Dental-craniofacial manifestation and treatment of rare diseases

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Rare diseases are usually genetic, chronic and incurable disorders with a relatively low incidence. Developments in the diagnosis and management of rare diseases have been relatively slow due to a lack of sufficient profit motivation and market to attract research by companies. However, due to the attention of government and society as well as economic development, rare diseases have gradually become an increasing concern. As several dental-craniofacial manifestations are associated with rare diseases, we summarize them in this study to help dentists and oral maxillofacial surgeons provide an early diagnosis and subsequent management for patients with these rare diseases.

In this review, we aim to summarize the related manifestations and treatment of dental-craniofacial disorders related to rare diseases, thus helping to improve understanding and certainly diagnostic capacity for dentists and oral maxillofacial surgeons.

DENTAL-CRANIOFACIAL DISORDER-RELATED RARE DISEASES

Tooth dysplasia

Congenital ectodermal dysplasia. Ectodermal dysplasias (EDs) are a group of more than 150 different genetic disorders deriving from ectodermal structural abnormalities.7,8 Ectodermal dysplasias have been described as ‘heritable conditions in which there are abnormalities of two or more ectodermal structures, such as the hair, teeth, nails, sweat glands, salivary glands, cranial-facial structure, digits and other parts of the body’.7 The abnormality in the development of tooth buds frequently results in congenital hypodontia (both primary and permanent dentitions) and/or changes in tooth morphology or size, such as peg-shaped or pointed teeth, taurodontism and enamel defects, including hypoplasia.9 The degree of tooth missing is always in the mild to moderate range, and a wide variation is observed regarding which teeth are missing; however, the most frequently reported missing teeth are the first molars, upper central incisors and canines10–14 (Table 1). Accordingly, composite restorations or crowns are almost always necessary for children as early as 2 years of age, and multiple denture replacements are often needed as the child grows, with dental implants providing a potential option in adolescence when the jaw is fully grown. The current option of extracting teeth and substituting them with dental implants is quite common.15 Additionally, orthodontic treatment is further necessary during the early teenage years as part of the best multidisciplinary approach. Furthermore, several studies have also reported reduced salivary secretion in ED patients, accompanied by a reduced buffering ability and increased bacterial counts.16–20 Therefore, a systematic preventive plan including fluoride use and...
The placement of fissure sealants will be especially important for patients with ED owing to their increased caries risk.

**Williams syndrome.** Williams syndrome (WS, also known as Williams–Beuren syndrome) is a multi-system genetic disorder that is caused by a genetic abnormality, especially the deletion of a specific region in chromosome 7q11.23 containing 26–28 genes, such as CLIP2, ELN, GTF2I, GTF2IRD1, BCL7A and LIMK1. Most of the gene deletions occur as a random event during the embryonic period rather than being inherited from affected parents, and different clinical features are believed to be linked to the loss of specific genes.26

WS is characterized by various degrees of intellectual disability or mental retardation, unique facial features, cardiovascular defects and hypersocial behaviour, and cardiovascular complications are the major cause of death in WS patients. Regarding physical disorders, ophthalmologic and auditory abnormalities (altered visual acuity, strabismus, sensorineural hearing loss and hyperacusis), cardiovascular diseases (vascular stenosis, hypertension and stroke), gastrointestinal diseases (constipation, colic, rectal prolapse and coeliac disease) and musculoskeletal disorders (joint laxity, joint contractures and scoliosis) are always observed in WS patients.21,27–29 Additionally, WS patients usually exhibit distinctive facial features, including a broad forehead, bitemporal narrowing, a short nose with a broad tip, malar flattening, full cheeks, a small jaw, a wide mouth with full lips and large ear lobes, as well as dental abnormalities, such as small or unusually shaped primary teeth, malocclusions with widely spaced teeth, hypodontia and tooth enamel defects.21,25,26 (Table 1) Regarding the nervous system, multiple missing genes result in several defects in the cerebellum, right parietal lobe, and left frontal cortical regions. Thus, patients with WS often present a cognitive impairment, poor balance and coordination, defects in language development and coordination of fine motor tasks, such as drawing and writing, and disabilities in visuospatial construction.21,33–37 Furthermore, WS patients consistently demonstrate an overly friendly, highly social and empathic personality, as well as excessive worry and fears, distractibility, and irritability. However, these patients have been reported to show a greater volume of memory and higher rhythm propensity and fondness of music.38–40 Concerning endocrine abnormalities, hypercalcaemia, diabetes mellitus, subclinical hypothyroidism and other endocrine disturbances are well described in some patients with WS, which might lead to muscle hypotonia, a diminished appetite, obesity and a below-average height and weight.21,24,41–43

Current management guidelines for WS have not been established, and the available treatments involve a combination of medical monitoring, pharmacotherapy, surgery, speech and behavioural treatments. Routine supervision of blood pressure, blood glucose and calcium levels are generally recommended for all patients with WS, surgery or stent insertion is the preferred method for moderate to severe aortic stenoses, an oral hypoglycaemic agent or insulin administration might be required for WS patients with potential diabetes mellitus, and bisphosphonate therapy is used for WS patients with significantly decreased bone density.21,44–46 Additionally, the application of anxiolytic and antipsychotic agents is occasionally prescribed, and behavioural treatments, such as the practice of relaxation, music treatment and social skill training, might be effective to channel the nature of affected patients.47 Other treatments, including dental restoration and orthodontic treatments, are administered depending on the patient's particular symptoms, and an early dental evaluation and dietary counselling are necessary to determine the presence of dental anomalies, such as caries and enamel structural defects.32

**Congenital erythropoietic porphyria.** Congenital erythropoietic porphyria (CEP), also known as Gunther's disease, is an autosomal-recessive genetic disorder resulting from a homozygous defect in uroporphyrinogen III cosynthase, located on human chromosome 10q25.2-q26.3, which leads to the overproduction and accumulation of porphyrin I and coproporphyrins I.48,49 The accumulation of porphyrins and the targets leads to a loss of membrane integrity, destruction of cellular organelles and cell apoptosis.51

Severe cutaneous photosensitivity due to massive porphyrin accumulation in the skin, characterized by subepidermal blistering with inflammatory infiltration, is the most frequently observed manifestation in CEP patients, although significant phenotypic variability in CEP has been reported.42 The severity of skin photosensitivity depends on the amount of porphyrin in the tissue,53 and thickened skin with hyperpigmentation as well as the rapid development of vesicles and bullae can be observed in any

| Rare diseases | Aetiology | Major manifestations | Dental-cranio-facial manifestations | Incidencea | Onset periodb |
|--------------|-----------|---------------------|------------------------------------|------------|--------------|
| Congenital ectodermal dysplasia | Abnormalities of the ectodermal structures | Excessively fragile and twisted hair; thick, brittle or discoloured nails; red or brown pigmentation on skin; overheating; respiratory infections | Hypodontia, peg-shaped or pointed teeth, taurodontism, enamel hypoplasia; Reduced salivary secretion | 80%; 30.2% | Childhood |
| Williams syndrome | Genetic deletion of chromosome 7q11.23 | Intellectual disability; cardiovascular defects; failure to thrive; lack of social inhibition | Small jaw, wide mouth with full lips; Malocclusions with widely spaced teeth, hypodontia and enamel defects | 75%–91%; 38%–93% | Childhood, Early childhood |
| Congenital erythropoietic porphyria | Genetic mutations in chromosome 10q25.2-q26.3 | Thickened skin with hyperpigmentation and bullae formation; anaemia; dry eyes | Facial scabs and scars; Teeth: reddish fluorescence, reddish-brown discoloration with a sharply defined margin; Sclerotic and osteolytic round lesions in the skull, maxilla, mandible | 47%; 73% | Early childhood, Childhood |

*aIncidence of dental-craniofacial manifestations  
*bOnset period of dental-craniofacial manifestations
sun-exposed areas. Additionally, facial scabs and scars and the destruction of auricular and nasal cartilages, cheeks, lips and forehead can be severe due to repetitive skin damage and bone resorption, resulting in unique facial features. The accumulation of excessive porphyrin in erythroid precursors, reticulocytes and erythrocytes can induce osmotic haemolysis and subsequently result in mild to severe anaemia, and haematological complications are the major predictors of a poor prognosis for patients with CEP. Dental disorders are also found in most CEP patients due to porphyrin deposition during tooth development, which exhibits fluorescence under long-wavelength ultraviolet light as well as visible reddish-brown discoloration with a sharply defined margin (Table 1). Ocular complications, including corneal scarring, loss of eyelashes and eyebrows, ulcerative keratitis and conjunctivitis, and skeletal abnormalities including fractures and a shortened stature, are also frequently observed in CEP patients.

Other symptoms, such as neurological manifestations, are not common, although Parkinson disease and corticobasal syndrome have been reported to be involved in some CEP patients. Multiple therapies have been proposed to treat CEP, such as the elimination of sun exposure using sunscreens (zinc oxide and titanium dioxide), oral β-carotene, or protective clothing; prevention of the reabsorption of porphyrins with oral activated charcoal and cholestyramine; erythrocyte transfusions; and hematopoietic stem cell transplantation (HSCT); HSCT is the only curative method for severe CEP. Other management strategies, such as topical lubrication for dry eyes, glucocorticosteroid therapy for anaemia and thrombocytopenia and splenectomy for splenomegaly, have also been reported for specific complications.

