INTRODUCTION

Bisphosphonates are anti-resorptive agents highly efficient in binding bone hydroxyapatite. They can modulate the bone turnover and increase its mineral density.\textsuperscript{1,2} Bisphosphonates are widely used to treat postmenopausal and corticosteroid-induced osteoporosis.\textsuperscript{3}

The use of bisphosphonates in osteoporosis has been shown to improve bone strength and decreases the risk of fracture.\textsuperscript{4} Moreover, bisphosphonates are used in Paget’s disease and fibrous dysplasia,\textsuperscript{5} to balance metastatic cancer (primarily breast and prostate) deposits in bone,\textsuperscript{6,7} to cure severe hypercalcemia\textsuperscript{8} and to treat the bone resorption defects of multiple myeloma.\textsuperscript{8} Some examples of widely used oral bisphosphonates are alendronate, risedronate, etidronate,

Abstract

Background: Oral bisphosphonates are widely used in the treatment of bone resorptive diseases. There is an evidence that oral bisphosphonates can exert adverse effects on the oral mucosa independently of their effects on the jaw bones.

Objective: To systematically map the literature on adverse effects of oral bisphosphonates on the oral mucosa of adults with bone resorptive diseases.

Design: Scoping review of the literature, including different study designs.

Methods: Systematic searches of the PubMed, LILACS, Google Scholar and EMBASE databases were conducted. Two independent reviewers screened titles and abstracts according to predetermined criteria.

Results: The search retrieved 26 unique articles, comprising 22 case reports, one case series and three reviews describing a total of 56 cases of oral adverse events related to oral bisphosphonates. 88% of the reported cases were female suffering from comorbidities other than metabolic bone diseases. The improper use of the oral bisphosphonate was the most suspected cause of the adverse effect on the oral mucosa. Its management mainly involved withdrawal of the medication.

Conclusion: Adverse effects on the oral mucosa can develop from using oral bisphosphonates. Standardised registration of these adverse effects in university clinics and private practises could provide additional information about their occurrence and severity.

KEYWORDS
bisphosphonates, erosive oral lesions, mouth ulcer, mucositis

1 | INTRODUCTION

Bisphosphonates are anti-resorptive agents highly efficient in binding bone hydroxyapatite. They can modulate the bone turnover and increase its mineral density.\textsuperscript{1,2} Bisphosphonates are widely used to treat postmenopausal and corticosteroid-induced osteoporosis.\textsuperscript{3}
TABLE 1 Eligibility criteria for studies

| Inclusion                                      | Exclusion                                                                 |
|------------------------------------------------|----------------------------------------------------------------------------|
| Clinical trials, cohort studies, case series, case reports, reviews reporting cases. | Studies reporting on MRONJ only.                                           |
| Adult patients taking oral bisphosphonates for bone resorptive diseases.          | Studies where the mucosal adverse effect is associated with pathology of the jaw structures. |
| Adverse effect of the bisphosphonates on the oral mucosa.                   | Reviews not reporting any original cases, surveys, comment letters, conference abstracts, letters to the editor. |
                                                                                   | In vitro and ex vivo studies.                                             |
                                                                                   | Animal studies.                                                           |

and minodronate. Although the most common oral formulation for these medications is tablets, a drinkable solution of alendronate and an alendronate oral jelly have been developed the last years.9,10

The therapeutic boundaries of bisphosphonates have been explored for several years and several adverse effects have been reported.11,12 Oral bisphosphonates, mainly used for the treatment of osteoporosis, have been linked to adverse events involving the upper gastrointestinal tract such as dysphagia, nausea, vomiting, dyspepsia, upper abdominal pain and discomfort.13-15 Moreover, oral bisphosphonates may cause erosive esophagitis, gastritis, duodenitis, delayed healing and haemorrhage due to oesophageal, gastric or peptic ulcers.16,17 Oral bisphosphonates may also cause musculoskeletal pain, ocular adverse events such as conjunctivitis and uveitis, hypocalcaemia and secondary hyperparathyroidism, atrial fibrillation and atypical fractures of the femoral diaphysis.11,12

