Therapeutic alternative for oral lichen planus treatment in an unusual location: case report

Thaís Torres Barros DUTRA1
Thâmara Manoela Marinho BEZERRA2
Filipe Nobre CHAVES3
Sthefane Gomes FEITOSA4
Fábio Willson Gurgel COSTA4
Karuza Maria Alves PEREIRA5

1Doutoranda, Departamento de Patologia Oral, Faculdade de Farmácia, Odontologia e Efermagem (FFOE), Universidade Federal do Ceará (UFC), 60430-355 Fortaleza - CE, Brazil
2Pós-Doutoranda, Departamento de Patologia Oral, Faculdade de Farmácia, Odontologia e Efermagem (FFOE), Universidade Federal do Ceará (UFC), 60430-355 Fortaleza - CE, Brazil
3Professor Adjunto, Mestrado em Ciências da Saúde, Curso de Odontologia, Universidade Federal do Ceará, Campus Sobral, 62010-820 Sobral - CE, Brazil
4Professor Adjunto, Departamento de Radiologia Odontológica, Faculdade de Farmácia, Odontologia e Efermagem (FFOE), Universidade Federal do Ceará (UFC), 60430-355 Fortaleza - CE, Brazil
5Professora Associada, Departamento de Morfologia, Faculdade de Medicina (FAMED), Universidade Federal do Ceará (UFC), 60430-160 Fortaleza - CE, Brazil

Abstract

Oral Lichen Planus (OLP) is considered a potentially malignant lesion (PML), although its rate of transformation is controversial. New treatments have been introduced recently, for example calcipotriene (D3 vitamin analog). Its incremental action with glucocorticoids is observed and isolated lip lesions may respond positively to combined topical therapy. Thus, objective of this case report is to show a therapeutic alternative for isolated and persistent OLP lip lesions. An 18-year-old teenager was referred to evaluate an erythematosus lower lip lesion with approximately 10 years of evolution. The inspection revealed multiple erythematous areas surrounded by thin whitish streaks located in the vermilion of the lower lip. No abnormalities in the skin, nails, scalp or other areas of oral mucosa were observed. The incisional biopsy was performed and the microscopic exam showed areas of basal layer degeneration with intense mononuclear inflammatory infiltrate banded and predominantly subepithelial. The therapeutic proposal was topical application of Daivobet®, a combination of topical corticosteroids and vitamin D derivative. The lesion had remission after the fifteen days, medication was suspended and indicated when there was a relapse. During follow-up, no recurrences or complications were observed. That combination therapy may be a new approach in treating OLP.

Descriptors: Lichen Planus; Lichen Planus, Oral; Lip; Pharmaceutical Preparations.

Resumo

O Líquen Plano Oral (OLP) é considerado uma lesão potencialmente maligna (LPM), embora sua taxa de transformação seja controversa. Novos tratamentos foram introduzidos recentemente, por exemplo, calcipotriene (análogo do vitamina D3). Lesões labiais isoladas podem responder positivamente a esta terapia tópica combinada, aumentando a efetividade do tratamento. Assim, o objetivo deste relato de caso é mostrar uma alternativa terapêutica para lesões labiais de LPO isoladas e persistentes. Um adolescente de 18 anos foi encaminhado para avaliar uma lesão eritematosa do lábio inferior com aproximadamente 10 anos de evolução. A inspeção revelou múltiplas áreas eritematosas cercadas por finas estrías esbranquiçadas localizadas no vermelho do lábio inferior. Não foram observadas anormalidades na pele, unhas, couro cabeludo ou outras áreas da mucosa oral. A biópsia incisional foi realizada e o exame microscópico mostrou áreas de degeneração da camada basal com intenso infiltrato mononuclear em faixas e predominantemente subepiteliais. A proposta terapêutica foi a aplicação tópica de Daivobet®, uma combinação de tópica de corteicoide e calcipotriene. A lesão teve remissão após os quinze dias, a medicação foi suspensa e indicada quando houvesse recidiva. Durante o acompanhamento, não foram observadas recorrências ou complicações. Essa terapia combinada pode ser uma nova abordagem no tratamento do LPO.

Descriptors: Líquen Plano; Líquen Plano Bucal; Lábio; Preparações Farmacêuticas.

