Abstract

**Introduction:** The aim of this review was to systematize known facts about the effects of adjuvants to local anesthetics for neuraxial and regional analgesia in order to determine the adjuvant with the best effect among all others. More precisely to consider the time to onset, the strength of the effect, duration of the motor and sensory block and some additional effects such as ‘marker of intravascular injection’, safety and toxicity profile. We aimed to find an ideal adjuvant which has all these properties to a good degree.

**Methodology:** For this narrative review we searched the information in Medline, PubMed, Scopus, and Embase databases. 105 articles were identified regarding the topic, published since 1989 to 2020. Data from 105 articles about adjuvants to local anesthetics was analyzed and synthesized in this review.

**Results:** Regional methods of analgesia are becoming a crucial part of anesthesiologists’ practice and the knowledge about adjuvants is developing alongside with it, so there are more and more studies devoted to it. All of them try to find the “ideal” adjuvant, having sufficient necessary effects, but we think that due to the difference in various classes of adjuvants, some may be better than others. However, use of combination of adjuvants is not desirable at all times.

**Conclusion:** A variety of adjuvants to local anesthetics are available now, yet the data about most of them remains inconclusive, so more studies are required to found out the best adjuvants with the most desirable profile and the least adverse effects

**Key words:** Local anesthetic; Adjuvants; Pain management; Regional anesthesia.

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1. **Introduction**

Modern toolkit of anesthesiologist includes a wide spectrum of methods of regional anesthesia, such as neuraxial and peripheral nerve blocks, which can also be used for various pain management procedures. Every method has its pros and cons, but regional methods are considered the safest one's now a days. There is a variety of interventions where regional analgesia can be used to achieve specific goals depending upon the specific circumstances in each case, especially in multiple organ dysfunction syndrome (MODS). These goals can be achieved by potentiating certain effects of local anesthetics or evading the adverse effects achieved by using adjuvants to local anesthetics.
A large number of clinical studies aimed to search an ideal adjuvant to local anesthetics. For example, in urgent surgery, a time to onset of sensorimotor block, but not the duration of postoperative analgesia, can be the primary effect. The need to start the surgery urgently after an urgent block in the stressful situation and lack of time may lead to inadvertent intravascular injection of local anesthetic or crossing the maximal safe dose.

The aim of this review is to systematize known facts about the effects of adjuvants to local anesthetics in order to determine an adjuvant with the most desirable effects among all others, including the time to onset, strength of the effect, duration of the block and some additional effects such as ‘marker of intravascular injection’, safety and the toxicity profile of the agent.

2. Methodology

For this narrative review we searched for information in Medline, PubMed, SCOPUS and Embase. 105 articles were included into this review, published since 1989 to 2020. Data from these articles about adjuvants to local anesthetics was analyzed and synthesized in this article. Key word for search were: ‘adjuvant’, ‘regional analgesia’, ‘neuraxial analgesia’. Articles that provided insufficient information or had no data available for extraction, ongoing studies, the articles published in foreign languages with no translation available, and articles not available for the review through library services, were not analyzed and included in this narrative review.

We also did not included case reports.

3. Discussion

In the following paragraphs main pharmacological agents used as adjuvants to local anesthetics out of each known groups will be discussed in brief.

3.1. Vasoactive agents

3.1.1. Alpha-2-receptors agonists:

Two most widely used alpha-agonists now a days are, clonidine and dexmedetomidine. Clonidine is a derivative of imidazole, which was earlier used as an adjuvant to local anesthetics. Clonidine is alpha-2-agonist, and has analgesic, hemodynamic and sedative effect during neuraxial and peripheral blocks.

3.1.2. Clonidine: is selective to alpha-2-adrenoceptors that are located on the primary afferent paths inside the spinal cord and brain stem. It also acts through the nucleotide-ion channels. It is considered that during peripheral nerve blocks clonidine blocks conduction in A and C fibers by increasing the potassium permeability.\(^3,4\) Clonidine can also produce local vasoconstriction, that can prolong action of local anesthetic by slowing its uptake from the place of action.\(^5\) A meta-analysis has shown an increase by 2 hours in the duration of action of moderate and long-acting local anesthetics when used with clonidine.\(^6\) Another systematic review by McCartney et al. has shown longer anesthesia with moderate-acting local anesthetic, when used with clonidine\(^6\).

