Continuous Nerve Block
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Abstract

Background: CPNB known as “perineural infusion of local anesthetic” is the catheter placement next to a peripheral nerve, preceded by local anesthetic administration through catheter, which gives anesthesia/analgesia for many days or even months. CPNBs may be delivered in a hospital setting, but mobile infusion using mobile, lightweight pumps is also a possibility. The most prevalent use of this technique is postoperative pain control. Treating intractable hiccups is considered one of the included other indications; initiating vasodilation to raise blood supply and sympathectomy after catastrophic vascularity, digit transfer/replantation, or limb salvage; relieving Raynaud’s disease vasospasm; and treating chronic pain and peripheral embolism, such as complex regional pain syndrome, phantom limb pain, trigeminal neuralgia, and cancer-induced pain. Perineural infusion may give analgesia after trauma while transit to a distant medical facility or while awaiting surgical repair. Catheter installation may be achieved utilizing various different methods, involving nerve stimulation, ultrasound guiding, paresthesia induction, fluoroscopic imaging, and basic tactile sensations (“facial click”). This article provides a review of the existing literature on continuous peripheral nerve blocks supported by evidence.

Keywords: Continuous; Nerve Block; Peripheral; Infusion

Anesthesiologists have spent over 25 years trying to ascertain if regional anesthesia, as opposed to general anesthetic, really enhances health outcomes after surgical treatment [1]. In 1946, Paul Ansbro delivered repeated supravacular injections of the brachial plexus to patients having surgery on their upper limbs to extend the duration of anesthesia. He was the one who originally characterized the CPNB [2].

CPNB is the percutaneous installation of a catheter near a peripheral nerve, followed by the injection of local anesthetic via the catheter. Consequently, “perineural local anaesthetic infusion” and CPNB are frequently used synonymously. Currently, the longest length of a peripheral nerve block with a single administration is between 8 and 24 hours. Accordingly, CPNB is a possibility whenever a sustained neural blockage is sought [3].

Installation of catheter may be achieved in a variety of ways. Depending on the technique of nerve localization, catheter implantation strategies might differ. The placement of the catheter near target nerve to offer adequate analgesia is one of the primary challenges offered by catheter methods. Confirmation (or verification) of a catheter tip’s precision. Ultrasound as a peripheral nerve block therapy is a revolutionary concept that is gaining popularity rapidly [4]. Infusion consists of a long-acting, diluted local anesthetic administered as a bolus alone, basal alone, or basal-bolus mixture [5].

Regional analgesia may be sufficient to justify its use over opioid-based analgesia due to its higher efficacy [6]. Considering the many recognised benefits of regional anaesthesia (RA), its use stays between 15 and 30 percent of current practise. This may be owing to the increased time and skill necessary for this technique, which often results in surgeons’ reluctance [7].

CPNB is utilized to extend intraoperative anesthesia. CPNB may give analgesia during trauma-related transportation or while awaiting surgical intervention. Nevertheless, the most prevalent use is postoperative analgesics [5]. This research aims to demonstrate the effectiveness of CPNB in extending the period of intraoperative nerve block anesthesia, as well as its use as a regional anesthetic modality and for postoperative analgesia. And to increase understanding of benefits and drawbacks as well as problems.

Anatomy of Peripheral Nerves

The central nervous system and the peripheral nervous system (PNS) are the two components of nervous system. CNS is composed of brain and spinal cord. PNS is primarily formed of cranial nerves, spinal nerves, their ramifications and their ganglia, that transport efferent and afferent neurones between the central nervous system and the entire body. In addition to the enteric nervous system, which comprises of nerve fibre plexuses and cell bodies in the gastrointestinal tract wall, it also contains the sympathetic trunks and ganglia [8].

Interscalene anatomy

Scalene muscles consist of three pairs of neck muscles positioned immediately laterally of the sternocleidomastoid, in front of the throat. There are anterior scalenus, medial scalenus and posterior scalenus [9].

