Personalized needle modification for CT-guided percutaneous infrrazygomatic radiofrequency ablation of the maxillary nerve through the foramen rotundum in order to treat V2 trigeminal neuralgia

Background: The computed tomography (CT)-guided radiofrequency ablation (RFA) of the maxillary nerve (V2) via foramen rotundum (FR) approach has been reported to offer the highest rates of pain relief in V2 trigeminal neuralgia (TN). However, the access to FR may be obstructed by the greater wing of the sphenoid bone.

Objectives: We report on an optimized CT-guided percutaneous infrrazygomatic of maxillary nerve through the foramen rotundum (FR) to treat V2 trigeminal neuralgia (TN) using personalized RFA needles based on patient’s individual CT-image parameters.

Patients and methods: 176 patients with isolated V2 TN were included. If the entry of the percutaneous needle into the FR canal was blocked by the greater wing of the sphenoid bone, straight RFA needles was bent at the tip with an angle $\alpha$ (the angle between the straight line from the external opening of FR to the skin entry point and the long axis of the FR canal). The maxillary nerve RFA was performed after confirmation with electrophysiological tests. Pain relief in the V2 territory and TN recurrence rate were followed for up to 60 months.

Results: Fifty-two patients (29.55%) required needle bending. The maxillary nerve thermal RFA resulted in analgesia in the V2 territory without affecting the V1 or V3 zone. TN recurrence rate at 6, 12, 24, 36, 48 and 60 months was 2.55%, 7.64%, 17.20%, 24.41%, 30.28% and 33.77%, respectively.

Conclusion: The personalized needle modification technique for maxillary nerve RFA through FR is safe and effective to treat V2 TN.

Keywords: trigeminal neuralgia, maxillary nerve, foramen rotundum, radiofrequency ablation

Introduction
Trigeminal neuralgia (TN) is a severe painful condition characterized by touch-evoked unilateral brief shock-like paroxysmal pain in one or more divisions (V1, V2, or V3) of the trigeminal nerve.\textsuperscript{1,2} TN pain can be very refractory to a number of therapies, including medications, nerve blocks, and microvascular decompression surgeries.\textsuperscript{3} Percutaneous radiofrequency thermocoagulation or ablation (RFA) has been reported to offer the highest rates of pain relief in TN.\textsuperscript{4} A number of research groups, amongst them ourselves, have reported that the computed tomography (CT)-guided RFA of the maxillary nerve (V2) via foramen rotundum (FR) approach can achieve selective analgesia to treat V2 TN.\textsuperscript{5,6} FR is a bony canal of 3 mm in diameter and 3–6 mm in length. The long axis of the canal runs in an obliquely
posterolaterally to anteroinferior direction, which may form an angle with the percutaneous needle trajectory. In addition, the external opening of the FR is located in the upper portion of the pterygopalatine fossa. Access to FR may be obstructed by the greater wing of the sphenoid bone. Under these circumstances, a straight RFA needle may not be able to reach FR, yielding a less satisfactory ablative effect. To address this issue, we have designed a modified RFA needle technique based on individual CT scans to improve the access of the FR canal for the treatment of V2 TN.

Methods
Design: retrospective chart review
Patient characteristics
From November 2012 to March 2017, 176 patients with primary isolated V2 TN were included (Figure 1). This study was conducted in accordance with the Declaration of Helsinki and was approved by the Ethics Committee of the First Affiliated Hospital of Jiaxing University. All cases were conducted in the same institution. The diagnosis of TN was confirmed according to the International Headache Society guidelines. Potential secondary causes of TN were ruled out. There were 74 males and 102 females patients with a mean age of 66.9±9.84 years (range 32–101 years). Seventy-seven patients had TN on the left side and 99 patients on the right side. The mean duration of TN pain was 2.4 years (range 0.5–13 years) and the mean pre-procedural numerical rating scale (NRS) pain score was 7.1 (range 6–10). All patients failed to respond to multiple pharmacological treatments and responded to diagnostic maxillary nerve block with >50% pain relief. Patients were counseled with risk and benefits of the RFA procedure and all patients signed a written informed consent (Table 1).

