Orthodontically Relevant Manifestations in People with Rare Diseases

Marcel Hanisch\textsuperscript{a, c}, Lale Hanisch\textsuperscript{b}, Johannes Kleinheinz\textsuperscript{a}, Gholamreza Danesh\textsuperscript{b}, Korbinian Benz\textsuperscript{c}, Joachim Jackowski\textsuperscript{c}

\textsuperscript{a}Research Unit Rare Diseases with Orofacial Manifestations (RDOM), Department of Cranio-Maxillofacial Surgery, University Hospital Münster, Münster, Germany; \textsuperscript{b}Department of Orthodontics, School of Dentistry, Faculty of Health, Witten/Herdecke University, Witten, Germany; \textsuperscript{c}Department of Oral Surgery and Dental Emergency Care, School of Dentistry, Faculty of Health, Witten/Herdecke University, Witten, Germany

Significance of the Study

- In the future, digital databases will gain in importance especially in the diagnosis of rare diseases. The orthodontist can ascertain several rare diseases on the basis of typical symptoms, such as oligodontia or dysgnathia, after which the patient can get more detailed information on the digital database.

Keywords

Rare diseases · Orofacial manifestations · Oral medicine · Orthodontics · Interdisciplinary dentistry

Abstract

Background: Approximately 15\% of all rare diseases occur with orofacial manifestations. Symptoms and manifestations of relevance to orthodontists represent a considerable proportion of these diseases and require appropriate strategies for their treatment. This article provides an overview of the orthodontically relevant manifestations of rare diseases.

Material and Methods: Overall, 3,639 rare diseases listed at the Orphanet, OMIM or Pubmed database were evaluated for orofacial manifestations. All rare diseases which were indicated with at least one orofacial manifestation were recorded in a database for rare diseases with orofacial manifestations called “ROMSE,” which was developed by the authors. All the rare diseases were analysed with regard to orthodontically relevant orofacial manifestations, such as dysgnathia, changes in the number of teeth, failures of eruption, pathologies of bone metabolism or orofacial clefts. For all rare diseases with orthodontic relevance, an exact analysis was undertaken. Results: The orthodontically relevant orofacial manifestation termed dysgnathia is described in 151 of 535 identified rare diseases (28.2\%). In these 151 rare diseases, 15 different subforms of dysgnathia, in the sense of skeletal misdevelopments of the jaws but without dental abnormalities, were described. Also changes in the number of teeth (17.9\%), orofacial clefts (27.6\%), failures of eruption (8.4\%) and pathologies of the bone (2.1\%) were described. Conclusions: Orthodontics play an important role in the diagnosis and treatment of orofacial manifestations in rare diseases. Databases such as ROMSE are a first step toward providing valid information in publicly accessible databases.
Introduction

Around 4 million people in Germany are affected by various diseases [1]. Studies have shown that approximately 15% of all rare diseases occur with orofacial manifestations [2–4]. Symptoms and manifestations of relevance to orthodontists represent a considerable proportion of these diseases and require appropriate strategies for their treatment [5]. Often, the cases treated are so comprehensive and complex that an interdisciplinary treatment plan becomes necessary [5]. The symptoms that are relevant for orthodontists are dysgnathia, changes in the number of teeth (anodontia, hypodontia, oligodontia), cleft anomalies and bone diseases such as cherubism [6]. On average, a rare disease is present for 7 years before it is correctly diagnosed [7].

According to a definition given by the European Union (EU), a disease is classified as “rare” when fewer than 1 in 2,000 people is affected by it. This definition applies to at least 30 million people in the 28 member states of the EU [8]. Since 2009, rare diseases in the EU have become the focus of public attention after the Council of the EU called upon member states to draw up plans and strategies for rare diseases at the appropriate level. These measures aim to ensure proper medical care for those affected. This should include diagnosis, treatment and rehabilitation measures [9]. For member states, the aim was to establish and accept a plan in this field by the end of 2013. Alongside this, results of a study commissioned by the German Ministry of Health were published in 2009 in which measures were presented to improve the situation relating to the health of people with rare diseases in Germany [10]. This report assessed the importance of rare diseases as being considerable. Among other aspects, research work was accorded a very high level of importance for the future care of patients with rare diseases. Moreover, the report considered it appropriate to have a “National Plan of Action for People with Rare Diseases” as drawn up by a National Action League. In October 2010 in Germany, the National Action League for People with Rare Diseases was subsequently set up with the aim of putting into practice the recommendations of the European Council. Finally, in 2013, the German Cabinet decided on a National Plan of Action for People with Rare Diseases for Germany, which ends in 2018. This plan comprises 7 fields of action and a total of 52 proposals [11]. The demands include the provision of sources of information and compiling epidemiological population-based patient registers [11].

Against this political background, a database for rare diseases with orofacial manifestations was first set up in 2013 [12]. The purpose of this article is to provide an overview of the orthodontically relevant manifestations of rare diseases described in Orphanet [13], OMIM [14],

---

**Fig. 1.** Subforms of dysgnathia described in the recorded rare diseases (n).

---
PubMed [15] and the ROMSE [16] database, which was developed by the authors based on these data, and to analyse them and present the problems associated with the correct diagnosis of rare diseases.

