Short Report

Homozygous null mutations in \textit{ZMPST24} in restrictive dermopathy: evidence of genetic heterogeneity


Restrictive dermopathy (RD) results in stillbirth or early neonatal death. RD is characterized by prematurity, intrauterine growth retardation, fixed facial expression, micrognathia, mouth in the ‘o’ position, rigid and tense skin with erosions and denudations and multiple joint contractures. Nearly all 25 previously reported neonates with RD had homozygous or compound heterozygous null mutations in the \textit{ZMPST24} gene. Here, we report three new cases of RD; all died within 3 weeks of birth. One of them had a previously reported homozygous c.1085dupT (p.Leu362PhefsX19) mutation, the second case had a novel homozygous c.1020G>A (p.Trp340X) null mutation in \textit{ZMPST24}, but the third case, a stillborn with features of RD except for the presence of tapering rather than rounded, bulbous digits, harbored no disease-causing mutations in \textit{LMNA} or \textit{ZMPST24}. In the newborn with a novel \textit{ZMPST24} mutation, unique features included butterfly-shaped thoracic 5 vertebra and the bulbous appearance of the distal clavicles. Skin biopsies from both the stillborn fetus and the newborn with c.1020G>A \textit{ZMPST24} mutation showed absence of elastic fibers throughout the dermis. This report provides evidence of genetic heterogeneity among RD and concludes that there may be an additional locus for RD which remains to be identified.

Conflict of interest

All the authors declare no conflict of interest.

Restrictive dermopathy (RD; online mendelian inheritance in man (OMIM) 275210) results in stillbirth or early neonatal death and is characterized by prematurity, intrauterine growth retardation (IUGR), fixed facial expression, micrognathia, mouth in the ‘o’ position, rigid and tense skin with erosions and denudations, and multiple joint contractures. Nearly all neonates with RD have homozygous or compound heterozygous null mutations in the zinc metalloproteinase (\textit{ZMPST24}) gene (1–7). \textit{ZMPST24} is critical for post-translational proteolytic cleavage of carboxy-terminal residues of prelamin A in two steps to form mature lamin A (8), an integral nuclear lamina protein encoded by lamin A/C (\textit{LMNA}) gene.

Thus far, 25 patients with RD have been reported to harbor \textit{ZMPST24} mutations (1–7, 9–11),
but many of them lacked detailed phenotypic information (4, 5). Here, we report clinical features and genotyping of three new cases of RD.

**Patients and methods**

A written informed consent was obtained from the parents of RD 600.3 and RD 500.3, and the study was approved by the Institutional Review Board at UT Southwestern. A written consent for autopsy was obtained from the parents of RD 200.3 at the Sanjay Gandhi Post Graduate Institute of Medical Sciences in Lucknow, India.

RD 600.3

This female infant was born via vaginal delivery at 34 weeks of gestation to non-consanguineous, healthy parents (Fig. 1) with one previous stillbirth who had features of RD. A prenatal diagnosis of 70% mosaic trisomy 20 was made. Prenatal ultrasound revealed a short chest and small stomach. Polyhydramnios was noted early on, but in the few weeks prior to delivery, oligohydramnios was present. Mother underwent oxytocin augmentation for non-reassuring fetal heart tracings. The birth weight was 1615 g. Apgar scores were 1 at 1 min and 5 at 5 min. The baby was immediately resuscitated, intubated and transferred to the neonatal intensive care unit.

Upon examination, she had tight, fragile skin with prominent superficial vasculature. An 8–9 cm laceration extended circumferentially around the neck from the left ear to past the right ear toward the back of the head. It was estimated to be 8–10 mm deep and had outward curling ragged edges along the entire length. The right external
jugular vein appeared exposed. There was minimal range of motion of the large and small joints. Her mouth was open and tightly fixed in an O-shaped position. There was almost no nasal bridge with a minimal pointed tip. Hands were small with relatively long fingers, and nails were unremarkable. She was also noted to have large fontanelles, low-set and posteriorly rotated ears, downslanting eyes, and micrognathia. An echocardiogram showed a small patent ductus arteriosus shunting left to right and radiographs showed short dysplastic clavicles. After a prolonged hospital course requiring mechanical ventilation, antibiotics for severe sepsis, total parental nutrition, and non-oliguric renal insufficiency, the infant died on day-of-life 16.

RD 500.3

This female infant was born via vaginal delivery at 33 weeks of gestation to consanguineous parents of Mexican origin (Fig. 1). The mother had preterm premature rupture of membranes. An abdominal sonogram showed IUGR. Labor was complicated by severe pre-eclampsia requiring magnesium sulfate therapy and oxytocin augmentation for non-reassuring fetal heart tracings. Upon delivery, the baby was cyanotic and hypotonic. The birth weight was 1192 g. Apgar scores were 1 at 1 min and 4 at 5 min. The baby was immediately resuscitated, intubated and transferred to the neonatal intensive care unit.