CT-guided percutaneous RFA of V2 nerve through FR
Patient was positioned in the supine position and under monitored anesthesia care. Access of FR was achieved as we described previously with individual modification. In short terms, a positioning grid was placed over the cheek of the affected side (Figure 2) and a semi-coronal CT scan was conducted. The specific CT-image frame (Figures 3, 5 and 6) that simultaneously captures the FR and its external and

Figure 1 Flow diagram.
internal opening was used to establish the needle trajectory. The built-in ruler of the CT instrument was used to measure the distance from the needle entry point to the external opening of FR. Using the CT instrument's image processing software, we drew a line from the midpoint of the FR canal (puncture target) against the lateral wall of the maxillary sinus to the

Table 1 Patient baseline characteristics (n=176)

<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>Time from diagnosis of V2 (year)</th>
<th>TN</th>
<th>NRS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male</td>
<td>Female</td>
<td></td>
<td>Left</td>
</tr>
<tr>
<td>66.9±9.84</td>
<td>74</td>
<td>102</td>
<td>2.4±1.4</td>
<td>77</td>
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</table>

Figure 2 The patient was positioned supine on the CT-fluoroscopy table. A roll placed under the shoulders and head taped at the chin to prevent unintentional movement. A positioning grid was placed over the cheek of the affected side.

Figure 3 Design of the RFA needle insertion route. (A) Along the lateral wall of the maxillary sinus, line 1 is drawn from the mid of the foramen rotundum (FR) canal to the skin entry point. Measure the needle insertion depth (distance, here showed 6.44 cm) and the puncture angle (the angle between line 1 and the sagittal plan, β=43). Since line 1 passes the greater wing of the sphenoid bone (arrow), a needle-bending technique is required. (B) Line 2 is drawn from the external opening of FR to the skin entry point and the distance (6.44 cm) measured. Puncture angle β1 between line 2 and the sagittal plane was about 43. Bending angle α between line 2 and the long axis of FR canal was about 151.
surface of the skin (needle entry point). If line 1 passes through the ipsilateral greater wing of the sphenoid bone (Figure 3A), line 2 is drawn from the external opening of the FR to the skin entry point. The angle \( \alpha \) between line 2 and the long axis of the FR canal (the degree at which the RFA needle is to be bent) and the angle between line 2 and the sagittal plane (puncture

**Figure 4** Making of a personalized RFA needle. Inserting the 10 cm 22 gauge straight RFA probe (upper panel) into a 16G sterilized piercing needle for about 5–8 mm (middle panel) and bend the RFA at the angle \( \alpha \) (bottom panel) based on the CT imaging measurements.

**Figure 5** The personalized RFA needle curved anterosuperiorly to pass the greater wing of sphenoid and reached the external opening of the FR canal (A). The personalized RFA needle was then turned inferoposteriorly to enter the FR canal (B).
angle $\beta_1$) were measured. The length of the segment of line 2 serves as puncture depth (Figure 3B). Then, the tip of a 10 cm 22 gauge straight RFA probe is inserted into a 16G sterilized piercing needle for about 5–8 mm and the RFA probe is bent to the degree of $\alpha$ (Figure 4). In this way, the RF probe becomes the patient’s individualized RFA needle. The skin and subcutaneous tissues were anesthetized with 2 mL of 1% lidocaine using a 27G intradermal needle. Under CT guidance, the personalized RFA needle was inserted infraygomatically and advanced based on predetermined parameters, including angle, path, and depth to reach the external opening of FR anterosuperiorly (Figure 5A). The needle was then turned inferoposteriorly using the individually made needle curvature to enter the FR canal (Figure 5B). If line 1 does not pass through the sphenoid bone, it can be set as the puncture path using the unmodified straight RFA needle at the angle between line 1 and the sagittal plane (puncture angle $\beta_2$) (Figure 6).

After confirming that there was no evidence of blood, cerebral spinal fluid, or paresthesia, a sensory test was performed by stimulating the RFA probe at 100 Hz with pulse width of 500 msec to generate paresthesia concordant to the patient’s usual TN pain at 0.1–0.5 V. A motor test was then performed by stimulating the probes at 2 Hz and 0.1–0.5 V to confirm that the probe was not in proximity to other adjacent nerves. 0.5 mL of 1% lidocaine was then injected 2 mins prior to ablation. Subsequently, continuous RFA was performed at 90°C for 120 s under intravenous propofol anesthesia. During the treatment, blood pressure, heart rate, electrocardiogram, and blood oxygen saturation were closely monitored. Rescue medications and equipment such as atropine, lidocaine, epinephrine, and airway devices were made immediately available. Oxygen was given via the nasal cannula. If the patient’s blood pressure rose by more than 20% of the baseline, Urapidil was administered at increments of 12.5 mg intravenously. After the procedure, the patient was transferred to the post-procedure recovery room where the NRS score was evaluated and recorded. The patient’s vital signs were monitored for at least four additional hours before being discharged. The intraoperative and postoperative complications were recorded and the immediate and long-term outcomes were evaluated during follow-up.

