The management of postoperative pain after musculoskeletal surgery – a narrative review

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Cost-effective care amidst the rapidly rising cost of medical services necessitates the implementation of a standardised multimodal analgesia plan to aid patient care. This review aims to address the physiology and pharmacological management of postoperative pain following musculoskeletal surgery.

Keywords: multimodal analgesia, musculoskeletal pain, postoperative management

Introduction

The healthcare environment functions on cost-effective care, and the rapidly rising cost of medical services is fast becoming a major consideration in patient management. The implementation of a standardised multimodal analgesia plan can streamline planned patient care since all role players know what to expect after a surgical procedure. This review will focus on strategies for use in the management of postoperative pain following musculoskeletal surgery.

The pathophysiology of musculoskeletal pain

No discussion of the treatment of pain is complete without a review of the physiology of nociception.

Ascending tracts

Whenever an injury occurs, peripheral nociceptors are activated. These receptors comprise i) high threshold mechanoreceptors (HTM) and ii) polymodal nociceptors (PMN – stimulated by serotonin, histamine, H+ ions, cytokines, bradykinins, prostaglandins and leukotrienes). Inflammatory mediators sensitise nociceptors and lower threshold for firing. Nociceptors are free nerve endings from which impulses are carried via primary afferent Aδ and C fibres. The release of glutamate and substance P here activates the second order neurons in the substantia gelatinosa of the dorsal horn of the spinal cord. These neurons, after crossing the midline, proceed to the brain via the spinothalamic tract which eventually terminate in the thalamus – the primary somatosensory processing area. From here, tertiary order neurons project to the primary and secondary somatosensory cortexes, the insula, anterior cingulate cortex (ACC) and the prefrontal cortex for conscious perception of pain, as well as interacting with the cerebellum and basal ganglia. Activation of the ACC, for instance, leads to processing of emotionally related stimuli which may go some way as to explain many of the psychological components of pain.

Descending tracts

The purpose of the descending pathways is in modulation of pain. Neurons from the higher centres as well as projections from the spinothalamic tract connect to the peri-aqueduct grey matter (PAG – with noradrenaline as transmitter) and from there to the nucleus raphe magnus (NRM – with serotonin as transmitter). From here, it proceeds down via the ipsilateral inhibitory dorsolateral columns to synapse with the substantia gelatinosa to accomplish three objectives: i) block transmission of nociceptive stimuli via the primary neurons, ii) direct inhibition of dorsal horn cells and iii) activate inhibitory dorsal horn neurons.

Each step of neurotransmission can be accessed and modified by the use of pharmacological agents in the management of pain.

Muscle pain

Besides the normal pain pathways, muscle pain is commonly associated with both abnormal muscle tension and trigger points. Understanding myofascial pain relates to these two concepts of trigger points and muscle tension (the product of viscoelastic tone and contractile activity). Viscoelastic tone is the elastic stiffness (related to the distance the contractile elements move) and viscoelastic stiffness (effect of velocity on the contractile elements) in the absence of electromyographic activity. Contractile activity is the sum of three elements: contracture (muscle activity independent of electromyographic action), electrogenic spasm (pathological contractions arising from electrical activity in the alpha motor neurons and endplates) and electrogenic stiffness (muscle tension derived from normal electrogenic muscle contracture in individuals who are not relaxed).

Trigger points are taut bands of muscle fibres producing pain in specific designated zones in a muscle group and can be identified as active or latent trigger points.

Other physiological mechanisms contributing to muscle pain may include (but is not limited to) increased metabolism and/or decreased perfusion in the muscle fibres contributing to ischaemia, peripheral and central sensitisation, autonomic hyperactivity as well as psychological influences.
Since pain after muscle injury is transmitted via the same pathways as other somatic pain, the altered sensory and sympathetic excitability due to the secreted peptides sensitise nociceptive receptor thresholds in a process called peripheral sensitisation. This is experienced as spontaneous pain and tenderness after injury. Further repetitive stimulation of second and higher order neurons may contribute to the development of central sensitisation. It has been shown that muscle injury contributes the major portion of summation of the central sensitisation process. Central sensitisation is implicated in chronic muscle pain syndromes and development of referred pain.