Several authors have shown that clonidine is capable of increasing block duration for two hours, and it was also reported, that effect was better in terms of quality of analgesia, when clonidine was used together with long-acting anesthetics. Authors also reported systemic adverse events such as hypotonia and bradycardia at high doses of clonidine.\(^5,7,8\)

3.1.3. Dexmedetomidine is an alpha-2-adrenoceptor agonist, which has hypnotic, analgesic, hypotensive effects and is capable of decreasing heart rate. Intrathecal administration of dexmedetomidine together with bupivacaine demonstrated an increase in motor and sensory block duration.\(^9\) Dexmedetomidine can prolong peripheral block for 200 min in the dose 1 mcg/kg. It prolongs duration not only of short-acting, but also long-acting local anesthetics. One of the meta-analysis have shown a presence of reversible bradycardia.\(^10\) Another meta-analysis reviewed 14 clinical studies with 848 patients, where perineural administration of dexmedetomidine vs clonidine was compared. It was shown, that dexmedetomidine prolonged motor and sensor block.\(^11,12\) Dexmedetomidine effect exceed such of clonidine in seven times.\(^13,14\) Another study by Andersen et al. compared perineural administration vs. intravascular use of dexmedetomidine. Both ways it prolonged action of ropivacaine, but perineural administration was more effective when compared to systemic use.\(^12\)

Dexmedetomidine has such adverse effects as hypotonia and bradycardia.\(^15\) There is no data about neurotoxicity of dexmedetomidine in humans and animals, but there is still discussion around the neurotoxicity of dexmedetomidine after its
perineural administration in patients with diabetic neuropathy. More studies should be performed to confirm or negate this hypothesis.16,17

3.1.4. Adrenaline is known as an adjuvant to local anesthetics for a long time and now a day is probably the most widely used adjuvant to local anesthetics for regional anesthesia. It can prolong the action of local anesthetics by decreasing systemic resorption of local anesthetic. This feature can also decrease systemic toxicity of local anesthetic, that can promote administration of higher doses and increase the block density.18,19 Adrenaline also has alpha-2 receptor mediated anti-nociceptive effect.

Negative hemodynamic reactions are rarely observed after adrenaline use as an adjuvant, but tachycardia and hypertension are possible after its intravascular injection which can be used as a marker of intravascular injection.20 Addition of adrenaline 5 mcg/ml to 1% lidocaine have shown an increase of block duration by 5 times. An increase of sensory and motor block was shown in the block of axillary nerves with lidocaine and brachial plexus with mepivacaine and bupivacaine. But addition of epinephrine does not change time of onset of action of ropivacaine.21,22

Despite some benefits of adrenaline as the adjuvant, it can significantly limit blood circulation in kidneys.23 This can have negative consequences in patients with diabetes mellitus with polyneuropathy. An increase of neurotoxicity has been shown in animal studies.24,25

3.2. Opioids

Opioids can be considered as the oldest adjuvants. Direct analgesic effects occurs after the action on antinociceptive structures, such as G-protein associated hyperpolarization of afferent sensor neurons. Opioids have a good credit in neuraxial blocks, but the data are controversial about their perineural administration.

3.2.1. Buprenorphine is a semisynthetic partial agonist of opioid receptors, which can also block voltage-dependent sodium channels. Intermediate metabolite of buprenorphine – norbuprenorphine act not only through mu-opioid receptors, but also through kappa and delta opioid receptors.26,27,28

Buprenorphine was used as an adjuvant for different regional techniques such as neuraxial, intravenous and peripheral nerve blocks.29 Studies30,31 have shown that a combination of buprenorphine and levobupivacaine can double the time of analgesia during brachial plexus block.