The scalenes extend to the sternocleidomastoid in depth (SCM). They extend from the cervical vertebra to the first and second ribs. The anterior scalene is almost vertical and its higher portion is hidden by the SCM, while its lower portion is concealed by the clavicle. Along its medial margin, the carotid artery flows. The internal jugular vein, intermediate omohyoid tendon, phrenic nerve, and transverse cervical and scapular arteries are all located between the anterior scalene and the sternocleidomastoid (in...
front of scalene behind the SCM). Between the muscle and clavicle, the subclavian vein is located. Its posterior border links the brachial plexus nerve roots that transverse it with medial scalene [9].

These muscles, in conjunction with the first rib, form the scalene triangle or interscalene triangle, through which the nerves and arteries of the brachial plexus transit. Moreover, the lung pleura and superior intercostal artery are positioned posterior to the anterior scalene [10].

**Anatomy of the brachial plexus**

The ventral rami from C5 to T1 nearly exclusively serve the upper limb, which arises at this level from the trunk. Motor fibers that emanate from the plexus serve the muscles of the upper limb. A number of these muscles are located in the back as part of a complex that connects the scapula to the vertebrae. Anatomists distinguish four areas of the plexus, from the ventral ramus to the designated nerve, extending outward from the ventral ramus. As shown in figure(1), these zones are roots, trunks, divisions, and cords [11].

The name of the cords of the plexus derives from their connection to the large axillary artery, which runs parallel to the upper limb. Multiple named nerves arise straight from the cords, as seen in figure [10].

**Anatomy of lower limb**

**Lumbar plexus**

The ventral rami formation of L1–4 that may be a T12 or L5 donation (in 50 percent of cases). The plexus forms inside the psoas major (in front of the L2–5 transversal divisions) [8]: L1 is separated into upper and lower halves. The iliohypogastric and ilioinguinal nerves are distributed via the upper division. The genitofemoral nerve is composed of the lower division and an L2 branching. L2–4 split into ventral and dorsal parts. Thigh lateral cutaneous nerve is supplied by L2 and L3, while L2–4 comprise femoral nerve. The obturator nerve is created when the ventral branches unite. In addition, the L4 and L5 branches combine forming sacroccygeal plexus lumbar sacral trunk.

**Sacrococcygeal Plexus**

Constitution considerable diversity is present. L4 to L5 and S1 to S3 form sacral plexus. S4, S5, and the coccygeal nerve combine to produce the coccygeal portion [10]. Lumbar sacral trunk is generated by L4 and L5 at the psoas major medial border. that connects margin of the pelvis to the S1 vertebra. The pelvic plexus is connected to ventral rami S1–4 and S5.

Between the nerve trunks of the sacral plexus are many veins. These vessels include superior gluteal, inferior gluteal, internal pudendal and iliolumbar vessels. The branches of essential nerve include [11]: Inferior gluteal nerve (L5, S1 and S2), Superior gluteal nerve (L4 and L5, S1), Perforating cutaneous nerve (S2 and S3), Pudendal nerve (S2–4) and Posterior femoral cutaneous nerve (S1–3). Together with the femoral nerve, sciatic nerve (S2–4) the largest nerve in the body is the sciatic and serves lower leg.

**Lateral femoral cutaneous nerve:** It is a sensory nerve that originates from the L2–L3 vertebral roots. It arises laterally to the psoas muscle and caudally to the ilioinguinal nerve in the pelvis. Parallel to the iliac crest and under the iliac fascia, its path is anterolateral. It arises from the pelvis under the inguinal ligament between anterior inferior the anterior superior and iliac spines to provide the lateral thigh with sensory innervation. [12].

**Femoral nerve**

It consists of the posterior divisions of the roots L2–L3–L4. In addition, it is situated in pelvis, laterally to psoas muscle, at cleavage between iliacus and psoas muscles. It lies superficial to iliopsoas muscle as it goes below inguinal ligament. Under inguinal ligament, the femoral nerve divides into three to four centimeter-long posterior and anterior branches. From anterior division, two sensory branches and two muscular branches service pectineus and sartorius muscles. The posterior division has a solitary sensory branch, the saphenous nerve, as well as muscular branches to the quadriceps [13].

