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Tinnitus and Temporomandibular Disorders
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Japan

1. Introduction

Temporomandibular disorders (TMDs) are a group of related disorders of the masticatory system (the masticatory musculature and the temporomandibular joint). The most frequent symptom is pain, usually localized in the muscles of mastication, the preauricular region, and the temporomandibular joint (TMJ). Patients often complain of jaw ache, earache, headache, and facial pain. In addition to pain, patients with these disorders frequently have limited or asymmetric jaw movement and joint sounds that are described as clicking or crepitus (McNeill, 1990). The causes of TMDs remain unclear, and numerous factors have been implicated.

The first description of the relationship between TMJ dysfunction and aural symptoms is thought to have been made by Costen in 1934. Costen reported various clinical cases of patients with ear and nasal symptoms; he summarized his findings by stating that hearing tests showed a mild type of catarrhal otitis with eustachian tube involvement (usually simple obstruction). The prognosis of such cases reportedly depended on these factors: (a) the accuracy with which refitted dentures relieved the abnormal pressure on the joint; and (b) the extent of the injury to the tube and to the condyle, meniscus, and joint capsule. A more general acknowledgement of the relationship between ear symptoms and TMJ dysfunction was subsequently made (Costen, 1934, 1936, 1944). Sicher reported that from an anatomical perspective, the eustachian tube cannot be compressed during the closure of the mandible; neither can the opening action of the tensor palati muscle on the tube be impaired in this condition, making Costen’s theory impossible (Sicher, 1948). Schwartz theorized that contractive muscle spasms may cause pain as a direct result of myofascial trigger mechanisms and by referred routes (Schwartz, 1956).

2. Tinnitus and treatment for TMDs

After Schwarzt’s proposal, Dolowitz et al. reported that muscle exercises were effective for the relief of aural symptoms, especially ear fullness (44 out of 47 diagnosed patients) and tinnitus (40 out of 43 diagnosed patients). Various therapies have since been tried for the relief of TMD symptoms, including aural symptoms (Dolowitz et al., 1964). These therapies include selective grinding of the teeth, bite plate, biofeedback training, thermotherapy and muscle training (Koskinnen et al., 1980; Principato & Barwell, 1978). Parker and Chole reported that tinnitus and vertigo were significantly more prevalent among a TMD group than among either of their control groups (Parker & Chole, 1995). Erlandsson et al. reported that stomatognathic treatment (occlusal splint, occlusal adjustment, and exercise therapy)
and biofeedback treatment seemed to have some positive effects in a subgroup of tinnitus patients (with normal hearing) (Erlandsson et al., 1991). Rubinstein performed extensive studies of TMD and tinnitus and made the following conclusions: (1) 46% of the patients who had received stomatognathic treatment reported no tinnitus or reduced tinnitus a few months after treatment. A two-year follow-up showed that the improvement persisted in most of those who had benefited from the treatment. (2) Frequent headaches, fatigue/tenderness in the jaw muscles, clinical findings of pain upon palpation of the masticatory muscles, impaired mandibular mobility, and signs of parafunctions were more prevalent among tinnitus patients than among an epidemiological sampling. (3) The awareness of diurnal bruxism and a feeling of jaw tenderness/fatigue may be related to fluctuating tinnitus, vertigo and hyperacusis. (4) The evaluation of treatment outcome showed some improvements at the group level: a decrease in tinnitus, mood improvement, and a reduction in clinical signs of the dysfunction of the masticatory system. And (5), a strong relationship existed between tinnitus complaints and several symptoms of TMD. In summary, a relatively low severity of tinnitus, normal hearing, and fluctuations in the tinnitus intensity were good predictors of the psychological, stomatognathic, and cervical treatment outcome (Rubinstein, 1993). Wright and Bifano reported that 38% of their clinic’s patients reported coexisting tinnitus; among the 93 subjects included in their study, 56% reported that their tinnitus resolved after TMD treatment, 30% reported a significant improvement, and 14% reported minimal or no change (Wright & Bifano, 1997). On the other hand, Vernon et al. divided 929 patients into two groups: a TMD group, consisting of 69 patients in whom no known cause of tinnitus was present other than one or more temporomandibular joint (TMJ) indicators (example: effect of jaw manipulation on tinnitus, incidence of pain in tinnitus ear, and sensation of fullness in tinnitus ear), and a comparison group with mixed etiologies. These two groups were then compared to determine the attributes of tinnitus that significantly separated these groups. They concluded by recommending that patients with tinnitus of unknown etiology who exhibited three or more TMJ indicators should be referred for TMD treatment (Vernon et al., 1992). Similarly, Chan and Reade reported that tinnitus may have a multifactorial etiology, similar to pain, for which the correction or elimination of any one of the factors may result in the complete resolution of symptoms (Chan & Reade, 1994). Thus, the absence of any significant or detectable auditory, genetic, drug-related, or trauma-related causes should be confirmed by otorhinolaryngologists before treating TMD patients with tinnitus.