Several studies have shown, that buprenorphine can significantly increase average duration of analgesia for the nerve blocks of upper and lower limbs. Candido et al. published several studies, where they showed an increase of duration in 1.5 to 3 times for analgesia of subclavicular, axillary and iliac nerves with the addition of buprenorphine.32,33 Available data have shown perspectives of buprenorphine used, but more studies are yet required. Buprenorphine is relatively safe drug for use, but several studies have shown an increase of nausea and vomiting after its administration.

3.2.2. Morphine: There is no doubt about good result of intrathecal and epidural administration of morphine,34,35 while its use as an adjuvant for the peripheral nerve blocks is doubtful and did not show any effect in most of the cases.36 No benefits of perineural administration of morphine were shown when compared to intramuscular use. Taking into account the absence of evidence it is not recommended for perineural blocks.

3.2.3. Fentanyl: In studies, that compared fentanyl with local anesthetics such as lidocaine, mepivacaine, ropivacaine for peripheral nerves blocks, the authors did not show any significant difference. But there was a prolongation of effect to 18 hours in a study, where a combination of bupivacaine, fentanyl and adrenaline was used for paravertebral blocks.37 There was also positive result of using a combination of bupivacaine, fentanyl and lidocaine for cervical plexus blocks.

3.2.4. Tramadol is a weak opioid, that also block sodium, potassium, noradrenergic and serotonin receptors.37,38,39 Use of tramadol as adjuvant to local anesthetics for peripheral blocks is doubtful.40,41 So its use is recommended only for the postoperative epidural infusion.42

3.3. NMDA antagonists

Ketamine and magnesium can be considered as NMDA antagonists in this context.

3.3.1. Ketamine has local analgesic effect and can decrease nerve conduction due to the block of NMDA receptors.20 Ketamine is used for the treatment of acute pain, palliative care and chronic neuropathic pain. Lower doses can be used for
postoperative analgesia or for the decrease of exogenous opioid hyperalgesia. Combination of ketamine and opioid significantly decrease pain scores, total morphine use and postoperative desaturation in patients after thoracic surgery.

Intravenous infusion of ketamine can be used for treatment of severe neuropathic pain, such as complex regional pain syndrome, postherpetic neuralgia, trauma of the spinal cord and phantom pain. There is very little data available regarding perineural ketamine use for regional anesthesia. Lee et al. showed that ketamine did not affect sensory and motor block at its perineural use and had high amount of complications. Adverse events of ketamine include hallucinations, memory defects, panic attacks, nausea, vomiting, sleepiness, sympathetic stimulation and cardiac depression. So, ketamine is not recommended as an adjuvant to local anesthetics.

### 3.3.2. Magnesium

Modulates potassium flow into the cell through NMDA receptors and has potential benefits as an adjuvant. It was shown that addition of magnesium prolongs the duration of peripheral nerve blocks with bupivacaine, procaine and levobupivacaine. There is also an information, that magnesium can increase activity of lidocaine through the increase of threshold of A-beta fibers, as shown in experimental rat models. An important benefit of magnesium is that none of the studies reported any adverse effects related to this adjuvant. Studies have shown that addition of magnesium can prolong femoral nerve block, interscalene and axillary blocks.

Doses higher than 150 mg can be associated with higher risks of nausea and vomiting. More studies regarding neurotoxicity of magnesium should be performed before its routine use for peripheral blocks.

### 3.3.3. Midazolam

Is a short-acting benzodiazepine, and acts as a GABA-A receptor agonist in central nervous system (CNS). There is some evidence, that GABA-A receptors are located not only in CNS, but there are not enough data on the mechanism of action of benzodiazepines in peripheral nervous system. Translocator protein (TSPO), which was previously known as benzodiazepine receptor (BDR), is considered a peripheral target of midazolam. TSPO is located throughout the body, were it functions as a protein, which binds cholesterol at the external mitochondrial membrane for the steroid production. It was also determined that TSPO plays a protective role in microglial cells after the nerve damage.