As with other anti-resorptive agents,18 bisphosphonate can cause medication-related osteonecrosis of the jaws (MRONJ) is the most well-known adverse effect in the field of oral and maxillofacial pathology.19 Bisphosphonates can promote avascular necrosis of bone tissue, especially in the maxilla and mandible.20 MRONJ appears either as exposed bone or as a non-healing extraction socket.21 The intravenous administration of nitrogen-containing bisphosphonates remains the most common etiologic factor for MRONJ,19 whereas the risk of MRONJ from oral bisphosphonates appears to increase when the therapy is continued for more than three years.11

Bisphosphonate toxicity to gastrointestinal cells is well documented.14,16,17 Recent in vitro studies provide evidence that high doses of oral bisphosphonates may also have a negative impact on the oral mucosa.21-24 Doneti et al (2014) suggested that alendronic acid may have a negative effect on keratinised oral human mucosa of osteoporotic patients without clinical signs of MRONJ,22 whilst recent experimental studies in animals have demonstrated that alendronic acid can induce an inflammatory process in the buccal and tongue mucosa.23,24 Furthermore, adverse effects on the oral soft tissues have been described in several case reports and are accompanied by pain and great discomfort for the patient, whilst a life-threatening situation has also been described.25 Hence, it is essential for the dental practitioner to overcome the diagnostic challenges deriving from the abundance of the various oral mucosal lesions, identify the effects on the oral mucosa and provide an effective treatment. Nevertheless, the existing evidence is limited to scattered case reports and reviews focussed on a specific type of mucosal lesion.26,27 The aim of the scoping review was to systematically map the literature to ascertain the adverse effects of oral bisphosphonates on the oral mucosa.

2 | METHODS

2.1 | Protocol

This scoping review was conducted and reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews (PRISMA-ScR) guidelines.28

2.2 | Eligibility criteria

Peer-reviewed journal articles were included if they involved adult human patients who used oral bisphosphonates as a therapeutic agent and had suffered from an adverse effect reported on their oral mucosa. Clinical trials, cohort studies, case series, case reports and systematic reviews reporting cases were included. Papers were excluded when they reported only MRONJ, mucosal adverse effects related with osteonecrosis of the jaws or did not report any cases. Surveys, comment letters, conference abstracts, ex-vivo and in-vitro studies and animal studies were also excluded (Table 1). There were no limitations concerning the language or publication status (accepted, in press, published).

2.3 | Information sources and search strategy

To identify potentially relevant articles, the following online databases were searched from 1969 to June 2020: PubMed, LILACS, Google Scholar and EMBASE (from 1969 onwards). The search was contacted in a period from March until May 2020. The whole search process was repeated in December 2020.

Additionally, all issues of four dental journals, that were deemed relevant to the topic of this review, were hand-searched for further potentially relevant publications: Journal of Oral Maxillofacial Surgery, Head and Neck Pathology, Journal of Oral Pathology and Medicine and Gerodontology. Grey literature was searched through Grey Literature report (http://www.greylit.org) and OpenGrey

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that was not possible for two studies. 30,31

screening resulted in the exclusion of 16 articles, so every possible
inging duplicates 60 unique articles were identified. Title and abstract
articles were recovered from the grey literature sources. After remov
The electronic search yielded 104 potential articles. No relevant ar
RESULTS
Citing these articles were hand-searched for titles not identified with

Abstracts were obtained for all the studies identified during the
electronic and hand-searches. In cases where an abstract was not availa
language other than English were translated.

Predetermined data (Table 2) were extracted from each study indepen
by two reviewers (CP and ZP) and arranged into data tables. The data abstrac
researchers, during a preliminary extraction of the data from ten articles. Disagreements were solved through discussion. Full-
text copies were obtained for all the selected studies. Articles in lan-

No quality assessment of the included articles took place, which was in accordance with available guidelines on scoping review.29

The electronic search yielded 104 potential articles. No relevant ar-
ticles were recovered from the grey literature sources. After remov-

text, that was not possible for two studies.30,31 After reading the full text,

2.4 | Study selection process

2.5 | Data charting process and synthesis

2.6 | Critical appraisal

3 | RESULTS

(www.opengrey.eu). Because of the challenges in finding papers with data on adverse oral effects not related to MRONJ, addi-
tional search strategies were also used. The reference list of all the full-
text articles selected after the screening and the list of articles citing these articles were hand-searched for titles not identified with the previous methods.