3. Treatments for TMDs

Although the causes of TMDs have not been clarified, various treatment modalities have been clinically attempted. The most prevalent treatment is occlusal treatment, including occlusal adjustment and occlusal reconstruction. The outcomes of occlusal adjustments, in particular, have been reported several times; however, a diverse range of outcomes has been noted, from very effective to only a placebo effect (Forssel et al., 1986, 1987; Forssel et al., 1999; Kopp, 1979; Kopp & Wenneberg, 1981; Tsolka et al., 1992; Tsolka & Preiskel, 1993; Vallon et al., 1991, 1995; Vallon & Nilner, 1997; Wenneberg et al., 1988). The use of occlusal adjustments has focused on the correction of a wide range of occlusal conditions (e.g., premature contact in the centric relation obtained by manipulation of the mandible, not by muscular directing; a slide between the intercuspal position [IP] and the centric relation [RCP]; and occlusal contact on the non-working side), rather than the elimination of a single
condition (e.g., only premature contact in the centric relation). Thus, the results of these treatments have been difficult to interpret. Egermark-Eriksson et al. performed a long-term epidemiologic study of the relationship between occlusal factors and TMD and reported that occlusal interference, as conventionally described (in an occlusal analysis based on the reference position of the RCP obtained by manipulating the mandible), does not seem to be of much help in explaining the development or persistence of TMD (Egermark-Eriksson et al., 1987). Pullinger et al. also reported that occlusion cannot be considered the unique or dominant factor in defining TMD populations (Pullinger et al., 1993). Since the relationship between these occlusal factors (premature contact in the centric relation obtained by manipulation, not a straight slide from the centric to intercuspal position) and TMD have not been scientifically demonstrated, the Technology Conference on the Management of Temporomandibular Disorders of the NIH/NIDR (held on April 29 through May 1, 1996, in the USA) concluded that no data exists to support many commonly held beliefs regarding TMD, nor does any data exist to support the superiority of any method of management compared with the effect of a placebo. The conference participants insisted that validated diagnostic methods, with respect to sensitivity and specificity, for the identification of TMD were lacking and noted the obvious need for basic research on TMD. They then stated their belief that occlusal adjustments are invasive and that the superiority of such treatment over other non-invasive therapies has not been demonstrated in randomized controlled prospective trials. Since this conference, many conflicting opinions have been reported (Dawson, 1999; Green et al., 1998), and most clinical researchers and basic researchers have become estranged from investigations of the relationship between occlusion and TMD. Therefore, studies on occlusion and TMD have been rarely performed since that time, and treatment for TMD has shifted to more conservative modalities (Tsukiyama et al., 2001). The outcomes of these conservative treatments for TMD have been symptomatic and temporary, not resulting in a cure, because the treatments have not been applied to patients based on evidence, as the causes of TMD remain unknown. Therefore, the effects of these treatments for tinnitus have also varied (Dolowitz et al, 1964; Erlandsson et al., 1991; Parker & Chole, 1995; Principato & Barwell, 1978; Schwartz, 1956). In 2005, Torii and Chiwata reported a highly significant relationship between the TMJ sounds and the discrepancy between the habitual occlusal position (HOP) and the bite plate-induced occlusal position (BPOP) (Torii & Chiwata, 2005). TMJ sounds are the most common and most relevant sign of TMD (Könönen et al., 1993; Köhler et al., 2009; Nydell et al., 1994). Thus, the possible effect of occlusal equilibration of the BPOP on TMD signs and symptoms has also been inferred. In their study, the BPOP was defined using a physiological muscular position as a reference position. The BPOP was obtained during voluntary jaw closing, while the patient was in an upright position and after wearing an anterior bite plate for a short period time (Fig. 1).

