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RESEARCH

Does articaine, rather than lidocaine, increase the risk of nerve damage when administered for inferior alveolar nerve blocks in patients undergoing local anaesthesia for dental treatment? A mini systematic review of the literature

P. Stirrup^{*1} and S. Crean²

Key points

Outlines the benefits and possible risks of using articaine rather than lidocaine.

Presents the results of a systematic search of the literature to determine the safety of articaine in inferior alveolar block anaesthesia.

Advises the profession following analysis of the results.

Aims This mini systematic review seeks to analyse the available literature and determine if a 4% articaine solution poses a greater risk of inferior alveolar and/or lingual nerve damage compared to that of 2% lidocaine, when administered for an inferior alveolar nerve block. **Results** After a mini systematic review of the published literature, seven suitable studies were identified: one double-blind random controlled trial (DBRCT) and six retrospective cohort studies. The DBRCT and two of the cohort studies concluded that 4% articaine poses no greater risk of nerve damage. The remaining four cohort studies suggested that caution should be exhibited when using a 4% local anaesthetic solution rather than a 2% solution. However, these studies also concluded that no evidence exists to explain the reasons for their results. **Discussion and conclusion** The included articles present no conclusive evidence to suggest that 4% articaine causes more nerve damage than 2% lidocaine, although some authors advise caution when using this agent. All studies conclude that further quality research is required, and it is therefore suggested that dental practitioners exhibit caution when choosing to use 4% articaine in an inferior alveolar nerve block until further scientific research has been performed.

Introduction

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Since 1949, lidocaine has been recognised as the 'gold-standard' of local anaesthetic (LA) agents.¹ However, the desire to develop fastacting agents with a short half-life that also produce profound anaesthesia has led to the development of other alternatives. One example is articaine, initially synthesised in 1969 and used for the first time in clinical dental practice in Germany in 1976.

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The reason for articaine's popularity appears to be due to its efficacy. Numerous studies have shown that articaine produces a more profound anaesthesia than that of lidocaine.2-8 Lidocaine is an amide compound, based on a benzene ring structure (C6H6). Articaine, in contrast, possesses a thiophene ring (C4H3), providing greater lipid solubility and an increased potency as a greater volume of an administered dose can enter the target neurons. Articaine's lipid solubility has been quoted at over four times greater than that of lidocaine.9 The same study confirmed that the onset of anaesthesia was achieved in 7.4 mins with articaine, as opposed to 8.7 mins with lidocaine.9 It has also been suggested that articaine provides a longer duration of anaesthesia due to its protein binding characteristics.10,11

With these attributes, it is perhaps not surprising that many studies have concluded that

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articaine is more efficient at producing profound anaesthesia than lidocaine.^{6,12-15} These papers include studies of both infiltration and nerve block anaesthesia. Other authors concluded that articaine has a faster onset than lidocaine,¹¹ and a meta-analysis has proved that articaine is 1.6–3.5 times more potent than lidocaine.² Several studies have concluded that articaine should be recommended for use over lidocaine.^{2,6,12,16} In 2007, Robertson *et al.* concluded that both the speed of onset and the anaesthetic efficacy of articaine were superior to those of lidocaine, when administered via a buccal infiltration technique in the posterior molar region.¹⁴

Another important attribute of a local anaesthetic agent is that of safety and this is perhaps where articaine compares less favourably. Since its introduction, several articles have been published warning of possible nerve damage when articaine

is administered in an inferior alveolar nerve block (IANB).^{17,18} These articles indicate a risk of causing temporary or permanent paraesthesia of the inferior alveolar nerve (IAN) but evidence also exists contradicting these claims.^{3,19,20}

It appears, therefore, that the dental profession faces a dilemma. Should the more efficient agent be used to achieve faster, more profound anaesthesia; or should the profession be wary of an agent that may have the potential to induce nerve damage?

A mini systematic review of the literature was performed by a single researcher with one, clearly focused question.²¹ The results of the study will hopefully provide advice to the dental profession, ensuring the continued provision of safe and effective local anaesthesia.

Methodology

The Scottish Intercollegiate Guidelines Network (SIGN) presents eight levels of evidence-based research. The SIGN tool was used in this study according to the criteria set out in Table 1.²² The development of the research question was aided using the PICOS method,²³ as described in Table 2. Inclusion and exclusion criteria were applied to the literature search as outlined in Tables 3 and 4. Basic search terms and medical sub-headings terms were developed and detailed in Boxes 1 and 2. Three electronic databases were chosen to systematically search the available literature:

- MEDLINE with Full Text
 Dentistry & Oral Sciences Source
- The Cochrane Library.
- 5. The Coefficient Library.

Quality assessment of studies

To ensure that the random controlled trials included in the review were accurately assessed against the inclusion and exclusion criteria, the risk of bias tool as described in the *Cochrane Handbook for Systemic Reviews of Intervention* was applied.²⁴

For the selected cohort studies, a methodology index for non-randomised studies (MINORS) was applied,²⁵ as described in Table 6. A record sheet was developed, and each study was subsequently scored as directed by Slim and Nini *et al.* 2003²⁵ as defined in Table 7.

Data extraction

Specifically designed data extraction forms were developed, allowing uniform data to be extracted under the following headings:

- Study design
- Study objectives
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- · Geographical origin of the study
- · Clinical setting for the study
- Study funding
- Study participants sex, age, numbers
- Type of anaesthetic agent used
- Study outcome-methods of recording and reporting nerve damage
- Comparison made between 'expected' and 'observed' outcomes
- Follow-up periods
- Attrition bias
- Data analysis of outcomes.

Results

Data extraction and results of the mini systematic review are detailed in Tables 8–15 and Figure 1.