A double-blind, prospective, randomized trial studied use of midazolam as an add-on during brachial plexus blocks with bupivacaine in 50 patients. Patients were divided into two groups: one group received 30 ml 0.5% bupivacaine with 2 ml of 0.9% NaCl or 30 ml 0.5% bupivacaine with 0.05 mg/kg of midazolam. Despite the small sample size, the results were significant for the average pain scores decreased and lower pain scores in 12 hours in the group receiving midazolam. The only adverse effect in midazolam group was a decrease of breathing rate 10-30 min after the injection. This decrease did not require any intervention except of addition of oxygen through the face mask.

A recent article studied efficacy of midazolam as an adjuvant during peribulbar block for cataract surgery. Midazolam in concentrations of 50 mcg/ml and 100 mcg/ml was combined with lidocaine and hyaluronidase. An average time of sensory block, but not the motor block, was significantly lower in midazolam groups. Total duration of motor and sensor analgesia was also longer in midazolam groups.

Data considering midazolam safety is limited and there is not sufficient amount of in vivo studies. Some in vitro studies indicate potential toxicity of peripheral use of midazolam, but this was not proved in in vivo studies. More randomized controlled trials are required before any recommendations considering midazolam use as the adjuvant to regional anesthesia.

### 3.4. Anti-inflammatory drugs

#### 3.4.1. Non-steroidal anti-inflammatory drugs (NSAIDs)

Act through the inhibition of COX-1, COX-2 or both enzymes. In one study ketorolac as an adjuvant to lignocaine improved analgesia in operations of the foot. In comparative study of ketorolac and dexmedetomidine ketorolac did not show benefits in the duration and onset time of the block. Animal models provide controversial conclusions regarding neurotoxicity of NSAIDs.

One study assessed the efficacy of combined use of lidocaine with dexketoprofen or acetaminophen as an adjuvant to regional intravenous analgesia. They found that both combinations decrease the time of onset and duration of sensory and motor block when
compared to control groups. There was no significant difference between paracetamol and dexketoprofen. There is yet not enough data for routine use of NSAIDs in practice for regional blocks.

**3.4.2. Dexamethasone** is a synthetic long-acting corticosteroid, with a broad spectrum of physiological effects. Except of direct anti-inflammatory action through the inhibition of COX-2, dexamethasone can act through the potassium channels of type C nociceptive fibers through the corticosteroid receptors, affecting activity of this fiber.58 A meta-analysis of placebo-controlled studies have shown, that dexamethasone can significantly prolong block of the brachial plexus. Dose of 4-10 mg can prolong action of local anesthetic from 730 min to 1306 min. Action of moderate-acting local anesthetic was increased by 175 min, motor block duration increased from 664 to 1102 min with average difference of 438 min.59

Farington et al. have shown, that the time of motor and sensor block onset in placebo and dexamethasone groups during the brachial plexus block with mepivacaine was similar, but duration of block increased from 228 to 332 min in dexamethasone group.60

*In vitro* studies and studies on mice have shown an increased risk of peripheral neurotoxicity and increased neurons dying rate. Dexamethasone as an adjuvant was studied in brachial plexus blocks, talocrural and paravertebral blocks, transversus abdominis plane (TAP) block.61 Several studies show positive results such as a decrease in pain score, decrease of postoperative opioid use and decrease of nausea and vomiting.15

A meta-analysis from 2011, that included 24 randomized controlled trials (RCT) on 2751 patients showed a decrease of postoperative pain after intravenous dexamethasone in dose of 0.1 mg/kg.62 A recent RCT studied systemic dexamethasone use together with paravertebral blocks during breast surgeries. Results were significant, and showed a decrease of postoperative pain up to 12 hrs after the operations in patients who received dexamethasone, when compared to control group.53

A decrease in pain severity score was observed after use of dexamethasone as adjuvant to blocks of anterior wall nerves at 2, 6 and 12 hours.63 DeeOliveira et al. in their meta-analysis have shown that intravenous administration of dexamethasone in dose of 0.1 mg/kg was associated with a decrease in pain score and opioid use.62

Another meta-analysis, that included 45 studies and 5796 patients, show lesser use of morphine 2-24 hrs after the operation in patients, who received 1.25 – 20 mg dexamethasone.64

But randomized clinical study, where 8 mg dexamethasone was used during 52 deliveries through cesarean section did not report a reduction of opioid use 24 hrs after the operation.66