Bone tissue abnormality

Osteogenesis imperfecta. Osteogenesis imperfecta (OI) is represented by a group of genetic disorders that mainly affect the bones, connective tissue and may increase skeletal fragility, also known as brittle bone disease. Among all cases, 85%–90% present a lack of type I collagen due to a mutation in the COL1A1 and COL1A2 genes that is inherited from the parents or develops de novo. With advances in genomic analysis and exome sequencing, several other gene mutations, such as CRTAP, LEPRE1, PPIB, SERPINH1, and SP7 mutations, which might result in defects in collagen post-translational modifications or osteoblast differentiation, have been found to be involved in the occurrence of OI.

OI had been recognized since the early 1980s. Fractures caused by mild trauma, bowing deformities of the long bones, and growth deficiency are the hallmark features, including macrocephaly and chest wall deformities. Additionally, typical extraskeletal manifestations can be associated variably with the disorder, including a dark or blue sclera, dentinogenesis imperfecta, pulmonary function impairment, the presence of wormian bones on skull radiographs, hyperlaxity of the ligaments and skin, and hearing impairment. Blue sclera and dentinogenesis imperfecta are always used as diagnostic signs of OI, and dentinogenesis imperfecta occurs more frequently in primary teeth than permanent teeth. Hearing loss is rare in the first 20 years of life, but half of patients aged more than 50 years report hearing loss. Radiological or histological examination can reveal generalized osteopenia and some combination of gracile ribs, long-bone bowing, and vertebral compression. Several clinical and genetic classifications have emerged to encompass the rare forms of osteogenesis imperfecta, beginning with David Sillence in 1971; however, they are associated with respective limitations. Additionally, in 2016, Forlino proposed a genetic-functional metabolic classification that is dependent on both the involved gene function and clinical features, updating several new types to classic Sillence types I–IV. The current classification of OI types is still debated.

There is no cure for OI, and all management strategies are symptom-based or complication-based. Regarding the medical management of OI, a multidisciplinary team is necessary. Physiotherapy, hydrotherapy and rehabilitation exercises focus on strengthening the muscles, restricting joint range of motion and improving the patients’ living ability. Audiology examination should be carried out in childhood, as patients with severe disease with pure conductive or sensory loss might require hearing aids or cochlear implants. Oral health management, including oral hygiene introduction (regular brushing and flossing to avoid tooth chipping and potential fracture during dental procedures), and crown restoration should be applied to prevent caries, periodontitis and to optimize aesthetics. Regarding drug therapies, bisphosphonates are antiresorptive drugs that are widely used in children with OI to increase the volume of bone, bone strength, restore vertebral shape and decrease fractures, although the newly formed bones still contain defective collagen, and patients who use bisphosphonates should be well-informed of bisphosphonate-related osteonecrosis of the jaws. Concerning orthopaedic surgery, lower extremity, upper extremity and spinal surgery are usually conducted, and combined with pre- and post-surgical rehabilitation, placement of an intramedullary telescoping rod can be applied after correction of the bone deformity to provide strength, alignment and to stabilize fractures. Hip and knee arthroplasty are frequently conducted for OI patients with joint osteoarthritis, and spinal fusion is generally used to correct the scoliosis in OI patients. Because pulmonary impairment is the leading cause of death in OI patients, potentially as a secondary effect of scoliosis and rib fracture, treatment for pulmonary complications, such as obstructive disease, should also be emphasized to avoid respiratory infections and insufficiency.

Hypophosphatemic rickets. Hypophosphatemic rickets is a genetic X-linked dominant form of rickets, also called X-linked hypophosphatemia (XLH), which is different from most cases of rickets in that the administration of vitamin D is not effective. XLH is caused by a loss-of-function mutation in the phosphate-regulating endopeptidase gene, X-linked (PHEx), and results in overactivity of fibroblast growth factor 23 (FGF23). The excess circulating FGF23 inhibits vitamin D 1α-hydroxylation and phosphate reabsorption by the kidneys, leading to hypophosphatemia and defective mineralization of the bones and thus facilitating rickets and osteomalacia.

The main manifestations of XLH patients are a disproportionately short stature, reduced growth rate and bone deformities, such as coxa vara, tibial torsion and lower limb bowing, which occur in children before the fusion of the epiphysis. Adult patients can present symptoms of osteomalacia, including myopathy, bone pain, neurologically complications and insufficiency fractures, due to enthesopathy and ectopic calcification. In addition, primary craniosynostosis has been reported to be present at or soon after birth, and mineralizing enthesopathy, osteoarthropathy and ossification of the spinal ligaments have also been reported during adulthood.

Along with skeletal abnormalities, dental implications are consistently observed, and some of the main manifestations are recurrent abscesses or sinus tracts associated with carious and trauma-free teeth of the primary and permanent dentition because of the dentin defect (wide predentin zones and tubular defects) and microdefects in the enamel, especially in the anterior teeth. The hypoplasia of the enamel and the lack of fusion of calcifications in dentin facilitates microbial penetration, leading to pulp infection, pulp necrosis and finally periapical periodontitis and abscesses. Other dental-related symptoms have also been reported, such as delayed tooth eruption, taurodontism (large pulp chambers, short roots, prominent pulp horns and a thin enamel layer), and a hypoplastic alveolar ridge. Early diagnosis and medical intervention for XLH is always associated with better therapeutic outcomes. The current standard treatment for XLH consists of activated vitamin D metabolites, oral inorganic phosphate salts and growth hormone
Table 2. Dental-craniofacial manifestations of bone tissue abnormality-related rare diseases

| Rare diseases                          | Aetiology                  | Major manifestations                                                                 | Dental-cranio-facial manifestations | Incidencea | Onset periodb |
|---------------------------------------|----------------------------|--------------------------------------------------------------------------------------|-------------------------------------|------------|---------------|
| Osteogenesis imperfecta               | Mutations in the COL1A1 or COL1A2 genes | Bones fracture easily; long bones deformity and small stature; loose joints; blue-grey colour of the sclera; loss of hearing | Facial deformities with high risk of fracture; Dentinogenesis imperfecta; Malocclusion and delayed tooth eruption | 60%        | Childhood; adulthood |
|                                        |                            |                                                                                      |                                      |            |               |
| Hypophosphatemic rickets              | Mutations in the phosphate-regulating endopeptidase gene                           | Disproportionate short stature; bone deformity; bone pain; hearing loss                 | Primary craniosynostosis; Recurrent abscesses with carious and trauma free teeth; Delayed tooth eruption, taurodontism | —          | At birth; Early childhood |
| Hypophosphatasia                     | Mutations in tissue non-specific alkaline phosphatase genes                        | Perinatal HPP: soft calvarium, deformed limbs, respiratory failure; Infantile hypophosphatasia: poor feeding, flail chest; Childhood hypophosphatasia: delayed walking, frequent fractures, open fontanels; Adult hypophosphatasia: painful feet, femoral pseudoarthroses, arthritis | Unossified calvarium with separated cranial sutures; Early loss of deciduous teeth, shell teeth, impaired dentogenesis, permanent dentition caries | 31%–40%    | At birth; Childhood |
| Marfan syndrome                      | Mutations in FBN1 gene            | Arachnodactyly, disproportionately long, slender limbs with thin, ectopia lentis; weak wrists, long fingers and toes; undue fatigue, shortness of breath, cold arms, hands, and feet | Long narrow skull, high arched palate, mandibular and maxillary hypoplasia; Crowded teeth and overbite | 63.6%      | Childhood |
| McCune–Albright syndrome             | Mutation in the gene GNAS        | Multiple bone fibrous dysplasia; café-au-lait skin pigmentation; endocrine diseases (Precocious puberty, testicular abnormalities and hyperthyroidism) | Facial asymmetry with expanding fibrous dysplasia lesion; Oral mucosal pigmentation; Dental malocclusion, dentin dysplasia, taurodontism and high caries index | 62%–100%   | Early childhood (3.4 years old) |
| Kallmann syndrome                    | Isolated Gonadotropin-Releasing Hormone Deficiency                                | Failure to start or fully complete puberty, primary amenorrhea or lack of testicle development; Total lack of sense of smell, hearing loss | Cleft palate, hare lip, high-arched palate; Hypodontia, malformed teeth and other dental abnormalities | 25%–30%    | At birth; Early childhood |
| Fanconi anaemia (FA)                 | Mutations in FA or FA-like genes                                               | Bone marrow failure, acute myeloid leukaemia; skin hyperpigmentation; short stature, abnormal thumbs, absent radii | Microcephaly, triangular face; Head and neck cancers | 51%        | Infanthishhood and Early childhood |

aIncidence of dental-craniofacial manifestations
bOnset period of dental-craniofacial manifestations

dentosis, to recreate a normal anatomic and mechanical axis. 102 Multiple osteotomies, and arthroplasty if necessary, are performed for deformed bones, and then the bone pieces are realigned in an improved position using external fixators or intramedullary nails to achieve a straightened configuration. 104, 106 The application of burosumab, a recombinant human IgG1 monoclonal antibody that targets FGF-23 and inhibits its activity, has been reported to increase renal tubular reabsorption of phosphate and thereby increase serum phosphate and serum 1,25(OH)2D in phase 1 and 2 clinical trials in both paediatric and adult patients with XLH. 107–109 Moreover, diverse dental treatments have been suggested for XLH patients, as well as routine radiographic control of the entire dentition, topical fluoride varnish and fissure sealing, stainless steel crowns or permanent crown restoration, and early non-surgical root canal treatment or surgical resection for teeth associated with apical periodontitis. 95 Teeth with severe periapical abscesses might require extraction followed by implant restoration.

