530 included mild myogenic involvement.
**Table 1. Patients with a rare mutation in a single known gene**

<table>
<thead>
<tr>
<th>Pedigree</th>
<th>Patient</th>
<th>Clinical findings</th>
<th>Gene</th>
<th>Variant</th>
<th>Novel variant</th>
<th>Zyg</th>
<th>Gene-Phenotype relationship(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOU1684 BAB4104</td>
<td>Contracts of elbows and knees, camptodactyly, syndactyly of left toes 3 and 4, overlapping fingers and toes, rocker-bottom feet, bilateral ptosis, micrognathia, short and webbed neck, hypospadias, and bilateral cryptorchidism</td>
<td>CHRNG</td>
<td>c.256C&gt;T; p.Arg86Gys</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2430 BAB6498</td>
<td>Contracts of hands, elbows, and knees; multiple pterygiums; short neck; TEV; DPF; small mouth; and delayed teeth</td>
<td>CHRNG</td>
<td>c.753_754 delCT; p.Val253fs</td>
<td>No</td>
<td>Hom</td>
<td>MPS, lethal type (OMIM 253290); Escobar syndrome (OMIM 265000)</td>
<td></td>
</tr>
<tr>
<td>HOU2606 BAB7080</td>
<td>Contracts of hands, elbows, and knees; multiple pterygiums; rocker-bottom feet; DPF; micrognathia; small mouth; ECHO: Atrial septal defect and tricuspid insufficiency</td>
<td>CHRNG</td>
<td>c.241C&gt;T; p.Gln81X</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
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<tr>
<td>HOU2331 BAB6209</td>
<td>Contracts of hands, elbows, and knees; multiple pterygiums; talipes valgus; small and low-set ears; micrognathia; and ambiguous genitalia</td>
<td>CHRNG</td>
<td>c.715C&gt;T; p.Arg239Gly</td>
<td>No</td>
<td>Hom</td>
<td></td>
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<tr>
<td>HOU2607 BAB7083</td>
<td>Contracts of elbows and knees, short neck, camptodactyly, TEV, micrognathia, and prominent ears</td>
<td>CHRNG</td>
<td>c.715C&gt;T; p.Arg239Gly</td>
<td>No</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU1530 BAB3934</td>
<td>Severe flexion contracture of knees, camptodactyly, normal feet, hip dislocation, capillary hemangioma, and EMC: mild myogenic lesions</td>
<td>EEE1</td>
<td>c.1147C&gt;T; p.Gln383X</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2432 BAB6500</td>
<td>Contracts of hands, hip-joint dysplasia, talipes valgus, facial dysmorphic features with a mask-like whistling appearance, and short neck</td>
<td>EEE1</td>
<td>c.1147C&gt;T; p.Gln383X</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2460 BAB6608</td>
<td>Contracture of elbows, camptodactyly, scoliosis, ptosis, flat and mask-like face, high-arched eyebrows, low-set ears, micrognathia, and cryptorchidism</td>
<td>EEE1</td>
<td>c.2023G&gt;A; p.Ala675Thr</td>
<td>No</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2530 BAB6827</td>
<td>Severe camptodactyly, hip dislocation, hypotonia, right TEV, developmental delay, thin corpus callosum on cranial MRI, mask-like face, high-arched eyebrows, bulbous and upturned nose, low-set ears, micrognathia, nephrocalcinosis, and cryptorchidism</td>
<td>EEE1</td>
<td>c.1210C&gt;T; p.Ala404Thr</td>
<td>No</td>
<td>Hom</td>
<td></td>
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<tr>
<td>HOU1522 BAB3903</td>
<td>Contracts of both hands, TEV, high-arched feet, and contracture of toes</td>
<td>COL6A2</td>
<td>c.289C&gt;A; p.Ala97Thr</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
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<tr>
<td>HOU1529 BAB3931</td>
<td>Contractures of elbows, ulnar deviation of hands, retromicrognathia, high-arched palate, crowded and delayed teeth, low-set ears, delayed bone age, TEV, and abnormal scar formation on left foot</td>
<td>CENP4</td>
<td>c.763A&gt;G; pThr255Ala</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
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<tr>
<td>HOU1539 BAB3955</td>
<td>Flexion contractures of hands, rocker-bottom feet, delayed neuromotor development, growth retardation, hypotonia, seizures, low-set ears, high-arched palate, laryngomalacia, AP-US: Hepatosteatosis and nephrocalcinosis, EEC: Hyparsphythytism, and EMC: Anterior horn cell involvement of spinal cord. Died at 1.5 years. Clinical diagnosis: Spinal muscular atrophy</td>
<td>ID5</td>
<td>c.613C&gt;A; p.Ala201Thr</td>
<td>Yes</td>
<td>Hem</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU1540 BAB3958</td>
<td>Severe contraction of knees, elbow contractures, ulnar deviation of hands, rocker-bottom feet, talipes valgus, short neck, and scoliosis</td>
<td>TPM2</td>
<td>c.180T&gt;G; p.Tyr60X</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU1541 BAB3960</td>
<td>Joint contractures, hypotonia, cleft palate, micrognathia, short neck, tall vertebral bodies on X-ray, mild talipes equinovarus, and pelvicaliectasis</td>
<td>GBET1</td>
<td>c.776C&gt;G; p.Ala259Gly</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2114 BAB5449</td>
<td>Elbow contractures, developmental delay, recurrent infections, swallowing difficulty, contractures of the facial muscles, chubby cheeks, blepharophimosis, epicanthal folds, synophrys, long eyelashes, hirsutism, prominent and wide nasal root, high-arched palate, small mouth, and severe dental caries</td>
<td>CRLF1</td>
<td>c.984_985insG; p.Ser328fs</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2683 BAB7305</td>
<td>Severe contraction of knees, contractures of elbows and hands, joint laxity, muscle hypoplasia, axillary and popliteal pterygium, delayed motor milestones, and ECHO: Localize septal hypertrophy</td>
<td>FH1</td>
<td>c.29C&gt;G; p.Ser10Cys</td>
<td>Yes</td>
<td>Hem</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HOU2789 BAB7707</td>
<td>Contractures of elbows and hands, patella agenesis, short stature, pectus carinatum, epiphelial and metaphysial dysplasia, mild TEV, and delayed motor milestones</td>
<td>SLC26A2</td>
<td>c.2057G&gt;A; p.Cys686Tyr</td>
<td>Yes</td>
<td>Hom</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Genes in which variant alleles were associated with traits suggesting phenotypic expansion. AD, autosomal dominant; AP-US, abnormally pelvic ultrasound; DPF, downsloping palpebral fissures; ECHO, echocardiogram; EEG, electroencephalography; Hem, hemizygous; Hom, homozygous; TEV, talipes equinovarus; Zyg, zygosity.*
The affected cousin and his parents were analyzed by Sanger sequencing, and the cousin was also found to be homozygous while the parents were heterozygous carriers for the same variant identified in BAB6500 (Figure 1A). After a careful search within the kindred in HOU2432, a third-degree cousin of BAB6500 was also found to have DA. The affected cousin and his parents were analyzed by Sanger sequencing, and the cousin was also found to be homozygous while the parents were heterozygous carriers for the same variant identified in BAB6500 (Figure 1A).