The muscular position has been reported to vary with the posture of the subject and is thought to be less reproducible than the manually obtained centric relation (which is guided by a dentist). However, Torii and Chiwata reported that the variation in the BPOP decreased after wearing an anterior bite plate, resulting in a reduction in the severity of TMD symptoms, compared with the symptoms experienced at the beginning of treatment (Fig. 2) (Torii & Chiwata, 2010). In addition, they reported the temporary effect of a bite plate on TMD and described the need for occlusal equilibration of the BPOP (Table 1, Fig. 3).
Fig. 1. **Anterior flat plane bite plate.** The plate covers the six upper anterior teeth and the first premolar teeth on both sides. The BPOP (physiological muscular position) is obtained during voluntary jaw closing after wearing the bite plate for a short period time.

Torii and Chiwata then performed a pilot study on the effect of occlusal equilibration using the BPOP on TMD and reported that the percentage of symptom-free patients at one year after treatment was 86% according to the Subjective Dysfunction Index (SDI) and 76% according to the Clinical Dysfunction Index (CDI); these changes in both indices after treatment were significant ($P < 0.01$) (Fig. 4 and 5) (Tables 1 and 2) (Torii & Chiwata 2010).
After occlusal adjustment

<table>
<thead>
<tr>
<th>Occlusal discrepancy</th>
<th>No discrepancy</th>
<th>Discrepancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discrepancy</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>No discrepancy</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

McNemar’s test: $X^2_{cal} = 18.05 > X^2_{(0.001)} = 10.83$, Significant. HOP: habitual occlusal position; BPOP: bite plate-induced occlusal position.

Table 1. Changes in the occlusal discrepancy between the HOP and BPOP before and after occlusal adjustment

<table>
<thead>
<tr>
<th>CDI</th>
<th>Myofascial</th>
<th>Disc disp.</th>
<th>Arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st Exam.</td>
<td>1 y</td>
<td>1 y</td>
<td>1 y</td>
</tr>
<tr>
<td>DiO</td>
<td>0%</td>
<td>80%</td>
<td>0%</td>
</tr>
<tr>
<td>DiI</td>
<td>0%</td>
<td>20%</td>
<td>36%</td>
</tr>
<tr>
<td>DiII</td>
<td>40%</td>
<td>0%</td>
<td>43%</td>
</tr>
<tr>
<td>DiIII</td>
<td>20%</td>
<td>0%</td>
<td>21%</td>
</tr>
</tbody>
</table>

CDI: Helkimo Clinical Index; 1st Exam.: first examination; 1 y: 1-year follow-up examination; Myofascial: Myofascial pain; Disc disp.: Disc displacement. DiO: no TMD; DiI: mild TMD; DiII: moderate TMD; and DiIII: severe TMD.

Table 2. Changes in Helkimo Clinical Index at first examination and after a 1-year follow-up examination for each TMD subtype

<table>
<thead>
<tr>
<th>NS</th>
<th>H1 B1</th>
<th>H2</th>
<th>H6</th>
<th>H7</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>↑ S</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NS</td>
<td></td>
<td>B2</td>
<td>H3</td>
<td>H4</td>
</tr>
<tr>
<td>Days</td>
<td>1</td>
<td>3</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Max.</td>
<td>40</td>
<td>42</td>
<td>42</td>
<td>42</td>
</tr>
</tbody>
</table>

NS: Not significant; S: Significant; Days: Days of visits; H1-10: Habitual occlusal position on different day; B1-10: Bite plate-induced occlusal position on different day; VAS: Score on a 10-point Visual Analogue Scale; Max.: Maximum unassisted opening (mm).

Fig. 3. HOP and BPOP positions and TMD symptoms on different days. The occlusal discrepancy between the HOP and the BPOP disappeared on day 6, and the disappearance was maintained between days 6 and 12. However, the discrepancy reappeared after 15 days. Therefore, equilibration of the occlusion was required to resolve the discrepancy. The occlusal equilibration was performed at 17 days.
Fig. 4. Dysfunction Index at first examination. Distribution of dysfunction indices before occlusal adjustment. Di O: no TMD; Di I: mild TMD; Di II: moderate TMD; Di III: severe TMD.