Hypophosphatasia. Hypophosphatasia (HPP) is a type of genetic bone metabolic disorder that results from a molecular defect in tissue non-specific alkaline phosphatase (TNSALP) genes, and a dominant effect of the mutated allele is usually suspected to be the cause of the disease. 10, 11 TNSALP is a membrane-bound glycosylated enzyme encoded by the ALPL gene, and its physiological function has been proposed to be involved in extracellular matrix mineralization, ATP hydrolysis and skeletal development. 112 The absence and reduced activity of TNSALP would result in increasing extracellular PPI in the bone matrix, an inhibitor of hydroxypapatite formation, which is an important component of bone and lead to rickets and osteomalacia. 113 In addition to hard tissues, such as bone and teeth, TNSALP is also essential for pyridoxal 5′-phosphate dephosphorylation and...
vitamin B6 production; thus, other organs, such as muscles, brain and liver, can also be affected in HPP patients.\textsuperscript{114} A small number of mutations are recurrently found, which result in a large number of compound heterozygous genotypes and a wide range of clinical symptoms. Based on the appearance of the first symptom, HPP is divided into several subtypes (during gestation or at birth; perinatal hypophosphatasia; before the first 6 months of life: infantile hypophosphatasia; onset ≥6 months to 18 years of age; childhood hypophosphatasia; after 18 years of age: adult hypophosphatasia), according to the classification proposed by Fraser et al.\textsuperscript{15} and Whyte et al.\textsuperscript{116} Additionally, odontohypophosphatasia refers to the phenotype when dental disease (including premature loss of deciduous teeth, especially the anterior teeth; large pulp chambers; impaired dentinogenesis; and rare enamel hypoplasia) is the only clinical abnormality, and no radiographic and histopathological evidence of rickets and osteomalacia can be observed (Table 2). Perinatal HPP is the most severe form of HPP, which is usually characterized by caput membrane, and deformed limbs, periodic apnoea with cyanosis and bradycardia resulting from chest deformities and lung hypoplasia, myelophthisic anaemia and intracranial haemorrhage at birth.\textsuperscript{118,119} Nearly all the bones appear to be completely unmineralized, which would quickly be fatal due to respiratory failure.\textsuperscript{120} Infantile HPP patients always show a failure to thrive as well as rachitic features, including leg bowing, joint enlargement, rib fracture, progressive deformity of the thorax and tracheomalacia.\textsuperscript{117,121,122} Other signs such as hypercalcaemia, muscle hypotonia, papilledema resulting from craniosynostosis and increased intracranial pressure can also be observed in some patients.\textsuperscript{123,124} Childhood HPP is the form with the greatest clinical variability. The infantile and childhood subtype are a continuum and are sometimes difficult to distinguish, with childhood HPP being more evident in most cases.\textsuperscript{123,125} Delayed walking, a waddling gait, chronic bone and joint pain, recurrent fractures, scoliosis, shell teeth, hypoplasia of the cementum and caries in the permanent dentition are characteristic manifestations and common in childhood HPP.\textsuperscript{126,127} Adult HPP is always associated with an early loss of permanent dentition and painless or due to femoral pseudofractures in the lateral cortices of the femora (Looser zones) and recurrent metatarsal stress fractures.\textsuperscript{128–130} Pyrophosphate arthropathy, pseudogout, spinal hyperostosis and calcific periartithitis are also observed in some adult HPP patients.\textsuperscript{129,131,132} There is confirmed medical management for HPP, but only therapeutic interventions that consist of symptom palliation, calcium balance maintenance and physical, occupational, dental and orthopaedic interventions, as necessary, to alleviate symptoms and reduce complications. Regarding tracheomalacia and pulmonary hypoplasia in infantile and childhood HPP, mechanical ventilation is necessary,\textsuperscript{133} and to avoid neurological complications, vitamin B6 or pyridoxine supplementation, and/or craniectomy if necessary, is recommended.\textsuperscript{134,135} Concerning pseudofractures or completed fractures in adult HPP, the use of teriparatide (recombinant human parathyroid hormone), load-sharing intramedullary fixations or ankle-foot orthoses is often applied, and naproxen might be used to diminish skeletal pain.\textsuperscript{136,130,137} It is worth noting that bisphosphonate treatment for ‘osteoporosis’ in HPP patients is ineffective, but PTH provides some mitigation.\textsuperscript{138} Additionally, dental prosthetic interventions (especially removable partial dentures in childhood) are well recommended to facilitate the normal development of speech and avoid abnormal transversal development of the jaw and related social problems.\textsuperscript{139} Currently, several newly proposed treatments in infants and young children, such as bone marrow and mesenchymal stem cell transplantation\textsuperscript{139} and enzyme replacement therapy (ERT, Asfotase alfa),\textsuperscript{133,140} have achieved promising results for the bones and lungs.

Marfan syndrome. Marfan syndrome (MFS) is a genetic autosomal dominant disorder of the connective tissue and is caused by mutations in the \textit{FBN1} gene, which encodes extracellular matrix protein fibrillin-1.\textsuperscript{141} Several studies have demonstrated that the transforming growth factor \(\beta\) (TGF\(\beta\)) signalling system may also be involved in the development of MFS.\textsuperscript{142,143} MFS can manifest either at birth or as a progressive disease that can be found as late as 30–40 years old.

Several signs and symptoms have been reported to be associated with MFS, especially the skeletal, cardiovascular and ocular systems.\textsuperscript{144} In the skeletal system, most individuals with MFS present arachnodactyly (positive Walker Murder sign and Steinberg sign), dolichostenomelia and deformity of the spine and chest wall, due to disproportionate linear overgrowth of tubular bones, and some MFS patients demonstrate osteopenia, ligament laxity, hindfoot valgus with footed abduction, a high risk of knee and ankle sprains, and protusio acetabuli. All these manifestations usually appear during childhood and may worsen during adolescence. Additionally, typical craniofacial deformities, such as a long narrow skull, high arched palate with crowing teeth, midface hypoplasia, mandibular retrognathia and malar hypoplasia, are also observed in MFS patients\textsuperscript{145,146} (Table 2). In the cardiovascular system, enlargement of the aortic root, pulmonary artery dilatation, valvulopathies, cardiomyopathy due to elastic fibre degeneration in the aorta, and subsequent abdominal aortic aneurysms and progressive myocardial dysfunction are common in MFS patients and are always associated with severe consequences for survival.\textsuperscript{147–150} In ocular and other systems, ectopia lentis, near-sightedness, intracranial hypotension-associated headache due to dural ectasia, spontaneous pneumothorax and striae atropeicae on the arms, hip and lower back, are very frequent in MFS patients.\textsuperscript{151–154}

Cardiac and pulmonary impactions are the most severe complications of MFS, while osteopenia and bone pain are the two most poorly managed features of MFS, particularly in elderly patients. Although there is no cure for MFS, life expectancy can be significantly increased after a life-style modification, such as reducing emotional stress and restricting physical activities\textsuperscript{155} and undergoing regular echocardiographic imaging assessments, pharmacological treatment (propranolol and other \(\beta\)-blockers, calcium channel blockers, angiotensin receptor blockers and angiotensin-converting enzyme inhibitors)\textsuperscript{157,158} and prophylactic surgery (replacement of the aortic valve and ascending aorta, vitreolensectomy with laser prophylaxis).\textsuperscript{159} For MFS patients with severe craniofacial deformities, combined orthognathic and orthodontic treatment is recommended to correct crowing teeth and mandibular retrognathia.