Discussion

Malamed and Gagnon's study of 1,325 participants enabled a statistical analysis of the results which indicated that the incidence of nerve damage was the same (1%) whether 4% articaine or 2% lidocaine was used as the LA agent. Indeed, this DBRCT concluded that articaine is a 'safe and effective' local anaesthetic agent.¹⁹

Both studies conducted by Pogrel,^{20,26} concluded that the incidence of nerve damage following the use of 4% articaine was in proportion to its market share. However, three of the studies indicated that the use of 4% articaine elicited more adverse outcomes than would be expected when compared to the agent's market share.^{17,27,28}

Limitations and characteristics of included studies

Several methodological inconsistencies exist throughout the included studies, making a direct comparison between the chosen articles difficult. When performing a study comparing two pharmaceutical agents, a true comparison can only be achieved with the knowledge of the relative use of the two drugs within the studied population. Haas and Lennon,¹⁷ Gaffen and Haas,²⁸ and Garisto, Gaffen *et al.*,²⁷ all used the 'null hypothesis' developed by Ronald Fisher.²⁹ However, the other included studies failed to indicate any comparison between expected and observed outcome events.

The creation of a 'barb' on the tip of the needle, resulting from contact with the bone, may also be a factor in the traumatic damage to both the IAN and lingual nerve (LN). However, whether or not this event occurred during any of the IANBs included in the studies, the resultant mechanical

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Box 1 Basic search terms

articaine
carticaine
septanest
ultracaine
septocaine
dental anaesthesia
ignocaine
lidocaine
xylocaine
paraesthesia
paresthesia
anaesthesia
anesthesia
dysaesthesia
dysesthesia
trigeminal nerve injuries
damage
njury
nferior alveolar nerve
nferior dental nerve
mandibular nerve
lingual nerve

Box 2 Medical sub headings terms (MeSH Terms)

articaine dental anaesthesia nerve injury

damage would be the same for both LA solutions.

Of the seven included papers, only one involves a DBRCT, three involve voluntary reporting of nerve damage, and the remaining three articles elicit their information from patients who have been referred to a specialist centre for the specific reason that they are experiencing some degree of nerve damage. This clearly results in a considerable degree of reporting bias.

With incidences of nerve damage ranging from 1:27,000 to 1:785,000,^{17,30} it is clear that this study's outcome is extremely rare. To obtain statistically significant results in a DBRCT would require a clinical trial on a very large scale. This could explain the existence of only one such study since 1976.¹⁹

Both Hillerup and Jensen,¹⁸ and Garisto and Gaffen,²⁷ make reference to the possibility of reporting bias in their papers, and Gaffen and Haas²⁸ admit that 'reported incidence numbers should be viewed cautiously.' In his 2007 paper, Pogrel²⁶ states that he estimates his study represents approximately 10% of all cases of nerve damage in the given population per year. However, reporting bias for patients referred to

a specialist centre would be the same for both LA solutions.

The only study that included a detailed physical examination of the patient was that of Hillerup and Jensen,¹⁸ using a 'standardised test of neurosensory functions' by a single operator to determine the presence and extent of any reported nerve damage.^{31,32} The remaining included studies merely noted the incidence of 'reported' nerve damage.

Pogrel's studies,^{20,26} using data from a specialist centre and Garisto and Gaffen's paper,²⁷ all failed to accurately examine the patient, relying instead on the patient's own descriptions and a log of reported cases to the adverse event reporting system (AERS). Pogrel's description of the patient 'examination' lacks sufficient detail to allow exclusion of detection bias.

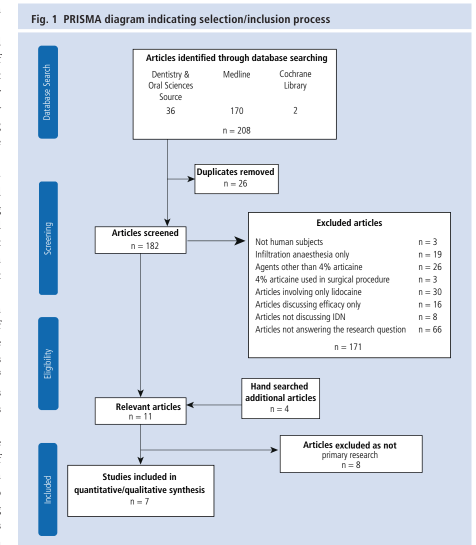
The description of the reporting of an 'electric shock' during the administration of the LA created notable discussion among the included authors. Four of the included papers noted the reporting of this phenomenon^{17,18,27,28} and all included these reports in their results as a 'nerve injury'. The remaining three papers failed to mention this possible event.^{19,20,26}

Interestingly, Hillerup and Jensen state that 'electric shock per se is probably of minor relevance for the aetiology of injection injuries.¹⁸ However, they then go on to question the cause of nerve injury, admitting that it is unknown as to whether the nerve is damaged via neurotoxicity or mechanically, via intra-fascicular injection.

Many authors are now advocating the use of 4% articaine in infiltration anaesthesia as an alternative to block anaesthesia due to the increased efficacy of this agent.^{33–36} The evidence presented in these studies indicates a clear efficacy advantage when using 4% articaine as a buccal infiltration compared to 2% lidocaine in an IANB. One author has even suggested that the IANB may now be an unnecessary procedure.³⁷

Concentration of the LA agent

Three of the chosen papers postulate that it may be the fact that, because articaine is administered in a 4% solution, it is the concentration of the LA solution rather than the actual pharmacology of the agent that causes damage to the nerve.^{17,27,28} This suggestion would appear to be confirmed by another study on rat sciatic nerves, which concluded that significantly more neurotoxic injuries were observed following the direct injection into the nerve of a 4% articaine solution compared to that of a 2% solution.³⁸



In a recent *in vitro* study, articaine proved to be less neurotoxic than lidocaine, mepivacaine and prilocaine.³⁹ Indeed, previous studies have concluded that no scientific evidence exists to confirm the suggestion that articaine causes increased paraesthesia and, to date, no causal relationship has been exhibited between an anaesthetic agent's concentration and neurological damage.^{40,41}

Implications for clinical research

This mini systematic review confirms that controversy still exists over the safety of 4% articaine and 1:100,000 adrenaline as a dental local anaesthetic agent.