It is interesting, that perineural administration decreases frequency of postoperative nausea and vomiting. Recent meta-analysis of 9 RCT’s with 575 patients, where dexamethasone was used as an adjuvant for TAP block, a decrease of postoperative nausea and vomiting was observed.66

Potential adverse effects of corticosteroids include immunity depression, delayed wound healing and hyperglycemia. However, two big meta-analyses did not find differences in the rate of wound infection and time of healing in study groups compared to control group.62,64 Data about hyperglycemia after dexamethasone use as an adjuvant is controversial with some studies indicating increased risk, while others show absence of any difference between the study groups.63,64 Tien et al. compared dexamethasone use in diabetic and non-diabetic patients. There was no significant change in glucose levels between 2 study groups.67

**3.5. Calcium channel blockers**

Verapamil is used as an adjuvant to local anesthetics during peripheral nerve blocks. It exerts its effect by decreasing the membrane permeability to calcium ions. Verapamil was studied as an adjuvant to bupivacaine in brachial plexus blocks and showed significant positive effect on the block duration.40

**3.6. Sodium bicarbonate**

It is an additional substance that can make the time of regional onset faster and increase the depth of block for different local anesthetics. Bicarbonate added to local anesthetics in order to make solution more alkaline, changes dissociation point and increases non-ionized part of the anesthetics, that passes well through the cellular lipid membrane, and makes the block onset faster.

Some anesthetics, such as bupivacaine and ropivacaine precipitate in the alkaline solution, so dose should be lower. Drug permeates lipid membrane faster in non-ionized state and block...
neuron depolarization. But data regarding this theory is controversial. pH of local aesthetic solution depends on the method of preparation, so it is impossible to calculate the exact amount of alkaline for the desired result.

Presence of adrenaline in the solution can alter action of sodium bicarbonate causing a decrease of the time of the onset, but also decreases the duration of anesthesia according to Sinnott et al.

3.7. Neostigmine

Neostigmine potentiates analgesic effect of local anesthetics by increasing acetylcholine levels in the nervous ends. Acetylcholine is a primary mediator in the spinal cord that participates in spinal analgesia. Neostigmine suppress afferent pain impulses to the posterior horns of the spinal cord through muscarinic M1 and M2 receptors. It is also considered that it potentiates anesthesia through increased excretion of nitrogen oxide in the spinal cord.

One of the studies have shown, that combination of 1, 2.5 or 5 mg of neostigmine and morphine led to the prolongation of anesthesia to 8 hours, that demonstrates potentiation of morphine analgesic effect, without any increase in adverse events rate.

Neostigmine as an adjuvant to intravenous analgesia for surgeries on upper limbs decreased time to block onset and prolonged time to first postoperative intake of analgesics. Postoperative bradycardia was more prominent after the administration of neostigmine. Neostigmine is less effective adjuvant when compared to dexamethasone and midazolam in brachial plexus blocks. A high amount of dose-dependent complications such as nausea and vomiting is observed during neostigmine administration.

Spinal neostigmine cause analgesia on animals and humans at a >100 mg doses, but related nausea and vomiting limited its use. Neurotoxicity of neostigmine has also been shown in its perineural use.

There is a big variety of adjuvants nowadays. They represent different classes of drugs, and so act through the different receptors and at different levels of pain conduction, so they have different set of properties, including adverse events. The number of studies regarding the adjuvant use in regional anesthesia techniques is limited, and the number varies for different adjuvants.

We tried to accumulate available data about different adjuvants according to their main properties described in the publications. Their use can be accompanied by some risk for healthcare professionals and patients. Some adjuvants have specific effects that make them different from all others. For example, main effect of sodium bicarbonate is to facilitate block onset. Physiological basis of this is well known, but preparation of working (alkaline) solution is problematic. This is particularly relevant for bupivacaine and ropivacaine, and usually anesthesiologists want to improve their action.