\textbf{McCune–Albright syndrome.} McCune–Albright syndrome (MAS) is a non-hereditary genetic disorder this is caused by a spontaneous postzygotic mutation of the gene \textit{GNAS}, which is involved in \(G\)-protein signalling and results in constitutive activation of \(G\)-protein as well as the production of excess \textit{CAMP}\textsuperscript{160,161}. The \textit{GNAS} mutation arises very early during embryogenesis and occurs only in the mosaic state, which leads to a variable pattern in affected tissues.\textsuperscript{162} MAS was initially defined as the triad of polyostotic fibrous dysplasia (FD) of the bone, café-au-lait skin pigmentation and precocious puberty,\textsuperscript{163} and over years it has been refined as a disorder involving at least one of the following clinical manifestations: café-au-lait macules, fibrous dysplasia, and autonomous endocrine hyperfunction.\textsuperscript{164} The light brown café au lait macules arising from the ectoderm are often the first observed feature of MAS patients, which usually appear at or shortly after birth. They are described as having an irregular border and mostly affect the midline of the body, such as the posterior neck, base of the spine and face.\textsuperscript{164} Additionally, oral mucosal pigmentation has been documented in a minority of MAS patients.\textsuperscript{165} FD arises from the mesoderm and can occur in one bone as well as a combination of craniofacial-axial bones and the appendicular skeleton as a hallmark of MAS. The affected bones are characterized by extensive lesions with a thin cortex and ‘ground glass’-like
intramandibular matrix.\textsuperscript{166} Craniofacial deformities may present as a facial asymmetry due to an expanding FD lesion, which may progress along with dental malocclusion, hearing impairment and vision changes.\textsuperscript{167,168} Craniofacial FD is also associated with tooth development and dental disorders, such as dentin dysplasia, taurodontism and a high caries index, and therefore, more frequent scaling and root planing as well as topical fluoride application are required to control dental plaque and caries in MAS patients.\textsuperscript{169,170} Jaw FD may contribute to aneurysmal bone cysts and osteosarcoma\textsuperscript{171,172} (Table 2). FD-involved sphenoid bone may lead to a pituitary adenoma, and conversely, excess growth hormone can worsen craniofacial bone disease.\textsuperscript{173,174} Limb deformities usually present both valgus and varus deformities in both knees, appendicular skeletal fractures as well as curved femurs.\textsuperscript{162,175} Additionally, scoliosis appears to occur frequently in MAS patients.\textsuperscript{176} Endocrine hyperfunction in MAS patients consists of several disorders, including precocious puberty, Cushing’s syndrome, excess growth hormone and prolactin, renal phosphate wasting, and hyperthyroidism, and it is more common in females than males. Thus, growth acceleration, testicular enlargement or Sertoli cell tumours, vaginal bleeding or spotting, hypophosphatemia-related rickets or fractures, advanced skeletal maturity and acromegaly may occur in most MAS patients.\textsuperscript{164,168} Additionally, hepatic and cardiac complications have also been reported in MAS patients.\textsuperscript{168}

The affected multi-organ clinical pattern makes the management of MAS complex and challenging. Regarding FD, physical therapy is used to maintain the strength and range of motion of affected bones.\textsuperscript{177} Surgical treatment involves orthopaedic surgery or orthognathic surgery combined with orthodontic therapy, lesion resection surgery and intramedullary device fixation, which are applied for FD associated craniofacial and limb deformities as well as fractures.\textsuperscript{170,178} and antiresorptive therapy using bisphosphonates is also advocated to relieve FD-related bone pain and reduce bone resorption.\textsuperscript{179,180} However, surgical management is not recommended in most patients, and there is no satisfactory treatment capable of altering the progress of FD in MAS patients. Thus, annual observation may be necessary. Concerning endocrinopathies, medical treatment is mainly used. Somatostatin analogues such as pegvisomant are used to treat excess growth hormone, the aromatase inhibitor letrozole combined with leuprolide or testosterone receptor antagonist is used for precocious puberty, and oral phosphorus and calcitriol supplementation is used for hypophosphatemia.\textsuperscript{181} In patients with FA, medical treatments, several surgeries, including thyroidectomy and hypophysectomy, have been reported as a potential option.\textsuperscript{168,182} Treatment for skin hyperpigmentation in MAS patients is not routinely conducted, although laser therapy can be applied with satisfactory results.\textsuperscript{182}

Kallmann syndrome. Kallmann syndrome (KS) is a form of a group of genetic disorders termed isolated gonadotropin-releasing hormone (GnRH) deficiency (IGD).\textsuperscript{183} Hypogonadotropic hypogonadism (HH) occurs with anoma and can be termed KS.\textsuperscript{184} Although the genetic defect in most IGD patients remains uncharacterized and involves a sequence variant of several different genes, the genes that regulate neurodevelopmental IGD pathways, including neural cell adhesion and axonal migration, have been reported to contribute to KS. Such genes include Kallmann 1, 2 (KAL1, 2), NDMA receptor synaptotagonin signalling and neuronal migration factor (NSMF), fibroblast growth factor receptors 1, 8 and 17 (FGF1, 8, 17), semaphorin 3A (SEMA3A), SRY Box 10 (SOX10) and many more.\textsuperscript{185}

The clinical features of KS can be divided into two parts: reproductive features and non-reproductive features. Among the reproductive features, KS patients are characterized by low blood levels of sex hormones, gonadotropins and subsequent failure of puberty onset, low libido and poor sexual function, as well as infertility. Other signs such as a micro penis and lack of testicular development or cryptorchidism in males and delayed menarche and failure to start menstruation in females have also been observed in some KS patients.\textsuperscript{184,186} Regarding non-reproductive features, a wide spectrum of clinical manifestations can be involved, most of which are associated with mutations in different genes. In addition to a partial or total lack of sense of smell due to olfactory placode cell migration from the nasal region to inside the hypothalamus, cleft palate, hare lip, a high-arched palate and other midline craniofacial defects, hypodontia, malformed teeth and other dental abnormalities, short metacarpals, scoliosis, neural hearing loss, bimanual synkinesis due to cerebellar ataxia, eye movement disorders, colour blindness, unilateral renal agenesis and other maladies are also frequently observed in KS patients (Table 2). In addition, as a result of a deficiency in either testosterone or oestrogen, which is important to maintain bone density, KS patients are exposed to a higher risk of developing secondary osteoporosis or osteopenia.\textsuperscript{193,194} Thus, an increased tendency toward fracture may be observed in these patients.

The treatment for KS patients aims first to initiate the development of secondary sexual characteristics and second, to develop fertility, sex hormone replacement in childhood and gonadotropin and GnRH pulsatile therapy in adolescence and adulthood are the most frequently applied treatments to stimulate virilization or oestrogenization and restore fertility.\textsuperscript{195} Early surgical corrections before 1 year of age are recommended for KS patients with cryptorchidism.\textsuperscript{196} Other skeletal phenotypes such as a cleft palate and lip and hypodontia require related repair surgery, and patients with tooth agenesis or hypodontia require dental restoration early in life.\textsuperscript{187}

Fanconi anaemia. Fanconi anaemia (FA) is a genetic autosomal recessive genetic disorder that results from several FA or FA-like genes, such as FANCA, -C, -D, -E, -F, -G, -J, -L, -M, -N, -P, -S and XPF, all which are involved in an impaired response to DNA damage.\textsuperscript{197–200} The proteins encoded by these genes and their complex have been reported to protect cells from oxidation-induced genotoxicity and to directly participate in STAT pathway activation and protein kinase pathway suppression, which is important in regulating the apoptosis of hematopoietic cells.\textsuperscript{200,201} Consequently, haematologic components, including blood cells and platelets, fail to develop, and bone marrow failure syndromes gradually develop.

Patients with FA are characterized by bone marrow failure, myelodysplastic syndromes/acute myeloid leukaemia (MDSs/AML), typical skeletal deformities and an increased incidence of solid tumours. Skin discolorations such as petechiae, bruises and cafe-au-lait spots are usually the first signs of haematologic problem in FA patients.\textsuperscript{202} Subsequently, FA patients may exhibit a pale appearance, feel tired, and develop infections, 20% of whom may develop MDSs/AML during their teens or young adulthood due to hypoplastic or aplastic anaemia or cytopenia, unexplained macrocytosis and bone marrow failure.\textsuperscript{202,203} Furthermore, facial and skeletal abnormalities are frequently observed in patients with FA, as well as short stature, fanconi facies (microcephaly, microphthalmia and ptosis, microphthalmia and triangular face), and abnormal radii and thumbs (clinodactyly and polydactyly)\textsuperscript{196,207–210} (Table 2). Other symptoms such as the onset of solid tumours or cancers (especially head and neck squamous cell carcinomas, HNSCCs), abnormal reproductive organs with reduced fertility, renal and urinary tract problems, heart defects and gastrointestinal problems have also been reported, but with a relatively lower incidence.\textsuperscript{199,211} Currently, the most common management for FA involves HSCT and the application of androgens and hematopoietic growth factors. HSCT is the only way to establish normal haematopoiesis and can significantly extend patients’ lifespan; however, not all patients are candidates for transplantation, and the high solid
and Kindler syndrome.218,220 Although the types differ in signs of junctional epidermolysis bullosa, dystrophic epidermolysis bullosa, skin mucosal, and soft tissue abnormalities. Hereditary epidermolysis bullosa (EB) comprises a group of rare genetic disease characterized by increasing skin fragility, which results in blisters or erosions of the skin and mucous membranes in response to minor mechanical injury, such as scratching.217 Mutations or errors in the genetic code lead to a defect in attachment between or within two layers of the skin-epidermis and dermis, thus resulting in extremely fragile skin.218 Currently, mutations in at least one of twenty different genes has been found to cause a large spectrum of phenotypes in EB patients, ranging from mild to lethal.219 Based on EB-related genes for proteins with different cellular localizations (intracellular, transmembrane or extracellular), EB is categorized into four major types: epidermolysis bullosa simplex, junctional epidermolysis bullosa, dystrophic epidermolysis bullosa, and Kindler syndrome.218,220 Although the types differ in signs and symptoms, the underlying mechanism is identical. These mutations of specific genes prevent the production of essential proteins to strengthen the skin or produce antibodies against structural components of the skin. Basal variants of epidermolysis bullosa simplex (EBS), the most common EB type, are frequently caused by dominant negative missense mutations in the KRT5 or KRT14 genes.221 These genes supply instructions for producing tough, fibrous proteins to provide resiliency to the epidermis. Mutations in either the KRT5 or KRT14 gene will lead to destabilization of the cytoskeleton and cytolysis upon mechanical stress. Similarly, junctional epidermolysis bullosa (JEB) results from mutations in the LAMA3, LAMB3, LAMC2, and COL17A1 genes, which lead to defects in the production of laminin 332, an important protein to help attach the epidermis to the dermis.