INTRODUCTION

Lichen Planus (LP) is a chronic, mucocutaneous, immunologically mediated disease of uncertain etiology1-4 which mainly affects the skin, scalp, nails and mucosa in the form of itchy papules with fine white lines in the surface (Wickham striae)5-7. Its terminology was given by the British physician Erasmus Wilson in 1869, possibly due to the similarity of clinical lesions to lichens growing on rocks. In the oral cavity, LP is difficult to diagnose, and clinical
and histopathological data must be combined, as there are six clinical subtypes of the disease (reticular, plaque-like, atrophic, erosive / ulcerative, papular and bullous) that may manifest in a combined manner, with periods of relapses and remissions\(^4\). In addition, Oral Lichen Planus (OLP) is considered a potentially malignant lesion (PML), although its rate of transformation is controversial\(^6\). The risk of malignant transformation increases with the presence of erosive and / or atrophic areas, tobacco and / or alcohol consumption, hepatitis C virus infection and when located in the tongue\(^10\). Due to anatomic, physiologic and functional peculiarities of the oral cavity, OLP requires specific evaluations for diagnosis and management\(^4\).

Management of OLP involves the treatment of atrophic and erosive / ulcerative lesions to relieve the symptoms as well as to reduce the potential risk of malignant transformation\(^1,11\). However, the protocol for treating OLP is not well defined\(^11\). Currently, the use of topical agents is preferred due to the few adverse effects. Topical corticosteroid is the most widely used drug because it has fewer adverse effects and a good patient response rate (between 30% and 100% of cases)\(^12,13\). Other topical agents that can be alternatively used are retinoids, topical aloe vera, biologics, oral curcuminoind, calcineurin inhibitors (e.g., tacrolimus, cyclosporine) and low intensity laser\(^14-16\).

New OLP treatments have been introduced recently. Calcipotriene, the D3 vitamin analog, was introduced in the United States in 1994 for the treatment of psoriasis vulgaris\(^17\). Its anti-inflammatory effects are inferior to those provided by glucocorticoids, but an incremental action is observed when the two drugs are combined\(^18\) and the cutaneous LP disease course seems to be similarly affected by both treatments\(^18\).

As lip OLP has a clinical course similar to cutaneous LP, isolated lip lesions may respond positively to combined topical therapy of glucocorticoids and calcipotriol. Thus, the objective of this case report is to show a therapeutic alternative for isolated and persistent OLP lip lesions.

**CLINICAL CASE**

An 18-year-old teenager was referred to the Oral Medicine Clinic to evaluate an erythematous lower lip lesion with approximately 10 years of evolution. The extraororal (Figure 1A and B) inspection revealed multiple erythematous areas surrounded by thin whitish streaks located in the vermilion of the lower lip. No abnormalities in the skin, nails or scalp were observed. The intraoral inspection shows continuity of lesions located in the vermilion of the lip to the labial mucosa (Figure 1C and D), with no other areas of oral mucosa involved.

The patient reported a burning sensation and mild intermittent pain, worsening when hot or spicy foods were ingested. There were no reports of systemic diseases, general or drug allergies, or use of tobacco or alcohol. In addition, no similar lesions appeared in other regions of the body during the 10 years of lip lesions onset. Several drug therapies (topical corticosteroids, topical antiretrovirals, vitamins (folic acid and B complex) and antibiotics) were previously administered by other professionals, but without remission of the clinical picture. His medical, social and cultural histories were not contributory.

Considering the clinical hypothesis of LP, an incisional biopsy was performed, and the specimen was sent for histologic examination. The specimen was fixed in 10% neutral formalin buffer and embedded in paraffin. Macroscopic examination revealed 3 soft tissue fragments, predominantly white, of irregular shape and surface, with rubbery and fibrous consistency, the largest fragment measuring 0.7 x 0.5 x 0.4 cm. The microscopic exam showed areas of basal layer degeneration with intense mononuclear inflammatory infiltrate banded and predominantly subepithelial (Figure 2A and B). The final diagnosis was LP.

The therapeutic proposal was topical application of Daivobet®, whose formulation contains calcipotriol hydrate 50mcg / g and betamethasone dipropionate 0.5mg / g, a combination of topical corticosteroids and vitamin D derivative. The patient was followed fortnightly and the lesion was remitted after the first fifteen days (Figure 3). The use of medication was suspended and indicated when
there was a relapse. During follow-up, no recurrences or complications were observed (Figure 4).

