Management of children with amelogenesis imperfecta: A Case series

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INTRODUCTION

Amelogenesis imperfecta (AI) is a group of hereditary conditions that primarily affect the quality and/or the quantity of dental enamel, with a variable occurrence of approximately 1/700-1/14,000.

In general, the inheritance pattern is usually autosomal dominant or recessive, or X linked. (Koruyucu M et al., 2014) (Leevailoj C et al., 2017) (Pathak A et al., 2011) (Reddy Ss et al., 2010) (Kala Vani S et al., 2012).

Witkop’s classification is the most frequently cited one in the literature. Four major categories based primarily on phenotype (hypoplastic (HP), hypomaturation (HM), hypocalcified (HC), HM/HP with taurodontism) subdivided into 15 subtypes based on clinical phenotype and mode of inheritance (Aldred M et al., 2003).

AI has been associated with different dental anomalies including discolored teeth, microdontia, congenitally missing teeth, taurodontism, pulp calcification, hypercementosis, crown and root resorption. In addition, AI was shown to be associated with malocclusions such as open bite and cross bite. (Leevailoj C et al.,2017) Apart from these difficulties, this condition also brought psychological problem and lack of self-confidence due to the appearance of their teeth. There are also other difficulties of managing AI including dental inexperience and anxiety of a child and parental expectation all of which further complicate its management.

Amelogenesis imperfecta affects deciduous dentition as well as a permanent dentition and requires a specific preventive and restorative care approach.

The aim of this article was to present the diagnostic elements of AI and the different therapeutic management.

CASE PRESENTATION

Case Report 1:

A 13-year-old boy was referred to the department of Pediatric and Preventive Dentistry, Monastir Dental Faculty, Tunisia, with an aesthetic chief complaint. Apart from the consanguineous marriage, his parents had past dental and medical history that were noncontributory. There was no family history of AI and his elder male siblings showed normal dental development.

Intraoral examination revealed yellow to yellowish-brown teeth with rough surfaces, conspicuous and irregular defects, a lack of contact points and gingival hypertrophy. The enamel alterations were
generalized affecting both lower and upper teeth (Fig1.a,b.c).

The panoramic radiography showed deciduous teeth and incomplete permanent dentition with a delayed eruption and several impacted teeth.

No density difference was observed between enamel and dentin. The molars presented intrapulpal calcifications (Fig1.d).

A diagnosis of hypoplastic AI was made, transmission of the genes took place by autosomal recessive mode. The patient was further investigated. Ultrasound of kidney, ureter, bladder showed medullary nephrocalcinosis with bilateral multiple calculi. Based on the systemic investigation, patient has been diagnosed to have bilateral medullary calcinosis (Fig1.e).

A treatment plan was developed which the main objective was to:
- reduce sensitivity
- improve esthetics
- Restore masticatory function.

First, the patient was advised to brush with warm water and to use a normal fluoride toothpaste.

Second, a gingivectomy and gingivoplasty were performed to adjust the height of the cervical line.

Finally, stainless steel crowns were made on the permanent molars to slightly increase and maintain vertical dimension. Polycarbonate crowns were made on the permanent incisors, canines and premolars (Fig1.f.g.h).

The Patient was addressed to The Nephrologist and the Orthodontist.

![Figure 1. Case 1 (a) intraoral frontal view before treatment (b) intaoral occlusal maxillary view before treatment (c) intraoral occlusal mandibular view before treatment (d) panoramic radiography (e) Ultrasound of kidney (f) intraoral frontal view after treatment (g) intraoral occlusal maxillary view after treatment (h) intraoral occlusal mandibular view after treatment](image-url)
Case Report 2:

A girl aged 14 years old, referred to the department of Pediatric and Preventive Dentistry, Monastir Dental Faculty, Tunisia, regarding the unsightly teeth and dental hypersensitivity.

The patient came from a consanguineous marriage and her sister had the same dental anomaly. Her general, physical, and clinical findings were normal.