Deep tissue pain does not have the plethora of scientific literature as seen with cutaneous pain models. Patients often describe it as a poorly localised, deep cramp-like pain. The convergent afferent inputs from skin, joints and viscera to the spinothalamic and other ascending tracts cause misinterpretation of information having their origin in Aδ and C fibres, resulting in referred pain. Hyperalgesia is much more likely to occur in smaller muscles or groups. It should be considered that a specific pain syndrome may have multiple origins (TMD may be myofascial or arthrogenic in origin).3

Perioperative pain management

It is safe to say that multimodal analgesia (MMA) has become the norm at most institutions. The multimodal approach consists of the use of different analgesia medication to target different receptors along the pain pathway1 to maximise analgesia but minimise side effects of any one group. Currently, the focus is on opioid-sparing techniques given the fact that opioid addiction has taken on epidemic proportions in large parts of the world. Postoperative musculoskeletal pain is a combination of pain subtypes (i.e. somatic sympathetic, neuropathic, psychogenic)6 which, with release of tumour promoting substances in turn will facilitate tumour growth. Decreased opioid consumption may favourably influence patient survival.9 Regional anaesthesia for analgesia in breast cancer surgery,10 prostate cancer11 and colon resection12 have all shown favourable outcomes.

For musculoskeletal surgery, upper and lower limb surgery is eminently suited for regional analgesia and can decrease reliance on opioids tremendously. The addition of dexamethasone or dexmedetomidine has been shown to increase the duration of nerve blocks considerably.13,14 Prolonging regional analgesia by continuous peripheral block by means of indwelling catheter is another option. It is safe if staff is trained to manage it and if the resources to maintain staffing is available. Although the addition of some adjuvants to nerve blocks have been shown to cause nerve damage in animal models, this has not been seen in human studies.15

Local anaesthesia infiltration into wound edges has been shown to be of benefit in caesarean section and laparoscopic cholecystectomy.16 The opioid-sparing effect, however, is short lived and of little clinical relevance. Where other regional techniques are not an option, infiltration is advantageous when used as part of an MMA plan. Injection into joint spaces provide analgesia comparable with regional techniques, albeit shorter lived.17 Periarticular cocktails differ from institution to institution but may include diluting local anaesthetics, steroids, opioids, nonsteroidal anti-inflammatory drugs (NSAIDs) and adrenaline.18

Intravenous lignocaine plays a role in MMA protocols at many institutions practicing according to the enhanced recovery after surgery (ERAS) guidelines. Recovery of bowel function, decreased opioid use and increased analgesia are seen after spinal surgery where intravenous (IV) lignocaine was part of the MMA protocol.19,20 The mechanism of action is thought to be that lignocaine attenuates the effect of a number of pro-inflammatory mediators.

The most obvious postoperative advantage of the use of regional nerve blocks are that mobilisation of the operated limb may commence within hours of the procedure.21 Though benefit is demonstrated in the immediate postoperative period, there seems to be little influence on later outcome, mostly due to a paucity of clinical studies.

Permanent incapacity due to neurological injury is the main fear in preventing the use of regional techniques in the perioperative period. Yet, permanent nerve damage after peripheral nerve block is extremely rare.22 The incidence of damage is 7.6:10 000 for epidural, 0–4.2:10 000 for spinals,23 and 0:1 416 for peripheral blocks.24 The advent of ultrasound-guided needle and catheter placement, as well as the use of nerve stimulation during placement has gone a long way in reducing the complication rate. Patient safety also relies on care and follow-up in the ward, in order to be alerted to the development of complications. A recognised postoperative analgesia protocol must be implemented to ensure safety at all time.
POX to the stable PGH2 – the precursor to other prostaglandins. Hydro peroxide prostaglandin, PGG2, which is then converted by COX to an unstable intermediate appropriately called prostaglandin H2 synthase. This enzyme twofold. What is referred to as cyclooxygenase (COX) is more effectively achieved within 5 minutes and peak effect is achieved within 40–60 minutes. Current evidence, however, do not universally support the routine ward use of intravenous acetaminophen over oral paracetamol, unless the duration of surgery precludes the effective use of the oral preparation.

Paracetamol does not decrease tissue inflammation and is therefore not a classic NSAID. The mechanism of action is twofold. What is referred to as cyclooxygenase (COX) is more appropriately called prostaglandin H2 synthase. This enzyme has two active sites – the COX and the peroxidase (POX) sites. Arachidonic acid is converted by COX to an unstable intermediate hydro peroxide prostaglandin, PGG2, which is then converted by POX to the stable PGH2 – the precursor to other prostaglandins (PG), leukotrienes and thromboxane. Paracetamol acts as a reducing agent on the POX site, decreasing the oxidising of COX needed for PG production. The peroxide dependant COX inhibition explains why paracetamol has a differential activity in the brain (where peroxide is low) and in the periphery (where peroxide levels are high due to cellular destruction). A second mechanism may be its direct inhibition on the COX-3 (thought to be a COX-1 splice variant) iso-enzyme, which is present in the central nervous system (CNS). This may explain the analgesic and antipyretic properties in the absence of anti-inflammatory effects. Activation of the descending serotonergic pathway and inhibition of endocannabinoid re-uptake is further postulated in the actions of paracetamol.