DISCUSSION

Epidemiological studies show a low OLP prevalence in the adult population (estimated at 0.5% to 2%)\(^8,20\). It preferentially affects women aged 30-60 years and presents itself clinically isolated or associated with cutaneous manifestations\(^{14,21,22}\). This case report presents an OLP restricted to the lower lip of a young man. OLPs that strictly affect the lip have a preference for males and are rare\(^{23}\). Two studies detected OLP in lip alone, with a prevalence of 6.3 and 8.9%\(^{23}\). However, when OLP affects other regions of the oral cavity, the lip is commonly affected\(^{20,24,25}\). OLP in lip vermilion has a different clinical aspect from skin LP due to peculiarities of this region, such as: thick squamous epithelium, abundant capillary supply within interdigitating rete ridges and dermal papillae, besides absence of follicular and salivary structures\(^{26}\). However, the lesions in the present case showed a classic appearance of an erosive OLP (Figure 1). OLP lesions, especially in the lip, are insidious and easily overlooked because they are asymptomatic in most cases, except for erosive and scarring forms due to severe local pain, lip atrophy and microlabial. The patient in this study reported that the lesions were present for 10 years and that no drug therapy was effective until then to eliminate the lesions definitively. In this case, incisional biopsy was an important tool to guide clinical treatment, since with the histopathological diagnosis of OLP associated with the clinical history of multiple treatments without remission, we can better direct the patient's treatment.

OLP etiology is controversial, but there is strong evidence that it is an immunologically mediated disease with antigen-specific and non-antigen-specific mechanisms involved in its pathogenesis\(^{27}\). Within the specific mechanisms, antigen expression occurs by basal layer keratinocytes and their consequent death by CD8 cytotoxic T cells. Non-specific mechanisms include mast cell degranulation and metalloproteinase matrix (MPM) activation\(^{27}\). The combination of these mechanisms causes T cell accumulation in the underlying lamina propria with consequent disruption of the basement membrane, T lymphocyte migration towards the epithelium and keratinocyte apoptosis. This process is perpetuated through the release of chemokines in the inflammatory site\(^{27}\). Based on this, corticosteroids, especially glucocorticoids, are the first line of choice in the treatment of OLP.

Midpotency corticosteroids such as triamcinolone, potent fluorinated corticosteroids such as fluocinolone acetonide and fluocinonide as well as superpotent halogenated corticosteroids such as clobetasol have been successfully applied in clinical practice\(^{28,29}\). The main form of administration of this drug for treatment of OLP is topical, since it promotes less adverse effects. However, side effects such as secondary candidosis, nausea, non-tolerated oral use, refractory response, mucosal atrophy, oral dryness, sore throat, bad taste, and delayed

Figure 2: Photomicrographs: A. Oral mucosa lining epithelium showing areas of atrophy and acanthosis. A large banded inflammatory infiltrate is observed on the lamina propria underlying mucosal lining tissue (H / E 100x). B. In greater detail is observed in the epithelium loss of clarity of the basal layer and exocytosis. The underlying lamina propria shows presence of intense lymphocyte inflammatory infiltrate (H / E 200x).

Figure 3. Extraoral and intraoral aspect of the patient after 15 days of Daivobet therapy: A. Frontal view with no skin abnormality; B. Approximate view of the lower lip showing remission of erythematous lesions and whitish striae; C. Evaluation of lower lip vermilion with no other areas of involvement.

Figure 4: Extraoral and intraoral aspect of the patient during follow-up: A. Frontal view with no skin abnormality; B. Approximate view of the lower lip showing remission of erythematous lesions and whitish striae; C. Evaluation of lower lip vermilion with no other areas of involvement.
healing have been observed\textsuperscript{30,31}. Calciplotriol (Daivonex / Dovonex) is an indicator of vitamin D and has been used to treat psoriasis vulgaris (PV) in the United States since 1994\textsuperscript{31}. Vitamin D analogs and topical corticosteroids are likely to have different modes of action. The calcipotriol suppresses lymphocyte proliferation by decreasing interleukin-1 and interleukin-8\textsuperscript{17} as well as induces terminal differentiation and inhibits proliferation of keratinocytes\textsuperscript{92}. Their anti-inflammatory properties are known to be inferior to those of glucocorticoids, but an incremental effect has been observed when these drugs are used in combination to treat PV\textsuperscript{17,18}, as there is better efficacy, superior tolerability and faster clinical action compared to individual drug use\textsuperscript{17,33,34}. In this case, we believe that calcipotriol when combined with corticosteroids promoted its best efficacy, since the onset of clinical improvement occurred after 14 days of combined use of calcipotriol and betamethasone. It is important to emphasize that the patient did not show improvement in clinical condition with the use of corticosteroids alone.