**Outcomes**

The NRS score before and after the procedure was assessed. The bending angle ($\alpha$ value), puncture depth, puncture angle ($\beta$ value), puncture time, and number of CT scans for the RFA were documented. V2 TN recurrence rate, defined as number of patient with returning V2 TN after RFA/total number of follow-up patients, was tracked for up to 48 months. Complications and adverse effects were recorded.

**Statistical analysis**

Data were presented as mean±standard deviations and analyzed using SPSS 16.0 software (SPSS, Inc., Chicago, IL, USA). Comparison of continuous data was performed by Student’s t-test. P-values lower than 0.05 were considered statistically significant.

**Results**

Infraygomatic percutaneous access to the FR canal was obstructed by the greater wing of the sphenoid in 52 of the 176 (29.55%) patients recruited in this study. The RFA needle was bent at an angle of 14–31º (19.6±4.3º) to facilitate the entry into the FR canal. In the other 124 patients, the RFA needle route was not blocked by the sphenoid wing so the access to the FR canal was gained using a straight probe. Sensory test at 0.1–0.5 mA produced paresthesia in the V2 territory. All patients obtained more than 95% V2 TN pain reduction in NRS score after RFA. There was no numbness in the territory of V1 or V3. The puncture depth, angle, and puncture time of the bent needle group and the straight needle group are compared in Table 2. The puncture
time and number of CT scans in the bent needle group were significantly greater than those in the straight needle group ($P<0.05$). Patients with V2 TN recurrence received repeated RFA. V2 TN total recurrence rate (Table 3) was 2.55%, 7.64%, 17.20%, 24.41%, 30.28%, and 33.77% at 6, 12, 24, 36, 48, and 60 months, respectively, while the recurrence rate of intervention group was 2.2%, 6.8%, 20.45%, 28.2%, 30%, and 33.3%. There was no statistical difference between the two groups.

There were no intracranial hemorrhage, trigeminal nerve V1 or V3 nerve injuries, cerebrovascular accidents, infections, or deaths recorded. All patients in the study suffered from varying degrees of numbness in the original pain area after treatment. There were 23 cases (non-intervention group/intervention group 19/4) of facial hematoma which resolved within 3 days after treatment with ice-bag compressions. Thirty-six patients of 167 (20%) required an average dose of $37.5$ mg Urapidil to help control an increase in blood pressure of more than 20% of the baseline. No significant decrease in heart rate occurred during the procedure.

**Discussion**

We reported a personalized needle modification technique for percutaneous infrrazygomatic CT-guided maxillary nerve RFA to treat isolated V2 TN. TN is one of the most common causes of facial pain and the treatment requires a multidisciplinary approach.\textsuperscript{10} For patients who could not tolerate or are not willing to undergo open craniotomy microvascular decompression surgery, percutaneous RFA provides sustained pain relief.\textsuperscript{9,11–13} A systematic review evaluated 166 studies and concluded that, comparing to glycerol rhizolysis, balloon compression of the trigeminal ganglion, and stereotactic radiosurgery, the RF thermocoagulation offers the highest rate of complete pain relief.\textsuperscript{4} The key for a successful RFA is the accurate placement of RFA electrode alongside the affected nerve. Traditionally, TN can be treated with RFA of the trigeminal Gasserian ganglion which can be accessed through the foramen ovale (FO) using the Härtel technique.\textsuperscript{14} CT fluoroscopy can be used to improve the visualization and access of FO.\textsuperscript{15,16} However, the FO approach is not optimal to treat sub-branch TN. It is often difficult to distinguish between V2 and V3 or V1 nerve once the needle passes the external opening of FO.\textsuperscript{5} Consequently, it often requires frequent adjustment of position of the needle to identify the V2 nerve. Performing these adjustments may increase the risk of dural puncture, infection, headache, and intracranial bleeding.\textsuperscript{17} Unintentional
lesioning of V3 or V1 may result in masseter muscle weakness or paralysis, diminished or absent corneal reflex, and keratitis. 9,13,18,19 Duration of pain relief for isolated V2 TN was significantly shorter than for V3 TN using the FO RFA technique20 likely because the V2 nerve is less successfully ablated using the FO approach. To address these concerns for isolated V2 TN, we amongst others have developed the FR approach at the pterygopalatine fossa to selectively block the V2 nerve.5,6,21 The V2 trigeminal nerve begins at the Gasserian ganglion and passes through the FR to exit the cranium.7 Blocking the V2 nerve at this peripheral location reduced risks of abovementioned complications related to the FO technique.5 As we performed more cases of V2 block/RFA using the FR approach, we have encountered a new challenge due to anatomical variations among individual patients. In some patients, percutaneous infrazygomatic access of FR was obstructed by the greater wing of the sphenoid bone (2). To overcome this obstacle, we have adapted the bent needle technique based on individual CT measurements and gained access to the FR canal. This individualized bent needle technique (with personalized angle for each patient) was required in 52 of 176 (29.55%) patients with isolated V2 TN. The average bending angle was 14–31° (19.6±4.3°), suggesting that a commercial RF probe bent at 20° would have a great potential for percutaneous infrazygomatic FR access under CT-guidance. The procedure time required by the bent needle technique was about 5 mins longer than using the straight needle (23.53±6.14 mins Vs 19.42±4.03 mins, P<0.05). On the other hand, the individualized bent needle technique allowed precise needle positioning in the FR canal and subsequently a highly selective RFA of the V2 nerve. The recurrence rate of V2 TN in our study at 24 months was 17.20%, lower than that (28.3%) reported in RFA of Gasserian ganglion,12 suggesting again that a selective V2 RFA is preferable to the ganglion RFA in treating V2 TN.