Fig. 5. Dysfunction Index at 1-year evaluation. Distribution of dysfunction indices after occlusal adjustment. Di O: no TMD; Di I: mild TMD; Di II: moderate TMD; Di III: severe TMD. None of the patients required an anterior bite plate during a one-year follow-up period after the completion of the occlusal adjustment. The number of visits varied between 2 and 34, with a mean of 11.0 ± 6.0 visits. Between 1 and 13 sessions of occlusal adjustment were performed, with a mean of 4.7 ± 3.5 sessions. The changes in the SDI and CDI were significant (p<0.01).
Fig. 6. **Mean HOP-BPOP difference before and after treatment.** Changes in the mean difference between the habitual occlusal position (HOP) and the bite plate-induced occlusal position (BPOP) before and after occlusal adjustment. x: mediolateral; y: anteroposterior; z: superoinferior. The changes were significant on the x-axis and y-axis (p<0.05), whereas the change on the z-axis was not significant (p>0.1).

Figure 6 and Table 1 indicate that a good outcome was obtained with resolution of the occlusal discrepancy by occlusal equilibration of the BPOP. Therefore, this treatment is a causal treatment, not a symptomatic treatment. In this pilot study, they reported that the otalgia of one patient and the tinnitus of another patient also completely disappeared after treatment. Table 3 shows the outcomes of the treatment of occlusal equilibration for TMDs and tinnitus in our practice.

**4. Causal treatment for TMDs**

Until now, occlusal discrepancy has been considered to be only one among several possible causes of TMD. At present, the only evidence-based causal treatment of TMD is occlusal equilibration, in which the habitual occlusal position (HOP) is aligned with the muscular position (BPOP). This treatment requires an articulator that can precisely reproduce the BPOP (voluntary or habitual closure movement). Torii observed various mandibular closures and found that the voluntary closing movement of the mandible consisted of a vertical movement against the occlusal plane at a point of less than 1 mm to the plane (Torii, 1989). He then developed an articulator that has a mechanism allowing vertical movement. Thus, the BPOP can be precisely reproduced using this articulator. When the upper and lower dental models are attached to the articulator with the BPOP wax record, the anteroposterior parallel space remains between both occlusal surfaces of the models after the wax record has been removed (Fig. 7).
### Table 3. Treatment outcome of TMD patients with aural symptoms

<table>
<thead>
<tr>
<th>No.</th>
<th>Patient number</th>
<th>Diagnosis</th>
<th>Treatment period</th>
<th>Outcome</th>
<th>Follow-up period</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>16 M</td>
<td>Limited opening</td>
<td>Disc disp.(R)</td>
<td>2 months resolved</td>
<td>5 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus (R), headache</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>2.</td>
<td>20 F</td>
<td>Limited opening</td>
<td>Disc disp.(B)</td>
<td>5 months resolved</td>
<td>3 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus(R) Myofascial</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td>22 F</td>
<td>Limited opening</td>
<td>Myofascial</td>
<td>5 months resolved</td>
<td>3 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Headache, tinnitus(L) Disc disp.(L)</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>4.</td>
<td>28 F</td>
<td>Limited opening</td>
<td>Myofascial (L)</td>
<td>3 months resolved</td>
<td>5 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus (L), fullness</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>35 F</td>
<td>Facial pain</td>
<td>Myofascial (L)</td>
<td>2 months resolved</td>
<td>3 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Headache, tinnitus (L)</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>37 F</td>
<td>Limited opening</td>
<td>Disc disp.(L)</td>
<td>3 months resolved</td>
<td>10 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus (L), headache</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>7.</td>
<td>42 F</td>
<td>Facial pain</td>
<td>Myofascial (L)</td>
<td>2 months resolved</td>
<td>5 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus (L), vertigo</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>8.</td>
<td>47 F</td>
<td>Limited opening</td>
<td>Disc disp.(L)</td>
<td>5 months resolved</td>
<td>2 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Tinnitus(L)</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>9.</td>
<td>73 F</td>
<td>TMJ pain</td>
<td>Arthralgia (R)</td>
<td>5 months resolved</td>
<td>7 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Vertigo, Tinnitus (R)</td>
<td>No recurrence</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. **Treatment outcome of TMD patients with aural symptoms**

No: Patient number; Visits: Number of visits; Period: Treatment period. Disc Disp.: Disc displacement; Myofascial: Myofascial Pain. M: Male ; F: Female. (R), (L) and (B): Mainly affected side on the (R): right; (L): left; (B): both sides.