Upon examination, she had thin, taught, translucent skin with prominent superficial vasculature. Extremities were rigid with flexural joint contractions at the elbows, fingers, knees, and ankles and extensor contractures of the wrists. Her mouth was in a fixed O-shaped position. Fingers showed rounded and bulbous tips consistent with acroosteolysis. She was also noted to have enlarged anterior fontanelles, low-set and hypoplastic ears, absent eyelashes and eyebrows, eyelids that did not close completely, small pinched nose, micrognathia, and multiple skin erosions on the chest and extensor surfaces.

The patient’s karyotype was 46, XX. Her hemoglobin and hematocrit were 12.0 g/dl (normal 14.5–22.5) and 36.1% (normal 45.0–67.0), respectively, requiring transfusion with packed red blood cells. Arterial blood pH was 6.96. The remainder of the blood chemistries and blood counts were normal at birth. Blood urea nitrogen was 18 mg/dl and glucose was 126 mg/dl. Her serum triglycerides were 130 mg/dl (normal 50–150) on day-of-life 4 and 51 mg/dl on day-of-life 5.

A chest radiograph showed butterfly vertebra at the thoracic 5 level (Fig. 2a). Clavicles were dysmorphic with a pseudoarthrosis at the juncture of the middle and distal one thirds with the distal portions bulbous in appearance (Fig. 2a). Radiograph of the skull revealed soft tissue swelling in the parietal regions, prominent sutures, micrognathia, and hypoplastic cervical vertebral bodies (Fig. 2b). Echocardiogram showed moderate patent ductus arteriosus and patent foramen ovale. A right thigh skin biopsy showed the epidermis and dermis to be slightly attenuated, and the dermis showed marked fibrosis and parallel collagen bundles (Fig. 2c). An elastic tissue stain, Verhoeff van Gieson, showed complete absence of elastic fibers throughout the dermis (Fig. 2d). The infant was maintained on mechanical ventilation, parenteral nutrition, antibiotics and pain medications and died on day-of-life 6.

RD 200.3

This was a stillborn male fetus born to Asian Indian primigravida mother at 32 weeks of gestation. Prenatal period was significant for oligohydramnios with premature rupture of membranes and preterm delivery. Upon birth, the fetus was noted to have tight skin with contractures in the neck, elbows, hip joints, and digits in hands and feet. Post-mortem examination showed the fetus weighed 750 g and was 48-cm long. Pinched nose, posteriorly rotated ears, tapering of the digits of hands and feet, superficial vasculature, absent eyelashes and eyebrows, and skin erosions were also noted (Fig. 3). The heart, lungs, liver, kidneys, gastrointestinal tract, and genitourinary tract looked normal. The brain tissue was autolyzed. The placenta was normal. Skin histology revealed thin epidermis and dermis. Epidermis displayed few variably sized keratohyaline granules. The dermis revealed loose fibrous connective tissue without any elastic fibers. The karyotype was 46, XY.

Methods

Mutational analysis

The exons and splice-site junctions of *LMNA* and *ZMPSTE24* were sequenced using genomic DNA as previously described (12, 13). In addition, approximately 2 kb of the promoter regions of *LMNA* and *ZMPSTE24* were sequenced in RD 200.3 (primers available on request).
Homozygous null mutations in ZMPSTE24 in RD

Fig. 2. Radiographs and histopathology of skin biopsy of restrictive dermopathy (RD) 500.3. (a) Anterior–posterior chest radiograph of patient RD 500.3 shows a butterfly vertebra at the thoracic 5 level (white arrow). Clavicles were dysmorphic with a pseudoarthrosis at the juncture of the middle and distal one thirds with the distal portions bulbous in appearance (black arrow). (b) Lateral radiograph of the skull of patient RD 500.3 reveals prominent sutures (gray arrow), micrognathia (white arrow), and hypoplastic cervical vertebral bodies (black arrow). (c) Hematoxylin and eosin stain of skin biopsy shows the epidermis and dermis to be slightly attenuated, and the dermis shows marked fibrosis and parallel collagen bundles. (d) An elastic tissue stain, Verhoeff van Gieson, shows complete absence of elastic fibers throughout the dermis.

Results

A homozygous mutation in ZMPSTE24, c.1085dupT (p.Leu362PhefsX19), was found in RD 600.3. Both the parents harbored heterozygous c.1085dupT mutation in ZMPSTE24. No disease-causing mutations were observed in sequencing of LMNA.