Traditionally, pulsed RF (PRF) has been used to treat TN.22 PRF is a nondestructive ablation that generates short bursts of RF current at 42°C with long pauses between bursts to allow heat to dissipate in the target tissue.23 The advantage of using PRF is probably due to the fact that the smaller demyelinated C fibers are selectively lesioned under a lower temperature without motor ablation. Chua et al recommended performing PRF prior to thermal RF for the purpose of avoiding the disturbing of sensory paresthesia and masseter paralysis.24 However, a recent randomized controlled trial comparing conventional thermal RF with PRF showed that conventional RF provided significant pain reduction and patient satisfaction improvement in patients with idiopathic TN whereas PRF failed to achieve these effects.25 Another randomized prospective study reported that better outcome was observed in the thermal treatment as compared to pulsed RF treatment of the Gasserian ganglion.26 Based on these findings, we performed the selective V2 RFA at 90°C for 120 s. All patient had immediate TN pain relief after the procedure and the V2 TN recurrence rate at 36 months in our cohort was lower than that reported in the studies using 75°C C for 4 mins (24.41% vs 36%),21 indicating that the temperature utilized for RFA may play an important role in patients with TN. Another study indicated that different temperature may cause different cardiovascular responses during RFA of trigeminal nerves.27 Significant elevation of the mean arterial blood pressure was observed in all patient studied (n=48) during RFA of the trigeminal ganglion using the FO approach. The correlation between the RFA stimulus and magnitude of the pressor was positive when the temperature used for RFA was below 75°C but it became negative when the temperature was above 75°C. These findings may indicate that responses of the trigeminocardiac reflex during trigeminal RFA are temperature dependent.28 We performed V2 RFA at 90°C and about 20% of our patients did require Urapidil to help control the elevation of blood pressure during the procedure. However, no significant tachycardia or bradycardia occurred during the RFA. We want to emphasize that vasoactive medications should be made immediately available during RFA of trigeminal nerves.

**Conclusion**

Nearly 30% of V2 TN patients in our cohort required the personalized needle-bending technique for percutaneous RFA of the maxillary nerve due to the obstruction of FR

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**Table 3 The V2 TN recurrence rate (patients or recurrence rate of intervention group)**

<table>
<thead>
<tr>
<th>Follow-up time period (month)</th>
<th>6 months</th>
<th>12 months</th>
<th>24 months</th>
<th>36 months</th>
<th>48 months</th>
<th>60 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of follow-up patients</td>
<td>157 (45)</td>
<td>157 (44)</td>
<td>157 (44)</td>
<td>127 (39)</td>
<td>109 (30)</td>
<td>77 (21)</td>
</tr>
<tr>
<td>Number of patients with V2 TN recurrence</td>
<td>4 (1)</td>
<td>12 (3)</td>
<td>27 (9)</td>
<td>31 (11)</td>
<td>33 (9)</td>
<td>26 (7)</td>
</tr>
<tr>
<td>V2 TN recurrence rate (%)</td>
<td>2.5 (2.2)</td>
<td>7.6 (6.8)</td>
<td>17.2 (20.45)</td>
<td>24.41 (28.2)</td>
<td>30.26 (30)</td>
<td>33.77 (33.3)</td>
</tr>
</tbody>
</table>
access by the ipsilateral greater wing of the sphenoid bone. We have optimized the maxillary RFA through the FR approach with precise localization of the entry of the FR canal. Highly selective thermocoagulation (90°C for 120 s) of the V2 nerve using this personalized technique is a safe and effective treatment and provides sustained pain relief and lower recurrence rate of isolated V2 TN pain.

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Author contributions
All authors contributed to data analysis, drafting or revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

References