Review Article

Dental implants in patients with oral autoimmune diseases

Theodora S. Tounta

Private Dentist, Athens, Greece

Abstract

Oral mucosal autoimmune diseases include a variety of disorders, like oral lichen planus (OLP), pemphigus vulgaris (PV), mucous membrane (MMP) and bullous pemphigoid (BP), epidermolysis bullosa acquisita (EBA), systemic lupus erythematosus (SLE) and Sjögren syndrome (SS) and can cause painful erosions, blisters and ulceration at the oral epithelium. Such diseases complicate dental hygiene and can lead to tooth loss. When natural teeth are missing, dental implants can improve quality of life for these patients. Osseointegration is necessary for dental implant success and is not contraindicated for patients with oral mucosal autoimmune diseases. However, the clinical dentist must consider the possible impact of oral autoimmune diseases on the oral epithelium that might affect implant success, also due to the difficulty of everyday oral hygiene, leading to bone absorption around the implant. Moreover, medication approved for the therapy of such diseases, such as corticosteroids, immunosuppressants and non-steroidal anti-inflammatory drugs (NSAIDs) could provoke osseointegration, as it compromises bone quality and affects the patient’s general health. However, the impact of these drugs on implant surgery depends on dose and duration of the drug and usually dental implantation is possible.

Keywords: Oral autoimmune diseases, Dental implants, Lichen planus, Pemphigus, Peri-implantitis

Introduction

An autoimmune disease is a condition in which the immune system reacts to antigens by producing autoantibodies or by activating cells such as lymphocytes, thereby producing a state of inflammation. Autoimmune diseases that affect the oral tissues are notably oral lichen planus (OLP), pemphigus vulgaris (PV), mucous membrane (MMP) and bullous pemphigoid (BP), epidermolysis bullosa acquisita (EBA) and systemic lupus erythematosus (SLE), which are characterized mainly by oral manifestations, such as erosions, blisters, papules and bullae and Sjögren syndrome (SS), which causes severe xerostomia. The lesions caused by these diseases are significantly painful and can be provoked by minimum pressure, as the patient talks, chews or performs dental hygiene. As a result, those patients avoid proper teeth brushing, in order to protect their wounded oral mucous membrane, or to avoid causing new lesions. Consequently, patients with autoimmune diseases that affect the oral cavity can easily lose their teeth by dental caries or periodontitis.

In the past, when the remaining teeth could not support a fixed prosthesis, the occlusion restoration could only be accomplished by removable partial or total dentures. However, the denture use is commonly problematic for patients with oral autoimmune diseases, as erosions, ulcerations, bulla formation and blisters cause additional mucosal discomfort and pain, due to the tissue pressure from the dentures. Therefore, in order to improve quality of life for these patients, fixed oral restorations have to be used, which leads to the necessity of dental implant rehabilitation. Dental implants can support fixed prosthesis or removable dentures, which are particularly stable, in contrast to simple dentures. However, autoimmune diseases with oral manifestations, as well as the medication intended for these patients, can affect the oral epithelium, as well as the bone quality, which are both important factors of implant osseointegration and long term success.

The author has no conflict of interest.

Corresponding author: Theodora S. Tounta, DDS, 29 K. Varnali Street, N. Erythrea, 14671, Athens, Greece

E-mail: theodoratounta@gmail.com

Edited by: Konstantinos Stathopoulos

Accepted 15 November 2018
Impact on periodontal and peri-implant tissues

In order to be successful, implant placement must be followed by osseointegration. Osseointegration is described by Branemark as the state when the implant and the alveolar bone are in contact, without the involvement of fibrous connective tissue. Osseointegration is affected, amongst other aspects, by the quality of the bone, as well as the atraumatic healing of the surgery site. Normally, the oral mucosa acts as a barrier and protects the alveolar bone from the oral cavity’s bacteria. However, if the oral mucosa is problematic, as when the patient suffers from autoimmune diseases with oral manifestations, the sealing may be disturbed and there is a higher possibility of tissue infection, which can undermine the long-term implant survival. Successful mucosal protection is achieved by the unhampered connection between the epithelial cells, as well as between the basement membrane and the connective tissue. The connection is accomplished by desmosomes in the former and hemidesmosomes in the latter.

However, oral autoimmune diseases cause alterations in the oral epithelium and connective tissue, which may also disturb the connection between the mucosa and the dental implant surface and lead to infection and implant loss.

Oral lichen planus

Oral lichen planus is a chronic autoimmune disease that affects the oral mucosa, the skin, the genitals, the nails and the scalp. The prevalence ranges between 0.5-2.6%, in several countries worldwide. White striations, blisters, erosions and atrophy are reported as clinical OLP features. Figures 1 and 2 show lichenoid lesions on the buccal mucosa.