2.4 | Study selection process

Abstracts were obtained for all the studies identified during the
electronic and hand-searches. In cases where an abstract was not available, full-text copies were obtained. Two reviewers (CP and ZP) screened titles and abstracts or full-text copies independently to eliminate articles that clearly failed to meet the eligibility criteria (Table 1). Any disagreements were settled through discussion. Full-
text copies were obtained for all the selected studies. Articles in lan-

data were described as “Not reported” (NR). Cases described in more than one studies were

Data charting process and synthesis

2.5 | Data charting process and synthesis

Predetermined data (Table 2) were extracted from each study indepen-
dently by two reviewers (CP and ZP) and arranged into data tables. The data abstraction table was pilot tested, by the two re-
viewers, during a preliminary extraction of the data from ten articles. Disagreements were solved through discussion between the two reviewers. When an article described multiple cases, only those that met the inclusion criteria were included in the review. Efforts were made to contact the author(s) and obtain additional information in cases when required data were not described in the articles. If this information could not be obtained, the data were coded as “Not reported” (NR). Cases described in more than one studies were identified (predominantly by using the extracted data) and excluded. Cases reported by individuals or organisations other than health care professionals (patients, marketing authorisation holder) were excluded.

Critical appraisal

No quality assessment of the included articles took place, which was

RESULTS

3 | RESULTS

The electronic search yielded 104 potential articles. No relevant ar-
ticles were recovered from the grey literature sources. After remov-
ing duplicates 60 unique articles were identified. Title and abstract screening resulted in the exclusion of 16 articles, so every possible effort was made to retrieve 44 articles in full text. Unfortunately,

Mapping of the evidence

3.1 | Mapping of the evidence

Most of the cases were female (n = 49). The age of the patients ranged from 48 to 96 years (median age 72 years, based on 53 pa-
tients). In three cases, the age of the patient was not mentioned.27,51

Ten cases involved patients who were residents of a nursing facility or were receiving 24-hour help at home.43,44,30,54-56,59

The major indication for prescribing bisphosphonates was osteo-

porosis (51/56). In one case, the bisphosphonate was administered
because of osteopenia, a preosteoporotic condition. Osteoarthritis was mentioned in one case. In another case, the bisphosphonate was administered to reduce the risk of osteoporosis secondary to polymyalgia rheumatica. In one case alendronate was prescribed after a knee replacement surgery. In one case alendronate was administered as a treatment to vertebral fracture.

The chief complaint of the patient was reported in 37 cases and mostly involved pain (24 cases) and difficulty in eating (13 cases). Almost half of the cases (25/56) involved patients who were suffering from diseases other than the metabolic bone diseases. In 22 cases, the medical history is not mentioned.

Dementia (11/56), Parkinson's disease (2/56) and cerebrovascular incidents (3/56) was reported in individual cases. One case reported a patient with Down Syndrome, whilst another case involved a legally blind patient suffering from schizophrenia. One case involved a patient with congenital deafness and another case was a non-verbal patient with learning difficulties. Cerebral palsy, hemiplegia, severe postural instability and a thoracic compression fracture were described in five individual cases.

Polypharmacy (the concurrent use of five or more medications) was reported in eleven cases (Table S3).

A previous history of adverse effects on the oral mucosa is not reported in any of the cases.

Nine patients were using a complete or partial denture. One case was edentulous. Dental status could not be verified for 46 cases.

Most of the cases involved patients who used oral bisphosphonates in the form of tablets, except for one case where the alendronate tablets had been recently replaced by a drinkable solution of alendronate.