The safety profiles of calcipotriene and betamethasone as monotherapies are well established\textsuperscript{35-37}. Adverse reactions to calcipotriene use are predominantly local and the most common are erythema or increased itchness\textsuperscript{38,39}. It is expected that the adverse effects will be similar to the adverse reaction profile of betamethasone, when using the drugs together\textsuperscript{17}. However, the patient in this study did not show any adverse effects during the combined therapy period, and there was a satisfactory and similar clinical response to PV patients. Alternative therapies for OLP treatment may be chosen through diseases that also have immune and inflammatory pathogenesis, such as PV.

In addition to the autoimmune nature, other factors are associated with the progression of OLP lesions, such as: allergic reactions to dental restorative agents (amalgam and gold), chronic irritants such as poorly fitted dental prostheses or pointed cusps (Koebner phenomenon), thyroid dysfunction, particularly hypothyroidism\textsuperscript{40} and hepatitis B or C virus infection. Genetic, immunological, psychological (stress and anxiety) factors, systemic conditions, and viral or bacterial infections may all play a significant role in the pathogenesis of OLP\textsuperscript{14}. Therefore, in addition to accurate history and pharmacological therapy, patient counseling to improve their lifestyle is indispensable. In this case report, no other contributing factors were associated with the development of the lesions.

It is important to stress the risk of malignant transformation of LP, since it is considered a lesion with potential for malignant transformation\textsuperscript{3,40,41}. It has been associated with tobacco use, alcohol and hepatitis C virus infection\textsuperscript{42}. Although these factors are not reported by the patient in this study, we encourage regular follow-up of patients with OLP. There is no consensus on the optimal number of annual dental visit returns, but two visits per year seem adequate and feasible\textsuperscript{43}. In this case, despite the absence of factors associated with malignant transformation, returns were scheduled every 6 months due to the new treatment of the lesion. After the remission of the clinical condition, visits to the dental office took place annually. Currently, the patient is 6 years free of the disease.

To our knowledge, this is the first case report of OLP treated by the combination of calcipotriene and betamethasone. This new therapeutic approach provides a safe and effective new alternative for treating lip OLP, providing remission of the lesion after conventional therapy has failed and a consequent improvement in the patient's quality of life. However, further prospective clinical studies involving larger groups of OLP patients are required to consolidate the pharmacological protocol implemented here and to investigate possible side effects of this treatment. We did not observe any side effects resulting from the pharmacological therapy adopted or any clinical changes in OLP during the recommended treatment.

CONCLUSION

LP may appear first and / or exclusively in the oral cavity. When in lip it presents itself clinically in different ways, having relevance in dental practice because it causes aesthetic impairment and often morbidity. Knowing the different forms of treatment available gives the choice of the right therapy for each case and leads to successful treatment and improved patient prognosis. We used calcipotriol and betamethasone combination therapy to treat a case of lip-restricted OLP, with complete remission after a long follow-up. This combination of drugs is widely accepted in the treatment of PV patients, with high success rates. We believe that the similar pathogenesis between the two lesions is the reason that leads to remission responses. We suggest that calcipotriol and betamethasone combination therapy may be a new approach in treating OLP.
REFERENCES


CONFLICTS OF INTERESTS
The authors declare no conflicts of interests.

CORRESPONDING AUTHOR
Thâmara Manoela Marinho Bezerra
Departamento de Patologia Oral, Faculdade de Farmácia, Odontologia e Enfermagem (FFOE)
Universidade Federal do Ceará (UFC)
60430-355 Fortaleza - CE, Brazil
E-mail: tmmbezerra@gmail.com

Received 07/04/2020
Accepted 23/10/2020