It has been suggested that osseointegration may be problematic in patients with oral lichen planus. Histologically, OLP causes degradation of the basic epithelial cells, as well as alternations of the basement membrane. As a result, the epithelium-connective tissue connection is affected, which leads to the creation of an empty space that can easily be infected. Moreover, T-cells are increased in number and excrete cytokines, which leads to further epithelial cell apoptosis. Therefore, it can be assumed that the implant survival is at stake if the implant is placed on damaged mucosal membrane. Moreover, Sugerman and Savage report increased levels of TNF-α in patients with oral lichen planus. However, TNF-α is a major provoking periodontitis and peri-implantitis factor. Consequently, it is possible that peri-implant inflammation is more acute at patients with OLP compared to healthy subjects and could lead to quicker bone deformation and thereby also implant loss.

However, the majority of clinical studies referring to implant placement in OLP patients demonstrates promising results. Czerninski et al studied 29 OLP patients who had received 54 dental implants. The authors suggested that there is no correlation between lichen planus and implant success. Similar findings are reported by Hernández et al. López-Jornet et al studied 32 OLP patients compared to 16 healthy people. The study involved a total amount of 106 dental implants. The writers suggest that there is no significant relationship between lichen planus and peri-implantitis. Reichart’s study referred to 3 OLP patients treated with 10 dental implants and suggests that dental
Dental implants in patients with oral autoimmune diseases

Implant rehabilitation is safe for lichen planus patients. Finally, Esposito et al placed 4 dental implants in 2 OLP patients with complete dentures and report better ability to masticate and fewer OLP flare-ups. All findings are additionally described in Table 1. A possible explanation of the high implant success rate may be that in most studies the autoimmune disease was under recession and the oral hygiene was excellent.

**Pemphigus vulgaris**

Pemphigus vulgaris describes a range of epithelial bullous diseases, in which IgG auto-antibodies are produced, provoking akantolysis. Pemphigus vulgaris affects skin and mucous membranes and appears in 0.1-0.5/100000 persons per year. In the early stages, patients may only present oral alternations. More specifically, intra-epithelial bullae appear, which rapidly grow in size, burst and leave painful erosions. Dental hygiene is particularly inflicted and the use of dentures causes constantly new bullae, because of the rubbing.

Patients with PV make auto-antibodies against the desmosomes that connect the epithelial cells. The epithelium rupture enables the bacterial infection, as the protective barrier no longer exists. The infection can then easily advance to the inner tissues and lead to bone loss and implant failure. Literature data referring to implant placement in patients suffering from PV, to the best of our knowledge, are scarce. Altin et al placed dental implants in a pemphigus patient in order to support overall denture. 32 months after, the results were still satisfying and the pemphigus symptoms were less intense after the implant rehabilitation. However, the surgery was performed only after the disease’s control with corticosteroid use for several months.

**Mucous membrane and bullous pemphigoid**

Mucous membrane and bullous pemphigoid are chronic autoimmune diseases that are characterized by the formation of sub-epithelial bullae, which burst and leave erosions and scars when healed. The prevalence is reported 1.3-2 and 13.4-21.7/100000 persons per year respectively. In both MMP and BP, auto-antibodies against hemidesmosomes - which are necessary for the bond formation between epithelial cells and the implant surface - are produced. As a result, hemidesmosome malfunction may prevent the formation of a protective epithelial barrier for the implant.

Moreover, in bullous pemphigoid patients’ serum, as well as the bullae liquid, an increase in cytokines, chemokines, interleukins and TNF-α has been found. These factors provoke bone absorption in active periodontitis and may also affect the bone tissue around the implants. References in the bibliography concerning the success of implantation in such patients are rare.

**Epidermolysis bullosa acquisita**

Patients who suffer from epidermolysis bullosa also present sub-epithelial blisters, as EBA leads to the formation of auto-antibodies against collagen type VII. Epidermolysis bullosa is a rare condition, with prevalence 0.2/100000. Usually, the bullae gradually decrease and leave erosions and scars, which may lead to microstomia. Collagen type VII is a major contributor to mucous membrane stability and well-functioning. As a result, collagen type VII destruction by the auto-antibodies may lead to problematic mucous membrane creation, which may enable infection by pathogens and prevent osseointegration.