The authors of all the included papers admit that, due to the extremely rare occurrence of the outcome, a carefully performed, high quality DBRCT would have to involve such vast numbers of participants that, logistically, such a study would pose certain problems.

It is generally accepted that 4% articaine exhibits greater lipid solubility, faster onset

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and increased duration of anaesthesia, more profound anaesthesia, and reduced toxicity than those of its counterpart, 2% lidocaine. With these favourable attributes, 4% articaine does indeed offer superior properties over 2% lidocaine but would a 2% articaine solution offer the same advantages?

Further research is required into the efficacy and safety of a 2% articaine solution. Indeed, a study in 2006 proved that the 4% articaine solution was not superior in its anaesthetic effect compared to 2% and 3% solutions of the same agent.⁴²

Implications for general dental practice

The highest level of evidence available to this study was that of Malamed and Gagnon's DBRCT in 2001.¹⁹ Although spread over 27 sites in two countries, this trial unfortunately exhibited several potential areas of bias. It did, however, conclude that there was no evidence to suggest that 4% articaine posed a greater risk of nerve damage than 2% lidocaine and that the

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use of 4% articaine in general dental practice can, therefore, be deemed safe and efficient.

Three further papers, not included in this study, also concluded that no conclusive evidence exists to suggest that 4% articaine poses a greater risk of nerve damage compared to other LA agents.^{3,10,12}

Conclusion

This mini systematic review of the literature has highlighted the fact that further research is required to determine the relative risks of using 4% articaine compared to 2% lidocaine in IANBs. Clearly, the use of 4% articaine is becoming increasingly popular as a means of achieving successful dental anaesthesia and, if current trends continue, this agent may become the number one anaesthetic of choice in the future. This steady increase in popularity is likely to be due to the proven efficacy of this LA agent, benefiting both the patient and the operator. Indeed, the incidence of inferior alveolar nerve damage may reduce in the future as more evidence emerges to support infiltration anaesthesia. With this in mind, and considering the contradictory evidence presented in this study, it is suggested that until factual evidence becomes available, dental practitioners should consider all the potential risks and benefits of a particular LA agent prior to its administration.

- Renton T. Oral surgery: part 4. minimising and managing nerve injuries and other complications. *Br Dent J* 2013; 215: 393–399.
- Brandt R G, Anderson P F, McDonald N J, Sohn W, Peters M C. The pulpal anesthetic efficacy of articaine versus lidocaine in dentistry: a meta-analysis. *J Am Dent Assoc* 2011; **142:** 493–504.
- Su N, Li C, Wang H, Shen J, Liu W, Kou L. Efficacy and safety of articaine versus lidocaine for irreversible pulpitis treatment: a systematic review and meta-analysis of randomised controlled trials. *Aust Endod J* 2016; 42: 4–15.
- Malamed S F, Gagnon S, Leblanc D. Efficacy of articaine: a new amide local anesthetic. JAm Dent Assoc 2000; 131: 635–642.
- 5. Powell V. Articaine is superior to lidocaine in providing pulpal anesthesia. *J Am Dent Assoc* 2012; **143**: 897–898.
- Paxton K, Thome D E. Efficacy of articaine formulations: quantitative reviews. *Dent Clin North Am* 2010; 54: 643–653.
- Haase A, Reader A, Nusstein J, Beck M, Drum M. Comparing anesthetic efficacy of articaine versus lidocaine as a supplemental buccal infiltration of the mandibular first molar after an inferior alveolar nerve block. *J Am Dent Assoc* 2008: **139**: 1228–1235.
- Kanaa M D, Whitworth J M, Meechan J G. A comparison of the efficacy of 4% articaine with 1:100,000 epinephrine and 2% lidocaine with 1:80,000 epinephrine in achieving pulpal anesthesia in maxillary teeth with irreversible pulpitis. *J Endod* 2012; **38:** 279–282.
- Bartlett G, Mansoor J. Articaine buccal infiltration vs lidocaine inferior dental block - a review of the literature. *Br Dent J* 2016; 220: 117–120.
- Yapp K E, Hopcraft M S, Parashos P. Articaine: a review of the literature. *Br Dent J* 2011; 210: 323–329.
- Costa C G, Tortamano I P, Rocha R G, Francischone C E, Tortamano N. Onset and duration periods of articaine and lidocaine on maxillary infiltration. *Quintessence Int* 2005; 36: 197–201.