RAMPANT DENTAL CARIES in patients with JEB seems to occur due to enamel hypoplasia, which decreases the intrinsic resistance of the tooth. However, despite normal dentition development, rampant dental caries are also frequently detected in DEB patients, which could be attributed to the special diet (soft) and limited oral clearance (limited tongue and cheek mobility) of EB patients (Table 3). Apart from easily blistering skin and mucous membranes, complications of epidermolysis bullosa may include the following: anaemia, muscular dystrophy, dysphagia, pyloric atresia, constipation, cardiomyopathy, renal insufficiency, syringomyelia, oesophageal atresia, and a high risk of cancer.220,225

EB has no effective therapy or cure at present. Although the longevity of patients with mild forms may not be affected significantly by EB, multiple interventions from a range of medical specialists are required to ensure quality of life. The

due to the severe cytotoxicity of chemoradiotherapy for FA patients, surgical resection is preferred, while HPV vaccination and frequent dental evaluations or bone marrow aspirates are recommended to rule out early cancers.199,211 Additionally, induced pluripotent stem cell transplantation and gene therapy for FA have recently been proposed with promising pre-clinical results.215,216

**Table 3.** Dental-craniofacial manifestations of skin, mucosa and soft tissue abnormality-related rare diseases

| Rare diseases                      | Aetiology                                                                 | Major manifestations                                                                 | Dental-cranio-facial manifestations | Incidencea | Onset periodb |
|-----------------------------------|---------------------------------------------------------------------------|--------------------------------------------------------------------------------------|-------------------------------------|------------|---------------|
| Hereditary epidermolysis bullosa  | Defect in attachment between the epidermis and dermis of the skin         | Hands and feet blisters at the site of rubbing                                        | Intraoral blistering with or without scar formation, oral vestibule; Enamel hypoplasia and/or caries | 38.6%–94.8% | Perinatal period |
| Peutz–Jeghers syndrome            | Mutations in the LKB1 gene                                                 | Benigne hamartomatous polypos in the gastrointestinal tract; skin hyperpigmented macules in hand and feet | Hyperpigmented macules in lip and oral mucosa (Gingiva, hard palate and inside of the cheek). | 18.1–100% | Early childhood |
| Mucopolysaccharidosis             | Absence of or malfunctioning of lysosomal enzymes                         | Developmental delay, intellectual disabilities, short stature; impaired motor function; hearing loss; respiratory distress, obstructive sleep apnoea; enlarged or diseased heart valves | High-arched palate, hypertrophy of the alveolar processes; Enlarged tongue, gingiva and associated anterior open bite; Delayed tooth eruption, impacted teeth | 56.3%–85.7% | Childhood |
| Mikulicz’s disease                | Abnormal IgG4 deposition and related inflammation                         | Continuous painless lachrimal gland swelling; pulmonary interstitial fibrosis           | Painless and persistent parotid, submandibular and sublingual salivary glands swelling | 70%–86.7% | Childhood |
| Primary light-chain amyloidosis   | Abnormal light chains deposition                                           | Renal failure; heart failure; enlarged liver                                           | Macroglossia, submandibular swelling | 54.5%–100% | Adulthood |

*aIncidence of dental-craniofacial manifestations

*bOnset period of dental-craniofacial manifestations
current treatment largely targets symptoms and focuses on
caring for and preventing the formation of new blisters. The
daily management available for EB patients includes wound
care, pain management, and protective bandaging. Attention
must be paid to improve nutrition and help with weight gain.217,222,228 For patients with severe EB, surgical treatment,
such as widening of the oesophagus, placement of a feeding
tube and skin grafting, is optional.227,228 Additionally, working
with a rehabilitation specialist may help relieve the limitations
on motion induced by scarring. Moreover, when performing
therapies, such as restorative dental treatment, local anæsthe-
sia, mucosal lubrication with hydrocortisone cream or triami-
nolone are recommended to relieve severe oral and perioral
scarring, microstomia, ankyloglossia or limited mouth opening.
Furthermore, instructions on toothbrushing and dietary habits
and a comprehensive assessment of caries risk or activity are
also strongly suggested to reduce the likelihood or severity of
caries disease.224,226

Peutz–Jeghers syndrome. Peutz–Jeghers syndrome (PJS) is an
autosomal dominant genetic disease that is mainly caused by
mutations (deletion, insertion, or single base pair substitutions) in the
LKB1 gene on chromosome 19p13.3, which encodes serine-threonine
kinase 11 (STK11) and may function as a tumour suppressor.229
STK11 has been reported to regulate cellular proliferation, apoptosis
and to play an important role in cell polarity, cell metabolism and
energy homeostasis.230–232 The function of LKB1 is complex and is
still being investigated, and no clear genotype-phenotype correlation
has been identified in PJS patients.231

PJS is characterized by intestinal benign hamartomatous polyps
combined with intermittent abdominal pain, as well as hyperpig-
mented macules that vary from 1 to 5 mm in size on the skin (nose,
periorbital, back of hands, and tips of toes and fingers), lip and oral
mucosa (gingiva, hard palate and inside the cheek are most
frequently involved).234 Mostly mucocutaneous pigmentation
appears early in childhood, before the onset of gastrointestinal
disease, and gradually disappears after puberty, but oral lesions
persist throughout life and are usually flat and painless.235 The size
and colour of the pigmentations are not affected by sunlight, unlike
regular ephelides (Table 3). Moreover, although hamartomatous
polyps have an extremely low potential for malignancy, patients with
PJS have an almost 15-fold higher tendency to develop cancer in
parts of the body, such as the pancreas, liver, lungs, breast, ovaries,
uterus, testes and other organs.235,236 Additionally, bowel obstruc-
tion, intussusception and iron-deficiency anaemia due to profound
gastrointestinal bleeding are also frequently reported in PJS patients.237,238

The management of PJS consists of the surveillance and treatment
of the hamartomatous polyps. Resection of the polyps is performed
only when serious bleeding occurs, and enterotomy is usually
performed to resect large nodules.238 As no standard treatment is
currently established for mucocutaneous pigmentation, cryosurgery,
electrodesiccation and the Q-switched ruby laser have been used to
remove these lesions but consistently result in unsuccessful removal
and scarring.239 Surveillance in PJS patients aims to detect sizable
gastroenterological polyps as well as cancer at an early stage to
reduce the likelihood of potential polyp-related complications, such
as intussusception, and improve the survival and cure rate of
potential cancer. Therefore, annual clinical examination and testicular
examination from birth until 12 years of age, colonoscopy and video
capsule endoscopy beginning at 8 years of age, and annual breast
MRI beginning at 25 years of age, among others, are recommended.233

Mucopolysaccharidosis. Mucopolysaccharidoses (MPS) are a
group of genetic metabolic disorders resulting from the absence
or malfunction of acid hydrolase, a kind of lysosomal enzyme that
is required to break down sulphated glycosaminoglycans
(GAGs).241 GAGs are series of long chains of sugar carbohydrates
that are present on both cell surfaces and in the extracellular
matrix in all tissues, such as heparan sulphate, keratan sulphate,
chondroitin sulphate and many more. Over time, due to the
deficiency of lysosomal enzymes, GAGs accumulate in the cells,
blood and connective tissues and subsequently result in cellular
damage and tissue and organ dysfunction.242 There are currently
11 known enzyme deficiencies resulting from more than 40
genetic disorders that account for 7 distinct MPS types.243

Different MPS types share several clinical phenotypes but have
various degrees of severity, and not all of the features may be
apparent at birth but may progress as GAGs start to accumulate.244
Each individual disorder has a wide spectrum of clinical features,
including skeletal dysostosis (prominent forehead, dwarfish with a
short neck and short trunk, claw hands, joint stiffness and spinal
dysostosis), neurologic manifestations (hearing loss, speech and
language delay, intellectual disabilities and learning deficiency,
declining neurological status), motor dysfunction (hyperactivity,
behavioural issues and motor impairment) and other somatic
symptoms (respiratory distress, hepatomegaly, hernias and excessive
body hair growth).245–251 Among the different types of MPS, type II
shows significant CNS involvement, type III shows severe motor
dysfunction, and type IV and VI shows more prominent skeletal
disorders, while the others present both.242 Additionally, distinct
facial manifestations have also been reported in most MPS patients,
especially in type I and VI, such as rough skin, bulging forehead,
depressed nasal bridge, enlarged mouth and thick lips.252 In the oral
cavity, GAG accumulation would contribute to an enlarged tongue,
gingiva and associated anterior open bite, high arched palate,
hypertrophy of the alveolar processes, delayed tooth eruption and
the formation of dental follicles.253–255 (Table 3).