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**Fig. 7. Mounting of upper and lower dental casts with BPOP wax record.** After the upper and lower casts are mounted with the BPOP wax record, and the wax record is removed, an anteroposteriorly and mediolaterally uniform space remains between both casts.

To interdigitate the teeth of the upper and lower models under this condition, a movement that is almost vertical to the occlusal plane of either model is necessary. If the models are allowed to make occlusal contact based on a hinge movement (like in a conventional articulator), the incisors and premolars make contact, and the molars will not occlude.
Therefore, to reproduce the occlusal relationship in the intercuspal position using an articulator, a mechanism that allows the upper model to be moved perpendicular to the occlusal plane is needed (Fig. 8).

![Vertically movable upper mounting ring.](image)

The procedure for occlusal analysis and occlusal equilibration in patients with TMDs is as follows.

1. After patient’s medical history has been taken and a clinical examination has been performed, an HOP record is taken using a polysiloxane material while the patient is in an upright position. While the HOP is being recorded, the patient is asked to close his or her mouth so as to achieve maximum intercuspsation and to hold that position until the material has set (approximately one minute). An anterior bite plate is then fabricated directly in the mouth using a self-curing acrylic resin material; the vertical dimension should be sufficient to produce a 1.0 mm-jaw separation at the second molars. The plate should cover the upper six anterior teeth and both first premolars. The occlusal surface of the plate should be flat and perpendicular to the mandibular incisors to allow free movement in all directions. The bite plate should be worn until the severity of symptoms is resolved, usually after one or two weeks; the bite plate is worn all day except while eating, speaking, or brushing one’s teeth. If the opening limitation and/or pain are not resolved using a bite plate, other symptomatic therapies, such as pharmacotherapy or physical medicine, may be necessary. Once the patient can freely open his or her mouth without pain, the treatment can proceed to the next step.

2. A BPOP record is then taken using the same material as that used to record the HOP. After the bite plate has been worn for 5 minutes, during which time the patient is asked to tap or slide his or her lower anterior teeth against the plate (muscular conditioning), the plate is removed and the recording material is applied to the occlusal surface using a syringe; the patient is then instructed to close his or her mouth until tooth contact is made and to hold that position. A BPOP wax record is then made using a wax registration material (Bite Wafer; Kerr USA, Romulus, MI, USA) and the same procedure mentioned above. The anterior part of the wafer should be cut off and softened in 60 °C water. After muscular conditioning using the bite plate has been performed, the softened wafer is placed on the premolars and molars on both sides. The patient is then instructed to bite the wafer gently, without biting through the wafer. The intercondylar distance is then measured using a caliper. Then, the distance between the condylar point and the lower incisal point in the occluded position is measured.
3. The intercondylar distance of the articulator is set to the same value as that of the patient. The gradation for zero on the scales on both sides indicates an intercondylar distance of 13 cm (the distance between the outsides of both condylar spheres). Therefore, when the condylar distance of the patient is 12 cm, the articulator should be reduced medially by 5 mm on each side. The articulator is then inverted, and the incisal point is marked on the mandibular cast table at a point the same distance from the condylar ball of the articulator as present in the patient. The lower cast is attached to the lower member of the articulator. The upper cast is then attached to the upper member using the BPOP wax record. At this time, a sheet of wrapping film is placed on the base of the upper cast, and some plaster is poured into the mounting ring of the upper member. The film will separate between the base of the upper cast and the poured plaster. Once the plaster has set, the film is removed and the upper cast and the poured plaster are adhered in place using $\alpha$-cyanoacrylate adhesive material. This mounting method prevents dimensional changes in the mounting plaster and subsequent inaccuracies in the articulated casts.

4. Next, the occlusal discrepancy is examined. Both condylar rods of the articulator are exchanged with the analyzing rods. Recording discs attached to graph papers are then inserted into the condylar shaft. An HOP polysiloxane record is placed between the upper and lower casts and the positions are tattooed on the graph papers using colored occlusal paper with recording needles. In the same manner, the BPOP is tattooed on the same graph paper using a different color occlusal paper. The difference between the HOP and the BPOP and the direction from the BPOP to the HOP are recorded (Fig. 9).