Intraoral examination showed generalized gingival inflammation, poor hygiene, anterior open bite deformity, yellow-brown rough fractured enamel of all the teeth and dilapidated maxillary permanent molar (16) (Fig.2.a.b.c).

Radiographic examination revealed no density difference between enamel and dentin, inclusion of the following teeth (32; 43; 23) and persistence of the left primary maxillary canine (63) (Fig.2.d)

The patient was diagnosed with AI (hypoplastic and hypomineralized type) via an X-linked-gene transmission mode.

Ultrasound of kidney, ureter, bladder were done and revealed no link between AI and nephrocalcinosis syndrome.

Treatment objectives taken into account were:
- preservation of tooth integrity and vitality of permanent teeth
- reducing sensitivity and improve esthetics
- normalization of oral functions

Treatment included oral prophylaxis, gingivectomy, and extraction of dilapidated permanent molar (16) and persistent primary canine (63).

Esthetic rehabilitation of decayed anterior teeth (incisors, canine and premolar) was done with polycarbonate crowns. Stain steel crowns was made on the permanent molars and removable partial denture for the extracted molar to obtain normal occlusion. (fig.2.e.f.g)

The patient was referred to the orthodontic department to correct the anterior open bite.

Figure 2. Case 2 (a) intraoral frontal view before treatment (b) intraoral occlusal maxillary view before treatment (c) intraoral occlusal mandibular view before treatment (d) panoramic radiography (e) intraoral frontal view after treatment (f) intraoral occlusal maxillary view after treatment (g) intraoral occlusal mandibular view after treatment
Case report 3:
An 11-year-old boy consulted the department of Pediatric and Preventive Dentistry, Monastir Dental Faculty, Tunisia, regarding the enhancing appearance of his teeth. He had a familial history of AI affecting her father, grandmother and several aunts and sister.

Oral examination revealed yellow-brown teeth with rough enamel; some enamel breaks in the anterior teeth generalized gingival inflammation, a lack of contact points and an anterior open bite (Fig.3.a.b.c)

Panoramic radiography showed a reduction in the enamel thickness, a similar X-ray density between enamel and dentin, and the absence of the right second mandibular premolar: 45. (Fig.3.d)

A clinical diagnosis of hypoplastic AI was made and the mode of gene-transmission took place by X.

The patient was further investigated. Ultrasound of kidney, ureter, bladder were normal: The diagnosis of nephrocalcinosis was eliminated.

A treatment plan was developed with the following objectives:
- to improve the esthetics
- to reduce the reported sensitivity of the teeth
- to restore the masticatory function.

Restoration of anterior teeth with polycarbonate crowns and posterior teeth with stainless steel crowns was planned (Fig.3.e.f.g).

The patient was referred to the orthodontic department to correct the anterior open bite.

![Figure 3. Case 3 (a) intraoral frontal view before treatment (b) intraoral occlusal maxillary view before treatment (c) intraoral occlusal mandibular view before treatment (d) panoramic radiography (e) intraoral frontal view after treatment (f) intraoral occlusal maxillary view after treatment (g) intraoral occlusal mandibular view after treatment](image)

Case report 4:
A 9 years old girl referred to the department of Pediatric and Preventive Dentistry, Monastir Dental Faculty, Tunisia, with a complaint of esthetic inadequacy of his teeth and dental hypersensitivity. Her family history confirmed the consanguinity between her parents and showed normal dental development.
The patient general condition was good but she showed a very low self-esteem because of her poor appearance.

Intraoral examination revealed mixed dentition, yellow permanent teeth with rough surfaces, a lack of contact points and gingival hypertrophy (Fig.4.a,b,c)

On the panoramic radiography, no density difference was observed between enamel and dentin. (Fig.4.d)

A diagnosis of hypoplastic AI was made with an autosomal recessive gene-transmission mode.

Treatment was discussed according to several objectives:
- functional restoration
- aesthetic restoration
- non-invasive rehabilitation that allowed evolution during growth

Treatment contained oral prophylaxis, gingivectomy, polycarbonate crowns on the incisors and canines to rehabilitate aesthetic of decayed anterior teeth and stainless steel crowns on the molars to obtain normal occlusion. (Fig.4.e,f)

**DISCUSSION**

Amelogenesis imperfecta is a developmental disturbance that interferes with normal enamel formation. (American Academy of pediatric Dentistry, 2013).

