Reduced Depth Technique with the Posterior Superior Alveolar Block

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Abstract

Purpose: The posterior superior alveolar (PSA) block is one of many techniques used to provide profound anesthesia for invasive dental procedures. This technique has a long history, with a high success rate, but is not without complication risks. The purpose of this study was to determine if pulpal anesthesia of the maxillary second molar could be achieved using a posterior superior alveolar block with a reduced depth of penetration of 10 mm compared to the current suggested depth of 16 mm.

Methods: Using a cold refrigerant, a thermal test was conducted using the buccal surface of a maxillary second molar of 43 participants. Positive neural responses were obtained from 100% of the participants (n=43) during the pretest. Each participant received a posterior superior alveolar block using a short (20mm), 27-gauge needle with the penetration depth reduced to 10mm. Post-test neural responses of these molars were evaluated using same cold thermal test technique.

Results: Study results demonstrated that the reduced depth technique for the PSA block was successful in 88% (n=38) of the participants; pulpal anesthesia of the maxillary second molar had been achieved. Furthermore, there were zero positive aspirations and zero hematomas observed in the participants.

Conclusion: The reduced needle depth technique showed promise in achieving desired results of pulpal anesthesia coupled with decreasing risk and complications associated with the PSA block. Additional blinded, randomized clinical studies are recommended to achieve evidence-based support for this reduced depth PSA block technique.

Keywords: local anesthesia, nerve blocks, pulpal anesthesia, clinical education

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Introduction

The use of local anesthesia is essential to facilitate many procedures in the dental field. There are a variety of target sites and techniques used to achieve patient comfort. The posterior superior alveolar (PSA) block is used to achieve pulpal and soft tissue anesthesia to the maxillary third molars, second molars, as well as the first molars, with the exception of the mesiobuccal root in some cases.\(^1\) When the middle superior alveolar (MSA) nerve is not present, as is the case with approximately 72% of the general population, the PSA nerve innervates the mesiobuccal root of the maxillary first molar and a PSA block will provide complete pulpal and soft tissue anesthesia here as well.\(^1\) The PSA nerve is the target area for the PSA block injection, which requires access via the height of the mucobuccal fold just distal to the apex of the maxillary second molar. The current recommended technique for accessing the PSA nerve for an “average-sized” adult is a depth of 16mm.\(^2\) There is allowance for the modification for “most smaller-skulled” patients to a penetrating depth of 10-14 mm.\(^2\) Student dental professionals are taught and tested using this recommended practice in their professional programs and on national and clinical board licensing examinations.\(^2,5\) This injection technique often evokes anxiety in some clinicians as there is no osseous contact alerting the clinician that the proper depth has been reached, thus over insertion is a possibility.\(^4\) The pterygoid plexus of veins is located in this area and the inadvertent penetration of this plexus and/or nearby maxillary artery can result in unpleasant complications for the patient.

The PSA block has a 3.1% positive aspiration rate, the second highest in the oral cavity, second only to the inferior alveolar block.\(^3\) The risk of causing a large hematoma often deters clinicians from utilizing this nerve block, while instead choosing a less suitable supraperiosteal injection, requiring multiple needle penetrations to the patient. A variety of PSA techniques have been explored, including a study by
One conservative insertion technique has been suggested in the literature in an attempt to minimize these risks.\textsuperscript{2,7} Given the depth of the target area of the PSA nerve as it exits the posterior superior alveolar foramina within the infratemporal fossa, it has been theorized that a shorter needle depth is sufficient for adequate anesthetic delivery while being far enough away from the pterygoid plexus of veins and maxillary artery to avoid puncture and hematoma risk.\textsuperscript{2,4}

Minimal literature exists however, to validate efficacy and hematoma risk reductions while delivering a PSA block with a reduced needle depth insertion technique. The purpose of this study was to determine if pulpal anesthesia of the maxillary second molar could be achieved using a PSA block with a reduced depth of penetration of 10 mm as compared to the standard suggested depth of 16 mm while minimizing complication risks.

**Materials**

This pilot study used a quasi-experimental design in which a single pre-test measurement (O1) was taken followed by an intervention (X) and finishing with a post-test measurement (O2).\textsuperscript{8} Investigators assessed whether a reduction in needle depth of the PSA block resulted in achieving pulpal anesthesia of the maxillary second molar. Since only approximately 28\% of the population has an MSA nerve to innervate the mesiobuccal root of the maxillary first molar, the second molar was selected as the test tooth to be studied.\textsuperscript{1-4} Approval for this study was granted through the University of New Mexico’s Institutional Review Board and the Human Research Protection Office (HRPO). Students enrolled in the undergraduate and graduate dental hygiene programs were recruited to participate. Informed consent was obtained from all participants and preliminary screening for eligibility was completed. The screening process included a review of health history, vital signs, and intraoral screening. Any participants indicating an allergy to lidocaine, blood clotting conditions, pregnant, or those taking anticoagulant medications or any type of analgesic within the last 12 hours were excluded from further participation in the study.

The intraoral screening was performed to evaluate teeth #2 and #15 to ensure they met the study criteria. Participants were immediately excluded from the study if they were missing both maxillary second molars. Each molar was assessed individually for any confounding features. Any maxillary second molar which had an amalgam, composite, crown or bridge, a root canal, an implant, frank decay or visible signs of active infection including an abscess or fistula in the maxillary molar area was not used in the study. Participants satisfying all criteria of the screening had a digital periapical radiograph of the qualified tooth taken as a final evaluation to confirm there were no radiolucent areas or visible abnormalities.