In contrast to the above, literature findings suggest that dental implantation is successful in such patients. Penarrocha et al placed 27 dental implants in 3 EBA patients, with 97.9% success. The writers suggest that dental implants are not contraindicated for patients suffering from epidermolysis bullosa, on the contrary implants can improve their quality of life. Letelier et al placed 11 dental implants in one EBA patient successfully and report better results than removable dentures. Penarrocha-Diago et al placed a total of 15 implants in

| Authors                  | Population                                                                 | Number of implants | Results                                |
|--------------------------|-----------------------------------------------------------------------------|--------------------|----------------------------------------|
| Czerninski et al.         | 14 OLP patients with implants and 15 OLP patients without implants            | 54                 | No correlation between OLP and implant success |
| Hernández et al.          | 18 OLP patients and 18 healthy people                                        | 56                 | No correlation between OLP and implant success |
| López-Jornet et al.       | 16 OLP patients with implants, 16 OLP patients without implants, 16 healthy people with implants | 56-50              | No correlation between OLP and peri-implantitis |
| Reichart                  | 3 OLP patients                                                              | 10                 | OLP patients can be treated with implants |
| Esposito et al.           | 2 OLP patients with complete dentures                                       | 4                  | Better ability to masticate and fewer OLP flare-ups |

Table 1. Relationship between oral lichen planus and dental implants.
4 patients and also suggest that dental implants can be a successful solution32. Penarrocha-Oltra et al placed 23 dental implants simultaneously with bone graft in 4 EBA patients and the results were successful33. Similarly, Larrazabal-Moron et al claim that simultaneous bone graft and implant placement is possible for EBA patients34. Table 2 summarizes findings from these studies.

**Sjögren syndrome**

Sjögren syndrome is a systemic disease that usually affects middle aged women. The prevalence of SS ranges between 0.5-1% of the population. Sjögren syndrome is characterized by mouth and eye dryness, caused by the presence of an inflammatory infiltrate of lymphocytes interfering with the function of the exocrine gland tissues3,35. Oral dryness causes numerous problems and leads to tooth loss, as the saliva protects from dental caries and periodontal and peri-implant diseases. The oral epithelium is characterized by painful erosions and sometimes colonization by Candida albicans36.

Numerous researchers refer to the use of dental implants in patients with Sjögren syndrome, as shown in Table 3. Korfage et al studied 140 cases of dental implants placed in 50 SS patients attending the University Medical Center Groningen. The writers concluded that the periodontal health indices reported were affected by the oral autoimmune disease, but the implant success rate was comparable to that concerning healthy people37. Binon placed 6 dental implants in one patient suffering from Sjögren syndrome and report successful rehabilitation at a follow up of 13 years38. Isidor et al placed a total of 90 implants in 8 women who suffered from Sjögren syndrome. All patients but one reported improvement in function and comfort 2 years after the surgery39. Similar results were reported by de Mendonça Invernici et al, who placed 3 dental implants in a Sjögren female patient that ameliorated the patient’s quality of life3. Finally, Chatzistavrianou and Shahdad report successful implant placement in two patients with SS after 18 months and two years follow-up respectively40.

| Authors                        | Population                                                                 | Number of implants | Results                                                                 |
|--------------------------------|----------------------------------------------------------------------------|--------------------|------------------------------------------------------------------------|
| Peñarrocha et al30             | 3 EBA patients, edentulous at the maxilla and/or the mandible              | 27                 | Dental implants are not contraindicated and improve quality of life    |
| Letelier et al31               | 1 31-year-old EBA patient with severe symptoms                              | 11                 | Dental implants are better than conservative removable dentures        |
| Peñarrocha-Oltra et al33       | 4 EBA patients                                                              | 23                 | Dental implants can be placed simultaneously with bone grafts in patients with EBA |
| Larrazabal-Moron et al34       | 1 52-year-old female EBA patient                                            | 2                  | Dental implants can be placed simultaneously with bone grafts in patients with EBA |
| Peñarrocha-Diago et al32       | 4 EBA patients, 1 man and 3 women, ages 26-35 years                        | 15                 | Dental implants can be successfully placed in patients with EBA        |

Table 2. Relationship between epidermolysis bullosa and dental implants.

| Authors                      | Population                      | Number of implants | Results                                                                 |
|------------------------------|---------------------------------|--------------------|------------------------------------------------------------------------|
| Binon38                      | 67-year-old male SS patient     | 6                  | Implant success after 13 years                                         |
| Isidor et al39               | 8 edentulous female SS patients | 90                 | Functional and aesthetic amelioration in 87.5% of patients             |
| Invernici et al3              | 58-year-old SS patient          | 3                  | Implant success 3 years after                                          |
| Chatzistavrianou and Shahdad40| Two female SS patients          | Not available      | Implant success after 18 months and 2 years                           |
| Korfage et al37              | 50 SS patients with dental implants | 140              | Impact on periodontal health but not on implant placement             |

Table 3. Relationship between Sjögren syndrome and dental implants.
Systemic lupus erythematosus

Systemic lupus erythematosus is an autoimmune disease with a wide range of symptoms, as most organs are affected. The prevalence ranges between 1.2-50/100000 worldwide41. The oral mucosa is affected in 40% of patients. Often, desquamative gingivitis, erosions and serious infections are present, as well as inflammation of the temporomandibular joint, that may complicate the opening of the mouth and pose difficulty in the implant surgery. Auto-antibodies against the nucleus and the cytoplasm antigens are created41. The immunological pathway of SLE resembles the one that develops during periodontitis and peri-implantitis42. Therefore, it is possible that SLE aggravates the periodontal and peri-implants diseases and may result in a higher possibility of implant loss compared to healthy people43.

