What is the recommended dental management for patients who are receiving oral bisphosphonate therapy?

**Background**

By decreasing osteoclastic activity, bisphosphonate drugs decrease rates of bone resorption, resulting in an increase in bone mass when given to patients with osteoporosis. They also have therapeutic effects for patients with rarer metabolic bone diseases such as Paget’s disease and osteogenesis imperfecta and for cancer patients with metastases to bone. Two forms of bisphosphonate treatment are currently available — oral (Table 1) and intravenous (see article on p. 618). Most patients receiving bisphosphonate therapy who are encountered in the general dental setting are receiving oral treatment, usually for osteoporosis.

**Management Advice**

It has been hypothesized, though not proven, that oral bisphosphonate therapy may be associated with osteonecrosis of the jaw. The scientific data for cases of bisphosphonate-related osteonecrosis of the jaw (BRONJ) are incomplete, and the vast majority of patients receiving oral bisphosphonate therapy do not experience any oral complications. As such, patients should be informed that the health benefits of oral bisphosphonate therapy far outweigh the minimal risk (if any) of BRONJ. In addition, good oral hygiene, accompanied by regular dental care, is the best way to minimize this risk, if it exists. Patients receiving bisphosphonate therapy should be advised to contact their dentist if any problem develops in the oral cavity. In general, patients who are taking oral bisphosphonates without other risk factors (Box 1) can be treated according to normal protocols and procedures, including surgery.

For patients receiving oral bisphosphonate therapy, dental treatment recommendations are similar to those for patients not taking the medication, as described in the following sections.

**Restorative and Prosthetic Dentistry**

All restorative procedures may be performed. At present, there is no evidence that malocclusion or occlusal forces increase the risk of BRONJ. Prosthodontic appliances should be adjusted for fit to avoid mucosal irritation.

**Periodontal Diseases**

Treatment protocols are similar to those for the general population (i.e., patients not taking the medication).

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**Table 1** Bisphosphonates currently available in Canada for oral administration

<table>
<thead>
<tr>
<th>Generic name</th>
<th>Brand name</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alendronate</td>
<td>Fosamax</td>
<td>Osteoporosis, Paget’s disease</td>
</tr>
<tr>
<td></td>
<td>Fosavance</td>
<td>Osteoporosis</td>
</tr>
<tr>
<td>Clodronate</td>
<td>Bonefos, Clasteon</td>
<td>Bone metastases of malignant tumours, hypercalcemia of malignancy</td>
</tr>
<tr>
<td>Etidronate</td>
<td>Didrocal</td>
<td>Osteoporosis</td>
</tr>
<tr>
<td></td>
<td>Didronel</td>
<td>Paget’s disease</td>
</tr>
<tr>
<td>Risedronate</td>
<td>Actonel</td>
<td>Osteoporosis, Paget’s disease of bone</td>
</tr>
<tr>
<td></td>
<td>Actonel Plus Calcium</td>
<td>Osteoporosis</td>
</tr>
</tbody>
</table>
Endodontics

If the tooth is salvageable, endodontic treatment is preferred to extractions or surgical manipulation. If extractions or surgical manipulations are necessary, such procedures should follow the recommendations discussed in the section “Oral and Maxillofacial Surgery” above.

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References


QUESTION 2

What is the recommended dental management for patients who are receiving intravenous bisphosphonate therapy?

Background

Since 2003, dentists have been observing osteonecrosis of the jaw as a potential complication of intravenously administered bisphosphonate therapy. Bisphosphonate-related osteonecrosis of the jaw (BRONJ) is defined as an area of exposed bone in the maxillofacial region that does not heal within 8 weeks after its identification by a health care provider, in a patient who is receiving or has previously been receiving bisphosphonates and who has not had radiation therapy to the craniofacial region.

The bisphosphonates currently available on the Canadian market for intravenous (IV) administration are listed in Table 1.

Management Advice

Dental Management of Patients Receiving IV Bisphosphonate Therapy

The risk of BRONJ appears to range between 1% and 10% in patients receiving IV bisphosphonate treatment. Any patient receiving such therapy should be informed of the signs and symptoms of BRONJ. In addition, before the IV bisphosphonate therapy is started, the patient should undergo a dental evaluation by a qualified dental professional, and dental recall examinations should be performed throughout the course of bisphosphonate therapy. The frequency of such examinations will be dictated by the patient’s clinical and dental status.