Forty-six patients used only alendronate (46/56), and one patient used alendronate in combination with etidronate and calcium.
Three patients used etidronate with calcium,27 and two patients used only etidronate.48 Two patients used minodronate,54,56 and two patients used risedronate.27,48

Of the 47 patients using alendronate, 21 patients were taking the medication on a weekly basis (44,7%), 16 patients were taking it daily (34%), whereas the dosage of alendronate was not specified for 10 cases. Etidronate was administrated daily in four cases,27,48 whilst in one case there was no information available concerning the dosage of etidronate.48 Minodronate was administered monthly.54,56 Risedronate was administered in one case weekly27 and in the other case daily.48

The duration of the bisphosphonate administration was specified for twenty-nine patients (51,8%) and ranged from 4 days32 to 10 years.58 Twelve patients used the bisphosphonate for one month or less; 11 patients had been using the medication for less than a year, and six patients had used the medication for a year or more.

Incorrect use of the bisphosphonate had been identified as the cause of the mucosal adverse effects in 30 of the cases (53,6%). In these cases, the bisphosphonate tablet was sucked, chewed, or allowed to dissolve on the oral mucosa before swallowing. In 10 cases incorrect use was attributed to physical or mental disability, whilst in other cases incorrect use was related with difficulty to swallow tablets (8/30), distraction (32,42) (2/30), habit (33,35) (2/30), wrong administration from the care giver (43,59) (2/30), or wrong instruction given by the general practitioner.57 (1/30). The difficulty to swallow tablets was related in two cases to cerebral palsy,43 or previous surgery for the removal of a thyroid tumour.47 No explanation for the improper use was proposed for six cases.

The diagnostic process was reported for 34 cases. Intraoral and extraoral examination took place in all these cases. A physical examination was performed in 14 cases, whilst in 13 cases, other mucosae were also inspected. Adjunctive visual tools were used in eight cases, whilst Nikolsky’s sign was used in three other cases. Histological examination of the mucosal lesions was used for 11 cases. Microbiological tests were used for 22 cases, whilst cytodiagnostics tests were performed in 10 cases. Finally, a CT scan was obtained in one case. A summary of the studies reporting a diagnostic process is presented in Table S2. In all of these cases, the diagnostic process was complex, involving a variety of the described tools.

The mucosal adverse effects were documented with respect to their location, size, shape and colour. Severe ulcerations of the oral mucosa, also described with the terms stomatitis and mucositis, were reported in most of the cases (80,3%). Erythema multiform minor52 and oral lichenoid reaction49 were described in two individual cases. Blister in and around the mouth48 (4/56), swelling48 (2/56), vesicles48 in the mouth (1/56) and petechiae48 (1/56) were also reported. In one case, the term mucosal alteration48 was used to describe the adverse effect on the oral mucosa, but no further explanation was provided (1/56). The most frequent location was the tongue and the lower lip, followed by the palate and the buccal mucosa.

The period from the onset of the mucosal adverse effect up to the day that the patient was reported by the health professional ranged from one day22,48 to three years,55 whilst it remains unknown for 16 cases. Twenty patients had suffered from the mucosal adverse effects for more than a month (20/40). In five of these cases, the patients had suffered from the adverse effects for more than a year35,39,50,55,57 (Table S4).

The differential diagnosis was reported for 34 cases and included other mucosal disorders, traumatic ulcerations and lesions caused by bacterial or viral infections, aphthous, thyroid abnormalities, syphilis, blood deficiencies, immunobullous diseases (so as pemphigus and pemphigoid), oral lichen planus, erosive oral candidiasis, autoimmune diseases and malignancy.

In most of the cases the management of the adverse effect included withdrawal of the oral bisphosphonate (85,7%); in three cases, this was followed by the administration of an alternative form of bisphosphonate.27,54,59 Adjunctive regimens included the local application of corticosteroid33,40,59 (4/48), or bicarbonate crème40 (1/48) and the use of benzydamine oromucosal spray and chlorhexidine mouthwash59 (1/48). In two cases, the dose of the bisphosphonate was reduced.27 In one case, there is no change reported concerning the bisphosphonate.27 In the rest of the cases, the administration of the bisphosphonate was corrected, and the patients were monitored ever since whilst taking the medication. The healing period ranged from 7 days32,34 to 13 months.44 The period required to achieve complete healing remains unknown for nine cases.

Previous unsuccessful treatment with analgesic and antiviral agents, topical and systemic corticosteroids, hyaluronic acid, topical and oral anti-mycotic agents, antibiotic agents and chlorhexidine (without modification of the oral bisphosphonate administration) was reported for 12 cases.