Nonsteroidal anti-inflammatory drugs

NSAIDs are the most frequently used pharmacological agents in the treatment of postoperative musculoskeletal pain. Their ability to inhibit COX, thereby interrupting the peripheral production of PGs from arachidonic acid, makes them suitable as analgesic drugs. PGs sensitise peripheral nociceptors so that even a small stimulus will now produce adequate pain impulse propagation to the CNS. The development of COX2-specific inhibitors were touted to be safer in that it circumvents the gastric mucosa erosion, potential renal impairment and platelet inhibition caused by inhibition of PG production via COX1, but the increased incidence of cardiac thrombotic events associated with long-term use has tempered its use (apart from patients with gastric ulcer disease). It is important to remember that many COX1 inhibitors are contra-indicated in patients with an allergy to sulphas. Concerns regarding NSAIDs increasing postoperative bleeding seem unfounded. In a recent meta-analysis examining inhibition of NSAIDs on bone healing, short-term NSAID use (<2 weeks) had no increased risk of non-union when compared to longer treatment periods (<4 weeks). Indomethacin seemed to have the greatest risk with an odds ratio (OR) of 1.66 to 9.03.

The predominant site of action for NSAIDs are peripheral, so unsurprisingly, topical administration of this group of drugs are very effective. NSAIDs have specific anti-nociceptive effects on painful conditions that involve muscles.

Systemic non-opioid analgesics

Systemic non-opioid analgesics remain the cornerstone of any MMA protocol. The opioid side effect profile is avoided and the effectiveness of these agents allow for faster mobilisation and rehabilitation. Table 1 summarises the commonly used non-opioid analgesic agents.

Paracetamol

Paracetamol is included in most MMA protocols because of its impressive safety profile (unless when administered in overdose), its cost and its opioid-sparing effect. Oral administration may cause bioavailability to vary (63–87%) and influence onset and duration of action. This is overcome by IV administration where onset of action is within 5 minutes and peak effect is achieved within 40–60 minutes. Current evidence, however, do not universally support the routine ward use of intravenous acetaminophen over oral paracetamol, unless the duration of surgery precludes the effective use of the oral preparation. Paracetamol does not decrease tissue inflammation and is therefore not a classic NSAID. The mechanism of action is twofold. What is referred to as cyclooxygenase (COX) is more appropriately called prostaglandin H2 synthase. This enzyme has two active sites – the COX and the peroxidase (POX) sites. Arachidonic acid is converted by COX to an unstable intermediate hydro peroxide prostaglandin, PGG2, which is then converted by POX to the stable PGH2 – the precursor to other prostaglandins (PG), leukotrienes and thromboxane. Paracetamol acts as a reducing agent on the POX site, decreasing the oxidising of COX needed for PG production. The peroxide dependant COX inhibition explains why paracetamol has a differential activity in the brain (where peroxide is low) and in the periphery (where peroxide levels are high due to cellular destruction). A second mechanism may be its direct inhibition on the COX-3 (thought to be a COX-1 splice variant) iso-enzyme, which is present in the central nervous system (CNS). This may explain the analgesic and antipyretic properties in the absence of anti-inflammatory effects. Activation of the descending serotonergic pathway and inhibition of endocannabinoid re-uptake is further postulated in the actions of paracetamol.

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**Table 1: Commonly used non-opioid agents**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Route*</th>
<th>Preoperative dose</th>
<th>Intraoperative dose</th>
<th>Postoperative dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paracetamol</td>
<td>IV/PO</td>
<td>1 000 mg (&gt; 50 kg)</td>
<td>1 000 mg</td>
<td>1 000 mg 6 hourly</td>
</tr>
<tr>
<td>Celecoxib</td>
<td>PO</td>
<td>400 mg</td>
<td>N/A</td>
<td>200 mg 12 hourly</td>
</tr>
<tr>
<td>Gabapentin</td>
<td>PO</td>
<td>300–1 200 mg</td>
<td>N/A</td>
<td>300–800 mg 8 hourly</td>
</tr>
<tr>
<td>Ketamine</td>
<td>IV</td>
<td>0.25–0.5 mg/kg</td>
<td>N/A</td>
<td>0.25 mg/kg/h IV</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>PO</td>
<td>600–800 mg</td>
<td>N/A</td>
<td>600 mg 6–8 hourly</td>
</tr>
<tr>
<td>Pregabalin</td>
<td>PO</td>
<td>75–150 mg</td>
<td>N/A</td>
<td>75 mg 12 hourly</td>
</tr>
</tbody>
</table>

*IV – intravenous, PO – orally, N/A – not applicable
has not produced consistent results as adjuvant in all MMA studies, but is synergistic to morphine and ketamine. Since it seems more effective when used intrathecally, it is thought that analgesia is (in part) due to direct NMDA blockade of the spinal nerves. Magnesium is a cheap addition to an MMA plan, especially when contraindications to the use of other analgesics are present.