Box 1

Risk factors for bisphosphonate-related osteonecrosis of the jaw in patients receiving oral bisphosphonate therapy

- Concomitant use of estrogen or glucocorticoids
- Comorbid conditions (e.g., malignancy)
- Poorly fitting dental appliances
- Intraoral trauma
- Presence of tori or other bony exostoses
- Pre-existing dental or periodontal disease
- Older age (> 65 years)
- Alcohol and/or tobacco use

Oral and Maxillofacial Surgery

Treatment protocols are similar to those for the general population (i.e., patients not taking the medication), unless other risk factors are present (Box 1). In such cases, conservative surgical technique, with primary tissue closure, should be considered when extractions or surgery are necessary (including elective dentoalveolar surgical procedures such as implant placement, reduction of tori or extraction of asymptomatic teeth).

Patients may use a chlorhexidine-containing rinse immediately before and after surgical procedures. Systemic antibiotic therapy may be considered for perioperative prophylaxis or if there is evidence of infection.

Questions

1. What is the recommended dental management for patients who are receiving intravenous bisphosphonate therapy?

2. What is the recommended dental management for patients who are receiving intravenous bisphosphonate therapy?
It is also important to identify patients with risk factors for BRONJ: dental extraction and/or oral bone surgery; poorly fitting dental appliances; intraoral trauma; presence of tori or other bony exostoses; pre-existing dental or periodontal disease; older age (> 65 years); prolonged exposure to bisphosphonate therapy; concomitant use of estrogen or glucocorticoids; comorbid conditions (e.g., malignancy); alcohol and/or tobacco use.1

Patients with cancer who are receiving IV pamidronate and/or zoledronic acid are at the greatest risk for BRONJ.1

If the patient’s situation permits, invasive dental procedures should be performed before the IV bisphosphonate therapy is started, with follow-up at 14–21 days to ensure complete healing at the surgical site. The following sections outline treatment recommendations for patients who are already receiving IV bisphosphonate therapy.1,3

Restorative and Prosthetic Dentistry

All restorative procedures may be performed. At present, there is no evidence that malocclusion or occlusal forces increase the risk of BRONJ. Prosthodontic appliances should be adjusted for fit to avoid mucosal irritation.

Periodontal Diseases

Nonsurgical therapy is preferred (such as scaling and root planing). Periodontal surgery is not recommended. When necessary, surgical treatment should be aimed primarily at obtaining access to root surfaces.

Oral and Maxillofacial Surgery

Whenever possible, nonsurgical endodontic or periodontal therapy is preferred to extraction, unless there is a risk of aspiration. Elective dento-alveolar surgical procedures, such as implant placement, reduction of tori and extraction of asymptomatic teeth, should be avoided. When an extraction or surgery is necessary, conservative surgical technique, with primary tissue closure, should be considered. The greater incidence of BRONJ in the mandible than the maxilla, especially in the posterior region of the mouth, must be taken into account in the decision to perform surgery.

Endodontics

For salvageable teeth, endodontic treatment is preferred to extractions or surgical manipulation. Manipulation beyond the apex should be avoided. Surgical procedures should be guided using the same recommendations mentioned in the section “Oral and Maxillofacial Surgery.”

Considerations for Any Surgical Procedure

Patients should use a chlorhexidine-containing rinse immediately before and after surgical procedures. Systemic antibiotic therapy may be considered for perioperative prophylaxis or if there is evidence of infection (and should follow the guidelines of the American Dental Association3) (Table 2).

<table>
<thead>
<tr>
<th>Generic name</th>
<th>Brand name</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clodronate</td>
<td>Bonefos, Clasteon</td>
<td>Bone metastases of malignant tumours, hypercalcemia of malignancy</td>
</tr>
<tr>
<td>Pamidronate</td>
<td>Aredia</td>
<td>Bone metastases of malignant tumours, hypercalcemia of malignancy, multiple myeloma, Paget’s disease</td>
</tr>
<tr>
<td>Zoledronic acid</td>
<td>Aclasta</td>
<td>Paget’s disease</td>
</tr>
<tr>
<td>Zometa concentrate</td>
<td></td>
<td>Bone metastases of malignant tumours, hypercalcemia of malignancy, multiple myeloma</td>
</tr>
</tbody>
</table>