Oral autoimmune diseases and peri-implantitis

As mentioned above, oral lesions caused by oral autoimmune diseases impair proper oral hygiene as it is painful and may provoke new manifestations. In addition, patients with oral autoimmune diseases often avoid dental appointments, as they fear that their symptoms may aggravate. As a result, it is possible for them to suffer from gingivitis, periodontitis and peri-implantitis due to the plaque accumulation.

Peri-implantitis is an inflammatory disease that affects the tissues around the dental implant and leads to bone absorption and eventually implant loss45. In order to prevent the severe peri-implantitis results, the dental clinician should pay close attention to patients with oral autoimmune diseases and implants. If peri-implantitis is identified at an early stage, it is easier to treat with the use of chlorhexidine, antibiotics and plaque removal46.

In the bibliography, many studies support the correlation between oral autoimmune and periodontal diseases47. Ertugrul et al showed that LP patients present a higher percentage of pathological bacteria in their saliva compared to healthy subjects, which may lead to more severe symptoms and quicker deterioration of periodontal and peri-implant disease48. In accordance with the above, the Wang et al survey suggests correlation between the periodontal diseases and OLP49. In addition, Thothar et al suggested that pemphigus vulgaris patients had higher plaque accumulation and pore pockets compared to healthy people. In fact, the pockets were deeper when the PV was more aggressive50. Tricamo et al found that patients with mucous membrane pemphigoid had statistically significant augmentation of gingival inflammation compared to the control group51.

Similarly, it seems that a correlation between systemic lupus erythematosus and periodontal diseases exists52,53.

Oral autoimmune diseases and soft tissue regeneration

In some cases of severe bone deficiency, in order to insert dental implants, it is necessary to perform periodontal plastic surgery and soft tissue regeneration, with the use of grafts and membranes53. However, oral autoimmune diseases may complicate such surgeries, as these specific disorders affect soft tissue healing. An additional problem for patients with oral autoimmune diseases is that the mucosa is thin and fragile and makes the surgery harder54.

Available data in the published literature are scarce. Katz et al claim lichen planus appearance after periodontal surgery and suggest avoiding traumatic surgery in patients with OLP55. In contrast, Toscano et al suggest that periodontal surgery may be performed after the proper pharmacological treatment of the oral autoimmune disease56. Furthermore, corticosteroid use, which is a common autoimmune disease treatment, impedes healing. However, Shin-Yu Lu supports that successful soft tissue regeneration with grafts and resorbable membrane is possible despite systemic corticosteroid use57.

Oral autoimmune diseases pharmacological therapy and implant rehabilitation

Systemic autoimmune diseases with oral involvement cause severe symptoms that range from discomfort and pain to organ failure and eventually death and require prompt action. Treatment is often challenging and medication may have severe side-effects and long duration. These drugs can also affect the implant osseointegration and long-term survival.

Corticosteroids

The most common therapeutic agent for the treatment of autoimmune diseases with oral manifestations is glucocorticosteroids, either topically or systematically5. In mild cases oral lesions can be treated with topical ointments9, whereas patients with more severe symptoms may need systematic administration with doses up to 0.5 mg/kg per day for oral lichen planus58 or even greater for pemphigus vulgaris (0.75-1.25 mg/kg/day)59. However, the use of corticosteroids in such doses affects bone quality and can result in glucocorticoid-induced osteoporosis60. More specifically, bone formation is diminished as osteoblasts are reduced in number and are less active and bone absorption is increased as osteocytes lead to apoptosis and osteoclasts are overactive61,62. Glucocorticoids also affect calcium metabolism, thereby interfering to mineralization of new bone63.

However, when bone healing and remodeling is affected, implant placement may be difficult for those patients. The
findings in the literature are controversial. Petsinis et al suggest that glucocorticosteroid intake for systemic diseases does not have a significant impact on the osseointegration and should not be considered as a contraindication for dental implant surgery. In accordance, Lu and Huang (2007) presented a successful implant placement in a chronic corticosteroid user and proposed antibiotic prophylaxis, strict oral hygiene and frequent recall appointments as special caution measures. However, in Keller’s survey (2004) osseointegration was affected in rabbits who received corticosteroids and it was suggested that this might jeopardize long-term implant stability. Similarly, Fujimoto et al worked with rabbits and showed that corticosteroids affect implant placement, but may have less impact on the osseointegration at the mandibular bone compared to the skeletal bone.

There is good evidence that glucocorticoids affect bone quality and can cause osteoporosis and implant failure. However, patients with autoimmune oral mucosa diseases differ amongst each other in medication agents, dose and duration, even when suffering from the same disease, so that care should be taken in an individual basis.