A homozygous missense mutation in ZMPSTE24, c.1020G>A (p.Trp340X), was found in RD 500.3. This patient also harbored a homozygous single nucleotide polymorphism (SNP), c.IVS5+18T>G (rs16827109), in ZMPSTE24. Both the parents had heterozygous c.1020G>A mutation in ZMPSTE24. No disease-causing mutations were observed in sequencing of LMNA.

No disease-causing mutations were observed in RD 200.3 upon sequencing of either LMNA or ZMPSTE24. The patient harbored a heterozygous c.1698C>T, p.His566His SNP in LMNA [rs4641 with minor allele frequency (MAF) in Asian subjects of 17–27%] and the following SNPs in ZMPSTE24: homozygous c.IVS5-50T>G (rs6677717, allele frequency not known), heterozygous c.651T>C, p.Asp217Asp (rs2076697, MAF 17–26%) and heterozygous c.IVS5+18T>G (rs16827109, MAF 9–27%). Sequencing of the 2.30-kb LMNA promoter region revealed the following variants: homozygous c.–1214T>C (rs2485661; MAF 46%) and homozygous c.–1242T>C (rs2485662; MAF 9–12%). The sequencing of the 2.15-kb ZMPSTE24 promoter region revealed the following variants: heterozygous c.–326G>A (rs3775483; MAF 11%), heterozygous c.–1269C>G (rs7548758, MAF 7–50%), heterozygous

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Fig. 3. Clinical features of restrictive dermopathy 200.3. (a) The infant had tight skin with skin breakdown in the neck, contractures in the elbows and digits in hands, and mouth in fixed ‘o’ position. A large circumferential neck laceration can be appreciated. (b) Prominent superficial vessels are seen in the posterior and lateral truncal region. There is normal number of digits on hands and feet, but fingers appear tapered. Scalp hair in the temporal and occipital regions appears normal.

c.–1355G>C (not listed) and homozygous c.–2036T>C (rs9326050, MAF 30–47%). All nucleotides are numbered from the first nucleotide of the translation start codon ATG.

Discussion

We describe detailed phenotype and genetic analysis of three cases of RD. Two newborns were found to have homozygous null mutations in ZMPSTE24, while one stillborn male did not have any disease-causing mutation in either ZMPSTE24 or LMNA. Most of the features of our cases, such as IUGR, joint contractures, skin breakdown, micrognathia, and acro-osteolysis, are typical of RD. Histologically, skin findings, including parallel collagen bundles and an absence of elastic fibers, are also consistent with previously reported cases of RD due to ZMPSTE24 mutations (9). However, butterfly thoracic 5 vertebra has not been previously noted. Also, although clavicles are often reported as hypoplastic or dysmorphic (2, 3, 9), the bulbous appearance of the distal portion of the clavicles in RD 500.3 is a novel observation.

Homozygous or compound heterozygous ZMPSTE24 mutations were first reported in 10 newborns with RD by Navarro et al. in 2005 (4). Since then, nearly all other cases of RD have also harbored null mutations in ZMPSTE24 (7). Interestingly, more than 50% of the patients with RD and ZMPSTE24 mutations harbored a homozygous c.1085dupT mutation (1, 3–5, 10) while an additional 17% were compound heterozygotes involving a c.1085dupT mutation on one allele (4, 5). One of our patients had homozygous c.1085dupT mutation while another had a novel homozygous c.1020G>A (p.Trp340X) mutation.

ZMPSTE24 mutations also cause autosomal recessive mandibuloacral dysplasia (MAD; OMIM 248370 and 608612) (13–16). In contrast to RD, MAD patients with ZMPSTE24 mutations harbor a null mutation on one allele and a missense mutation on the other allele (13–16). We have previously documented, using a yeast halo assay, that null mutations have no ZMPSTE24 activity, whereas the missense mutants have partial loss or nearly normal activity (13, 16, 17). Thus, the variable manifestations of the two disorders could be explained by varying amounts of prelamin A accumulation.

Only eight cases of MAD as a result of ZMPSTE24 mutations have been reported thus far (13–16, 18). Both MAD and RD patients with ZMPSTE24 mutations manifest prematurity, micrognathia, small pinched nose, sparse or absent hair, enlarged fontanelles, dysplastic clavicles and
Homozygous null mutations in **ZMPSTE24** in RD

Table 1. Comparison of clinical features of previously published and our patients with RD and mandibuloacral dysplasia as a result of **ZMPSTE24** mutations