**Table 2** Proposed antibiotic therapy1

<table>
<thead>
<tr>
<th>Patient’s penicillin status</th>
<th>Suggested antibiotic</th>
<th>Oral regimen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not allergic to penicillin</td>
<td>Amoxicillin</td>
<td>500 mg t.i.d. for 14 days</td>
</tr>
<tr>
<td></td>
<td>May be combined with metronidazole</td>
<td>250 mg t.i.d. for 14 days</td>
</tr>
<tr>
<td>Allergic to penicillin</td>
<td>Clindamycin</td>
<td>300 mg t.i.d. for 14 days</td>
</tr>
<tr>
<td></td>
<td>Azithromycin</td>
<td>250 mg t.i.d. for 10 days</td>
</tr>
</tbody>
</table>
Dental Management of Patients with BRONJ

If BRONJ is suspected but not yet confirmed (e.g., duration of unhealed exposed bone less than 8 weeks; Fig. 1), the patient should be followed carefully. Additional common findings include pain, swelling, paresthesia, suppuration, soft-tissue ulceration, intraoral or extraoral sinus tracks, and loosening of teeth. Radiographic findings can vary from changes in bone density to no obvious alteration to the bone pattern.

The differential diagnosis for BRONJ includes gingivitis, periodontal diseases (e.g., necrotizing ulcerative periodontitis), osteomyelitis, sinusitis, temporomandibular disorder, trauma, periapical lesions, osteoradionecrosis, bone tumours and metastatic lesions.

Standard radiography such as panoramic and periapical radiography may help in the detection of BRONJ in the early stages. Computed tomography may also be considered. No imaging is required for patients with established clinical evidence of BRONJ.

The dental professional should alert the patient’s physician to the diagnosis and should report cases of BRONJ to the appropriate agencies, such as the manufacturer of any agent implicated. There is no published evidence to suggest that discontinuation of bisphosphonates will promote resolution of BRONJ.

If pain is present, it should be managed appropriately with nonsteroidal anti-inflammatory drugs or narcotic analgesics. The patient should be advised to use chlorhexidine (0.12%) or another similar oral antimicrobial rinse, and systemic antibiotic therapy may be prescribed if there is evidence of secondary infection. Establishing and maintaining good oral hygiene is essential.

Any patient with established BRONJ needing surgical procedures should be referred to an oral and maxillofacial surgeon, who may consult other qualified specialists as appropriate. Any dentoalveolar surgical procedure (i.e., extractions, implants or apical surgery) should be avoided since the surgical sites will likely result in additional areas of exposed necrotic bone. However, loose teeth should be removed from the exposed bone if there is a danger of aspiration. Similarly, loose segments of bony sequestra should be removed, but without exposing uninvolved bone. Sharp bone edges should be removed, to prevent trauma to the adjacent soft tissues. Segmental jaw resection may be required for symptomatic patients with large segments of necrotic bone or pathologic fracture.

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References


Further Reading


A patient brochure on bisphosphonates produced by McGill University’s faculty of dentistry is available online at www.cda-adc.ca/jcda/vol-74/issue-7/617.html.

QUESTION 3

Which digital intraoral sensor is better?

Background

There are 2 types of intraoral sensors: direct sensors and storage phosphor sensors. Direct sensors, whether they use charge-couple-device or complementary metal oxide semiconductor technology, are equivalent in terms of image quality. Image display is instantaneous as these sensors are connected to a computer. The storage phosphor sensor is a plate, with dimensions comparable to those of conventional film; images are obtained when the plate is inserted into and read by a scanner.

Several experts believe that today’s sensors are reaching their technological limits. Both direct and storage phosphor sensors are capable of producing diagnostic images for the tasks dentists perform daily, such as diagnosing caries, identifying periapical lesions and evaluating periodontal bone loss (Figs. 1–4).

Digital Sensor Characteristics

The characteristics of digital sensors that have an impact on image quality are contrast resolution, spatial resolution, latitude and sensitivity.

Contrast resolution is the ability to detect differences between shades of grey. Theoretically, a sensor capable of capturing more shades of grey (greater bit depth) is better. However, because computer monitors display only 8-bit images, in practice there will be no difference between intraoral sensors that capture 8-bit images (256 levels of grey), 12-bit images (4,096 levels of grey) and 14-bit images (16,384 levels of grey). In addition, the number of grey shades differentiated by the human eye is between 32 and 60.