Individual BAB6608 from family HOU2460 had contractures of the elbows, camptodactyly, scoliosis, pterygium, and rocker-bottom feet (33). During pregnancy, and postnatal death.

Table 2. Clinical and genotype summary of the patients with potential oligogenic or mutational burden model

<table>
<thead>
<tr>
<th>Pedigree</th>
<th>Patient</th>
<th>Clinical findings</th>
<th>Gene</th>
<th>Variant</th>
<th>Zyg</th>
<th>Gene-Phenotype relationship(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOU2163</td>
<td>BAB5611</td>
<td>Terminated pregnancy at 24 weeks of gestation.</td>
<td>CHRN</td>
<td>c.715C&gt;T; p.Arg239Cys</td>
<td>Hom</td>
<td>MPS, lethal type (OMIM 252390), Escobar syndrome (OMIM 26500)</td>
</tr>
<tr>
<td>HOU2432</td>
<td>BAB6018</td>
<td>Contractures of knees and elbows, developmental delay, hip dislocation, torticollis, scoliosis, pes planus, micrognathia, anteverted nostrils, high-arched palate, prominent ears, brain MRI: ventricular dilatation and brain stem atrophy</td>
<td>COL6A3</td>
<td>c.7720delC; p.Leu2574fs</td>
<td>Hom</td>
<td>Bethlem myopathy (OMIM 158810); Ullrich congenital muscular dystrophy (OMIM 254090)</td>
</tr>
<tr>
<td>HOU2278</td>
<td>BAB6019</td>
<td>Mild joint contractures, developmental delay, hip dislocation, scoliosis, flat occiput, micrognathia, and rocker-bottom feet</td>
<td>COL6A3</td>
<td>c.7720delC; p.Leu2574fs</td>
<td>Hom</td>
<td>Bethlem myopathy (OMIM 158810); Ullrich congenital muscular dystrophy (OMIM 254090)</td>
</tr>
<tr>
<td>HOU2332</td>
<td>BAB6212</td>
<td>Unclassified DA, narrow thorax, polyhydramnios during pregnancy, and postnatal death</td>
<td>GPR126</td>
<td>c.19C&gt;T; p.Arg7X</td>
<td>Hom</td>
<td>Recently reported in 3 families with lethal arthrogryposis multiplex congenita (29)</td>
</tr>
<tr>
<td>HOU2682</td>
<td>BAB7302</td>
<td>Contractures of distal joints, delayed neuromotor development, hypotonia, swallowing difficulty, pyloric stenosis, tibia vara, enlarged metaphyses, and multiple milia</td>
<td>LIFR</td>
<td>c.274C&gt;T; p.Gln92X</td>
<td>Het</td>
<td>SWS (OMIM 601559)</td>
</tr>
<tr>
<td>HOU2616</td>
<td>BAB7125</td>
<td>Contractures of hands and elbows, severe developmental delay, swallowing difficulty, high-arched palate, pectus excavatum, scoliosis, hypertrophy in calf muscles, and mild mitral valve insufficiency</td>
<td>MYBPC2</td>
<td>c.707C&gt;T; p.Thr236Ile</td>
<td>Het</td>
<td>Recently associated with perisylvian polymicrogyria, cerebellar hypoplasia and arthrogryposis (33)</td>
</tr>
<tr>
<td>HOU2620</td>
<td>BAB7140</td>
<td>Preterm birth, delayed motor milestones, contractures of hands and elbows, webbed neck, high-grade rotoscoliosis, congenital ptosis, vertebral segmentation and fusion defects</td>
<td>MYO18B</td>
<td>c.6322C&gt;T; p.Arg2108X</td>
<td>Hom</td>
<td>Recently reported in 2 families with Klippel-Feil anomaly, myopathy, and DA (32)</td>
</tr>
<tr>
<td>HOU2621</td>
<td>BAB7143</td>
<td>Contractures of hands and feet (DA), delayed motor milestones, myotonia, microcephaly, difficulty in swallowing solid food, constipation, congenital ptosis, hip dislocation, TEV, ECHO: Bicuspid aorta, MRI: Cerebellar vermian hypoplasia and Dandy-Walker malformation</td>
<td>VPS8</td>
<td>c.3130G&gt;A; p.Val1044Ile</td>
<td>Hom</td>
<td>Recently reported in 2 families with Klippel-Feil anomaly, myopathy, and DA (32)</td>
</tr>
<tr>
<td>HOU2791</td>
<td>BAB7713</td>
<td>Rupture of the patellar ligament</td>
<td>RIPK4</td>
<td>c.1681G&gt;A; p.Val561Met</td>
<td>Hom</td>
<td>Patellar pterygium syndrome 2, lethal type (OMIM 263650)</td>
</tr>
</tbody>
</table>

