Perineural dexamethasone in ultrasound-guided interscalene brachial plexus block with levobupivacaine for shoulder arthroscopic surgery in the outpatient setting: randomized controlled trial*

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Abstract  
Background and objectives: In shoulder arthroscopy, on an outpatient basis, the patient needs a good control of the postoperative pain that can be achieved through regional blocks. Perineural dexamethasone may prolong the effect of these blocks. The aim of this study was to evaluate the effect of perineural dexamethasone on the prolongation of the sensory block in the postoperative period for arthroscopic shoulder surgery in outpatient setting.

Methods: After approval by the Research Ethics Committee and informed consent, patients undergoing arthroscopic shoulder surgery under general anesthesia and ultrasound-guided interscalene brachial plexus block were randomized into Group D – blockade performed with 30 mL of 0.5% levobupivacaine with vasoconstrictor and 6 mg (1.5 mL) of dexamethasone and Group C – 30 mL of 0.5% levobupivacaine with vasoconstrictor and 1.5 mL of 0.9% saline. The duration of the sensory block was evaluated in 4 postoperative moments (0, 4, 12 and 24 hours) as well as the need for rescue analgesia, nausea and vomiting incidence, and Visual Analog Pain Scale (VAS).

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Results: Seventy-four patients were recruited and 71 completed the study (Group C, n = 37; Group D, n = 34). Our findings showed a prolongation of the mean time of the sensitive blockade in Group D (1440 ± 0 min vs. 1267 ± 164 min, p < 0.001). It was observed that Group C had a higher mean pain score according to VAS (2.08 ± 1.72 vs. 0.02 ± 0.17, p < 0.001) and a greater number of patients (68.4% vs. 0%, p < 0.001) required rescue analgesia in the first 24 hours. The incidence of postoperative nausea and vomiting was not statistically significant.

Conclusion: Perineural dexamethasone significantly prolonged the sensory blockade promoted by levobupivacaine in interscalene brachial plexus block, reduced pain intensity and rescue analgesia needs in the postoperative period.

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Introduction

Shoulder arthroscopy is currently one of the most frequent orthopedic procedures and is mostly performed in the outpatient setting. Roughly 30% of patients submitted to shoulder arthroscopy report strong intensity pain on the first postoperative day, frequently requiring opioids for analgesia.1–4 It is, therefore, a challenge to anesthesiologists, given appropriate pain management during the initial postoperative period is essential for the success of surgical outpatient treatment.1

Regional blockades with local anesthetics are usually used to reduce moderate to strong intensity postoperative pain,5,6 providing efficacious analgesia with minimum side effects when compared to opioids,2,7 albeit limited duration when given in single injections.2,3,6,7 Several drugs such as single injection block adjuvants like epinephrine,5,9 opioids,10,11 ketamine,11,12 and clonidine have been studied aiming to extend the analgesic effect.13,14

Dexamethasone, a long-action and highly potent glucocorticoid, has been shown to prolong the effects of peripheral nerve blocks when added to local anesthetics.2,7,9,15–23 This effect seems to be mediated by several mechanisms of action, such as local vasoconstriction and/or direct effect on peripheral nerves,4 although, its use in outpatient surgery...
has not been well established, given most studies assessed patients during longer inpatient stay periods.\(^7\)\(^,\)\(^{20}\)

The assumption of the present study was that dexamethasone changes the duration of interscalene brachial plexus block when added to local anesthetics. Thus, the objective of the study was to assess the effect of perineural dexamethasone on extending the duration of the sensorial block in the postoperative period of outpatient arthroscopic shoulder surgery in comparison to placebo.

**Method**

The prospective, randomized, parallel, double blind and controlled clinical trial was approved by the local Research Ethics Committee, and was conducted according to the Declaration of Helsinki and registered on Plataforma Brasil with the Ethical Appreciation Presentation Certificate (CAAE n° CAAE5759816.7.0000.5704). The study was also registered in the Brazilian Clinical Trials Registry (ReBEc) with protocol n° RBR-5v2xr. All participating patients signed a consent form. Data were collected between January and July 2017 at Complexo Hospitalar Santa Genoveva and at the Uberlândia Medical Center, both located in the city of Uberlândia, Minas Gerais. The study protocol, design, and report followed the recommendations established by the Consolidated Standards of Reporting Trials – CONSORT).

**Inclusion and exclusion criteria**

Patients submitted to shoulder arthroscopy to repair rotator cuff tear in the outpatient setting were recruited for the study. Additional inclusion criteria were adult age group (aged between 18 and 65 years), American Society of Anesthesiology physical status 1 and 2 (ASA P1 and P2), Body Mass Index (BMI) < 30 kg.m\(^{-2}\), and both genders.