**Materials and Methods**

The ethical approval for this study was obtained from the ethical review committee (ref. No. 2017-374-f-N), Ethikkommission der Ärztekammer Westfalen-Lippe und der Westfälischen Wilhelms-Universität, Münster, Germany. Overall 3,639 rare diseases listed at the Orphanet Classification of Rare Diseases [17] were evaluated for orofacial manifestations in the Orphanet [13], OMIM [14] and Pubmed [15] databases between November 2012 and January 2018. All rare diseases indicated with at least one orofacial manifestation were incorporated in the specialized database for Rare Diseases with Orofacial Manifestations “ROMSE” (http://romse.org), which was developed by the authors [12, 16]. All the recorded rare diseases were analysed with regard to orthodontically relevant orofacial manifestations, such as dysgnathia, changes in the number of teeth, failures of eruption, pathologies of bone metabolism or orofacial clefts. For all rare diseases with orthodontic relevance, such as dysgnathia and the classification of the various forms that it takes, an exact analysis was undertaken.
Results

Overall 535 rare diseases with orofacial manifestations had been recorded and could be incorporated in the specialized database ROMSE. The orthodontically relevant orofacial manifestation termed dysgnathia is described in 151 of these 535 identified rare diseases (28.2%). Under the main category “dysgnathia,” in the sense of skeletal misdevelopments of the upper or lower jaw but without dental abnormalities, 15 subforms were described in these 151 rare diseases. These include among others: microgenia \((n = 65)\), micrognathia \((n = 17)\), high arched palate \((n = 33)\), mandibular prognathism \((n = 22)\), retrognathia \((n = 7)\) or hyperplasia in the maxilla \((n = 4)\). Figure 1 provides an overview of the subforms of dysgnathia described in the recorded rare diseases. Dental abnormalities comprise subforms including failures of eruption (8.4%) such as tooth retentions \((n = 13)\), dentitio tarda \((n = 30)\) or dentitio praecox \((n = 2)\). Changes in the number of teeth are described in a total of 96 of the rare diseases (17.9%) and these concern hyperdontia \((n = 9)\), hypodontia \((n = 59)\), oligodontia \((n = 20)\), anodontia \((n = 6)\) or hyperhypodontia \((n = 2)\). Further orthodontically relevant orofacial manifestations described in rare diseases are cleft lips and palates in 27.6% \((n = 148)\) and pathologies of the bone in 2.1% \((n = 11)\).

Discussion

The results presented above underline the high proportion of orofacial manifestations that occur in rare diseases and that can be categorized under “Orthodontics.” Dysgnathia, for example, is described in almost one third of the rare diseases with orofacial manifestations. If the changes in the number of teeth are added, their proportion rises even further. In many cases, however, the orofacial manifestations in rare diseases have not been described by dentists or orthodontists but are based on
medical or genetic publications [12]. This is attributable to the scarcity, or even lack, of publications relating to dental/orthodontic practice or oral surgery, and it has resulted in terminology for orofacial findings that is sometimes incorrect. Thus, in the case of rare diseases with no recorded orthodontic findings, no assumptions can be made with any certainty as to whether, for example, the disease involves a genuine mandibular prognathism or a maxillary retrognathism. We suggest that a structured standardized recording should be undertaken in collaboration with university hospitals or within the “European Reference Networks” [18].

On average, a rare disease is present for 7 years before it is correctly diagnosed [7], whereas oral manifestations can serve as an important indicator of the underlying disease [19, 20]. Especially in patients with orthodontically relevant symptoms, such as dysgnathia or oligodontia, these not only can give a clue to the underlying disease, but also indicate serious impact on the oral health-related quality of life [21]. Research has also shown that patients with syndromic oligodontia which had not been linked to a rare disease in the first place reported substantially worse outcomes [22]. Especially when it comes to the combination of the symptoms oligodontia and dysgnathia, as found for example in odonto-onychodermal dysplasia, the orthodontist has an important role to play in recognizing this in the context with a rare disease (Fig. 2–10). In particular, these externally otherwise inconspicuous forms of ectodermal dysplasia are often not recognized, resulting in years of uncertainty for the persons concerned [22].

Dental and medical background information, in particular concerning orofacial manifestations, plus material for further reading, can be retrieved via the ROMSE database mentioned above, although its limitations have to be taken into account. At present, this database is available only in German. An English version is currently being set up so that the database can be used globally. In the future, however, computer-based systems will offer health professionals passive decision support. On the basis of symptoms, structured, weighted lists of possible differential diagnoses are presented to the user. By using such tools diagnostic quality could be enhanced and diagnostic findings accelerated [23]. Databases, such as ROMSE, are thus a first step to provide the necessary data for the construction of an intelligent software system [23]. Furthermore, the demand for freely accessible information is high. This is shown by the large number of websites on the Internet run by patients’ organizations [24]. Therefore, databases like ROMSE should be available as a source of information for patients and their relatives.

The orofacial manifestations of rare diseases, such as dysgnathia or changes in the number of teeth (e.g. oligodontia, hypodontia), presented here make clear the important role that orthodontics play in the early diagnosis and treatment of rare diseases. Genetic diseases such as ectodermal dysplasias with WNT 10A mutations can
Orthodontists have not yet produced structured findings relating to such rare diseases, of which there are many. Databases, such as ROMSE, are a first step toward providing valid information in publicly accessible databases.

Disclosure Statement

The authors declare no conflict of interests.