A baseline neural response was obtained using a thermal test by applying a large cotton pellet with a refrigerant, 1,1,1,2 Tetraflouroethane (Endo-ice®, Coltène/Whaledent Inc; Cuyahoga Falls, Ohio), to the middle third of the buccal surface of the tested tooth. Investigators noted either a positive or negative response to the test. After confirmation of positive response, a cotton tip applicator with 5\% lidocaine topical anesthetic ointment was applied to the site of tissue penetration for 2 minutes. Participants were asked to close their mouth slightly, and shift their mandible towards the test side. They received the reduced depth PSA block using a 27-gauged, 20 mm short needle, angled 45 degrees posteriorly, 45 degrees superiorly and 45 degrees medially to the point of insertion. The needle was inserted at the height of the mucobuccal fold slightly distal to the second maxillary molar and advanced to a depth of 10 mm. All PSA blocks were completed either by the investigator or co-investigator. The left-handed investigator completed the PSA blocks used to test tooth #15, and was observed by the co-investigator. PSA blocks used to test tooth #2 were completed by the right-handed co-investigator and were observed by the left-handed investigator. Both investigators were present for each injection to ensure proper technique with the reduced needle depth for the PSA block was achieved.

Once it was agreed upon by the investigator/observer that the depth of 10 mm had been reached at the proper angle, the investigator/operator aspirated in two planes and administered one full cartridge (~1.8 mL) of lidocaine 2\% with 1:100,000 epinephrine. At 10 minutes, the thermal test was conducted again to assess the neural response of the test tooth. The same refrigerant and technique previously described was used. Investigators noted either a positive or negative response to the test for each subject with a negative response indicating pulpal anesthesia had been achieved.

**Results**

A total of 49 participants completed consent and enrolled in the pilot study however, after completing the screening process 6 participants were excluded as they either failed to satisfy minimum tooth requirements on tooth #2 or #15 or did not meet health history requirements. A total of 43 subjects, 39 females and 4 males, were eligible and participated in the study.

The pretest yielded a one hundred percent (n=43) “positive” baseline neural response when exposed to the refrigerant. Post-test results revealed an 88\% (n=38) negative response, indicating no neural response was felt and pulpal anesthesia had been achieved on the majority of participants. This compared to 12\% (n=5) of participants who still
indicated a positive response on the post-test and did not achieve pulpal anesthesia. Figure 1 illustrates pre-test and post-test results of neural response of the tested teeth.

Investigators evaluated the injections administered to tooth #2 and tooth #15 individually. Twenty subjects received the modified PSA injection on tooth #2 and twenty-three subjects received it on tooth #15, resulting in a 95% and 83% negative neural response to the post-thermal test, respectively. Table I and Figure 2 illustrate the results for teeth #2 and #15 individually.

A Fisher’s Exact test was performed against the null hypothesis statement that there was no difference between the two groups. The test revealed (p=0.219 and α-value of 0.05 ) that the null hypothesis could not be rejected. A Chi-squared test was also performed and confirmed previous findings. No positive aspirations or hematoma were observed as a result of the procedure.

Table I. Pulpal anesthesia achieved with reduced depth PSA technique for individual tooth

<table>
<thead>
<tr>
<th>Tooth Number</th>
<th>Number of injections</th>
<th>Pulpal anesthesia achieved using modified PSA</th>
<th>Pulpal anesthesia not achieved</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>20</td>
<td>19 (95%)</td>
<td>1 (5%)</td>
</tr>
<tr>
<td>15</td>
<td>23</td>
<td>19 (83%)</td>
<td>4 (17%)</td>
</tr>
</tbody>
</table>

Discussion

Historically, it has been taught that in order to achieve profound anesthesia, the proper depth of penetration for the PSA block is 16mm. This depth of penetration is widely accepted among clinicians and educators, and is considered the measurable standard on local anesthesia clinical board examinations providing the rationale that a control was unnecessary for the purposes of this pilot study.

The traditional PSA block has a success rate of 95%. Complications of the PSA block are well documented in the literature and may range from local minor irritation at the injection site, trismus, or hematoma to more severe complications such as paresthesia and potential permanent eye complications. This pilot study yielded an overall success rate of 88% in achieving pulpal anesthesia on the second molar, using the reduced depth technique. Additional successes of the study included no positive aspirations and no hematomas. Results of this study indicate that more conservative injection techniques could be explored to decrease the complication risks. Perhaps clinicians would then be less fearful of causing unsightly hematomas and utilize this effective nerve block to achieve profound anesthesia.

As hypothesized, the majority of participants achieved full pulpal anesthesia with the modified technique, however lack of randomization and the convenience sample used limits the generalizability of these results to a larger population. It is also recognized that the majority of participants were women, and while still considered to have an “average” size skull, it is accepted that women generally have a smaller skull size. This could lead clinicians to think that the success rate of the reduced depth technique was influenced; however, investigators believe that results from this study provide support of the effectiveness of anesthesia through a PSA block at a reduced needle depth.
side for purposes of providing the best viewing conditions of the needle position and depth. A study conducted by Khan et al. concluded that there was no difference in periodontal assessments based on the clinician's “handedness” and investigators believe the study results would have been replicated if only one investigator administered the injections on both sides of the mouth.

The limitations of a quasi-experimental pilot study are acknowledged. However, this quasi-experimental design was chosen intentionally for this pilot study to determine the logistical feasibility of conducting a larger randomized, blinded clinical study.

Conclusion

The reduced needle depth technique showed promise in achieving desired results of pulpal anesthesia coupled with decreasing risk and complications. Additional randomized, controlled, blinded clinical studies are recommended to achieve evidence-based support for the academic and dental communities to assess replacing the current recommended PSA block technique with the modified PSA block with a reduced needle depth.

Disclosure

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