Patients were excluded from the study if they presented one or more of the following criteria: skin infection on the puncture site, coagulopathy, brachial plexus neuropathy, use of systemic corticoids, routine use of opioids, diabetes, severe lung disease, psychiatric disorder and intolerance to one or more of the study drugs.

**Randomization and masking**

The Microsoft Excel® 2016 software Mersenne Twister (MT19937) algorithm was used to create random numbers to randomize the sequence between Group C (control) and D (dexamethasone) in the ratio of 1:1. Each number was printed and put into an opaque and sealed envelope.

The allocation to each group was determined upon admission to the operating room after the envelope was opened. All patients participating in the study were registered on a confidential table, in which each line was represented by one patient and pertaining medical record number, study allocation number and group to which he or she belonged to. The anesthesiologist responsible for admission, opening the envelope, and filling out the registration form was the same who prepared study medication. The anesthesiologist who performed the block was not aware of the group to which each patient belonged to and also was responsible for assessing the outcomes of the study. Patient participants were not aware to which group they belonged to.

**Intervention protocol**

Patients were monitored with 5-lead electrocardiography, pulse oximetry, non-invasive arterial pressure, gas analyzer, capnography and anesthesia depth monitoring using bispectral index (BIS\(^TM\), Medtronic, MN, US). General anesthesia was induced with 2 mg.kg\(^{-1}\) propofol, 2 mcg.kg\(^{-1}\) fentanyl, 0.6 mg.kg\(^{-1}\) rocuronium and 1 mcg.kg\(^{-1}\) lidocaine without vasoconstrictor. Target-controlled infusion of propofol was maintained (plasma concentration between 1 and 4 mcg.mL\(^{-1}\)), as was continuous infusion of remifentanil (0.1 to 0.3 mcg.kg\(^{-1}\).min\(^{-1}\)). Patients were intubated and ventilated with an oxygen/nitrous oxide mixture (50/50), with 1 L.min\(^{-1}\) fresh gas flow.

After induction of general anesthesia, antisepsis of puncture site was performed with 2% alcohol chlorhexidine. Ultrasound-guided location of interscalene brachial plexus was performed with a 13 MHz linear transducer (SonoSite NanoMaxx™, Bothel, WA – US), puncture with a 22G A50 needle (Stimuplex™, B. Braun, Melsungen AG – Germany), and injection of 30 mL of perineural anesthetic solution (2) and 1.5 mL of the study drug prepared in a separate 3 mL syringe according to randomization. In Group D, 30 mL of 0.5% levobupivacaine (Novabupi™, Laboratório Cristália, RJ, Brazil) was injected with vasoconstrictor (adrenaline 1:200.000) + 6 mg (1.5 mL) of dexamethasone (Decadron™, Aché Laboratórios Farmacêuticos S.A., SP, Brazil), and in Group C 30 mL of 0.5% levobupivacaine with vasoconstrictor (adrenaline 1:200.000) + 1.5 mL of 0.9% saline.

In order to perform shoulder arthroscopy, patients were placed in the beach chair position. During the intraoperative period, patients were medicated with 4 mg of dexamethasone and 4 mg of ondansetron for nausea and vomiting prophylaxis. To provide multimodal postoperative pain control, we associated the regional block with 2 g of dipyrrone and 100 mg of ketoprofen administered via systemic route. After finishing the arthroscopic procedure, patients were extubated and sent to the Post-Anesthesia Care Unit (PACU).

**Postoperative and outcome assessment**

Patients were assessed four times during the postoperative period: in the Postoperative Care Unit (PACU) and 4, 12 and 24 hours after the procedure, and upon hospital discharge. The duration of the sensorial block (in minutes) was established as the primary outcome and tested with cotton soaked in alcohol solution. Tactile or temperature sensations at the shoulder area were tested, and an affirmative answer to at least one of them led to considering the sensorial block had ended. When there were complaints of pain in the interval between assessments, this time was considered as the end of the sensorial block. The end of the local anesthetic solution injection was the starting time for the assessment of block duration.
The secondary outcomes assessed were: intensity of pain measured by the Visual Analogical Scale (VAS), incidence of Postoperative Nausea and Vomiting (PONV), need for rescue analgesia in the initial 24 hours (beginning with dipyrone and ketoprofen and, if required, tramadol), and rate of block failure index, characterized by complaint of pain or unchanged sensitivity on the limb operated on at the PACU. Patients were oriented to ask for analgesics as soon as they began feeling pain.