- Katyal V. The efficacy and safety of articaine versus lignocaine in dental treatments: a meta-analysis. J Dent 2010; 38: 307–317.
- Srinivasan N, Kavitha M, Loganathan C S, Padmini G. Comparison of anesthetic efficacy of 4% articaine and 2% lidocaine for maxillary buccal infiltration in patients with irreversible pulpitis. Oral Surg Oral Med Oral Pathol Oral Radiol Endod 2009; 107: 133–136.
- Robertson D, Nusstein J, Reader A, Beck M, McCartney M. The anesthetic efficacy of articaine in buccal infiltration of mandibular posterior teeth. J Am Dent Assoc 2007; 138: 1104–1112.
- Darawade D A, Kumar S, Budhiraja S, Mittal M, Mehta T N. A clinical study of efficacy of 4% articaine hydrochloride versus 2% lignocaine hydrochloride in dentistry. *J Int Oral Health* 2014; 6: 81–83.
- Balto K. Administration of articaine anesthesia may lead to superior profound pulpal anesthesia compared with lidocaine in adult patients. *J Evid Based Dent Pract* 2011; 11: 183–184.
- Haas D A, Lennon D. A 21 year retrospective study of reports of paresthesia following local anesthetic administration. *J Can Dent Assoc* 1995; **61:** 319–320, 323–326, 329–330.
- Hillerup S, Jensen R. Nerve injury caused by mandibular block analgesia. Int J Oral Maxillofac Surg 2006; 35: 437–443.
- Malamed S F, Gagnon S, LeBlanc D. Articaine hydrochloride: a study of the safety of a new amide local anesthetic. *J Am Dent Assoc* 2001; **132**: 177–185.
- Pogrel M A. Permanent nerve damage from inferior alveolar nerve blocks: a current update. J Calif Dent Assoc 2012; 40: 795–797.
- 21. Griffiths P. Evidence informing practice: introducing the mini-review. *Br J Community Nurs* 2002; **7**: 38–39.
- Petrisor B A, Keating J, Schemitsch E. Grading the evidence: levels of evidence and grades of recommendation. *Injury* 2006; **37:** 321–327.
- Makela M, Witt K. How to read a paper: critical appraisal of studies for application in healthcare. *Singapore Med J* 2005: 46: 108–114.
- Higgins J P T, Altman D G, Gøtzsche P C *et al.* The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. *BMJ* 2011; **343:** d5928.
- Slim K, Nini E, Forestier D, Kwiatkowski F, Panis Y, Chipponi J. Methodological index for non-randomized studies (minors): development and validation of a new instrument. ANZ J Surg 2003; 73: 712–716.
- Pogrel M A. Permanent nerve damage from inferior alveolar nerve blocks--an update to include articaine. J Calif Dent Assoc 2007; 35: 271–273.
- Garisto G A, Gaffen A S, Lawrence H P, Tenenbaum H C, Haas D A. Occurrence of paresthesia after dental local anesthetic administration in the United States. J Am Dent Assoc 2010; 141: 836–844.
- Gaffen A S, Haas D A. Retrospective review of voluntary reports of nonsurgical paresthesia in dentistry. J Can Dent Assoc 2009; 75: 579.
- Saxon E. Defining the null hypothesis. *BMC Biol* 2015; 13: 68.
- Pogrel M A, Thamby S. Permanent nerve involvement resulting: from inferior alveolar nerve blocks. JAm Dent Assoc 2000; 131: 901–907.
- Robinson P P. Observations on the recovery of sensation following inferior alveolar nerve injuries. Br J Oral Maxillofac Surg 1988; 26: 177–189.
- Robinson P P, Smith K G, Johnson F P, Coppins D A. Equipment and methods for simple sensory testing. Br J Oral Maxillofac Surg 1992; 30: 387–389.
- Corbett I P, Kanaa M D, Whitworth J M, Meechan J G. Articaine infiltration for anesthesia of mandibular first molars. *J Endod* 2008; 34: 514–518.
- Kanaa M D, Whitworth J M, Corbett I P, Meechan J G. Articaine buccal infiltration enhances the effectiveness of lidocaine inferior alveolar nerve block. *Int Endod J* 2009; 42: 238–246.
- 35. Poorni S, Veniashok B, Senthilkumar A D, Indira R, Ramachandran S. Anesthetic efficacy of four percent articaine for pulpal anesthesia by using inferior alveolar nerve block and buccal infiltration techniques in patients with irreversible pulpitis: a prospective randomized double-blind clinical trial. J Endod 2011; 37: 1603–1607.
- Leith R, Lynch K, O'Connell A C. Articaine use in children: a review. *Eur Arch Paediatr Dent* 2012; 13: 293–296.