AD, autosomal dominant; AR, autosomal recessive; ECHO, echocardiogram; Het, heterozygous; Hom: homozygous; TEV, talipes equinovarus.
and cryptorchidism. Individual BAB6827 from family HOU2530 had severe camptodactyly of the hands, bilateral hip dislocation, right talipes equinovarus, hypotonia, had developmental delay, cryptorchidism, nephrocalcinosis detected in the sonogram of the abdomen, and thin corpus callosum. He also had distinctive facial features of a mask-like face, high-arched eyebrows, a bulbous and upturned nose, low-set ears, and micrognathia (Figure 1B). Moreover, given the consanguinity between the parents in these 4 families, we examined the absence of heterozygosity (AOH) regions in all probands with ECELI mutations. Haplotype block analysis based on single nucleotide polymorphism data culled from exome sequencing (i.e., B-allele frequency) revealed that the maximal shared region of AOH in the 4 families was approximately 840 Kb on the distal long arm of chromosome 2, including the ECELI gene. Since there are 3 different mutations, the actual haplotypes in these AOH blocks are presumed to differ (Figure 1C).

In 6 patients, we identified 4 different homozygous variants in CHRNG, a known causal gene for lethal and nonlethal MPS. The identified variants include 1 stop-gain (c.241C>T; p.Gln81X), 1 frameshift deletion (c.753_754delICT; p.Val253Alafs), 1 missense mutation (c.256C>T; p.Arg86Cys), and 1 recurrent missense mutation (c.715C>T; p.Arg239Cys), which was observed in 3 patients from reportedly unrelated families (Figure 2A). The common findings in these patients were prenataly detected restriction of movements; craniofacial dysmorphisms, including down-slanting palpebral fissures, low-set ears, and micrognathia; and multiple contractures of the limbs (Figure 2B). The distribution of CHRNG mutations detected in our study is shown in Figure 2, C and D. Two of the identified CHRNG mutations have not been reported, while the other mutations, including the recurrent mutation, have been reported previously in MPS patients (20, 36). In 1 of 3 patients with the same homozygous CHRNG mutation (BAB5611), we additionally identified another homozygous deleterious mutation in a different arthrogryposis gene, ERC22 (c.1775C>G; p.Arg592His) in 3 affected kindred with antecubital pterygium syndrome (COFS) type 2 (OMIM 610756), and 2 other allelic disorders (trichoheridystrophy [OMIM 601675] and xeroderma pigmentosum, group D [OMIM 278730]).

Other identified variants in known genes are delineated in the following sections. Although 17 of these families are singleton cases with different mutations in different known genes (Table 1), we observed 8 families with an arthrogryposis gene mutational load, as we observed in family HOU2163 with CHRNG and ERC22 mutations (Table 2). These families have another homozygous or compound heterozygous predicted deleterious variant in an additional known gene or in a novel gene that cannot be excluded from having a potential clinical contribution because of its function in arthrogryposis formation or close interactions with known arthrogryposis gene products.

After exclusion of known genes, we examined patient genomes for novel genes, investigating all potential inheritance models. Our strategy of choosing best-candidate genes was based on function of the gene; interactions with other known arthrogryposis genes; functional prediction scores calculated by different computational algorithms (PolyPhen-2, MutationTaster, SIFT, LRT, and Phylop); review of our internal database (~4,800 exomes), including the Turkish subpopulation (~630 exomes), or other available databases such as the 1,000 Genomes Project (http://www.1000genomes.org), the Exome Variant Server, NHLBI GO Exome Sequencing Project (ESP; Seattle, Washington, USA; http://evs.gs.washington.edu/ EVS/), the Atherosclerosis Risk in Communities Study (ARIC) database (http://drupal.cscs.unc.edu/ aric/), and the Exome Aggregation Consortium (ExAC, http://exac.broadinstitute.org/) databases to examine the rarity of the identified variants; and segregation in the family studied. Using this approach, we identified compound heterozygous variants in FBN3 (c.1883C>T; p.Ser628Phe and c.8357G>A; p.Gly2786Asp) in a patient with contractions of the elbows and knees, compound heterozygous variants in MYO9A (c.6845G>A; p.Gly2282Glu and c.608A>G; p.Tyr203C) in a patient with multiple limb contractures, and a heterozygous missense PSD3 variant (c.437T>C; p.Leu146Thr) in 3 affected kindred with antecubital pterygium syndrome (APS) (OMIM 178200) (Table 3).

### Discussion

In our study, we define a general approach to investigate the molecular etiology underlying arthrogryposis by using WES in a patient cohort of subjects with different clinical subtypes of arthrogryposis. We identified known and novel variants in known genes and variants in potential novel candidate genes, and we provide evidence for phenotypic expansion in a few cases. Moreover, some subjects were found to have evidence for an apparent mutational load with disease-associated variants in more than one arthrogryposis gene locus and to be manifesting a more severe clinical phenotype.