4 | DISCUSSION

The purpose of this scoping review was to collate the available information on the adverse effects for the oral mucosa by adults using oral bisphosphonates. The nature and the rare occurrence of these adverse effects preclude observational prospective or retrospective studies. As a result, the available evidence is limited to the uncontrolled information presented in case reports. Case reports are a low level of evidence, and they are prone to various types of bias.60 Another limitation of case reports is that they cannot ascertain the cause of the adverse effects on the oral mucosa. The low prevalence of these adverse effects is the main reason why cases described in pharmaceutical databases27,48 were included in our study despite their limited documentation.

The pathophysiology underlying the adverse effects on the oral mucosa remains unknown at the present time. Alendronic acid has been shown to initiate inhibition of oral keratinocyte proliferation, alteration of the progress of epithelial terminal differentiation and changes in molecular composition and desmosomal morphology in the human oral mucosa of osteoporotic women without clinical signs of medication-related osteonecrosis of the jaws.22 There is also evidence that alendronic acid has a cytotoxic effect on direct contact
with periodontal ligament fibroblasts. According to another study, bisphosphonates have a strong negative effect on human oral keratinocytes by reducing cell viability, migration ability and accelerating the apoptosis rate of the oral mucosa. In a recent in vitro study, it has been suggested that the decrease in the number of hemidesmosomes noticed after administration of alendronic acid could indicate the formation of ulcerous lesions in murine buccal mucosa. Consequently, bisphosphonates may interfere in regenerative, apoptotic and ultrastructural cellular processes, leading to the development of mucosal adverse effects.

The majority of the patients in this study were older females, reflecting the higher prevalence of osteoporosis in older women. Two-thirds of the patients (66%) expressed pain and great discomfort in eating. These symptoms are of great significance for health care professionals, especially when treating older adults. In ageing population, dysphagia (swallowing difficulty), is a growing health concern, contributing to a variety of negative health status changes; such as, increased risk of malnutrition, dehydration, weight loss and aspiration pneumonia. Obviously, initiating a communication between the health care professionals can higher the awareness for the mucosal adverse effects and attribute to their early identification and management.

With respect to the type of oral bisphosphonate, alendronate was used in 84% of the cases, which reflects the wide use of this medication in patients with osteoporosis. Amongst the mucosal adverse effects of interest in this review, the most common one (80%) was oral ulceration. Erythema multiforme minor was also described in one case. Nevertheless, we made no attempt to differentiate the erythema multiforme minor from oral ulcerations as a toxic reaction to the oral bisphosphonate. A lichenoid reaction following treatment with oral bisphosphonates was also described in one case. A lichenoid reaction is associated with a greater likelihood of medication errors was reported in 19 per cent of the patients. Twelve per cent of the patients might have experienced difficulty to maintain an upright position, due to Parkinson’s disease, cerebral palsy, postural instability or compression fracture of the vertebra, whilst five per cent suffered from comorbidities (cerebrovascular incidents) that may impair the postural balance. Finally, twenty-five per cent of the patients experienced difficulty in swallowing, which is a common problem in older adults and may be attributed to neurologic disorders, such as transient ischaemic attacks and strokes, and neurodegenerative diseases, such as Parkinson’s disease. All of these conditions are associated with a greater likelihood of dosing errors and the subsequent complications, so a detailed medical history prior to the prescription of oral bisphosphonates is indispensable. It is noteworthy that the current European guidelines for the treatment of osteoporosis do not consider possible cognitive, postural, or swallowing impairment of the patient. Because incorrect administration can cause adverse effects on the oral mucosa, clear instructions on how to take oral bisphosphonates should be provided to all bisphosphonate users and, when applicable, their caregivers. When the administration of bisphosphonate tablets is contra-indicated other types of bisphosphonate formulation can be used, such as intravenous bisphosphonates and denosumab. Intravenous bisphosphonates in the treatment of osteoporosis have a proven efficacy, a better bioavailability than oral bisphosphonates and a good safety profile. Moreover, the long dosing intervals improve patient convenience and help to overcome adherence to drug therapy issues, whilst the risk of developing MRONJ in osteoporotic patients after exposure to intravenous bisphosphonates or denosumab remains very low. With regard to denosumab, there are no published clinical data suggesting that this medication can cause adverse effects on the oral mucosa that do not relate to MRONJ. In addition, a drinkable form of alendronate could also be used. Furthermore, an alendronate jelly administered once per week per os may be an effective alternative to tablets for older adults with swallowing difficulties. Some limitations of this review are apparent, with regard to the use of the alendronate oral drinkable solution and the alendronate sodium hydrate (oral jelly). Firstly, there are no published clinical data concerning the occurrence of adverse effects on the oral mucosa. Secondly, alendronate sodium hydrate is approved and used only in Japan. Regarding intravenous bisphosphonates and denosumab, there is limited data comparing their use with oral bisphosphonates in postmenopausal women with osteoporosis. These comparative studies include a small number of patients and have a relatively short term follow-up. As mentioned in the summary of the European guidance for the diagnosis and management for osteoporosis in postmenopausal women, oral bisphosphonates, when tolerated, remain the initial pharmacological intervention in the majority of the cases.
Oral mucosal lesions can develop as a result of a local factor as trauma or mechanical irritation or malignancy. They can be caused by a systemic disease or an auto-immune disorder, a viral, bacterial or mycotic infection, a toxic reaction on medication or a vitamin deficiency. Hence, identifying an oral mucosal lesion can be a challenging process. The diagnosis should be based on a thorough investigation of the patient including extraoral, intraoral and mucosal lesion assessments combined with a comprehensive medical history. Identifying the side-effects of the used medication can often provide valuable information. Additionally, diagnostic tests, including histological and microbiological samples, are necessary to establish a proper diagnosis.