- Malamed S F. Is the mandibular nerve block passé? J Am Dent Assoc 2011; 142 (Spec Iss): 3S–7S.
- Hillerup S, Bakke M, Larsen J O, Thomsen C E, Gerds T E. Concentration-dependent neurotoxicity of articaine: an electrophysiological and stereological study of the rat sciatic nerve. *Anesth Analg* 2011; **112**: 1330–1338.
- Malet A, Faure M, Deletage N, Pereira B, Haas J, Lambert G. The comparative cytotoxic effects of different local anesthetics on a human neuroblastoma cell line. *Anesth Analg* 2015; **120**: 589–596.
- Malamed S F. Local anesthetics: dentistry's most important drugs, clinical update 2006. J Calif Dent Assoc 2006; 34: 971–976.
- Malamed S F. Articaine versus lidocaine: the author responds. J Calif Dent Assoc 2007; 35: 383–385.
- Hintze A, Paessler L. Comparative investigations on the efficacy of articaine 4% (epinephrine 1:200,000) and articaine 2% (epinephrine 1:200,000) in local infiltration anaesthesia in dentistry-a randomised double-blind study. *Clin Oral Investig* 2006; **10**: 145–150.
- Petrisor B, Bhandari M. The hierarchy of evidence: Levels and grades of recommendation. *Indian J Orthop* 2007; 41: 11–15.
- Aguiar J, Chebroux A, Martinez-Taboada F, Leece E A. Analgesic effects of maxillary and inferior alveolar nerve blocks in cats undergoing dental extractions. *J Feline Med* Surg 2015; **17:** 110–116.
- Hung P, Chang H, Yang P, Kuo Y, Lan W, Lin C. Comparison of the Gow-Gates mandibular block and inferior alveolar nerve block using a standardized protocol. *J Formos Med Assoc* 2006; **105**: 139–146.
- Potocnik I, Tomsic M, Sketelj J, Bajroviä F F. Articaine is more effective than lidocaine or mepivacaine in rat sensory nerve conduction block in vitro. *J Dent Res* 2006; 85: 162–166.
- Sisk A L. Evaluation of the Akinosi mandibular block technique in oral surgery. J Oral Maxillofac Surg 1986; 44: 113–115.
- Baroni D B, Franz-Montan M, Cogo K *et al.* Effect of articaine on mental nerve anterior portion: histological analysis in rats. *Acta Odontol Scand* 2013; **71**: 82–87.
- Batista D S, Berto L A, Volpato M C *et al.* Anesthetic efficacy of articaine and lidocaine for incisive/mental nerve block. *J Endod* 2010; **36**: 438–441.
- Chopra R, Jindal G, Sachdev V, Sandhu M. Double-blind crossover study to compare pain experience during inferior alveolar nerve block administration using buffered two percent lidocaine in children. *Pediatr Dent* 2016; 38: 25–29.
- Danielsson K, Evers H, Holmlund A, Kjellman O, Nordenram A, Persson N E. Long-acting local anaesthetics in oral surgery. Clinical evaluation of bupivacaine and etidocaine for mandibular nerve block. *Int J Oral Maxillofac Surg* 1986; **15**: 119–126.
- Rood J P. Inferior alveolar nerve blocks. The use of 5 per cent lignocaine. *Br Dent J* 1976; **140**: 413–414.
 Ahmad Z H, Ravikumar H, Karale R, Preethanath R S,
- Ahmad Z H, Ravikumar H, Karale R, Preethanath R S, Sukumaran A. Study of the anesthetic efficacy of inferior alveolar nerve block using articaine in irreversible pulpitis. *J Contemp Dent Pract* 2014; 15: 71–74.
- Kambalimath D H, Dolas R S, Kambalimath H V, Agrawal S M. Efficacy of 4 % articaine and 2 % lidocaine: a clinical study. J Maxillofac Oral Surg 2013; 12: 3–10.
- Moorthy A, Stassen L F. The occurrence of paraesthesia of the maxillary division of the trigeminal nerve after dental local anaesthetic use: a case report. *J Ir Dent Assoc* 2015; 61: 34–35.
- Choi E, Seo J, Jung B, Park W. Diplopia after inferior alveolar nerve block anesthesia: report of 2 cases and literature review. Oral Surg Oral Med Oral Pathol Oral Radiol Endod 2009; 107: e21–e24.
- Al-Sandook T, Al-Saraj A. Ocular complications after inferior alveolar nerve block: a case report. J Calif Dent Assoc 2010; 38: 57–59.
- Wyman R J. Nerve injury following a mandibular block: a case report. *Dent Today* 2010; 29: 14.
- 59. Pedlar J. Prolonged paraesthesia. Br Dent J 2010; **195**: 119.
- Fowler S, Reader A. Is a volume of 3.6 mL better than 1.8 mL for inferior alveolar nerve blocks in patients with symptomatic irreversible pulpitis? *J Endod* 2013; **39**: 970–972.
- Steinkruger G, Nusstein J, Reader A, Beck M, Weaver J. The significance of needle bevel orientation in achieving a successful inferior alveolar nerve block. J Am Dent Assoc 2013; 137: 1685–1691.

Table 1 The hierarchy of evidence. Adapted from the Scottish Intercollegiate Guidelines Network (SIGN)⁴³

Level of evidence	Description of evidence
1++	High quality meta-analysis, systematic reviews of RCTs or very low risk of bias RCTs
1+	Well conducted meta-analysis, systematic reviews of RCTs or very low risk of bias RCTs
1-	Meta-analysis, systematic reviews of RCTs or RCTs with a high risk of bias
2++	High quality systematic reviews of cohort or case-control studies or high quality cohort or case-control studies with a very low risk of confounding bias or chance and a high probability that the relationship is causal
2+	Well conducted cohort or case-control studies with a low risk of confounding, bias or chance and a moderate probability that the relationship is causal
2-	Cohort or case-control studies with a high risk of confounding, bias or chance and a significant risk that the relationship is not causal
3	Non-analytical studies. Case reports and case series
4	Expert opinion

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Table 2 PICOS parameters applied to the studyPICOSSearch strategy applicationPopulationPatients receiving IANBs with either 4% articaine hydrochloride + 1:100,000 adrenaline or patients receiving IANBs with 2% lidocaine + 1:100,000
adrenaline. Males and females. All agesInterventionStudies involving the administration of an IANB with 4% articaine + 1:100,000 adrenalineComparisonStudies involving the administration of an IANB with 2% lidocaine + 1:100,000 adrenalineOutcomePost injection nerve damage indicated by prolonged temporary or permanent anaesthesia, paraesthesia or dysaesthesia in both the intervention and comparison groupsStudiesRandomised controlled trials comparing 4% articaine + 1:100,000 adrenaline + 2% lidocaine + 1:100,000 adrenaline in IANBs. Cohort studies investigating
the use of 4% articaine + 1:100,000 adrenaline as a dental local anaesthetic agent in IANBs.

Table 3 Search inclusion criteria

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Inclusion criteria	Reason for inclusion			
English language papers	No translation facility. Author only speaks English.			
Papers published since 1976	Articaine's first use in clinical dentistry			
Human subjects only	Relevant to general dental practice			
Male and female subjects	Maximum number of participants			
Global participation	Maximum number of participants			
Subjects of all ages	Maximum number of participants			
Articles involving IANB anaesthesia	Specific to study question			
LA agents, lidocaine and articaine only	Specific to study question			
Inferior alveolar and/or lingual nerve damage	Anatomical possibility of damage to either nerve during the administration of an IANB.			
Permanent and/or temporary nerve damage	Both indicators of nerve damage			
Suitable ethical approval obtained	Ethical and moral issues relating to research			
Random controlled trials	Good quality evidence			
Cohort studies	Large number of subjects			

Table 4 Search exclusion criteria **Exclusion criteria Reason for exclusion** Articles describing only infiltration anaesthesia Administration of a nerve block is postulated as a cause of nerve damage Articles describing the use of anaesthetic agents other than articaine or lignocaine Other anaesthetic agents not widely used in general dental practice Studies investigating the use of articaine for 'surgical dentistry' Possible surgical cause of nerve damage Both recognised causes of possible inferior alveolar and lingual nerve paraesthesia Studies investigating the use of articaine for removal of lower third molars and placement of mandibular implants 'Sponsored' articles, unless a conflict of interest is declared Author bias Case studies Poor quality evidence Letters to editors Personal opinions