The most common known arthrogryposis-associated genes identified in our cohort were ECELI and CHRNG. While the phenotypes associated with these variants were overall consistent with those reported, some patients had novel clinical features. For instance, BAB6500 had a distinct whistling face, which is commonly observed with MYH3 variants in Freeman-Sheldon syndrome (also known as DA type 2A) (OMIM 193700) (9). MYH3 encodes one of the contractile proteins of skeletal muscle, while ECELI mutations result in perturbed formation of the neuromuscular junction. Although developmental pathways that cause DA type 2A and DA type 5D are different, the observed whistling-face finding in an ECELI mutated case not only shows the complexity of the phenotypic spectrum seen in this heterogeneous subgroup of arthrogryposis, but also underscores the limitation of clinical characterization of arthrogryposis subtypes. Patient BAB6827 had hypoplasia of the corpus callosum (HCC), in addition to DA. Given the functional and expression patterns of ECELI in conjunction with neurologic findings in the patient (i.e., developmental delay and hypotonia), it is feasible to suggest that HCC may be part of the phenotype. Alternatively, there may be a yet unidentified second molecular diagnosis in this patient.

Families consistent with a potential oligogenic or mutational burden model. In patient BAB5611, a homozgyous variant in ERC22 was detected, in addition to a homozygous variant in CHRNG (Figure 2A). Mutations in ERC22 have previously been reported in patients with COFS type 2 (37), a progressive neurodegenerative and arthrogryposis-related disorder. Vogt et al. and Morgan et al. reported a family with a homozygous missence CHRNG mutation; 2 affected siblings died in early infancy, while the third
Figure 1. Segregation analyses, photographs, and AOH regions of the patients with ECEL1 mutations. (A) Detailed pedigrees of the families and their corresponding Sanger-sequencing chromatograms showing the segregation analyses. (B) Photographs of the patients. Note the whistling-face appearance of BAB6500, which is commonly observed in DA type 2A rather than ECEL1-associated DA type 5D. (C) AOH study of the patients based on calculated B-allele frequency data culled from WES analysis. Gray shaded areas indicate AOH regions. Note the approximately 840-Kb overlap between the AOH regions, which includes the ECEL1 gene.
Figure 2. Segregation results, photographs, and variant distributions of the patients with CHRNG mutations. (A) Pedigrees and Sanger sequencing analyses of the patients that represent the proper segregation of the WES-detected CHRNG variants. One of the patients with more severe phenotype (BAB5611) has another homozygous variant in a known arthrogryposis gene (ERCC2), which also segregates in the family. (B) Photographs of the probands showing the major clinical features, including joint contractures, multiple pterygiums, and distinct facial dysmorphism. (C and D) Schematic representations of gene and protein structures of CHRNG and localization of the identified variants. Neur_chan_LBD, neurotransmitter-gated ion-channel ligand binding domain; Neur_chan_memb, neurotransmitter-gated ion-channel transmembrane region.
affected sibling had nonlethal MPS (15, 20). Moreover, the same family had 4 additional cousins with nonlethal MPS phenotype and 1 deceased cousin. They concluded that, since the family represented a single observation, the significance of the clinical finding of variability of expression was unclear. We suggest that mutational burden or genetic load may contribute to clinical variability and disease severity, as has been shown in patients with Charcot-Marie-Tooth disease, wherein genetic burden contributes to phenotypic variability and complex neuropathy (38).

Variants in both MYO18B and MYH7B that were identified in BAB7140 (Figure 3A) were recently reported in 2 separate studies in patients with common clinical findings of myopathy and vertebral anomalies. Alazami et al. identified a null homozygous mutation in MYO18B in 2 patients from 2 unrelated families that shared a similar phenotype of Klippel-Feil anomaly, myopathy, arthrogryposis, microcephaly, and distinctive facies (32). Esposito et al. identified 2 homozygous mutations in MYH7B and ITGA7 in a patient with congenital myopathy, scoliosis, and cardiomyopathy, and they concluded that a synergistic effect of these 2 mutations results in a severe phenotype (39). Our patient was a 14-year-old female and presented with more severe clinical findings, suggesting a blended phenotype (Figure 3B, Table 4, and ref. 40). The most remarkable finding in our patient was a high-grade thoracolumbar rotoscoliosis with cervical vertebral segmentation and fusion defects (Figure 3B). We compared the clinical features and identified mutations of our patient with those reported in 2 studies (Table 4). Our data strongly suggest that the mutational burden from the variants observed at the 2 genetic loci, MYO18B and MYH7B, leads to a more severe and blended phenotype of arthrogryposis and vertebral anomalies, including high-grade scoliosis and fusion defects. Additionally, AOH findings based on data culled from WES showed that both MYO18B and MYH7B variants are surrounded by AOH blocks on chromosome 22 and 20, respectively. Interestingly, the AOH data detected on chromosome 20 revealed a large AOH block (53.7 Mb) that nearly encompassed the entire chromosome (~85%), suggesting a possible uniparental disomy (Figure 3C), as has been recently reported with clinical exome sequencing for other recessive disease traits. Apart from the blended phenotype outcome of different homozygous variants, the predicted structural similarity between these 2 myosin genes leads us to speculate that variants observed in these 2 genes in the same patient may induce a more severe disruption of the myosin gene network and cause a much more severe phenotype.

In family HOU2278, which had 2 affected siblings, we identified a homozygous frameshift mutation in COL6A3 in both patients. An additional homozygous missense mutation in BICD2 was also observed in the more severely affected elder brother, while the sister was a heterozygous carrier for this variant (Table 2). Mutations of both COL6A3 and BICD2 genes have previously been reported in muscle diseases comprising arthrogryposis (41, 42). However, while homozygous mutations of COL6A3 were reported in patients with Ullrich congenital muscular dystrophy (OMIM 254090), only heterozygous mutations of BICD2 were reported in patients with spinal muscular atrophy, lower extremity–predominant, type 2 (OMIM 615290). Our data suggest that the more severe phenotype observed in the brother might be a consequence of the mutational burden at the 2 loci and the biallelic homozygous inheritance of the BICD2 mutation.