Spatial resolution is the ability to capture details and is measured in line-pairs per millimetre (lp/mm). Film achieves a resolution of up to 20 lp/mm. Newer sensors with a pixel size of 20 µm are able to resolve 25 lp/mm. Storage phosphor systems achieve a lower resolution than direct sensors. Most dentists can perceive 6 lp/mm and up to 10–12 lp/mm with magnification; images magnified above that become pixilated and non-diagnostic. Digital sensors available today have a resolution of 7 lp/mm or more.

Latitude is the ability of digital receptors to provide diagnostic images
with a range of exposures. A disadvantage of conventional film is that it is easily overexposed or underexposed. Although the latitude of direct sensors is comparable to that of film, storage phosphor sensors have a greater latitude and, under normal conditions, images are unlikely to be overexposed or underexposed. The downside is the greater dose of radiation that patients will receive if greater exposure is used consistently.

Sensitivity is the amount of exposure required to produce an image. The more sensitive the receptor, the less exposure is required. One well-known advantage of intraoral digital radiography is the lower dose of radiation to which patients are exposed. The most sensitive intraoral film available is F-speed. Storage phosphor systems can produce images using half the exposure necessary with F-speed film. Direct systems require more exposure than storage phosphor systems, but less than F-speed film.

All imaging software products offer a range of tools for dentists to use to enhance their images. However, the goal is to acquire good-quality diagnostic images that require no enhancement, as modifying images may have deleterious effects.

Clear task-specific indications for the various enhancement tools have yet to be developed.

**Management Advice**

**Direct Systems**

**Advantages**
- Instantaneousness
- Additional images can be obtained without removing the sensor from the mouth
- Spatial resolution superior to storage phosphor

**Disadvantages**
- Sensors are expensive and fragile
- Physical properties of the sensor: thick, rigid, attached cable. Positioning devices are available for all direct sensors to allow the device to be placed parallel to the teeth. However, this technique is not always possible, particularly for patients with a narrow palate. Reverting to the bisecting technique is more frequent than with film. Missed apices are a common problem, particularly for new users of this technology (Fig. 5). The presence of the cable makes obtaining an image of vertical bitewings almost impossible.
- More than one size sensor will be needed. Most companies offer size 1 and 2 sensors whose active areas are smaller than their film counterparts. Some companies now offer a size 0 sensor for pediatric applications. Size 2 sensors are required for interproximal examinations to view the bone level, but obtaining a distal image of the canines with these large sensors is challenging (Fig. 3).
- More exposures are required compared with film because of the smaller active surface area of direct sensors and difficulties in positioning (Fig. 6).
- The learning curve is greater than with storage phosphor sensors.

**Storage Phosphor Systems**

**Advantages**
- Latitude superior to direct sensors and film
- Sensitivity superior to direct sensors and film
- Sensor thickness and flexibility are comparable to those of film
- Plates available in sizes 0 to 4
- Plates compatible with standard positioning devices for obtaining periapical, horizontal and vertical interproximal radiographs
• Transition from film to storage phosphor is simple

Disadvantages
• Spatial resolution inferior to that of direct sensors
• Scanning of exposed plates is required. Scanning time increases with the size and number of plates and required resolution
• Space for the scanner is required, preferably in a dimmed environment as exposed plates are sensitive to light
• With handling, plates become scratched and damaged at the edges (Fig. 7) and must be replaced regularly

Lighting Requirements
The lighting conditions under which images are interpreted must be considered. Dental operators are generally equipped with high ambient light; this must be reduced to create an environment suitable for analysis of digital images. Adjusting the contrast and brightness of monitors will also improve image quality. Cathode ray tube monitors tend to lose brightness with time.

Transition Period
Regardless of the system selected, expect a transition period to adapt to looking at digital images, which appear to have less detail. The evidence shows that the information needed to make common diagnoses is there. The medical profession adopted digital radiology to replace conventional plain films before the dental profession, possibly because radiologists were used to reading computed tomography and magnetic resonance imaging scans on monitors. However, as stated by Ludlow and Mol, “It is no longer a matter of if but rather when the majority of dental offices will use digital imaging.”

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References
Is the choice of attachment for implant overdentures influenced by the angulation of the implant?