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Table 5 Search strategy, 18 November 2016	
Search no.	Search term
S1	(MM 'carticaine')
S2	septanest
\$3	articaine
S4	ultracaine
S5	septocaine
S6	(MM 'anesthesia, dental+')
S7	lignocaine
58	lidocaine
S9	xylocaine
S10	S1 or S2 or S3 or S4 or S5 or S6
S11	S7 or S8 or S9
S12	paraesthesia
S13	paresthesia
S14	anaesthesia
S15	anesthesia
S16	dysaesthesia
S17	dysesthesia
S18	(MM 'trigeminal nerve injuries+')
S19	damage
S20	injury
S21	inferior alveolar nerve
S22	inferior dental nerve
S23	mandibular nerve
S24	lingual nerve
S25	S12 or S13 or S14 or S15 or S16 or S17 or S18 or S19 or S20
S26	S21 or S22 or S23 or S24
S27	S10 and S11 and S25 and S26

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Table 6 Methodology index for non randomised studies (MINORS) ²⁵				
Methodological items for non-randomised studies	Item description			
Clearly stated aim	Relevant and precise study question, relating to available literature			
Inclusion of consecutive patients	All eligible participants included in study			
Prospective collection of data	Data collected as per guidelines established prior to study commencement			
Endpoints appropriate to study aim	Clear, quantifiable outcome addressing study question			
Unbiased endpoint	Blind assessment of endpoint			
Review period appropriate to aim	Review period sufficient to allow outcome occurrence and measurement			
Attrition bias less than 5%	All patients should be reviewed			
Prospective calculation of study size	Information regarding study population size necessary to achieve 95% confidence interval and level of statistical signific			
Additional items for use in comparative studies	Item description			
Suitable control	'Gold-standard' as per available information			
Contemporary groups	Groups studies during the same time period			
Baseline equivalent groups	Group criteria similar at start point			
Statistical analysis	Suitable statistics with confidence intervals or relative risk			

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Table 7 MINORS criteria scores			
Item score	Reason		
0	Not reported		
1	Reported but inadequate		
2	Reported and adequate		

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Table 8 Search strategy and results (performed on 30 December 2016)					
Search no.	Search term	Dentistry & oral science	Medline	Cochrane	
S1	(MM 'carticaine')	2	303	3	
S2	septanest	2	4	1	
S3	articaine	216	398	3	
S4	ultracaine	4	47	9	
S5	septocaine	6	3	1	
S6	(MM 'Anesthesia, Dental+')	1,277	5,827	9	
S7	lignocaine	332	2,405	11	
58	lidocaine	561	25,426	47	
S9	xylocaine	306	713	1	
S10	S1 or S2 or S3 or S4 or S5 or S6	1,429	6,139	9	
S11	S7 or S8 or S9	592	26,463	55	
S12	paraesthesia	117	1,134	195	
S13	paresthesia	31	7,415	50	
S14	anaesthesia	6,591	65,803	1078	
S15	anesthesia	6,591	200,202	334	
S16	dysaesthesia	24	265	23	
S17	dysesthesia	61	1278	13	
S18	(MM 'trigeminal nerve injuries+')	84	833	13	
S19	damage	3,284	433,750	2,568	
S20	injury	9,260	549,161	2,570	
S21	inferior alveolar nerve	1124	2,102	13	
S22	inferior dental nerve	78	142	18	
S23	mandibular nerve	568	3,556	36	
S24	lingual nerve	269	1,298	18	
S25	S12 or S13 or S14 or S15 or S16 or S17 or S18 or S19 or S20	18,767	1,145,705	4,497	
S26	S21 or S22 or S23 or S24	1,492	5281	55	
S27	S10 and S11 and S25 and S26	36	170	2	

Table 9 Included studies 'SIGN' Level Type of study of evidence Title and author(s) Year A 21-year retrospective study of reports of paresthesia following local anesthetic administration. Hass and Lennon¹⁷ 2-1995 Retrospective cohort 2-Retrospective review of voluntary reports of nonsurgical paresthesia in dentistry. Gaffen and Haas²⁸ 2009 Retrospective cohort Nerve injury caused by mandibular block analgesia. Hillerup and Jenson¹⁸ 2006 2-Retrospective cohort Permanent nerve damage from inferior alveolar nerve blocks – an update to include articaine. Pogrel²⁶ 2007 2-Retrospective cohort 1-Articaine hydrochloride: a study of the safety of a new amide local anesthetic. Malamed, Gagnon et al.¹⁹ 2001 Random controlled trials Occurrence of paresthesia after dental local anesthetic administration in the United States. Garisto, Gaffen et al.²⁷ 2-2010 Retrospective cohort 2012 2-Permanent nerve damage from inferior alveolar nerve blocks: a current update. Pogrel²⁰ Retrospective cohort

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Table 10 Examples of excluded studies				
Article(s)	Reason for exclusion			
Aguiar, Chebroux <i>et al.</i> ⁴⁴ Hung, Chang <i>et al.</i> ⁴⁵ Potocnik, Tomsic <i>et al.</i> ⁴⁶ Sisk ⁴⁷ Baroni, Franz-Montan <i>et al.</i> ⁴⁸ Batista, Berto <i>et al.</i> ⁴⁹	Incorrect population. $n = 6$ Studies on rats and cats Studies using Cow–Gates and Akinosi IANB Studies of mental and incisive nerve blocks			
Chopra, Jindal <i>et al.</i> ⁵⁰ Danielsson, Evers <i>et al.</i> ⁵¹ Rood ⁵²	Incorrect intervention. $n = 48$ Studies comparing lidocaine, etidocaine and bupivacaine			
Rood ⁵²	Incorrect comparator. n = 1 5% lidocaine solution used in study			
Ahmad, Ravikumar <i>et al.</i> ⁵³ Kambalimath, Dolas <i>et al.</i> ⁵⁴ Moorthy, Stassen ⁵⁵ Choi, Seo <i>et al.</i> ⁵⁶ Al-Sandook, Al-Saraj ⁵⁷	Incorrect outcome. n = 42 Studies measuring articaine's efficacy only Studies detailing damage to nerves other than IAN and/or LN			
Choi, Seo <i>et al.</i> ⁵⁶ Wyman ⁵⁸ Pedlar ⁵⁹	Incorrect studies. n = 8 Case reports and letters to editors			
Fowler, Reader ⁶⁰ Steinkruger, Nusstein <i>et al.</i> ⁶¹	Articles not answering study question. n = 66 Studies comparing volume of anaesthetic agent and injection technique			