![Fig. 9. Use of the articulator as an occlusal position analyzer.](image)

The analyzing rods are then replaced with the condylar rods, the BPOP wax record is placed between the upper cast and the lower cast, and all the adjustable parts are set. The BPOP wax record is then removed, and the upper cast is moved downward until tooth contact has been made. Premature occlusal contact is located on the casts by marking and pulling an occlusal tape (Occlusion Foil; Coltene/Whaledent GmbH/Co. KG, Langenau, Germany). One of several therapies (selective tooth grinding, prosthodontic and orthodontic treatments, or orthognathic surgery) may then be selected for the occlusal equilibration of the BPOP, according to the degree of the discrepancy between the HOP and the BPOP.
5. The patient should be sitting in an upright position without a headrest support. The bite plate is worn in the patient’s mouth. After muscular conditioning has been performed by tapping and sliding the lower anterior teeth against the plate for 5 minutes, the plate is removed from the patient’s mouth. The operator holds Miller ribbon holders (Buffalo Dental Manufacturing Co., Inc. Brooklyn, New York, USA) for the right and left quadrant recordings with red occlusal foils and asks the patient to close his or her jaw until tooth contact, then to hold that position. True occlusal contact should be confirmed by pulling the foil laterally. The premature contact should be compared with the marked contact on the casts of the articulator.

6. The premature contact on the cast is removed using a small pear-shaped carbide bur with a slow speed handpiece. The ground spot is marked with a pencil. The incisal pin of the articulator is then removed from the upper member and the articulator is closed to make occlusal contact. The incisal pin must be removed to enable a hinge movement of the articulator, because during mandibular closure, the condition in which the condyle is suspended in the capsule changes to a condition in which the condyle is contracted against the disc and the eminence, resulting in a hinge movement. The location where the next premature contact appears should be confirmed by marking and pulling with an occlusal foil. The contacts should then be recorded.

7. The patient is then reclined and the premature contact is ground down using a diamond point of a similar size and shape and a high speed handpiece. Then, the patient is raised to an upright position and the bite plate is placed in the mouth. After muscular conditioning has been performed, the patient is asked to close his or her mouth until tooth contact has been made and to mark the premature contact with an occlusal foil. If the new contact is located at the same position as on the articulator, the articulator is regarded as precisely reproducing the BPOP and it can be reliably used in the following steps during future appointments. After the new contact has been confirmed, the contact is ground down. The impressions of the upper and lower dental arches are taken in the mouth. After the patient wears the bite plate in the mouth for 5 minutes, the bite plate is removed, and a BPOP wax record is then taken for the next appointment.

8. New upper and lower casts are made and attached to the articulator in the previously described manner. Steps 6 and 7 are repeated. The occlusal adjustments are continued in the mouth by referring to the contacts marked on the casts. The occlusal adjustment is completed by confirming the occlusal contacts on the premolars and molars on both sides of the casts attached to the articulator and in the mouth. Generally, two types of premature occlusal contact occur in TMD patients: one is a premature contact at the most posterior tooth with the occlusion opened anteriorly (often seen in young patients), the other is a premature contact at the anterior tooth with the occlusion opened posteriorly (often seen in elderly patients). The former type should be treated using occlusal adjustments. The latter should be treated by restoration of the teeth or using a fixed, or removable prosthesis. After completing the occlusal equilibration, HOP and BPOP polysiloxane records are taken and both positions are confirmed to be consistent with each other on the articulator.

5. Possible mechanism of Tinnitus related to TMD

Table 3 indicates that the side affected by TMD coincides with the side affected by tinnitus. Figure 10 shows the shifts recorded on both condylar areas of the articulator in case number...
9 shown in Table 3. This case has been previously reported (Torii & Chiwata, 2007). The patient complained of pain in her right ear and TMJ as well as tinnitus in her right ear. The shift on the right side was larger than that on the left side. The shifts noted after the resolution of her symptoms indicated that a displacement of the mandible from its correct position (muscular position) to a deflected position (habitual occlusal position) existed before treatment.