The surgeon saw patients seven days after the procedure when the presence of any neurological alteration that could be attributed to regional anesthesia was assessed.

Sample calculation and statistical analysis

The sample was calculated considering the mean duration of the block as 730 minutes with long action local anesthetic, and a prolongation effect of 75% of the sensorial block with the addition of dexamethasone, according to findings in a previous publication, and alpha error of 5% and test power of 80%. Toward that end, 34 participants per group were required for bicaudal analysis. Taking into account losses and withdrawals of 10% of participants, the sample was calculated as 74 participants.

The D’Agostino test was used to assess the normality of the distribution of the variables studied. We used the Student t-test for independent samples to compare the duration of the sensorial block, weight, age and body mass index. For VAS, comparison between groups was performed using the adjustment in a generalized linear model with Poisson distribution, followed by the Wald multiple comparison test. The Chi-Square or exact Fisher test were used to compare sex ratio, ASA classification, incidence of PONV and rescue analgesia. Quantitative variables are presented as means ± standard deviation. Nominal categorical variables are presented in absolute figures (valid percentages). We considered p < 0.05 as the significant statistical level. SAS software, version 9.3, was used for statistical analysis of data.

Results

Seventy-four patients were randomized and 71 completed the study, 34 patients in group D and 37 patients in Group C. Three patients of Group D were excluded from the analysis due to protocol violation. The details of the conduction of the study are presented in Figure 1. Patient baseline demographic data are presented in Table 1 and there were no differences between groups.

Univariate analysis was performed, and we used multivariate analysis for variables that presented significance (p < 0.01). The procedure was aimed at controlling possible result confounders. In the model studied, the only significant variable was duration of sensorial block (r² = 0.56, p < 0.001). We calculated the Odds Ratio of 0.008 with a Confidence Interval (CI) of 0.0005–0.1408 (p < 0.001) to assess dexamethasone as a protective factor for pain.

The findings of the study showed an extension of the mean duration of sensorial block in Group D (p < 0.001) (Table 2; Fig. 2). Group C presented a higher pain score mean at the 24-hour assessment (p < 0.001) (Table 2; Fig. 3). No patients complained of pain at the 0, 4 and 12-hour assessments. The incidence of PONV was higher in Group C, but not statistically significant (p = 0.16) (Table 2). In Group C, 24 patients (68.4%) required at least one dose of rescue analgesia, while in Group D, none of the patients asked for postoperative analgesia (p < 0.001) (Table 2).

There were no block failures among patients in both Groups. After seven days, at the surgeon follow-up appointment, none of the patients had neurological abnormalities or complaints that could be attributed to regional anesthesia.

Discussion

The results of the present study have shown that perineural dexamethasone extends significantly mean duration of sensorial block promoted by levobupivacaine for interscalene brachial plexus block, which corroborates similar findings of
Table 1  Baseline characteristics of the population studied.

| Variables            | Group C (n = 37) | Group D (n = 34) | p-value |
|----------------------|-----------------|-----------------|---------|
| Age (years)          | 47.2 ± 13       | 50.7 ± 11       | 0.25^a  |
| Gender (F/M) %       | 46/54           | 59/41           | 0.70^g  |
| Weight (kg)          | 70.4 ± 11       | 65.4 ± 11       | 0.16^a  |
| BMI                  | 27.2 ± 4.0      | 27.4 ± 5.2      | 0.73^a  |
| ASA Physical Status  |                 |                 | 0.97^a  |
| I                    | 23 (62.1)       | 21 (61.7)       |         |
| II                   | 14 (37.9)       | 13 (38.3)       |         |

Values presented as means ± standard deviation and numbers (%). BMI, Body Mass Index; ASA, American Society of Anesthesiologists; F, female; M, male; Group C, Control Group; Group D, Dexamethasone Group.

^a Student t-Test for independent samples was used to compare groups.

^b Chi-square test was used to compare groups.

Table 2  Distribution of outcomes studied among groups.

| Outcomes                | Group C (n = 37) | Group D (n = 34) | p-value |
|-------------------------|-----------------|-----------------|---------|
| Sensorial Block Time (min) | 1267 ± 164      | 1440 ± 0        | < 0.001^a |
| 24-hour VAS             | 2.08 ± 1.72     | 0.02 ± 0.17     | < 0.001^b |
| Rescue analgesia        | 24 (68.4)       | 0 (0)           | < 0.001^c |
| PONV                    | 2 (5.4)         | 0 (0)           | 0.16^d   |

Values presented as means ± standard deviation and numbers (%). VAS, Pain Visual Analog Scale; PONV, postoperative nausea and vomiting; Group C, Control Group; Group D, Dexamethasone Group.