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Table 11 MINORS checklist for included studies							
Criteria	Haas & Lennon ¹⁷	Gaffen & Haas ²⁸	Hillerup & Jenson ¹⁸	Pogrel ²⁶	Malamed & Gagnon ¹⁹	Garisto & Gaffen ²⁷	Pogrel ²⁰
Clearly stated aim	2	2	2	2	2	2	2
Inclusion of consecutive patients	1	2	2	2	1	2	2
Prospective collection of data	2	2	2	2	2	2	2
Endpoint appropriate to study	2	2	2	2	2	2	2
Unbiased assessment of endpoint	1	1	1	1	2	1	1
Appropriate follow-up period	0	1	2	2	1	1	2
Loss to follow-up less than 5%	1	0	0	0	0	0	0
Prospective calculation of study size	0	0	0	0	0	0	0
Adequate control group	N/A	N/A	N/A	N/A	2	N/A	N/A
Contemporary groups	N/A	N/A	N/A	N/A	2	N/A	N/A
Baseline equivalence groups	N/A	N/A	N/A	N/A	2	N/A	N/A
Adequate statistical analysis	N/A	N/A	N/A	N/A	1	N/A	N/A
Total score	9	10	11	11	17	10	11

Table 12 Risk of assessment bias ²⁴				
Bias	Malamed and Gagnon ¹⁹			
Random sequence generation (selection bias)	Low risk 'There were no statistically significant differences in the studies between the articaine and lidocaine treatment groups with respect to age, sex, weight, race distribution or the proportion of subjects undergoing simple or complex procedures'			
Allocation concealment (selection bias)	Unclear risk. Not mentioned in methodology			
Blinding of outcome assessment (detection bias)	Unclear risk. 'Randomised, double-blind' mentioned in methodology but no other details			
Participant awareness (performance bias)	Unclear risk. Not mentioned in methodology			
Incomplete outcome data (attrition bias)	High risk. No mention of attrition at 24 hour and 7 day follow-up interviews			
Sponsorship (funding bias)	Low risk. 'The manufacturer of the drug products used in the three trialsproviding materials and funding.' The same company manufactures both the intervention and comparator drugs			
Selective reporting (reporting bias)	Unclear risk. 'The vast majority of these events are related by (telephone interviews with) patients and are alleged not confirmed'			
Overall risk of bias	Unclear risk.			

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Table 13a Data extraction					
Study	Haas & Lennon ¹⁷	Gaffen & Haas ²⁸	Hillerup & Jensen ¹⁸		
Study publication date	April 1995	October 2009	May 2006		
Study design	Retrospective cohort	Retrospective cohort	Retrospective cohort		
Study objectives	Prolonged paraesthesia following LA in dentistry	Prolonged paraesthesia following LA in dentistry	Prolonged paraesthesia following LA in dentistry		
Geographical origin	Ontario, Canada	Ontario, Canada	Denmark		
Study setting	Not stated	Not stated	'All dental practitioners'		
Study funding	Not stated	'no declared financial interests'	Not stated		
Eligible study participants	143, male and female, all ages	172, male and female, 11–80 years	52, male and female, 24–81 years		
LA agents used	Lidocaine, articaine, prilocaine, mepivacaine, bupivacaine	Lidocaine, articaine, prilocaine, mepivacaine, bupivacaine	Lidocaine, articaine, prilocaine, mepivacaine		
Outcome reporting and recording	Voluntary reports to PLP	Voluntary reports to PLP	Telephone call to GDP. Type and volume of LA used. Electric shock experienced? Written questionnaires and patient interviews		
Comparison made between 'expected' and 'observed' outcomes	Yes	Yes	No		
Study period	21 years, 1973–1993	10 years, 1999–2008	8 years, 1997–2004		
Attrition bias	Not stated	Not stated	30 patients (58%) lost to follow up after 12 months		
Data analysis of outcomes	Chi: square analysis	Chi: square analysis	Chi: square analysis		
Ethical approval	Not stated	Stated obtained	Not stated		