In patient BAB6212, we identified a homozygous stop-gain mutation in GPR126, which was recently reported as a novel gene in 3 families with arthrogryposis (29). However, in this patient, WES data also revealed compound heterozygous variants in another novel gene, MYBPC2 (Figure 4A). MYBPC2 mutations were not previously associated with any Mendelian disease; however, mutations of MYBPC1, another myosin-binding protein, have previously been associated with DA type 1B (OMIM 614335) and lethal congenital contracture syndrome 4 (OMIM 614915) (10, 43). MYBPC proteins play structural and regulatory roles in muscle function. These proteins provide thick filament stability and modulate contractility through dynamic interactions with the head region of myosin and actin (44). Apart from these functional and molecular data, the mutated amino acids Thr236 and Ser255 are highly conserved in vertebrates (Figure 4B), and MYBPC2 has close interactions with various known arthrogryposis genes (Figure 4C). Although GPR126 mutations were recently associated with arthrogryposis, MYBPC2 may also be considered as a contributory gene to the patient’s phenotype due to the gene function and interactome structure of its protein product.
In 2 patients (BAB7302 and BAB7125), we identified different homozygous stop-gain mutations in LIFR, consistent with Stuve-Wiedemann syndrome (SWS) (OMIM 601559) (45). In both cases, we also identified potential contributory variants in other known genes (Table 2). Patient BAB7302 had compound heterozygous mutations in PI4KA, a gene recently associated with brain malformation and arthrogryposis (33). Patient BAB7125 had compound heterozygous variants in MYH14 (Table 2), which encodes a member of the nonmuscle myosin II family of ATP-dependent molecular motors that interact with cytoskeletal actin and regulate cytokinesis, cell motility, and cell polarity (46). Heterozygous mutations of MYH14 were reported in a family with a complex phenotype of peripheral neuropathy, myopathy, hoarseness, and hearing loss (47). Our patient was a 15-month-old male and had severe developmental delay, multiple joint contractures, pectus excavatum, scoliosis, hypertrophy in calf muscles, and mild mitral valve insufficiency. Although our patient had a normal hearing test and creatine kinase level, the neurologic findings and calf enlargement, which is a typical clinical feature in neuromuscular diseases, suggest that the compound heterozygous mutations of MYH14, in addition to LIFR homozygous mutation, may contribute to the severe phenotype observed in our patient due to a potential mutational burden effect.
In patient BAB7143, a homozygous missense variant in a novel gene, VPS8, was identified (Figure 5A). The patient had multiple joint contractures, delayed motor milestones, craniofacial dysmorphism, and cerebellar vermis hypoplasia detected in cranial magnetic resonance imaging (MRI) (Figure 5B and Table 2). VPS8 is one of the Vps proteins that functions in endosomal biogenesis and Golgi-to-lysosome trafficking in yeast (48, 49). Homozygous or compound heterozygous mutations in other Vps genes, VPS33B and VPS53, were previously associated with arthrogryposis–renal dysfunction–cholestasis syndrome (ARCS) type 1 (OMIM 208085) and pontocerebellar hypoplasia type 2E (PCH2E) (OMIM 615851), respectively (50, 51). Joint contractures and cerebellar anomalies were previously defined in patients with PCH2E (52). Moreover, VPS8 has close interactions with VPS33B and VPS53, as well as VIPAS39 (Figure 5C), and mutations of VIPAS39 were associated with ARCS type 2 (OMIM 613404) (53). Thus, VPS8 is a potential candidate gene for the arthrogryposis and MRI findings observed in our patient. Since the observed brain anomalies in the subject are not common findings in disorders with arthrogryposis, we further examined the exome data for an additional variant that may explain the MRI findings, and we identified another homozygous missense variant in POLR3A. POLR3A is not a known gene for arthrogryposis, but homozygous mutations in POLR3A have been associated with hypomethylating leukodystrophy-7 (HLD7) (OMIM 607694) (54). HLD7 is an autosomal recessive neurodegenerative disorder characterized by childhood onset of progressive motor decline manifest as spasticity, ataxia, tremor, cortical and cerebellar atrophy, as well as mild cognitive regression. The cerebellar component of HLD7 suggests that the identified POLR3A variant may have a role in the intracranial findings in this patient. The missense changes in VPS8 and POLR3A both affect highly conserved valine residues on the respective proteins (Figure 5D).

In another patient (BAB7713) with arthrogryposis and cardiomyopathy findings, we identified homozygous mutations in 2 different known genes, RIPK4 and LMNA. The major clinical features of the subject were contractures of hands and feet, myopathy, hypotonia, and dilated cardiomyopathy detected by echocardiogram. RIPK4 is a known gene for popliteal pterygium syndrome type 2 (OMIM 263650), and LMNA mutations were reported in several allelic disorders, including dilated cardiomyo-

Table 4. Comparison of the genotype and clinical findings in patient BAB7140 harboring MYO18B and MYH7B mutations with those of recently reported patients that represent similar findings