**Immunosuppressive drugs**

Very often immunosuppressants such as cyclosporine, azathioprine, mycophenolate and methotrexate are used for the treatment of autoimmune diseases with oral manifestations, especially when remission occurs. However, to the best of our knowledge there are no published data concerning the impact of immunosuppressive drug agents, used for autoimmune disease treatment, on dental implant surgery in humans. The only studies found referred to animals and suggest controversial results.

Therefore, there are no clear instructions on oral surgery for patients under such immunosuppressive agents. The clinician must bear in mind that soft tissue and bone healing are affected to some extent and the patient is more prone to infections.

**Non-steroidal anti-inflammatory drugs**

Patients who suffer from lupus erythematosus are sometimes recommended to use non-steroidal anti-inflammatory drugs (NSAIDs), in order to control inflammatory arthralgia and myalgia, as well as fever. However, NSAIDs and especially COX-2 inhibitors are thought to impair osseointegration, as they delay bone healing, particularly when NSAIDs are used in large doses and for long periods. This can be explained by the fact that non-steroidal anti-inflammatory drugs inhibit the COX-2 activity, which results in reduced prostaglandins and regulation of the inflammation. However, prostaglandins and especially E2 affect bone metabolism and may lead to problematic osseointegration. With regard to the above, caution should be taken when deciding if implant rehabilitation is suitable for patients with lupus erythematosus who use non-steroidal anti-inflammatory drugs for long periods of time.

**Conclusion**

To sum up, the use of dental implants in patients with autoimmune diseases with oral manifestations is still currently under investigation. It is clear that the severity of the disease is the major factor that the dental clinician will take into consideration in order to decide the implant solution. Patients with severe symptoms are rather unlikely to refer to the dentist for dental implant rehabilitation, as it is not their priority. In mild cases, where the drug treatment may be exclusively local, or for short periods and in low dose, the implant success rates are similar to these of healthy people. On the contrary, patients with average disease severity should be treated on an individual basis. Implant placement should always be attempted after flare ups of the disease have been managed. The implant placement spot on the mucosa should be lesion free. The drug duration and dose is of paramount importance for the implant survival, as augmentation in the drug dosage impairs the bone and soft tissue quality.

Finally, dental clinicians should place emphasis on oral hygiene, which is often neglected by patients with oral autoimmune diseases. Successful oral hygiene ameliorates the soft tissue lesions, reduces the infection possibility and protects from periodontal and peri-implant diseases, that could lead to implant loss. Teeth brushing should be performed by applying low pressure and using a soft toothbrush and mild antimicrobial mouthwashes are recommended.

In the future, more studies concerning dental implant placement in patients with oral autoimmune diseases are needed, as well as studies discussing the potential and limitations of regenerative periodontal tissue techniques in these patients.

**Acknowledgements**

I wish to thank Dr Kartanos Euaggelos and Dr Foteinopoulou Elpida for their valuable donation of photographic material from their personal archive. I would also like to thank Mrs Katri Lois for proofreading the article.

**References**

1. Zeher M, Szegedi G. Types of autoimmune disorders. Classification. Orv Hetil 2007;8;148(1):21-4.
2. Schürer M, Yeoh SC, Coleman H, Georgiou A. Oral mucosal diseases: the inflammatory dermatoses. Aust Dent J 2010;55(Suppl 1):23-38.
3. de Mendonça Invernici M, Finger Stadler A, Vale Nicolau G, Naval Machado MÁ, Soares de Lima AA, Compagnoni Martins M. Management of Sjögren’s Syndrome Patient: A Case Report of Prosthetic Rehabilitation with 6-Year Follow-Up. Case Rep Dent 2014;2014:761251.
4. Mustafa MB, Porter SR, Smoller BR, Sitaru C. Oral mucosal manifestations of autoimmune skin diseases. Autoimmun Rev
Dental implants in patients with oral autoimmune diseases

2015;14(10):930-51.

5. Bränemark P. Osseointegration and its experimental background. J Prosthet Dent 1983;50(3):399-410.

6. Atsuta I, Ayukawa Y, Kondo R, Oshiro W, Matsuura Y, Furushashi A, Tsukiyama Y, Koyano K. Soft tissue sealing around dental implants based on histological interpretation. J Prosthodont Res 2016;60(1):3-11.

7. Stevens A, Lowe JS. Iatrologia του ανθρώπου. Τρίτη έκδοση; εκδόσεις Π.Χ. ΠΑΝΑΧΩΛΗ.

8. Steinbacher M, Zorn C, Steinhöfel K, Götzfried S, Auinger K, Buser D. Implant-supported fixed complete denture in a patient with Sjögren’s syndrome. J Oral Implantol 2016;42(6):498-505.

9. Penarrocha-Diago M, Penarrocha-Diago A, Penarrocha-Diago M. Oral rehabilitation with implant-supported fixed complete denture in a patient with Sjögren’s syndrome: a clinical report. J Oral Maxillofac Surg 2009;67(7):1499-502.