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Table 13b Data extraction

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Study	Pogrel ²⁶	Malamed, Gagnon et al. ¹⁹	Garisto, Gaffen et al. ²⁷	Pogrel ²⁰	
Study publication date	April 2007	February 2001	July 2010	October 2012	
Study design	Retrospective cohort	3 double blind random controlled trials	Retrospective cohort	Retrospective cohort	
Study objectives	Prolonged IAN/LN paraesthesia following LA in dentistry	Direct comparison of efficacy and safety between 4% articaine and 2% lidocaine	Record incidence of nerve dam- age after LA in dentistry	Prolonged IAN/LN paraesthesia following LA in dentistry	
Geographical origin	Maxillofacial Dept, UCSF, USA	27 sites, 8 in the UK and 19 in the USA	USA	Maxillofacial Dept, UCSF, USA	
Study setting	Primary and secondary dental care	No stated	Voluntary reports to FDA's AERS	Primary and secondary dental care	
Study funding	Not stated	'Materials and funding' provided by manufacturers of the LA agents	No 'disclosures' reported by authors	Not stated	
Eligible study participants	57, sex and ages not stated	1325, male and female, aged 4–80 years	226, male and female, 15–78 years	38, sex and ages not stated	
LA agents used	Lidocaine, articaine, prilocaine, mepivacaine, bupivacaine	2% Lidocaine, 4% articaine,	Lidocaine, articaine, prilocaine, mepivacaine, bupivacaine	Lidocaine, articaine, prilocaine, carbocaine	
Outcome reporting and recording	Examination of patient at UCSF. Details of examination not stated	Interviews and telephone calls to the patients. No further details of examination	Voluntary reports to FDA's AERS. Duration of paraesthesia noted	Examination of patient at UCSF. Details of examination not stated	
Comparison made between 'expected' and 'observed' outcomes	Yes	No	Yes	Yes	
Study period	3 years. 01/01/03-31/12/05	Not stated	11 years, November 1997– August 2008	6 years, 01/01/06-31/12/11	
Attrition bias	Not stated	3 patients lost to follow up (0.23%)	Not stated	Not stated	
Data analysis of outcomes	Narrative	Narrative	Descriptive statistical analysis	Narrative	
Ethical approval	Not stated	Stated as obtained in UK and USA	Stated as obtained and approved by University of Toronto	Not stated	

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Table 14	Summary	of outcome charad	cteristics of included	studies

Study	Design	Number of eligible participants with outcome *	Number of participants with outcome following intervention (articaine)	Number of participants with outcome following comparison (lidocaine)	Reported outcomes
Haas & Lennon ¹⁷	Retrospective cohort	143 <u></u> *	50	5	Paraesthesia following the injection of LA in non-surgical dentsistry
Gaffen & Haas ²⁸	Retrospective cohort	172*	109	23	Non-surgical paraesthesia
Hillerup & Jensen ¹⁸	Retrospective cohort	52 *	29	10	Non-surgical IAN or LN injury following a unilateral IANB
Pogrel ²⁶	Retrospective cohort	57 *	17	20	Damage to IAN or LN following an IANB
Malamed, Gagnon <i>et al.</i> ¹⁹	Double-blind random controlled trial	13	8	5	'Numbness or tingling 4 – 8 days after the procedure'
Garisto, Gaffen <i>et al.</i> ²⁷	Retrospective cohort	226 *	116	11	Oral paraesthesia following dental treatment
Pogrel ²⁰	Retrospective cohort	38 *	14	10	Damage to IAN or LN following an IANB

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*In all the included studies except Malamed, Gagnon et al., agents other than articaine and lidocaine were also studied and included in the study results. The inclusion of prilocaine, mepivacaine, bupivacaine and carbocaine explains the discrepancy between the sum of the intervention (articaine) and comparison (lidocaine) participants and that of the number of eligible participants in each study.

Table 15a Summary of study findings					
Study	Haas & Lennon ¹⁷	Gaffen & Haas ²⁸	Hillerup & Jensen ¹⁸		
Number of incidences of IAN damage with articaine	Not reported	Not reported	5		
Number of incidences of LN damage with articaine	Not reported	Not reported	24		
Number of incidences of IAN and/or LN damage with articaine	50 (33.6%)	109 (59.9%)	29 (54%)		
Number of incidences of IAN damage with lidocaine	Not reported	Not reported	3		
Number of incidences of LN damage with lidocaine	Not reported	Not reported	7		
Number of incidences of IAN and/or LN damage with lidocaine	5 (3.4%)	23 (12.6%)	10 (19%)		
Expected frequency of IAN and/or LN damage with articaine $\!$	5.3	26.5	Not reported		
Observed frequency of IAN and/or LN damage with articaine	10	42	Not reported		
Expected frequency of IAN and/or LN damage with lidocaine $\!$	3.7	23.8	Not reported		
Observed frequency of IAN and/or LN damage with lidocaine	0	6	Not reported		
*Expected frequencies calculated using the 'null hypothesis'. ²⁹					

Table 15b Summary of study findings					
Study	Pogrel ²⁶	Malamed, Gagnon <i>et al.</i> ¹⁹	Garisto, Gaffen <i>et al.</i> ²⁷	Pogrel ²⁰	
Number of incidences of IAN damage with articaine	Not reported	Not reported	Not reported	Not reported	
Number of incidences of LN damage with articaine	Not reported	Not reported	Not reported	Not reported	
Number of incidences of IAN and/or LN damage with articaine	17 (29.8%)	8 (1%)	116 (51.3%)	14 (37%)	
Number of incidences of IAN damage with lidocaine	Not reported	Not reported	Not reported	Not reported	
Number of incidences of LN damage with lidocaine	Not reported	Not reported	Not reported	Not reported	
Number of incidences of IAN and/or LN damage with lidocaine	20 (35%)	5 (1%)	11 (4.9%)	10 (26%)	
Expected frequency of IAN and/or LN damage with articaine	Not reported	Not reported	32	Not reported	
Observed frequency of IAN and/or LN damage with articaine	Not reported	Not reported	116	Not reported	
Expected frequency of IAN and/or LN damage with lidocaine	Not reported	Not reported	130	Not reported	
Observed frequency of IAN and/or LN damage with lidocaine	Not reported	Not reported	10	Not reported	
*Expected frequencies calculated using the 'null hypothesis'. ²⁹					

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Appendix 1 Glossary of abbreviations

AERS: Adverse Event Reporting System

DBRCT: Double Blind Random Controlled Trial

IAN: Inferior Alveolar Nerve

IANB: Inferior Alveolar Nerve Block

LA: Local Anaesthetic

LN: Lingual Nerve

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MeSH: Medical Sub Headings

MINORS: Methodological Index for Non-Randomised Studies

PICOS: Population, Intervention, Comparator, Outcome, Studies

PRISMA: Preferred Reporting Items for Systematic reviews and Meta-Analyses

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SIGN: Scottish Intercollegiate Guidelines Network

UCSF: University of California, San Francisco

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