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Alazami et al. (ref. 32)</th>
<th>Esposito et al. (ref. 39)</th>
<th>Present study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Identified gene(s)</td>
<td>MYO18B</td>
<td>MYO18B</td>
<td>MYH7B</td>
</tr>
<tr>
<td>Variant type</td>
<td>Nonsense</td>
<td>Nonsense</td>
<td>Missense</td>
</tr>
<tr>
<td>Zygosity</td>
<td>Homozygous</td>
<td>Homozygous</td>
<td>Homozygous</td>
</tr>
</tbody>
</table>

| Ethnic origin | Arab | Arab | Italian | Turkish |
| Consanguinity | Yes | Yes | Yes | Yes |
| Sex | Male | Female | Female | Female |
| Premature delivery | Yes | No | N/A | Yes |
| Delayed motor development | Yes | Yes | Yes | Yes |
| Hypotonia | Yes | Yes | Yes | Yes |
| Short stature | Yes | Yes | N/A | Yes |
| Microcephaly | Yes | Yes | N/A | No |
| Ptosis | Yes | Yes | No | Yes |
| Arched eyebrows | Yes | Yes | No | Yes |
| Hypertelorism | Yes | Yes | No | Yes |
| Bulbous nose | Yes | Yes | N/A | Yes |
| Low-set ears | No | Yes | No | Yes |
| Micrognathia | No | Yes | Yes | Yes |
| Low posterior hair line | Yes | Yes | No | Yes |
| Webbed neck | No | Yes | No | Yes |
| Arthrogryposis | Yes | No | No | Yes |
| Vertebral fusion defect | Yes | Yes | No | Yes |
| Scoliosis | No | Yes | Yes (High grade) | Yes |
| Hip deformity | No | Yes | Yes | No |
| Creatine kinase | Normal | Normal | Normal | Normal |
| Electromyography | Myopathy | N/A | N/A | N/A |
| Muscle biopsy | Pathologic | N/A | Pathologic | N/A |
| Echocardiography | N/A | N/A | Cardiomyopathy | Normal |
deviation of hands, talipes equinovarus, and delayed bone age
measured via hand X-ray (Supplemental Figure 3). Besides these,
our patient was not microcephalic, and his growth parameters
were in the normal range when measured at 6 years of age, unlike
typical SCKL patients. Additionally, another remarkable finding
of our patient was abnormal scar formation with wound healing
observed on his left foot (Supplemental Figure 3), which sug-
gests possible connective-tissue involvement in his phenotype.
Further analysis of the exome data did not reveal a reasonably
plausible variant that might explain the abnormal scar formation
in the patient. Clinical features of our patient overlapping with
SCKL and distinct findings including arthrogryposis and abnor-
mal wound healing suggest that the homozygous mutation of
CENPJ observed in our patient may cause another allelic disor-
der apart from SCKL type 4 and MCPH-6 or may represent phe-
notypic expansion of CENPJ-related diseases.

In one male patient (BAB3955), we identified a hemizygous
variant in IDS, which is a known gene for mucopolysaccharidosis
type 2 (MPS2) (OMIM 309900). MPS2 is an X-linked metabolic
disorder caused by deficiency of the lysosomal enzyme iduronate
sulfatase that leads to progressive accumulation of glycosaminoglycans in nearly all cell types. The typical clinical features include severe airway obstruction; skeletal deformities, including joint
cartilages; cardiomyopathy; and neurologic decline (57). The

Patients that represent phenotypic expansion. In 3 patients, we
identified novel homozygous variants in known genes; however,
the clinical findings we observe in association with the newly
described variant alleles suggest phenotypic expansion for trait
manifestations previously attributed to this gene. In patient
BAB3931, we identified a novel homozygous missense mutation in
RIPK4, the causal gene for Seckel syndrome (SCKL) type 4
(OMIM 613676) and autosomal recessive microcephaly type 6
(MCPH-6) (OMIM 608393). Arthrogryposis is not a typical find-
ing of SCKL type 4, but elbow-flexion contracture, hip disloca-
tion, and talipes deformities were defined in SCKL type 1 (OMIM
210600). Moreover, Sarici et al. reported a Turkish patient
with SCKL, accompanied by semilobar holoprosencephaly and
arthrogryposis (56). The clinical findings of our patient over-
lapping with SCKL and distinct findings including arthrogryposis and abnor-
mal wound healing suggest that the homozygous mutation of
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contractures; cardiomyopathy; and neurologic decline (57). The
hepatosplenomegaly, and liver cirrhosis leading to death in early childhood (58). However, GSD4 can show extreme clinical heterogeneity, and different clinical manifestations may present, such as nonprogressive hepatic form and neuromuscular forms. Arthrogryposis is one of the characteristic findings of the congenital neuromuscular form of GSD4, which is mostly seen with hydrops fetalis and perinatal death. There are also juvenile and adult neuromuscular forms that are dominated by myopathy, and the classic clinical manifestation of liver cirrhosis generally is not present in these neuromuscular forms (59–61). The diagnosis of GSD4 can be confirmed by the determination of the branching enzyme activity in affected tissues. However, the commonly used assays are indirect and not sensitive enough for accurate evaluation of low levels of branching activity (60, 62). In our 2-year-old male patient, the presence of joint contractures was compatible with the congenital neuromuscular form, but the clinical manifestation was not severe. Additional patient findings suggesting phenotypic expansion were cleft palate, pterygium colli, pelvicaliectasis, cryptorchidism, and tall vertebral bodies detected on X-ray, which, to our knowledge, had not been reported previously.