Pediatric dentists are often the first to encounter children with AI and it is imperative that treatment requires an overall comprehensive plan that will also include a rough draft of future treatment needs (Markovic D et al., 2010).

Early identification of the disorder and preventive measures are crucial for children with AI (Markovic D et al.,2010).

Diagnosis is based on the family history, pedigree plotting, meticulous clinical and radiological observation. Dental radiographs of AI teeth provide important information to the clinician with respect to the degree of enamel mineralization to design an appropriate treatment plan (Reddy Ss et al.,2010)(Crawford PJ et al, 2007)

However, this could be challenging to achieve aesthetic improvement and functional rehabilitation in
AI patients (Koruyucu M et al., 2014) (Markovic D et al., 2010).

Poor aesthetics, worn dentition, and interocclusal space problems (both insufficient space and open bite) complicate the treatment course of AI patients (Markovic D et al., 2010).

Oral hygiene can be particularly difficult for these patients due to the sensitivity while brushing. The use of warm water for tooth brushing will go some way towards relieving symptoms while rinsing. Also, regular use of fluoride mouthwashes can help to reduce sensitivity and prevent caries. (Sabandal MMI et al., 2017) (Crawford PJ et al., 2007)

The progression of treatment during childhood has been described as a temporary phase followed by a transitory phase.

In infancy, the primary dentition is protected by the use of preformed metal crowns on posterior teeth. Either polycarbonate crowns or composite restorations are used on anterior teeth. (Crawford PJ et al., 2007) (Athanasiou A et al., 2014)

In the mixed dentition, stainless steel crowns, onlays for permanent molars, composite resin or glass ionomer cement for primary and permanent teeth. (Crawford PJ et al., 2007) (Athanasiou A et al., 2014).

For permanent incisors, direct or indirect composite veneers will improve aesthetics, reduce sensitivity and reduce incisal wear. Good periodontal health and eruption process are important for long term success of restorative treatments (Crawford PJ et al., 2007).

However, there is still no standard formula, current protocol or guideline for successful treatment. Initial treatment should be planned depending on AI type, severity and oral health habits of the patient. (Sabandal MMI et al., 2017) (Athanasiou A et al., 2014)

The success of bonding of composite resins to teeth affected by AI type, is related to how the enamel responds to acid etching. High failure rates with adhesive restorations in patients with AI have been described, especially with HC AI. In this type of AI, the enamel has a higher protein content, which might reduce the bonding of restorations. Although deproteinization in HC AI has been reported to enhance the bonding of composite restorations, recent studies concluded that the procedure of deproteinization had no significant effect on the success of the adhesive restorations. (Neto NL et al) (Saroglu Sonmez, 2008)

In some of the previous case reports, it has been suggested that children with apparently autosomal recessive AI should, at least, have a renal ultrasound examination to exclude the association with nephrocalcinosis syndrome. (Kala Vani S et al., 2012) (Hunter L et al., 2007) (Patel A et al., 2015)

In fact, since nephrocalcinosis is often asymptomatic and can be associated with impaired renal function, pediatric-dentists who see children with a generalized and thin hypoplastic AI should consider a renal ultrasound scan and referral to a Nephrologist. Children with nephrocalcinosis should also be considered for a dental check. (Kala Vani set al., 2012) (Hunter L et al., 2007) (Patel A et al., 2015)

**CONCLUSION**

AI is a disorder that should be diagnosed in an early stage to minimize the destructions of the teeth and to avoid its deleterious consequences.

The treatment plan should be based on patient’s age, type of defects and individual needs of the patients.

An interdisciplinary approach should be used to treat the patient starting at the first stage, not only due to functional and aesthetic reasons, but also for the positive psychological impact on young patient.

**REFERENCES**


7. Koruyucu, M., Bayram, M., Tuna, E. B., Gencay,