An important limitation of the current review was the incomplete reporting of the cases in the included articles. Missing information about the medical history (39%), the exact number and type of medications taken (34%), the dental history (82%), the dosage of the bisphosphonates (19%), the administration period (48%), the onset and duration of the adverse effects (28.5%) and the healing period (16%) hindered mapping of the evidence. Evidently, incomplete reporting would be an obstacle for any systematic review on this topic. Hence, future case reports should adhere to more strict reporting standards and provide a more detailed description of the cases, since other types of clinical studies are not feasible. Standardised registration of oral mucosal adverse effects in university clinics and private practices could also provide additional information on these adverse effects.

Bisphosphonates are highly effective for the prevention of subsequent fractures in patients with osteoporosis or osteoporosis related fragility fractures. After discovering bisphosphonates in the nineties, 31 million prescriptions were dispensed in the United States only in 2008. Although the number of prescriptions is declining every year due to adverse effects, in 2012 still over 15 million prescriptions are taken (34%). The dental history (82%), the dosage of the medications taken (34%), the administration period (48%), the onset and duration of the adverse effects (28.5%) and the healing period (16%) hindered mapping of the evidence. Evidently, incomplete reporting would be an obstacle for any systematic review on this topic. Hence, future case reports should adhere to more strict reporting standards and provide a more detailed description of the cases, since other types of clinical studies are not feasible. Standardised registration of oral mucosal adverse effects in university clinics and private practices could also provide additional information on these adverse effects.

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5 | CONCLUSION

Adverse effects on the oral mucosa can arise from using oral bisphosphonates. The most common adverse effect is oral ulceration. Half of the cases were related to improper administration of the medication. Health care professionals and oral health professionals, in particular, should consider the possibility of adverse effects in older adults who are prescribed oral bisphosphonates, especially when complications in administration can be expected due to mental or physical difficulties. Standardised registration of these adverse effects in university clinics and private practices, could provide additional information about their prevalence and severity.

CONFLICT OF INTEREST

There are no conflicts of interests in connection with this article.

AUTHOR CONTRIBUTIONS

CP conceived and designed the study. CP and ZP conducted the study with the support of HCW, WJK and CDvdMW. HCW offered her expertise in geriatrics and osteoporosis. HCW, WJK and CDvdMW provided critical feedback and CP wrote the manuscript with input of all the authors. CDvdMW had the final planning and supervision of the study. All authors read and approved the final manuscript.

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