We identified a homozygous novel variant in GBEI in patient BAB3960. The mutations in GBEI were previously associated with glycogen storage disease type 4 (GSD4) (OMIM 232500). Glycogen branching enzyme deficiency is causative of GSD4 that is characterized by the accumulation of polyglucosan in almost all tissues. Clinical features of the typical GSD4 are failure to thrive, hepatosplenomegaly, and liver cirrhosis leading to death in early childhood (58). However, GSD4 can show extreme clinical heterogeneity, and different clinical manifestations may present, such as nonprogressive hepatic form and neuromuscular forms. Arthrogryposis is one of the characteristic findings of the congenital neuromuscular form of GSD4, which is mostly seen with hydrops fetalis and perinatal death. There are also juvenile and adult neuromuscular forms that are dominated by myopathy, and the classic clinical manifestation of liver cirrhosis generally is not present in these neuromuscular forms (59–61). The diagnosis of GSD4 can be confirmed by the determination of the branching enzyme activity in affected tissues. However, the commonly used assays are indirect and not sensitive enough for accurate evaluation of low levels of branching activity (60, 62). In our 2-year-old male patient, the presence of joint contractures was compatible with the congenital neuromuscular form, but the clinical manifestation was not severe. Additional patient findings suggesting phenotypic expansion were cleft palate, pterygium colli, pelvicaliectasis, cryptorchidism, and tall vertebral bodies detected on X-ray, which, to our knowledge, had not been reported previously.
The best of our knowledge, were not reported in GSD4 patients previously. Also, the general hypotonia detected in the subject was compatible with the juvenile form. We suggest that our data support the clinical heterogeneity of GSD4 and expand the phenotypic spectrum with the novel clinical findings.

Patients with rare deleterious variants in a single novel candidate gene. We identified compound heterozygous variants in FBN3 in patient BAB7826 (Figure 6A). The subject had multiple joint contractures with severely affected elbows and knees (Figure 6B). FBN3 is one of the fibrillin genes that function in the structural architecture of connective tissues; mutations of FBN3 have not been reported in any Mendelian disorder. Other fibrillin genes, FBN1 and FBN2, were previously associated with disorders comprising arthrogryposis, but FBN3 was not. Mutations of FBN1 were described in autosomal dominant Weill-Marchesani syndrome (OMIM 277600), and FBN2 mutations are associated with DA type 9, also known as Beals syndrome (OMIM 121050). Our data lead us to speculate that mutations of FBN3 may also cause arthrogryposis, as found for other fibrillin gene mutations.

The exome analysis of BAB6499 revealed 2 heterozygous variants in MYO9A, one of which is de novo and the other inherited from the mother (Figure 7A). The subject had DA with predominant involvement of the fingers and feet (Figure 7B). Mutations of MYO9A were not previously associated with any human disease. MYO9A is one of the unconventional myosins that are members of the myosin superfamily that display the general domains of conventional myosins (63, 64). In a mouse study, Northern blot analysis detected mouse Myo9a expression in limb buds, and in situ hybridization to limb buds suggested that Myo9a may be localized to the precartilagenous mesenchyme (65). Also in the same study, Myo9a was shown to be expressed in the skeletal and the nervous systems, 2 important systems where dysfunction may play a role in the formation of arthrogryposis and consistent with the hypothesis that mutations of MYO9A may cause arthrogryposis. Moreover, the mutated amino acids Gly2282 and Tyr203 are highly conserved in vertebrates (Figure 7C), and MYO9A has interactions with some other myosins that have previously been associated with arthrogryposis disorders (Figure 7D). The segregation pattern of identified variants in our patient suggests that deleterious de novo mutations or compound heterozygous mutations of MYO9A may cause arthrogryposis.

In family HOU2061 with 8 affected individuals diagnosed with APS, we applied WES to 3 affected cousins, and a heterozygous variant in PSD3 was detected and shown to cosegregate with autosomal dominant arthrogryposis. To confirm this identified variant, we analyzed 12 individuals from 4 generations in the pedigree with available DNA (8 affected, 4 unaffected) by Sanger sequencing. We observed that all affected individuals were carriers for the same mutation, while the unaffected kindred were normal for the same allele (Figure 8A). Additionally, we calculated the logarithm of odds (LOD) score, and we found that, based on the family size, our data (θ = 0, LOD score = 1.806) supports the potential linkage of PSD3 to the phenotype. The function of PSD3 is not well established, and mutations of PSD3 were not reported in any Mendelian disorder; however, the association of PSD3 with systemic sclerosis—which is a heterogeneous disorder characterized by extensive skin fibrosis, microvascular changes, and autoimmunity—was reported (66). The skin fibrosis component of systemic sclerosis might be a clue for explaining the phenotype observed in APS patients with PSD3 mutation by reason of the fact that the main phenotype observed in APS is skin webbing around the elbow joint (Figure 8B). Nevertheless, the biological mechanism underlying APS and molecular pathway of PSD3 needs to be investigated further.

In summary, beside the known and novel variants in known genes, we identified variants in 5 potential novel candidate genes (FBN3, MYBPC2, MYO9A, PSD3, and VPS8) in patients with different arthrogryposis subtypes. Three of 5 novel genes were observed as a single causal gene, while variants in 2 novel candidate genes were identified as a second locus in addition to a known gene that may affect the phenotype with a mutational load or may cause a blended phenotype. Moreover, we identified a potential mutational burden in 6 patients with homozygous or compound heterozygous mutations in 2 different known genes. In 4 of 6 patients, both identified genes were previously associated with an arthrogryposis disorder, and the mutational burden may have contributed to the clinical severity observed (38); meanwhile, in 2 of 6 patients, the identified genes—in addition to known arthrogryposis genes—were associated with different disorders without arthrogryposis and resulted in an apparent blended phenotype with clinical manifestations of both disorders (40, 67). Arthrogryposis is observed in a broad group of genetically heterogeneous disorders with unknown molecular etiology in the considerable majority. In our study, the WES method enabled us to investigate potential oligogenic or mutational burden models, as well as to
identify novel candidate genes in arthrogryposis patients. We suggest that using genomic sequencing methods in patient populations with arthrogryposis disorders may identify novel arthrogryposis genes and provide further molecular etiological insights into this phenotype, as well as provide opportunities to uncover novel genetic mechanisms, including potential contributions of multilocus variation.