10. Sugerman PB, Savage NW. Oral lichen planus: causes, diagnosis and management. Aust Dent J 2002;47(4):290-9.

11. Aboushelib MN, Elsafi MH. Clinical Management Protocol for Dental implants in patients with oral autoimmune diseases JRPMS 2013;5(15):7.

12. Ding C, Ji X, Chen X, Xu Y, Zhong L. TNF-α gene promoter polymorphisms contribute to periodontitis susceptibility: evidence from 46 studies. J Clin Periodontol 2011;44(8):748-59.

13. Czeminski R, Eliezer M, Wilensky A, Sokolsky A. Oral lichen planus and dental implants-a retrospective study. Clin Implant Dent Relat Res 2013;15(2):234-42.

14. Hernández G, Lopez-Pintor RM, Arriba L, Torres J, de Vicente JC. Immunohistological study of the human dermis and hypodermis. Dermatol Venereol 2013;27(424-27).

15. Heath W, Pringle D, McClelland A, Heseltine J, Daffern M. Clinical practice guidelines for the management of cutaneous lupus erythematosus. J Dtsch Dermatol Ges 2009;7:434-40.

16. Reichart PA. Oral lichen planus and dental implants. Report of 3 cases. J Oral Maxillofac Surg 2003;61(1):6-10.

17. Esposito SJ, Camisa C, Morgan M. Implant retained overdentures for patients with severe lichen planus: a clinical report. J Prosthet Dent 2003;89(1):6-10.

18. Kasperkiewicz M, Ellbrecht CT, Takahashi H, Yamagami J, Zillikens D, Payne AS, Aroagi M. Pemphigus. Nat Rev Dis Primers 2017;13(10):273-85.

19. Altin N, Ergun S, Kat J, Sancakli E, Koray M, Tanyeri H. Implant supported oral rehabilitation of a patient with pemphigus vulgaris: a clinical case report. J Prosthodont Res 2013;57(4):1-6.

20. Xu HH, Werth VP, Parisi E, Sollecito TP. Mucous membrane pemphigoid. Dent Clin North Am 2013;57(4):611-30.

21. Schmidt E, Zillikens D. Pemphigoid diseases. Lancet 2013; 381(9863):320-32.

22. Ahmed AR, Kurgis BS, Rogers RS. Cicatricial pemphigoid. J Am Acad Dermatol 1991;24:987-1001.

23. Bertram F, Brocker EB, Zillikens D, Schmidt E. Prospective analysis of the incidence of autoimmune bullous disorders in Lower Franconia, Germany. J Dtsch Dermatol Ges 2009;7:434-40.

24. Gudi VS, White MI, Cruickshank N, Herriot R, Edwards SL, Nimmo F, Ormerod AD. Annual incidence and mortality of bullous pemphigoid in the Grampian region of north-east Scotland. Br J Dermatol 2005;153(424-27).

25. Joly P, Barcault S, Sparsa A, Bernard P, Bédane C, Duvert-Lehembre S, Courville P, Bravard P, Rémond B, Dofoel-Hantz V, Bénichou J. Incidence and mortality of bullous pemphigoid in France. J Invest Dermatol 2012;132:1998-2004.

26. Yucel-Linberg T, Bägle T. Inflammatory mediators in the pathogenesis of periodontitis. Expert Rev Mol Med 2013;15(15):7.

27. Kim JH, Kim SC. Epidermolysis bullosa acquisita. J Eur Acad Dermatol Venereol 2013;27(10):1204-13.

28. Anerio G, Perez-González J, Páramo-Fernández D, Catalán A, Sturrock RD. Autoimmune bullous dermatoses: the biology of the autoimmune bullous pemphigoid. Eur Dermatol 2011;20(7):482-9.

29. Csorba K, Chiriac MT, Florea F, Ohnajzerova B, Saidov A, Afsa A, Vuta V, Botana A. Blister-inducing antibodies target multiple epitopes on collagen VII in mice. J Cell Mol Med 2014 Sep 18;17(1):111-27.

30. Penarrocha M, Rambía J, Balaqu J, Serrano C, Silvestre J, Baigán JV. Placement of implants in patients with pemphigoid bullosa. J Oral Maxillofac Surg 2007;65(7 Suppl 1):103-6.

31. Leteiler MG, Jara CC, Peñarrocha-Oltra S, Gomar-Vecher S, Diogo MP. Fixed Implant-Supported Full-Arch Prosthesis in Epidermolysis Bullosa With Severe Symptoms. J Oral Implantol 2016;42(6):498-505.

32. Peñarrocha-Diago M, Serrano C, Sanz JM, Silvestre FJ, Baigán JV. Placement of endosseous implants in patients with oral epidermolysis bullosa. Oral Surg Oral Med Oral Pathol Oral Radiol Endod 2000;90(5):587-90.