When femoral nerve passes beneath inguinal ligament, as seen in the figure (2), Medial to the nerve is the femoral artery, whereas the femoral vein is medial to the artery (VAN from medial to lateral). The nerve is surrounded by the iliac fascia, which separates it from the circulatory systems, as well as the superficial deep fascia of thigh (fascia lata), rectus femoris muscular branch stabilizes joint of hip, whereas three vasti muscular branches stabilize joint of knee [11].

**Obturator nerve**

L2-L3-L4 roots anterior divisions often give birth to a mixed (sensory and motor) nerve. It enters pelvis on psoas muscle medial side, travels anteriorly parallel to lateral pelvis to obturator foramen, and exits posteriorly. The nerve separates into anterior and posterior branches at the thigh. The anterior division descends anterior to the adductor brevis and obturator externus muscles and posterior to the pectineus and adductor longus muscles [13].

The gracilis, adductor brevis, adductor longus, and sometimes the pectineus are innervated by this nerve. Additionally, it supplies the hip joint with articular branches. Posterior division penetrates obturator externus and descends under adductor brevis and in front of adductor magnus to supply medial part of leg with skin. It contains adductor magnus, knee joint and obturator externus [11].

**Sciatic nerve**

At its origin, the sciatic nerve is 2 cm diameter and the body's biggest nerve. It exits the pelvis by the greater sciatic foramen beneath piriformis, travels between the greater trochanter and ischial tuberosity, and divides into the tibial and common fibular nerves at various levels proximal to the knee. It is separated from the gluteus maximus, to which it joins superiorly at posterior ischial surface, by nerve of quadriceps femoris [12].

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The tendon then attaches to the obturator internus, gemelli, and quadratus femoris, which separate it from the obturator externus and hip joint. The posterior femoral cutaneous nerve and the inferior gluteal artery accompany it medially. It is posterior to the adductor magnus, and the long head of the biceps femoris crosses it posteriorly. A line is drawn from medially to the midpoint of the ischial tuberosity and greater trochanter to the top of the popliteal fossa. On occasion coming straight from the sacral plexus, proximal articular branches supply the hip joint via its posterior capsule. Muscular branches arise from the biceps femoris, semitendinosus, semimembranous, and ischial portion of the adductor magnus [13].

**Tibial nerve**

Ventral branches (anterior divisions) of the fourth and fifth lumbar and the first to third sacral ventral rami form tibial nerve. From the posterior thigh and popliteal fossa, it falls to the distal edge of the popliteus. After an anterior soleus arch crossing, the popliteal artery enters the leg. It is proximally protected by the hamstring muscles in the thigh, but in the popliteal fossa it becomes much shallower and runs laterally to the popliteal arteries. Tibial nerve becomes shallow to popliteal vessels and passes to the artery medial side at knee. In the distal popliteal fossa, two heads intersection of gastrocnemius covers it [14].

Between the medial malleolus and the heel, posterior tibial vessels and tibial nerve travel in lower leg. Its proximal part is deep to gastrocnemius and soleus; however, fasciae and skin only cover its distal third, but hallucis longus sometimes covers it. Initially medial to posterior tibial arteries, the posterior tibial artery then passes behind them and descends laterally to them till it bifurcates. With the exception of its distal connection to posterior tibial surface, it rests on tibialis posterior throughout vast majority of its length. Tibial nerve split and terminates into lateral and medial plantar nerves under flexor retinaculum [12].

**Branches**
The articular, muscular, sural, medial calcaneal, and medial and lateral plantar nerves are branches of the tibial nerve [15].

**Articular branches**
The superior, inferior medial genicular arteries, and the middle genicular artery are accompanied by articular branches that supply the joint of knee. They create a plexus with an obturator nerve branch that supplies the oblique popliteal ligament. The branches that accompany the superior and inferior genicular arteries likewise feed the capsule’s medial portion. Just before to the bifurcation of the tibial nerve, the ankle joint is supplied [14].

**Muscular branches**
The gastrocnemius, plantaris, soleus, and popliteus muscles are fed by branches that grow between the heads of the gastrocnemius. The soleus nerve reaches the muscle’s superficial area. The popliteus branch descends obliquely over the popliteal vessels before curving anteriorly along the muscle’s distal border. In addition to supplying proximal tibiofibular joint, the tibia and tibialis posterior, it sends out an interosseous branch that drops near the fibula to the distal tibiofibular joint [15]. Individually or through a shared trunk, tibialis posterior, flexor digitorum longus, soleus and flexor hallucis longus are supplied by muscle branches in the leg. The flexor hallucis longus branch is associated with the fibular vessels [12].