Methods

Human subjects and sample collection. Fifty-one Turkish patients and 1 Arab patient with arthrogryposis were evaluated by 1 or more pediatricians and/or clinical geneticists. Genomic DNA was extracted from whole blood using the Gentra Puregene Blood Extraction Kit per the manufacturer’s protocol (QIAGEN). All the genomic studies were performed on the DNA samples extracted from whole blood.

WES and data analysis. Samples from all patients underwent WES at Baylor College of Medicine Human Genome Sequencing Center through the Baylor-Hopkins Center for Mendelian Genomics research initiative. Genomic DNA samples obtained from patients processed according to protocols previously described (68). Briefly, DNA sample was prepared into Illumina paired-end libraries and underwent whole-exome capture using BCM-HGSC core design (52 Mb, NimbleGen, Roche Sequencing), followed by sequencing on the Illumina HiSeq 2000 platform with an approximately 150× depth of coverage. Data produced was aligned and mapped to the human genome reference sequence (hg19) using the Mercury pipeline. Variants were called using the ATLAS (an integrative variant analysis pipeline optimized for variant discovery) variant calling method and SAMtools (The Sequence Alignment/Map) and annotated using the in-house–developed Cassandra annotation pipeline that uses ANNOVAR (http://annovar.openbioinformatics.org/en/latest/) (69–71).

The findings of WES studies were deposited into the NCBI’s database of Genotypes and Phenotypes (dbGaP) archive (phs000711.v3.p1) (http://www.ncbi.nlm.nih.gov/gap/).

PCR validation and segregation analyses. To confirm the identified WES-detected variant, we applied Sanger sequencing following PCR amplification, and segregation analyses were performed on all DNA-available members of the families. Same pairs of primers are...
particular homozygous/hemizygous deletion was ≥0.5% in the whole cohort — and calls from consecutive exons were merged.

RPKM thresholds were determined based on the analysis of distribution of RPKM values in previously identified and confirmed homozygous deletions. Finally, we filtered out homozygous CNVs that did not overlap with larger (≥1 Kb) AOH regions. RPKM values were also used for further visualization of detected deletions.

Interaction network analysis. As a part of our approach to identify novel candidate genes, we performed interaction network analyses. The predicted interaction network had been based on 6 data sources, including coexpression, colocalization, pathways, physical interactions, predicted interactions, and shared protein domain data sources. Being predicted to be included in a potential interaction network with known disease genes may further support the novel gene association with the disease.

Study approval. This study was approved by the Institutional Review Board at Baylor College of Medicine. Informed consent was obtained from all subjects prior to enrollment in the project. Additionally, written consent was obtained from patients or patients' parents to publish patient photos.

utilized for both PCR amplification and Sanger sequencing. PCR reactions were performed following the protocols for HotStarTaq DNA Polymerase (QIAGEN).

AOH and copy number variation (CNV) analysis. To examine AOH regions surrounding candidate variants, we calculated B-allele frequency using WES data as a ratio of variant reads to total reads. These data were then processed using the circular binary segmentation algorithm to identify AOH regions (72).

To identify heterozygous CNVs, we used WES data. WES data were processed using CoNiFER software (73) and HMZDelFinder (https://github.com/BCM-Lupskilab/HMZDelFinder). HMZDelFinder is an in-house–developed algorithm implemented in the R programming language (R Core Team 2014, http://www.R-project.org). First, for every individual, we computed the total number of reads in each exon and normalized read-depth values (reads per kilobase per million [RPKM] mapped reads) using the utility provided with CoNiFER (73). Next, we identified homozygous deletions by analyzing exons for which the RPKM value was lower than 0.5. Then, low-quality samples (5% of individuals with the highest number of deletions) were removed. Next, low-quality/common deletions were removed— if the frequency of a particular homozygous/hemizygous deletion was ≥0.5% in the whole cohort — and calls from consecutive exons were merged.

RPKM thresholds were determined based on the analysis of distribution of RPKM values in previously identified and confirmed homozygous deletions. Finally, we filtered out homozygous CNVs that did not overlap with larger (≥1 Kb) AOH regions. RPKM values were also used for further visualization of detected deletions.

Interaction network analysis. As a part of our approach to identify novel candidate genes, we performed interaction network analyses. The predicted interaction network had been based on 6 data sources, including coexpression, colocalization, pathways, physical interactions, predicted interactions, and shared protein domain data sources. Being predicted to be included in a potential interaction network with known disease genes may further support the novel gene association with the disease.

Study approval. This study was approved by the Institutional Review Board at Baylor College of Medicine. Informed consent was obtained from all subjects prior to enrollment in the project. Additionally, written consent was obtained from patients or patients’ parents to publish patient photos.
Author contributions

YB participated in study design, analyzed the WES data, and wrote the manuscript. EK, TH, and DP contributed to clinical assessment of the patients and WES analyses. ZCA and TG participated in computational and bioinformatics design and analysis. ZCA performed copy number variation and interaction network analyses. EOY, GAT, HA, DT, STB, AG, SI, AWEH, WLC, AK, TC, OOV, TY, IAB, NE, and BT participated in clinical evaluation of the patients and sample collection. MMA participated in DNA extraction, PCR validation, and figure production for the manuscript. SNJ, DMM, EB, and RAG organized the WES data analyses. JRL participated in study design, data organization, management and analyses, and writing of the manuscript. All coauthors reviewed the manuscript.

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