^a Student t-Test for independent samples was used to compare groups.

^b Multiple comparison Wald Test was used to compare groups.

^c Chi-square test was used to compare groups.

^d Fisher Exact Test was used to compare groups.

Figure 2  Postoperative duration of sensorial block. Group C, Control Group; Group D, Dexamethasone Group. Student t-test was used for independent samples to compare groups (p < 0.001).

Figure 3  Pain Visual Analog Scale (VAS) in the initial 24 hours. Group C, Control Group; Group D, Dexamethasone Group. Comparison between groups was performed using the adjustment in a generalized linear model with Poisson distribution, followed by the Wald multiple comparison test (p < 0.001).

We also observed reduction in pain intensity, assessed by VAS, and lower consumption of rescue analgesia in the dexamethasone group during inpatient stay. These findings are in agreement with the literature, in which the use of dexamethasone was associated with reduction in the cumulative use of opioid^5,7 and lower pain scores^5,7,22 during the initial 24 postoperative hours.

Improvement in analgesia with dexamethasone for shoulder surgery^5,7,9,15-24 has been observed regardless of
intravenous or perineural administration route of the drug. In the present study, the lower intensity pain scores in the dexamethasone Group may be attributed both to the higher cumulative dose of dexamethasone used in the group and the increased duration of the sensory block. However, the increased duration of the sensory block, tested by the tactile and thermal test, cannot be explained by possible systemic effects of dexamethasone, underscoring an actual effect of the drug in prolonging the effect of local anesthetics.

To date, dexamethasone seems to be the best adjuvant method to prolong the duration of sensory blocks, superior to clonidine, epinephrine or midazolam. Moreover, its safety profile is promising, with a low risk of neurotoxicity. Perineural use in diabetic patients does not significantly change blood glucose levels, given hyperglycemia induced by steroids has only been confirmed in high dose regimens of intravenous dexamethasone. None of the patients in our study presented signs or symptoms of neurotoxicity attributed to dexamethasone, although the sample size was insufficient to detect rare results, and patients were not followed-up beyond seven days, date, of the last surgeon follow-up assessment.

Regarding PONV, studies have shown that after 24 post-operative hours, the complication is reduced in patients in whom dexamethasone was used as a perineural adjuvant. In the present study, the incidence of PONV in Group D was lower, in agreement with the literature, albeit there was no statistical significance due to its low occurrence. The effect can be attributed to a higher cumulative dose of dexamethasone in the intervention group, although the study was not designed to detect such a difference.

The major limitation of the study was restricting patient outcome follow-up to a 24-hour period because of the outpatient nature of the surgical procedure. In this scenario, all patients in the dexamethasone group still presented the sensory block when discharged, totaling a period of 1,440 minutes (24 hours). The fact prevented the assessment of the actual duration of the sensory block in patients who received perineural dexamethasone, which could have been longer than the observed. Another limitation of the study was the concomitant use of a 4-mg intravenous dose of dexamethasone in both groups as a strategy to reduce nausea and vomiting. The management may have increased the duration of analgesia in the control group and, consequently, falsely increased the mean duration of the sensory block in the same group, given that, within the interval between assessments, the sensory block was considered extinguished at the time when the patient asked for rescue analgesia.

In addition to extending the duration of the sensory block, the present study showed that adding perineural dexamethasone to single-shot block techniques is an interesting strategy to avoid using rescue analgesics in outpatient surgery, given that none of the patients of the perineural dexamethasone group required medication to control pain during inpatient stay. The absence of signs of neurotoxicity or complications associated with dexamethasone corroborates the safety of its use in this scenario. This study also tested 6 mg, a novel dose of perineural dexamethasone, which had not been described in the literature until the time of our study, and that also enabled to administer a diminished prophylactic intravenous dose of dexamethasone to patients for nausea and vomiting.

Despite the short follow-up, the present study is a baseline for future investigations in outpatient surgery, to determine exactly how long the sensory block duration is prolonged by perineural dexamethasone, the reduction in costs associated with less consumption of rescue analgesia and the incidence of possible side effects, such as late neurotoxicity.

**Conclusion**

Perineural dexamethasone, as an adjuvant to levobupivacaine, proved to be capable of prolonging the duration of the sensory component of the brachial plexus block for outpatient shoulder arthroscopy surgery. Perineural dexamethasone reduced pain intensity and patient requirement for rescue analgesia during the postoperative period.

**Conflicts of interest**

The authors declare no conflicts of interest.

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