33. Peñarrocha-Diago A, Aloy-Prósper A, Ata-Ali J, Peñarrocha-Diago M, Peñarrocha-Diago M. Implants placed simultaneously with particulated bone graft in patients diagnosed with recessive dystrophic epidermolysis bullosa. J Oral Maxillofac Surg 2012;70(1):51-7.

34. Larrazabal-Morón C, Boronat-López A, Peñarrocha-Diago M, Peñarrocha-Diago M. Oral rehabilitation with bone graft and simultaneous dental implants in a patient with epidermolysis bullosa: a clinical case report. J Oral Maxillofac Surg 2009;67(7):1499-502.

35. Mavragani CP, Mouftopoulos GM. Sjögren’s syndrome. Ann Rev Dermatol 2014;9:273-85.

36. Alsakran Altamimi M. Update knowledge of dry mouth- A guideline for dentists. Afr Health Sci 2014;14(3):736-42.

37. Kortlage A, Rahrooer OM, Arends S, Meiners PM, Visser A, Kroese FGJ, Boestma H, Vissink A. Dental Implants in Patients with Sjögren’s Syndrome. Clin Implant Dent Relat Res 2016;18(S5):937-45.

38. Binon PP. Thirteen-year follow-up of a mandibular implant-supported fixed complete denture in a patient with Sjögren’s syndrome: a clinical report. J Prosthodont 2005;94(5):409-13.

39. Isidor F, Brandum K, Hansen HJ, Jensen S, Sindet-Pedersen S. Outcome of treatment with implant-retained dental prostheses in patients with Sjögren’s syndrome. Int J Oral Maxillofac Implants 1999;14(5):736-43.

40. Chatzistavrianou D, Shahdad S. Implant Treatment in Patients with Sjögren’s Syndrome. Clin Implant Dent Relat Res 2016;18(5):937-45.

41. Deloro G, Bootsma H, Vissink A. Dental Implants in Patients with Sjögren’s Syndrome: A Review of the Literature and Two Clinical Case Reports. Eur J Prosthodont Restor Dent 2016;24(1):40-6.

42. Alibila JB, Lam RK, Clowie CM, Sánchez GK. Systemic lupus erythematosus: a review for dentists. Afr Health Sci 2014;14(3):736-42.

43. Kunz M. Lupus erythematosus. Part I: epidemiology, genetics and immunology. J Dtsch Dermatol Ges 2013;11(8):709-19.

44. Sete MR, Figueredo CM, Stjahnke F. Periodontitis and systemic lupus erythematosus. Rev Bras Reumatol Engl Ed 2016;56(2):165-70.

45. Ergun S, Kat J, Ciffer ED, Koray M, Esen BA, Tanyeri H. Implant supported oral rehabilitation of a patient with systemic lupus erythematosus: case report and review of the literature. Quintessence Int 2010;41(10):863-7.

46. Koyanagi T, Sakamoto M, Takeuchi Y, Maruyama N, Ohkuma M, Izumi M, Peñarrocha-Diago M, Peñarrocha-Diago M. Oral rehabilitation with bone graft and simultaneous dental implants in a patient with epidermolysis bullosa: a case report. J Oral Maxillofac Surg 2009;67(7):1499-502.
Azizi A, Rezaee M. Comparison of periodontal status in gingival oral lichen planus patients and healthy subjects. Dermatol Res Pract 2012;2012:561232.

Ertugrul AS, Arslan U, Dursun R, Hakki SS. Periodontopathogen profile of healthy and oral lichen planus patients with gingivitis or periodontitis. Int J Oral Sci 2013;5(2):92-7.

Wang H, Luo Z, Lei L, Sun Z, Zhou M, Dan H, Zeng X, Chen Q. Interaction between oral lichen planus and chronic periodontitis with Th17-associated cytokines in serum. Inflammation 2013;36(3):696-704.

Thorat MS, Raju A, Pradeep AR. Pemphigus vulgaris: effects on periodontal health. Int J Oral Sci 2010;5(3):449-54.

Tincamo MB, Rees TD, Hallmon WW, Wright JM, Cueva MA, Plemons JM. Periodontal status in patients with gingival mucous membrane pemphigoid. J Periodontol 2006;77(3):398-405.

Zhang, Zhang X, Feng G, Fu T, Yin R, Zhang L, Feng X, Li L, Gu Z. Periodontal disease in Chinese patients with systemic lupus erythematosus. Rheumatol Int 2017;37(8):1373-9.

Resnik RR. Surgical Techniques to Increase Bone Augmentation Success. Dent Today 2015;34(11):114-9.

Lorenzana ER, Rees TD, Hallmon WW. Esthetic management of multiple recession defects in a patient with cicatricial pemphigoid. J Periodontal 2001;72(2):230-7.