<table>
<thead>
<tr>
<th>Finding</th>
<th>Previously reported RD cases (n = 25) (1–7, 9–11)</th>
<th>Our cases</th>
<th>ZMPSTE24 MAD cases (n = 8) (13–18)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IUGR</td>
<td>12/14 (86%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Birth weight: normal/low (&lt;2500 g)/very low</td>
<td>0:0:7:0 (7)</td>
<td>1615 g</td>
<td>1192 g</td>
</tr>
<tr>
<td>extremely low (&lt;1500 g)/extremely low</td>
<td></td>
<td>750 g</td>
<td>2:2:0:0 (4)</td>
</tr>
<tr>
<td>Birth at ≤33 weeks of gestation</td>
<td>16/17 (94%)</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Tight, thin, translucent, or shiny skin</td>
<td>20/20 (100%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Skin erosions</td>
<td>19/19 (100%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Micrognathia</td>
<td>17/17 (100%)</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>Small pinched nose</td>
<td>7/7 (100%)</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>Mouth in the ‘o’ position</td>
<td>19/19 (100%)</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>Sparse or absent hair, eyelashes or eyebrows</td>
<td>4/4 (100%)</td>
<td>NA</td>
<td>+</td>
</tr>
<tr>
<td>Low-set dysplastic ears</td>
<td>6/6 (100%)</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>Enlarged fontanelles</td>
<td>7/7 (100)</td>
<td></td>
<td>NA</td>
</tr>
<tr>
<td>Contractures and rigid posture</td>
<td>20/20 (100%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Prominent superficial vasculature</td>
<td>6/6 (100%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Dysplastic/hypoplastic clavicles</td>
<td>4/4 (100)</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>Bulbous appearance of distal clavicle</td>
<td>0</td>
<td></td>
<td>NA</td>
</tr>
<tr>
<td>Butterfly vertebrae</td>
<td>0/1 (0)</td>
<td></td>
<td>NA</td>
</tr>
<tr>
<td>Death in neonatal period</td>
<td>14/16 (88%)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Partial lipodystrophy</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

IUGR, intrauterine growth retardation; MAD, mandibuloacral dysplasia; NA, information not available; RD, restrictive dermopathy.

acro-osteolysis (Table 1). However, MAD as a result of **ZMPSTE24** deficiency is not lethal within the newborn period with death reported at ages 3, 28, and 37 years (13–15). MAD patients do not present with IUGR, fixed facial expression, mouth in the ‘o’ position, or skin erosions and denudations (Table 1). Development of contractures and joint stiffness is also delayed in MAD patients.

Navarro et al. (19) have reported heterozygous abnormal splice-inducing mutations (c.IVS11+1 G>A and c.1824C>T) the **LMNA** in two patients aged 5 and 6 months with presumed diagnosis of RD. However, both these mutations cause Hutchinson–Gilford progeria syndrome (20, 21) and therefore, these may have been misdiagnosed. More recently, a newborn with RD was reported to carry a heterozygous c.1821G>A (p.Val607Val) **LMNA** mutation (22); however, whether this mutation induces an alternative splicing was not demonstrated.

The phenotype of the stillborn fetus was consistent with a diagnosis of RD except for the presence of tapering rather than rounded and bulbous digits. The lack of mutations in **LMNA** and **ZMPSTE24** in this fetus suggests additional loci for RD. The presence of two heterozygous SNPs in **ZMPSTE24** reduces the likelihood of a large deletion of one allele. Regardless, a heterozygous deletion or c.‐1355G>C promoter region variant in **ZMPSTE24**, without a concomitant null mutation, is not sufficient to cause RD by themselves. Furthermore, we did not find any disease-causing nucleotide alterations in the proximal (approximately 2 kb) promoter regions of **LMNA** and **ZMPSTE24**. However, lack of RNA precludes us to determine if this patient harbored any homozygous cryptic intronic mutation in **ZMPSTE24**. We were also unable to exclude the possibility of small deletions in **LMNA**.

There is a significant risk of recurrence of RD in subsequent pregnancies. As illustrated in our cases, early diagnosis of RD is difficult given that most affected fetuses have an unremarkable prenatal course. Ultrasound findings of polyhydramnios, decreased fetal movements, and growth retardation are suggestive of RD, but are non-specific and late findings. A small fixed open mouth on ultrasound is also a late finding (23) and as such is not useful in prenatal genetic counseling. Skin biopsy at 20 weeks of gestation has been attempted (24), but failed to predict the presence of RD probably because fetal skin development is not complete by then. Thus, the best option for early prenatal diagnosis is genetic testing of DNA obtained via chorionic villus sampling or amniocentesis. This would be mostly done in families who have previously had newborns with RD.
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case (2), prenatal diagnosis revealed the presence of a homozygous ZMPSTE24 mutation; the pregnancy was subsequently terminated.

In conclusion, we report a novel and a previously reported homozygous null mutation in ZMPSTE24 in two newborns with RD. We also report a stillborn fetus with features of RD but no mutations in LMNA or ZMPSTE24, suggesting an additional locus may exist for RD.

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References