Alpha-2-adrenoceptor agonists significantly improve block duration, but do not affect the time of onset. Adrenaline have a clear markers of intravascular injection and can prolong action of short- and moderate-acting local anesthetics, but does not prolong action of long-acting anesthetics (the data regarding long-acting anesthetics is controversial). Adrenaline can also prevent toxic action of local anesthetics.

Most of the opioids have some benefits regarding their analgesic effect, when compared systemic and perineural administration. Buprenorphine can be an exclusion with its additional effect on sodium channels. Intrathecal use of opioids is well known, but tramadol is recommended only for epidural infusion.

Regional methods of analgesia are becoming a crucial part of anesthesiologists’ practice and the knowledge about adjuvants is developing along with it, so there are more and more studies devoted to it.

4. Conclusion

This is a developing area of modern pain management and has more questions than answers. There is a huge need of new high quality studies, including randomized-controlled studies, in this field for available adjuvants, where precise and unified methods should be used for the assessment of different effects of each particular adjuvant on different block features. Also, precise documentation of adverse effects or systemic effects is required for each adjuvant. Combination of adjuvants is not positive at all times.
Table 1: Summary table of adjuvant to local anesthetics (data in the table is based on the data from the references)

| Adjuvant            | Neuraxial blocks | Perineural | Facilitates block onset | Prolongs block | Increase block strength | Marker of I/V injection | Adverse events                                                                 | Source                      |
|---------------------|------------------|------------|-------------------------|----------------|-------------------------|--------------------------|-------------------------------------------------------------------------------|-----------------------------|
| Adrenaline          | +                | +          | -                       | +              | +++                     | +++                      | Hypertension, tachycardia, arrhythmia                                         | 20, 23, 24, 25, 18, 19, 22, 80 |
| Dexamethasone       | ?                | +          | -                       | ++             | +                       | -                        | Immunity suppression, increase risk of infections, wound healing problems, hyperglycemia | 14, 15, 60, 61, 62, 63, 64, 65, 66, 67, 80 |
| Ketamine            | +                | -          | -                       | -              | -                       | -                        | Hallucinations, memory defects, nausea, vomiting, sleepiness                  | 20, 40, 43, 44, 45          |
| Morphine            | +                | -          | -                       | -              | -                       | -                        | ?                                                                              | 34, 35, 36                  |
| Buprenorphine       | +                | +          | +                       | +              | +                       | -                        | ?                                                                              | 26, 27, 28, 29, 30, 31, 32, 33, 81, 85, 83, 84, 85                             |
| Fentanyl            | +                | -          | -                       | -              | -                       | -                        | ?                                                                              | 37                          |
| Tramadol            | +                | -          | -                       | -              | -                       | -                        | ?                                                                              | 40, 37, 38, 39, 40, 41, 42 |
| Magnesium           | +                | +          | -                       | +              | +                       | -                        | Not observed                                                                  | 17, 46, 47, 48, 49          |
| Clonidine           | +                | +          | -                       | +++            | +\+                     | +\+                      | Hypotonia, bradycardia                                                        | 3, 4, 5, 6, 7, 8            |
| Dexametomidine      | +                | +          | -                       | +++            | +\+                     | +\+                      | Hypotonia, bradycardia                                                        | 10, 11, 12, 13, 14, 15, 16, 17, 86 |
| Sodium bicarbonate  | -                | +          | +++                     | -              | -                       | -                        | ?                                                                              | 11, 12, 16, 17              |
| Neostigmine         | +                | +          | +                       | -              | -                       | -                        | Nausea, vomiting, bradycardia                                                 |                              |
| Midazolam           | -                | +          | +                       | +              | +                       | -                        | Potential neurotoxicity                                                        | 50, 51, 52, 53, 54          |
| Verapamil           | ?                | +          | ?                       | +              | -                       | -                        | ?                                                                              | ?                           |

+++ adjuvant with the best effect amongst others; ++ adjuvant with good effect; + adjuvant that have this effect; [ ] adjuvant without effect; “?” no data available
5. Conflict of interest

None declared by the authors

6. Authors’ contribution

DD: Concept, study work, manuscript editing
NO, DK, LE: Study work, manuscript writing
ZO: Manuscript editing

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