**Background**

The treatment of edentulous patients with implant-retained or implant-supported removable prostheses yields satisfactory results in terms of both masticatory function and patient satisfaction. However, maintenance is a concern, especially during the first year of denture wear, regardless of the type of attachment (bar, magnets or stud). Some researchers have reported the need for frequent reactivation of loose components and replacement of fractured matrices and patrices. Manufacturers recommend specific positioning and angulation of implants planned for overdentures to ensure predictable retention of the attachment and to prevent premature wear and fatigue, thereby reducing maintenance.

In determining implant angulation, an anatomic plane such as Camper’s plane, the Frankfurt plane or the mandibular plane can be chosen as a reference. The occlusal plane of the existing or planned prosthesis would also be an adequate reference for measuring the angulation of implants or planning their ultimate position.

It has been suggested that implants should be placed as parallel to the path of insertion of the overdenture and as perpendicular to its occlusal plane as possible. Positioning the implants in accordance with these references facilitates denture insertion and avoids excessive nonaxial loading.

Because of differences in the size and shape of the residual ridge and the maxillomandibular relation, ideal angulation of the implants during placement can be difficult or impossible to achieve. In a preliminary study, 19% of the implants evaluated had an angulation of 90° to the reference plane, 11% had a lingual inclination and 70% were inclined labially. Figure 1 illustrates divergent inclines with 2 implants.

Other studies have shown that between-implant divergence or convergence of about 10° is technically manageable, but excessive wear of the attachments has been described with greater angulation. Labial inclination superior to 6.5° and lingual inclination superior to 6° in relation to the sagittal plane were associated with more repairs, whereas there was no difference in the incidence of adjustments or repairs associated with angulations projected on the frontal plane. Treatment failure can also be related to mechanical weakness of the attachment system employed.

**Assessing Implant Angulation**

**Intraoral Assessment**

Plastic or metallic extension pins can be connected to the implants to allow angulation to be evaluated directly, with the implant in the patient’s mouth. Angulation of the implants is then estimated by visually comparing the direction of the extension pins and standard marks on an instrument with preset angu-
Minor Discrepancies in Between-Implant Angulation

In cases of minor or mild discrepancies in between-implant angulation, the following principles should be observed:

- The matrix parts of ball attachments should be oriented according to a common path of insertion for the individual abutments before the application of acrylic.

- Matrix components with special designs to tolerate between-implant angulation discrepancies are commonly available. Plastic parts with specific levels of resiliency, which are designed to tolerate divergence or convergence between implants of up to 40°, are available for some systems, particularly the cylindrical designs.

It should be emphasized that the amount of wear in cases of between-implant angulation discrepancies is directly related to the magnitude of the angle (Fig. 7).

Laboratory Assessment

The device described above can also be used to estimate implant angulations on a dental cast, obtained by pouring an impression of the arch with dental stone (Figs. 3 and 4). Alternatively, a protractor can be used, which will yield a more accurate measurement of the angle.

Radiographic Assessment

Panoramic radiography (Fig. 5) and teleradiography (Fig. 6) permit evaluation of the direction of the implants relative to a reference plane, such as an anatomic or denture occlusal plane, and measurement of between-implant angulation.

Choice of Attachment System

When the location and alignment of the implants are adequate, the choice of attachments should be based on clinical criteria such as the degree of retention required, the number of implants and the available prosthetic space.
place, which adds flexibility in the search for an ideal attachment position.

- Magnet or bar attachments can be used in selected cases. For the magnetic type, lack of a mechanical engagement between matrix and patrix prevents problems related to implant angulation, providing a workable solution in even the most severe cases. However, less reliable retention and maintenance problems, such as wear of components and corrosion of the magnetic alloys, have been frequently reported as the main disadvantages of these systems. Bar-clip attachments are another option when angulation of implants is excessive. An adequate path of insertion and adequate retention for overdentures can be easily achieved by splinting the implants with a metallic bar, although this type of attachment usually requires more vertical and between-implant space. It also costs more, requires extra laboratory and chairside time, and is often more difficult for patients to clean than the stud type of attachment.

Conclusions
The angulation of dental implants can have a clinically relevant effect on the attachment system for implant overdentures. To prevent excessive wear of the attachment components, loss of retention, maintenance problems and unnecessary costs, the most effective system should be chosen after careful assessment of implant angulation.

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References