Katz J, Goultshcn J, Benoliel R, Rotstein I, Psanty S. Lichen planus evoked by periodontal surgery. J Clin Periodontol 1988;15(4):263-5.

Toscano NJ, Holtzclaw DJ, Shumaker ND, Stokes SM, Meehan SC, Rees TD. Surgical considerations and management of patients with mucocutaneous disorders. Compend Contin Educ Dent 2010;31(5):344-50, 52-9.

Lu SY, Huang CC. Resolution of an active peri-implantitis in a chronic steroid user by bone augmentation with PepGen P-15 and a barrier membrane. J Oral Implantol 2007;33(5):280-7.

Manousakis I, Manousakis K, Petsch WK, Schneider SW. Individualizing treatment and choice of medication in lichen planus: a step by step approach. J Dtsch Dermatol Ges 2013;11(10):981-91.

Tsuruta D, Ishii N, Hashimoto T. Diagnosis and treatment of pemphigus. Immunotherapy 2012;4(7):35-45.

Cho P, Schneider GB, Kellogg B, Zanhanis R, Keller JC. Effect of glucocorticoid-induced osteoporotic-like conditions on osteoblast cell attachment to implant surface microtopographies. Implant Dent 2006;15(4):377-85.

Whittier X, Saag KG. Glucocorticoid-induced Osteoporosis. Rheum Dis Clin North Am 2016;42(1):177-89.

Defranco DJ, Lian JB, Gowacki J. Differential effects of glucocorticoid on recruitment and activity of osteoclasts induced by normal and osteocalcin-deficient bone implanted in rats. Endocrinology 1992;131(1):114-21.

Canalis E, Mazziotti G, Giustina A, Bilezikian JP. Glucocorticoid-induced osteoporosis pathophysiology and therapy. Osteoporos Int 2007;18(10):1319-28.

Petsinis V, Kamarogou C, Alexandridi F, Alexandridis K. The impact of glucocorticosteroids administered for systemic diseases on the osseointegration and survival of dental implants placed without bone grafting: A retrospective study in 31 patients. J Craniofac Surg 2017;45(8):1197-200.

Keller JC, Stewart M, Roehm M, Schneider GB. Osteoporosis-like bone conditions affects osseointegration of implants. Int J Oral Maxillofac Implants 2004;19(5):687-94.

Fujimoto T, Nimm A, Sawai T, Ueda M. Effects of steroid-induced osteoporosis on osseointegration of titanium implants. Int J Oral Maxillofac Implants 1998;13(2):183-9.

Tunicliffe DJ, Singh-Grewal D, Kim S, Craig JC, Tong A. Diagnosis, Monitoring, and Treatment of Systemic Lupus Erythematosus: A Systematic Review of Clinical Practice Guidelines. Arthritis Care Res (Hoboken) 2015;67(10):1440-52.

Sakakura KE, Maromar N, Holzhausen M, Nociti FH Jr, Alba RC Jr, Marcantonio EJ Jr. Influence of cyclosporin A therapy on bone healing around titanium implants: a histometric and biomechanic study in rabbits. J Periodontol 2003;74(7):976-81.

Carvas JB, Perera RM, Bonfá E, Silveira CA, Lima LP, Caparbo Vde F, Mello SB. No deleterious effect of low dose methotrexate on titanium implant osseointegration in a rabbit model. Clinics (Sao Paulo) 2011;66(6):1055-9.

Gordon C, Amissah-Arthur MB, Gayed M, Brown S, Bruce IN, D'Cruz D, Empson B, Griffiths B, Jayne D, Khamesha M, Lightstone L, Norton P, Norton Y, Schreiber K, Isenberg D, The British Society for Rheumatology guideline for the management of systemic lupus erythematosus in adults. Rheumatology (Oxford) 2018;57(1):1-45.

Gomes Fi, Aração MG, Pinto VD, Condram DV, Barbosca FC, Silva AA, Bezerra MM, Chaves HV. Effects of nonsteroidal antiinflammatory drugs on osseointegration: A review. J Oral Implantol 2015;41(2):219-30.

Bourinos LA, Karachalios T, Poultsides L, Malizos KN. Do steroids, conventional non-steroidal anti-inflammatory drugs and selective Cox-2 inhibitors adversely affect fracture healing? J Musculoskeletal Neuronal Interact 2009;9(1):44-52.

Thomas MV, Puleo DA. Infection, inflammation, and bone regeneration: a paradoxical relationship. J Dent Res 2011;90(9):1052-61.

Kalogrou EM, Sklavounou A. Is Dental Implantation Indicated in Patients with Oral Mucosal Diseases? Balk J Dent Med 2017;21:83-92.

Reichart PA, Schmidt-Westhausen AM, Khongkhunthian P, Strietzel FP. Dental implants in patients with oral mucosal diseases - a systematic review. J Oral Rehabil 2016;43(5):388-99.