The current treatments include allogeneic HSCT and enzyme
replacement therapy, which is only available for MPS type I, III and VI.
ERT can easily reach the reticuloendothelial organs and significantly
reduce GAGs and related inflammatory cytokines, but it provides
limited improvement of joint pain stiffness and other skeletal
changes.256,257 Thus, ERT is considered to be effective in reducing
the development of pathology but not reversing established
deformities. HSCT is only recommended for young children with
severe MPS, as the application of HSCT can promote the establish-
ment of intellectual development and functions but shows limited
efficacy in preventing osteoarticular deformities.258–261 Additionally,
other therapies, including the application of active site-specific
chaperones that increase residual enzyme activity or genistein that
inhibits GAG synthesis (substrate reduction therapy), nonsteroidal
drug administration for anti-inflammatory treatment, surgery inter-
ventions for correcting deformities or respiratory status, and gene
therapy referring to inserting the wild copy of the defective genes,
have recently been proposed and shown promising results in clinical
or pre-clinical trials.262–266 For patients with severe occlusal
characteristics, such as a marked overjet, anterior open bite and
mandibular protrusion, orthodontic with or without orthognathic,
surgery should be considered to improve quality of life after ERT.253

Mikulicz’s disease. Mikulicz’s disease (MD) was first described in
1892 by Mikulicz as a developmental swelling disorder of the lacrimal
and salivary glands.267 In recent past, MD has been considered
a part of Sjögren syndrome, but with the development of
specific laboratory examinations, it is now accepted as part of
IgG4-related disease (IgG4-RD) and distinct from Sjögren syn-
drome.268 IgG4-RD is a chronic inflammatory condition that is
characterized by Th2 and regulatory immune reactions, manifest-
ing as high serum levels of IgG4 and tissue infiltration with
lymphocytes and IgG4-secreting plasma cells.269,270 However, the
physiological role of IgG4 in IgG4-RD remains unclear.
Clinically, MD patients present dry eye, exophthalmos and dry
mouth, as well as continuous and painless bilateral and
symmetrical enlargement of the related glands, while most MD
patients do not have keratoconjunctivitis sicca. MD usually involves the parotid, submandibular and occasionally sublingual salivary glands, but minor salivary gland involvement can also be observed. Nelson reported a MD patient with only nonulcerated, painless, irregular swelling of the left hard palate in 1963 (Table 3). Additionally, MD occasionally appears in combination with other IgG4-RD, including autoimmune pancreatitis, Riedel’s thyroiditis, Kütten’s tumour and other extra salivary and lacrimal gland lesions.

Currently, MD is mainly treated with immunosuppressive therapy, and it shows a good response to the administration of steroids, including a rapid improvement in glandular swelling and salivary secretion. However, relapse frequently occurs in the absence of therapy.

Primary light-chain amyloidosis. Primary light-chain amyloidosis (AL) is caused by the developmental amyloid deposition of abnormal protein fibres (misfolded free immunoglobulin or \( \lambda \) light chains), which can lead to structural and functional damage of different organs, such as the heart, kidneys, liver and brain. Skin and mucous membrane changes, including purpura, petechiae, ecchymoses and bullous lesions, are often the first features of primary AL patients, while weight loss and severe fatigue usually occur in most primary AL patients. Additionally, other symptoms are also frequently reported, such as heart failure, renal failure or end-stage renal disease, hepatomegaly without scan defects and postural hypotension, and cardiac involvement is the main determinant of survival. Macroglossia, submandibular swelling, alopecia and shoulder pain are not common in primary AL patients, while weight loss and severe fatigue usually occur in most primary AL patients. Additionally, in severe patients with heart or renal failure, renal and cardiac transplantation may improve quality of life and prolong the lifespan.

Others

Angelman syndrome. Angelman syndrome (AS) is a genetic disorder resulting from a new maternal mutation, deletion or imprinting defect in chromosome 15q11-q13, which contains the UBE3A and OCA2 gene, rather than being inherited from the parents. Because the paternal inherited allele of UBE3A is epigenetically silenced in most neurons, maternal deletions of UBE3A result in a nearly total selective loss of brain UBE3A function. The UBE3A gene encodes a HECT (homologous to E6-associated protein C terminus) domain E3 ubiquitin ligase, which ubiquitinates protein substrates such as p53, p27, Pbl/Ect2, Ephexin5 and many more, and leading to their degradation. Multiple mutations in UBE3A contribute to a defective catalytic function and result in disorders when regulating neuronal apoptosis, differentiation and axon outgrowth.

AS is characterized by childhood epilepsy and severe developmental delay with or without mental retardation. Developmental delay in AS individuals is usually observed before the first year of life, which manifests as decreased sleep; speech impairment; poor oral-motor functions, such as sucking and chewing; movement or balance disorders, such as tremors and ataxia; and behavioural uniqueness, such as a happy demeanour, excitability or hyperactivity. Seizures usually start between 1 and 3 years of age, which first only present as mild myoclonic jerks and spells of atypical absences and then become more frequent in early childhood, tending to decrease in adolescence. A typical pattern of large-amplitude slow-spike waves can be noted and used for diagnosis using an electroencephalogram. Additionally, a tendency to develop flexion contractures and valgus deformity of the feet, scoliosis and imperfect manual function are also observed with age in some individuals with AS. Several dental-craniofacial deformities in AS patients have also been reported in several studies, although with a reduced incidence, including a flat occiput, protruding tongue, mandibular prognathia, wide mouth and wide-spaced teeth (Table 4).

Currently, there is no specific treatment for AS, and didactic-supportive therapy is generally applied with the aim of promoting mental development, such as educational training, speech training (the use of communication devices, adapted pictogram and modified sign language), and antiepileptic drug therapy, including sodium valproate and benzodiazepines for idiopathic generalized epilepsies. The remainder of treatment is applied according to the specific manifestations in individuals with AS: disordered sleep is treated with behavioural interventions and melatonin administration; behavioural modification is recommended for hypermotoric or hyperactive behaviours rather than drug therapy; special adaptive chairs or positioners and physiotherapy are provided for ataxic children; orthotic bracing or surgery for subluxed or pronated ankles, thoraco-lumbar jackets

| Table 4. Dental-craniofacial manifestations of other related rare diseases |
| --- |
| **Rare diseases** | **Aetiology** | **Major manifestations** | **Dental-cranio-facial manifestations** | **Incidence** | **Onset period** |
| Angelman syndrome | Genetic mutations in chromosome 15q11-q13 | Developmental delay; movement and balance disorder; behavioural uniqueness (excitability or hyperactivity); seizures; abnormal EEG | Unique behaviour: happy demeanour, poor oral function; Mandible prognathia, protruding tongue, wide mouth and wide-spaced teeth | 75% | Early childhood |
| Langerhans-cell histiocytosis | Abnormal proliferation of Langerhans type cells | Lytic bone lesions; fever; diabetes insipidus (Hand-Schüller-Christian triad); scaly skin lesions in scalp, ear canals, and abdomen (Letterer-Siwe disease) | Ulcers, scabby and granuloma with pain and swelling at oral mucosa; Bone defect in craniofacial bones with punched-out appearance; Gingival necrosis with the movement of teeth | 55%–80% | Childhood or adulthood |

EEG, electroencephalogram

aIncidence of dental-craniofacial manifestations

bOnset period of dental-craniofacial manifestations
for severe scoliosis, and related orthodontic treatment, orthognathic surgery as well as dental restoration for correction of open bite, prominent diastema between the central incisors and other dental-craniofacial deformities are also supplied.295 Additionally, education about oral hygiene, such as regarding brushing techniques, should also be emphasized to maintain a caries-free state in the affected child.

Langerhans cell histiocytosis. Langerhans cell histiocytosis (LCH) is a developmental histiocytosis syndrome characterized by an abnormal proliferation of histiocytes. LCH is marked by excessive proliferation of Langerhans-type cells, which have immunophenotypic and ultrastructural similarities (CD1a antigen) to antigen-presenting Langerhans cells.297 The cause and pathogenesis of LCH remains a matter of debate concerning whether it is a kind of reactive or neoplastic process since Alfred Hand first described and misdiagnosed it as tuberculosis in 1983.298 The cytokine storm in the lesion region and high rate of spontaneous remissions support that LCH is an exaggerated physiological response of Langerhans cells.299,300 while the monoclonal proliferation of pathologic cells and a newly discovered potential somatic mutation of an oncogene, the BRAF gene, in LCH patients provides solid evidence that LCH is a malignancy.301–303