![Recorded mandibular shift from the habitual occlusal position to the correct occlusal position.](image)

Fig. 10. Recorded mandibular shift from the habitual occlusal position to the correct occlusal position. In this case, the patient was diagnosed as having Ménière disease by two independent otorhinolaryngologists and was treated with isosorbide, etc. However, when the drugs failed to improve her severe symptoms, she visited our clinic complaining of pain in the right TMJ region. An occlusal analysis was performed using dental models mounted on an articulator after relieving the painful symptoms using appliance therapy, and an occlusal discrepancy was identified. Occlusal equilibration of the BPOP was then performed (see Table 3).

This displacement seems to have distorted the normal relationship between the disc and the condyle in the TMJ, since the correct relationship between the disc and the condyle is thought to be fundamentally established during the development of the function of the masticatory muscles (with regard to the muscular position, that is the BPOP) before tooth eruption. On the other hand, the existence of an occlusal discrepancy between the HOP and the BPOP means that as the mandible voluntarily closes, the elevator muscles require additional activity to adapt the mandible from the BPOP to the HOP as multiple teeth come in contact with a stable position during the change from isotonic muscle contraction to isometric. This condition may cause muscle tension, muscle fatigue and muscle spasm, resulting in muscle pain. Therefore, when considering the mechanism of tinnitus related to TMD, the displacement of the TMJ and muscle tension or spasm should be taken into account. First of all, regarding the displacement of the TMJ, Pinto found that a tiny ligament connects the neck and anterior process of the malleus to the medioposterosuperior part of the TMJ capsule, the interarticular disc, and the sphenomandibular ligament. He also described that the fibrous layer of the tympanic membrane seemed to be continuous with this tiny ligament and that the tiny ligament had an embryologic origin common with that of the malleus and incus (Pinto, 1962). However, Komori et al. performed an anatomical study and concluded that no evidence was found to support the “tiny ligament theory,” as the ligament did not show any visible movement in response to a given strain. (Komori et
al., 1986) Rodriguez-Vásquez et al. reported that the discomalleolar ligament is a capsular structure similar to an intrinsic ligament that is formed by the association of capsular fibers coming from the posterosuperior and medial end of the articular disc. They then described that these fibers belong to the upper lamina of the bilaminar zone (Rodríguez-Vásquez et al., 1998). Eckerdal observed the petrotympanic fissure and reported that the dimensions of the fissure and the morphologic arrangement of the intrafissural and extrafissural soft tissues do not corroborate the theory that forces generally may be conducted through the fissure from the TMJ to the ear ossicles (Eckerdal, 1991). Parker and Chole reported that although the discomalleolar ligament and the mallear portion of the sphenomandibular ligament exist and could conceivably transmit mechanical energy to the malleus, they can not account for high frequency subjective tinnitus from local perturbations of the position of the malleus (Parker & Chole, 1995). In summary, the direct transmission of force arising from the displacement of the TMJ seems unlikely to be capable of causing ear symptoms. However, Johansson et al. described in their radiographic and histologic study that a medially displaced TMJ disc may exert traction or friction or may rub against the auriculotemporal nerve trunk. Since the auriculotemporal nerve innervates the tympanic membrane, the irritation of the nerve medially or posteromedially to the TMJ, before the nerve divergence, may give rise to symptoms from the ear region (Johansson et al., 1990). Williamson reported that noxious pain stimuli to the peridiscal tissues (similar to condyle or disc displacement) could result in the constriction of the internal auditory and posterior auricular arteries. Such constriction may result in a decrease in the blood supply to the middle and inner ears. Thus, the displacement of the TMJ may cause tinnitus (Williamson, 1990). Myers reported that beginning with the appearance of a reciprocal clicking in the TMJ, the disk is forced anteromedially with full intercuspation of the teeth. This change is accompanied by the stretching of the posterior laminae and the stretching or tearing of the lateral collateral ligament. As the laminae are stretched, vascular soft tissue is pulled forward and tends to become trapped between the head of the condyle and the roof of the fossa. This injury recurs each time the teeth are intercuspated, with displacement of the disk. This causes a chronic inflammation with extension of the inflammatory area, especially along three pathways. Among these three pathways, the second pathway is located in the carotid sheath area. In the case of a displaced disk specimen, the arteries, veins and nerves are impossible to dissect because of the fibrous material adhering to them and the surrounding structures. Tension would thus be produced on the whole carotid sheath in response to any movement of the neck or jaw. This tension of the carotid sheath fibrosis would be transmitted to the jugular foramen, making the saccus endolymphaticus unable to expand to perform any possible regulation of the endolymphatic pressure; as a result, the increased endolymphatic pressure on the hair cells of the cochlea could cause tinnitus (Myers, 1988).