**Alpha-2 agonists**

Dexmedetomidine and clonidine have been investigated as part of MMA protocols. These agents act as agonists on α₂ receptors in peripheral and central neuronal connections. Dexmedetomidine is an ideal opioid replacement agents since it has no respiratory depression effects. A number of studies have shown benefit in the use of dexmedetomidine as part of patient-controlled analgesia, decreasing reliance on opioids for pain relief while providing analgesia on par with opioids.44

**Systemic opioids**

Opioids have been the mainstay of postoperative analgesia for various reasons; including, simplicity of use, predictability and clinical familiarity. The presence of opioid receptors in the periphery is well documented but the efficacy of opioids are still attributed to actions within the CNS. Recently, the opioid epidemic and heightened awareness of opioid side effects have shifted the focus away from opioids. Although the updated World Health Organization (WHO) ladder for pain management reserve opioids for moderate to severe pain, or where more simple analgesics have failed, it cannot be eliminated completely from an MMA plan for procedures like hip or knee arthroplasty.45

Oral opioid preparations include hydrocodone and oxycodone – both available in a variety of combination preparations. Tramadol is a binary analgesic in that it possesses both opioid and non-opioid-related mechanisms of action. It binds to mu receptor with an affinity of about 1/6 000 of morphine, but also acts as serotonin and noradrenaline reuptake inhibitors. The total analgesic effect is due to the summation of these effects.46 Serotonin syndrome is a life-threatening drug interaction when tramadol is administered with mono-amino oxidase (MAO-I) inhibitors, selective serotonin reuptake inhibitor (SSRI) and serotonin noradrenaline reuptake inhibitor (SNRI) antidepressants. The incidence of nausea and vomiting is increased, and in Caucasians, 10% of patients will have reduced effect due to an inherited enzyme defect (needed to activate the prodrug). Furthermore, tramadol use is associated with increased opioid dependence and emergency unit attendance.47

**Skeletal muscle relaxants**

Muscle relaxants are classified into two groups: antispasmetics (dantrolene and baclofen) and antispasmodics (benzodiazepines, cyclobenzaprine, tizanidine, carisoprodol). The mechanism of action of antispasmodics (apart from benzodiazepines) is not fully understood – central NMDA blockade has been suggested for cyclobenzaprine, while tizanidine had α₂ agonist properties. There is precious little evidence to suggest the efficacy of any of the antispasmodics in acute settings, but cyclobenzaprine has been incorporated in a few MMA protocols as a preoperative adjuvant.48 Antispastics have not been used in the acute setting.

Studies has shown cyclobenzaprine to be an effective adjuvant in the management of acute lower back pain49,50 while being the least sedative of all the muscle relaxants. There is however a notable paucity in evidence for its efficacy in the acute postoperative setting (apart from incorporation into preoperative MMA protocols). Since it is structurally related to tricyclic antidepressants, its side effect profile and contra-indications are similar to these agents.

**Facing the challenge**

The implementation of clinical guidelines can go a long way to standardise practice. These guidelines should follow established evidence-based strategies to be able to champion this burden. Lin et al.51 reviewed how clinical practice guidelines (CPG) may benefit patients and the healthcare service providers in the management of musculoskeletal pain conditions. They suggested 11 recommendations and their best practice of care for musculoskeletal pain.51 Although the recommendations aim to encompass both acute and chronic pain management, the suggestions applicable to the acute setting, are listed below:

1. Care must be patient-centred. It needs to speak to the specific patient context, and allow shared decision making and effective communication.
2. Assessment of psycho-social factors are important.
3. A physical exam including complete neurological exam with assessment of mobility and muscle strength is important.
4. Patients should be educated and informed about their condition and available management options.

**Conclusion**

The need for postoperative analgesia following musculoskeletal surgery is a given. Although the ideal pharmacological management programme is not quite established, a multidisciplinary multimodal analgesia plan may go a long way in making the surgical experience far more satisfying for the patient.

**Declaration**

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**Conflict of interest**

The author declares no conflict of interest.

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