LCH consists of a wide spectrum of clinical disorders that vary from isolated bone lesions to multiple skeletal or visceral lesions, with or without lymph node involvement. Several clinical investigations have demonstrated that the peak incidence of LCH occurs between 1 and 3 years of age and that multiple-organ systematic diseases mostly begins before 2 years of age.304 At present, LCH is usually divided into two types according to the lesion scope: localized LCH (also called eosinophilic granuloma) and disseminated LCH (including Letter-Siwe disease and Hand–Schüller–Christian disease). Localized LCH consists of a simplex rash without affecting organs, simplex bone damage and multiple bone damage with or without diabetes insipidus.305 Diabetes insipidus is one of the most common manifestations in LCH patients due to the involvement of the central nervous system, including the hypothalamus and pituitary.306 Regarding rashes, invasive nodules and plaques or generalized seborrhoeic dermatitis-like rashes of the scalp, skin folds and retroauricular area are commonly reported. Additionally, LCH may also manifest as ulcers, scabs and granuloma with pain and swelling of the oral and genital mucosal region.307 Bone damage mainly involves the axial skeleton, such as the skull bones (orbit and temporal bone), mandible, pelvis and spine. Bone lesions of LCH demonstrate a combination of bone destruction and adjacent soft tissue mass, which can be easily observed using CT and MRI.308 The bone defect in skull bones usually displays a typical punched-out appearance with a scalloped or irregular margin and varies from poorly to well-defined. The supralateral and frontal part of the orbit and the tympanic, mastoid, and squamosal portions of the temporal bone are most frequently affected location in the skull bones.309,310 Multiple bone defects in the mandible can occur and temporal bone are most frequently affected location in the skull orbit and the tympanic, mastoid, and squamosal portions of the temporal bone.308,310 Multiple bone defects in the temporal bone are most frequently affected location in the skull bones.309,310 Multiple bone defects in the mandible can occur and temporal bone are most frequently affected location in the skull orbit and the tympanic, mastoid, and squamosal portions of the temporal bone.

Treatments for LCH are variable depending on the localization and number of lesions, and the aim of treatment is to correct the organ dysfunction as well as to limit the spread of disease. For localized LCH, surgical resection and curettage with or without bone grafting combined with topical steroids or vinblastine are the first-line therapy.311 For disseminated LCH, systemic therapy consisting of steroids and vinblastine is recommended312; cytarabine or HSCT are used as second-line therapy in patients who do not respond to the steroids and vinblastine.314,315 Furthermore, chemotherapy and radiotherapy can also be used alone or in combination for disseminated LCH, but this treatment paradigm is controversial, and strong chemoradiotherapy is not recommended to avoid severe toxic and side effects.316 In addition, a BRAF inhibitor, vemurafenib, was recently approved by the US Food and Drug Administration to treat severe BRAF mutation-positive LCH patients and showed substantial clinical and biological improvements without severe toxicities or adverse events.317

CONCLUSION
Approximately 7,000 rare diseases have been identified around the world, which far exceeds the recently defined 121 rare diseases in China. However, not all rare diseases have effective diagnosis or treatment regimens, and in consideration of the limited resources, it is much more meaningful to focus on the rare diseases with relatively a higher incidence as well as feasible management or precaution tactics. Based on the established ‘Rare Diseases List in China’, it is important for our clinicians to fully comprehend the related clinical features and potential treatments for early diagnosis and management as well as for improving the patient’s prognosis. Moreover, several rare diseases show unique dental-craniofacial manifestations in the early disease period, such as congenital erythrophagocytic porphyria, hypophosphatasia, Marfan syndrome, and Peutz–Jeghers syndrome, among others (Tables 1–4); thus, dentists and oral maxillofacial surgeons could be the first clinicians to identify these rare diseases at an early stage, making it particularly necessary for them to adequately grasp the related clinical features.