Regarding masticatory muscle fatigue or tension, Fuchs observed the masticatory muscle activity during sleep at night and concluded that the results of the observed activity supports the suggestion that dysfunctional patients without biteplates have a higher muscular activity during sleep than healthy persons (Fuchs, 1975). In addition, Mao et al. reported that the near absence of type II A fibers (fast-contracting, fatigue-resistant fibers) from human jaw muscles seems to suggest that these fibers may be readily susceptible to fatigue during sustained efforts at stronger force levels, since fast, fatigue-susceptible (type IIB) fibers must be recruited at these levels (Mao et al., 1993). Myrhaug reported that bite deformity is known among otologists to be a cause of tinnitus, and this type of tinnitus
arises as an autogenous vibration in the sound-conducting system, possibly because of
tremor or myoclonus (myorhythmia) in the tensor muscle of the middle ear. This condition
represents a fatigue reaction resulting from prolonged irritation and stress of the muscles
innervated by the trigeminal nerve (masticators). This form of fatigue reaction apparently
only affects the small muscles in this group because of their anatomical relationship. Hence,
the two tensor muscles are in a vulnerable position in bite anomalies (Myrhaug, 1964).
Koskinnen et al. and Parker and Chole were skeptical about Myrhaug’s theory (Koskinnen
et al., 1980; Parker & Chole, 1995). However, Watanabe et al. reported that tinnitus occurred
whenever a certain mimic facial muscle contracted voluntarily or involuntarily. Under the
operating microscope, the contraction of the stapedial muscle synchronous with the
contraction of the mimic facial muscle was observed. The tinnitus disappeared completely
immediately after the tendon of the stapedial muscle was sectioned (Watanabe et al., 1974).
Myrhaug described a matter similar to that in the report of Watanabe et al. in which
contractions of the tensor tympany muscle were viewed directly under a microscope during
voluntary stimulation by grinning and clenching movements of the jaw.
Taking into account the facts that myofascial pain, disc displacement, tension-type
headaches and tinnitus were resolved after occlusal equilibrating treatment for the BPOP
(Table 3), tinnitus related to TMD may be caused by the spasmodic synkinesis of the tensor
tympani and the stapedial muscle with muscle spasm of the masticatory muscles (including
relative muscles), or by a decrease in the blood supply to the middle and inner ear with
noxious stimuli to the peridiscal tissue, or by increased endolymphatic pressure with injury
to the peridiscal tissues of the TMJ.

6. Conclusion
First, an otolaryngologist should confirm the absence of any significant or detectable
auditory, genetic, drug-related, or trauma-related causes of tinnitus. Then, in cases with
suspected TMD, the patient should be referred to a dentist with experienced treating TMD.
The dentist should perform the examination based on the RDC/TMD (Dworkin &
LeResche, 1992), with particular attention to the occlusal discrepancy on an articulator.
Occlusal equilibration of the BPOP should be performed to resolve the main symptoms of
TMD (TMJ clicking, TMJ pain, muscle pain and difficulty in mandibular movements), since
the mechanism of tinnitus related to TMD remains uncertain. Patients are fortunate if the
tinnitus related to TMD disappears as a result of the treatment. In most cases, tinnitus
related to TMD is likely to disappear together with the other TMD symptoms after the
occlusal equilibration of the BPOP. In the absence of TMD, further medical treatments (e.g.,
psychiatric, neurological and otolaryngological) may be indicated.

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Up to Date on Tinnitus encompasses both theoretical background on the different forms of tinnitus and a detailed knowledge on state-of-the-art treatment for tinnitus, written for clinicians by clinicians and researchers. Realizing the complexity of tinnitus has highlighted the importance of interdisciplinary research. Therefore, all the authors contributing to the this book were chosen from many specialties of medicine including surgery, psychology, and neuroscience, and came from diverse areas of expertise, such as Neurology, Otolaryngology, Psychiatry, Clinical and Experimental Psychology and Dentistry.

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