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ADDITIONAL INFORMATION
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REFERENCES
1. Dawkins, H. J. S. et al. Progress in rare diseases research 2010-2016: an IRDRC perspective. Clin. Transl. Sci. 11, 11–20 (2018).
2. Aydmé, S. & Schmidtke, J. Networking for rare diseases: a necessity for Europe. Bundesgesundheitsbl. Gesundheitsforsch. Gesundheit 50, 1477–1483 (2007).
3. Azie, N. & Vincent, J. Rare diseases: the bane of modern society and the quest for cures. Clin. Pharmacol. Ther. 92, 135–139 (2012).
4. Schieppati, A., Henter, J.-I., Daina, E. & Aperia, A. Why rare diseases are an important medical and social issue. Lancet 371, 2039–2041 (2008).
5. Melnikova, I. Rare diseases and orphan drugs. Nat. Rev. Drug. Discov. 11, 267–268 (2012).
6. Ekins, S. Industrializing rare disease therapy discovery and development. Nat. Biotechnol. 35, 117–118 (2017).
7. James, W. D., Elston, D. M. & Berger, T. G. Ectodermal dysplasias: a clinical classification. Am. J. Med. Genet. A. 162 (1994).
8. Pinheiro, M. & Freire-Maia, N. Ectodermal dysplasias: a clinical classification and a causal review. Am. J. Med. Genet. A. 53, 153–162 (1994).
9. Halai, T. & Stevens, C. Ectodermal dysplasia: a clinical overview for the dental practitioner. Dent. Update 42, 779–790 (2015).
10. Lexer, M. O., Bardow, A., Hertz, J. M., Nielsen, L. A. & Kreiborg, S. Anomalies of tooth formation in hypohidrotic ectodermal dysplasia. Int. J. Paediatr. Dent. 17, 10–18 (2007).
11. Leao, J. C., Ferreira, A. M. C., Bandeira, V., Figueredo, F. V. & Porter, S. R. Anhydrotic ectodermal dysplasia (Christ-Siemens-Touraine syndrome). A case report. Int. Dent. J. 55, 89–92 (2005).
12. Crawford, P. J., Aldred, M. J. & Clarke, A. Clinical and radiographic dental findings in X linked hypohidrotic ectodermal dysplasia. J. Med. Genet. 28, 181–185 (1991).
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75. Land, C., Rauch, F., Montpetit, K., Ruck-Gibis, J. & Glorieux, F. H. Effect of intra-venous pamidronate therapy on functional abilities and level of ambulation in osteogenesis imperfecta. J. Pediatr. 148, 456–460 (2006).
76. Esposito, P. & Plotkin, H. Surgical treatment of osteogenesis imperfecta: current concepts. Curr. Opin. Pediatr. 20, 52–57 (2008).
77. Krishnan, H. et al. Primary and revision total hip arthroplasty in osteogenesis imperfecta. Hip Int. 23, 303–309 (2013).
78. Topouchian, V., Finidori, G., Glorion, C., Padovani, J. P. & Pouliquen, J. C. Posterior spinal fusion for kyphoscoliosis associated with osteogenesis imperfecta: long-term results. Rev. Chir. Orthop. Reparatrice. Appar. Mot. 90, 525–532 (2004).
79. Janus, G. J. M., Finidori, G., Engelbert, R. H. H., Pouliquen, M. & Pruijs, J. E. H. Operative treatment of severe scoliosis in osteogenesis imperfecta: results of 20 patients after halo traction and posterior spondylodesis with instrumentation. Eur. Spine J. 9, 486–491 (2000).
80. McAllion, S. J. & Paterson, C. R. Causes of death in osteogenesis imperfecta. Bone 28, 51–56 (2001).
81. Krishnan, H. et al. Primary and revision total hip arthroplasty in osteogenesis imperfecta. Hip Int. 23, 303–309 (2013).
82. Murayama, T., Iwatsubo, R., Akiyama, S., Amano, A. & Morisaki, I. Familial dental problems in hypophosphatemic rickets. J. Clin. Invest. 124, 1587–1594 (2014).
83. Imel, E. A. et al. Prolonged correction of serum phosphorus in adults with X-linked hypophosphatemia using monthly doses of KRN23. J. Clin. Endocrinol. Metab. 100, 2565–2573 (2015).
84. Linglart, A. & Biosse-Duplan, M. Hypophosphatasa. Curr. Osteoporos. Rep. 14, 95–105 (2016).
85. Lia-Baldini, A. et al. A molecular approach to dominance in hypophosphatasia. Hum. Genet. 109, 99–108 (2001).
86. Millan, J. L. The role of phosphatases in the initiation of skeletal mineralization. Calcif. Tissue Int. 93, 299–306 (2013).
87. Millan, J. L. & Plotkin, H. Hypophosphatasia - pathophysiology and treatment. Actual. Osteol. 8, 164–182 (2012).
88. Silva, L., Castello, W., Mateus, M. & Branco, J. C. Childhood hypophosphatasia with myopathy: clinical report with recent update. Acta Reumatol. Port. 37, 92–96 (2012).
89. Fraser D. Hypophosphatasia. Am. J. Med. 22, 740–746 (1957).
90. Whyte, M. P. et al. Hypophosphatasia: validation and expansion of the clinical nosology for children from 25years experience with 173 pediatric patients. Bone 55, 229–239 (2015).
91. Whyte, M. P. Hypophosphatasia — aetiology, pathology,ogenesis, diagnosis and treatment. Nat. Rev. Endocrinol. 12, 233–246 (2015).
92. Shohat, M., Rimoin, D. L., Graz, H. E. & Lachman, R. S. Perinatal lethal hypophosphatasia: clinical, radiologic and morphologic findings. Pediatr. Radiol. 21, 421–427 (1991).
93. Silver, M. M., Vilos, G. A. & Milne, J. K. Pulmonary hypoplasia in neonatal hypophosphatasia. Pediatr. Pathol. 8, 483–493 (1988).
94. Whyte, M. P. et al. Aftosia alfa treatment improves survival for perinatal and infantile hypophosphatasia. J. Clin. Endocrinol. Metab. 101, 334–342 (2016).
95. Bianchi, M. L. Hypophosphatasia: an overview of the disease and its treatment. Osteopors. Int. 26, 2743–2757 (2015).
96. Whyte, M. P. Hypophosphatasia: an overview for 2017. Bone 102, 15–25 (2017).
97. Hofmann, C. et al. Clinical aspects of hypophosphatasia: an update. Clin. Rev. Bone Miner. Metab. 11, 60–70 (2013).
98. Mornet, E. Hypophosphatasia. Best Pract. Res. Clin. Rheumatol. 22, 113–127 (2008).
99. Taketani, T. et al. Clinical and genetic aspects of hypophosphatasia in Japanese patients. Arch. Dis. Child. 99, 211–215 (2015).
100. Whyte, M. P. Physiological role of alkaline phosphatase expressed in hypophosphatasia. Ann. N. Y. Acad. Sci. 1192, 190–200 (2010).
101. Arun, R., Khazim, R., Webb, J. K. & Burn, J. Scisosis in association with infantile hypophosphatasia: a case study in two siblings. Spine 30, E471–E476 (2005).
102. Sutton, R. A. L., Mummm, S., Coburn, S. P., Ercion, K. L. & Whyte, M. P. “Atypical femoral fractures” during bisphosphonate exposure in adult hypophosphatasia. J. Bone Miner. Res. 27, 987–994 (2012).
103. Khandwala, H., Mummm, S. & Whyte, M. Low serum alkaline phosphatase activity and pathologic fracture: case report and brief review of hypophosphatasia diagnosed in adulthood. Endocr. Pract. 12, 676–681 (2006).
104. Coe, J. D., Murphy, W. A. & Whyte, M. P. Management of femoral fractures and pseudofractures in adult hypophosphatasia. J. Bone Jt. Surg. 88, 981–990 (1988).
105. Guallar, L. N. et al. Calcific periarthritis as the only clinical manifestation of hypophosphatasia in middle-aged sisters. J. Bone Miner. Res. 29, 929–934 (2014).
106. Beck, C., Morbach, H., Richl, P., Stenzel, M. & Grischik, H. J. How can calcium pyrophosphate crystals induce inflammation in hypophosphatasia or chronic inflammatory joint diseases? Rheumatol. Int. 29, 229–238 (2009).
107. Rodriguez, E. et al. Respiratory mechanics in an infant with perinatal lethal hypophosphatasia treated with human recombinant enzyme replacement therapy. Pediatr. Pulmonol. 47, 917–922 (2012).
108. de Roo, M. G. A. et al. Infantile hypophosphatasia without bone deformities presenting with severe pyridoxine-resistant seizures. Mol. Genet. Metab. 111, 404–407 (2014).
109. Baumgartner-Sigl, S. et al. Pyridoxine-responsive seizures as the first symptom of infantile hypophosphatasia caused by two novel missense mutations.
260. Parini, R. et al. Open issues in mucopolysaccharidosis type I-hurler. Orphanet J. Rare Dis. 12, 112 (2017).
261. Kubaski, F. et al. Hematopoietic stem cell transplantation for patients with mucopolysaccharidosis II. Biol. Blood. Marrow Transplant. 23, 1795–1803 (2017).
262. Fan, J.-Q. A counterintuitive approach to treat enzyme deficiencies: use of enzyme inhibitors for restoring mutant enzyme activity. Biochem. Biophys. Res. Commun. 389, 1–11 (2008).
263. Pirottonvka, E. et al. Two-year follow-up of sanfilippo disease patients treated with a genistein-rich isoflavone extract: assessment of effects on cognitive functions and general status of patients. Med. Sci. Monit. 17, CR196–CR202 (2011).
264. Giuigliani, R., Harmatz, P. & Wrash, J. E. Management guidelines for mucopolysaccharidosis VI. Pediatrics 120, 405–418 (2007).
265. Ohn, H. & Lee, J. Current and potential therapeutic strategies for mucopolysaccharidoses. J. Clin. Pharm. Ther. 39, 215–224 (2014).
266. Sawamoto, K. Chen, H. H., Almeicia-Diaz, C. J., Mason, R. W. & Tomatsu, S. Gene therapy for mucopolysaccharidosis. Mol. Genet. Metab. 123, 59–68 (2018).
267. Mikulicz. J. Über Eine Eigenartige Symmetrische Erkrankung der Trunen und Mundspeicheldrusen (Stuttgart: Theodor Billroth, 1892).
268. Yao, Q., Wu, G. & Hoschar, A. IgG4-related Mikulicz’s disease is a multigland lymphoproliferative disease distinct from Sjögren’s syndrome: a Caucasian patient and literature review. Clin. Exp. Rheumatol. 31, 289–294 (2013).
269. Umemura, H. et al. Comprehensive diagnostic criteria for IgG4-related disease (IgG4-RD). 2011. Mod. Rheumatol. 21, 23–30 (2012).
270. Maehara, T. et al. Interleukin-21 contributes to germinal centre formation and immunoglobulin G4 production in IgG4-related dacrocyoadenitis and salivary gland, so-called Mikulicz’s disease. Ann. Rheum. Dis. 71, 2011–2020 (2012).
271. Bhatti, R. M. & Stelow, E. B. IgG4-related disease of the head and neck. Arch. Otolaryngol. Head Neck Surg. 135, 885–893 (2009).
272. Lim, J. & Feldman, B. Current and potential therapeutic strategies for IgG4-related diseases. Front. Immunol. 9, 125 (2018).
273. Yamamoto, M., Takahashi, H., Sugai, S. & Imai, K. Clinical and pathological characteristics of Mikulicz’s disease (IgG4-related plasmacytic exocrinopathy). Autoimmun. Rev. 4, 195–200 (2005).
274. Nelson, W. R. Kay, S. & Salley, J. J. Mikulicz’s disease of the palate. Ann. Surg. 157, 152–156 (1963).
275. Hamano, H. et al. High serum IgG4 concentrations in patients with sclerosing pancreaticitis. N. Engl. J. Med. 344, 732–738 (2001).
276. Nied, G. et al. Inflammatory lesions of the lung, submandibular gland, bile duct and prostate and patients with IgG4-associated multifocal systemic fibro-sclerosis. Respir. Res. 12, 455–457 (2007).
277. Kitagawa, S. et al. Abundant IgG4-positive plasma cell infiltration characterizes chronic sclerosing sialadenitis (Kuttner’s tumor). Am. J. Surg. Pathol. 29, 783–791 (2005).
278. Abe, A. et al. The clinical characteristics of patients with IgG4-related disease with infiltration of the labial salivary gland by IgG4-positive cells. Mod. Rheum. Mat. 24, 949–952 (2014).
279. Himi, T., Takano, K., Yamamoto, M., Naishiro, Y. & Takahashi, H. A novel concept of Mikulicz’s disease as IgG4-related disease. Auris Nasus Larynx 39, 9–17 (2012).
280. Moriyama, M. et al. Clinical characteristics of Mikulicz’s disease as an IgG4-related disease. Clin. Oral Investig. 17, 1995–2002 (2013).
281. Merlini, G. & Belotti, F. Molecular mechanisms of amyloidosis. N. Engl. J. Med. 349, 583–596 (2003).
282. Borowicz, J., Gillespie, M. & Miller, R. Cutaneous amyloidosis. Skinmed 9, 96–100 (2011). quiz 101.
283. Merlini, G., Comenzo, R. L., Seldin, D. C., Wechalekar, A. & Gertz, M. A. Immunoglobulin light chain amyloidosis. Expert. Rev. Hematol. 7, 143–156 (2014).
284. Caccialanza, R. et al. Nutritional status of outpatients with systemic immunoglobulin light chain amyloidosis. Am. J. Clin. Nutr. 83, 350–354 (2006).
285. Milani, P., Merlini, G. & Palladini, G. Light chain amyloidosis. Mediters. J. Hematol. Infect. Dis. 10, e2018022 (2018).
286. Nelson, L. M., Gustafsson, F. & Gimsing, P. Characteristics and long-term outcome of patients with systemic immunoglobulin light chain amyloidosis. Acta Haematol. 133, 336–347 (2014).
287. Prokaeva, T. et al. Soft tissue, joint, and bone manifestations of AL amyloidosis: clinical presentation, molecular features, and survival. Arthritis Rheum. 56, 3858–3868 (2007).
288. Palladini, G. Association of melphalan and high-dose dexamethasone is effective and well tolerated in patients with AL (primary) amyloidosis who are ineligible for stem cell transplantation. Blood 103, 2936–2938 (2004).
289. Merlini, G., Wechalekar, A. D. & Palladini, G. Systemic light chain amyloidosis: an update for treating physicians. Blood 121, 5124–5130 (2013).
290. Gertz, M. A. Immunoglobulin light chain amyloidosis: 2014 update on diagnosis, prognosis, and treatment. Am. J. Hematol. 89, 1132–1140 (2014).
291. Clayton-Smith, J. & Laan, L. Angelman syndrome: a review of the clinical and genetic aspects. J. Med. Genet. 40, 87–95 (2003).