**Saphenous nerve**

Saphenous nerve of saphenous is the biggest and tallest of cutaneous branches of femoral nerve. In the femoral triangle, it descends lateral to femoral artery entering adductor canal, at which it passes beside artery to lay medial to it. At canal’s distal end, it exits artery and rises with saphenous branch of descending genicular artery through the aponeurotic covering [15].

**Common peroneal nerve**

Originating from sciatic nerve in lower thigh region. It travels through popliteal fossa lateral section prior to twisting around fibular neck. The nerve after that split into 2 branches, deep and superficial peroneal nerves (deep to peroneus longus). It supplies [14]. There are no muscular branches, however there are sensory branches to the calf lateral cutaneous nerve and the sural nerve. Superior peroneal (previously musculocutaneous) nerve – lies. Along the interosseous membrane anterior to the tibia at the ankle, deep peroneal nerve (previously anterior tibial nerve) - runs [16].

**Pharmacology and Infusion Regimen of Local Anesthetics**

By blocking sodium ions from reaching neuronal membranes via channels or ionophores, local anesthetics inhibit neural propagation. Sodium ions are generally prevented access during the resting state of these channels. When a neuron is triggered, the channel opens or activates, allowing sodium ions to enter the cell and causing depolarization. After this abrupt shift in membrane potential, the sodium channel enters an inactive state in which further sodium ion input is blocked and active transport mechanisms return sodium ions to the cell’s outside. This repolarization restores the channel’s original state. Understanding these sodium channel states helps explain the vulnerability of diverse nerve fiber types to local anesthetics [17].

**General properties of local anesthetics**

All local anaesthetics have three main components in their molecules [18]: Intermediate ester or amide linkage, Tertiary amine, Lipophilic aromatic ring. All of these constituents confer unique therapeutic features on the molecule.

**Anesthetic potency**

Local anesthetics vary in potency, often allowing for doses between 0.5 and 4 percent. This is mostly the consequence of changes in lipid solubility (table 1). This increases diffusion across nerve sheaths and neural membranes. The aromatic ring and its substitutions, as well as those supplied to the tertiary amine, control this feature [19].
The addition of large alkyl groups boosts the potency of a parent molecule (compare tetracaine to procaine or bupivacaine to mepivacaine) (compare tetracaine to procaine or bupivacaine to mepivacaine). Several parameters, including size, type, and myelination of fiber influence the minimal dosage of local anesthetic necessary to inhibit nerve impulse transmission; pH (acidic pH inhibits blockage), frequency of nerve stimulation, and electrolyte levels are among variables that influence neuron blockage (hypokalemia and hypercalcemia antagonize blockade) \[20\].

Table (1) A general guideline for safe dosage and expected durations of various local anesthetics for the average adult patient \[21\]

<table>
<thead>
<tr>
<th>Local anesthetic preparations</th>
<th>Maximal single dosing by mass or Volume (mg/kg)</th>
<th>Maximal Dose (mg)</th>
<th>Max Volume (ml)</th>
<th>PNB (min)</th>
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<tr>
<td>Lidocaine plain</td>
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<td>0.5% (5mg/ml)</td>
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<td>Lidocaine plus epinephrine there is 5 mcg/mL of epinephrine in commercial preparation lidocaine + epi</td>
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<td>0.5% (5mg/mL) + epinephrine</td>
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<td>1% (10mg/mL) + epinephrine</td>
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<td>Bupivacaine</td>
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<td>0.25% (2.5 mg/ml)</td>
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<td>0.5% (5mg/ml)</td>
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<tr>
<td>0.25% (2.5 mg/ml) + epi</td>
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<tr>
<td>Ropivacaine/levobupivacaine</td>
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<td>0.5% (5mg/mL)</td>
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<tr>
<td>Ropivacaine</td>
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<td>0.5% + epi</td>
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<td>levobupivacaine 0.5%</td>
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Duration of action
Local anesthetics vary in their duration of action due primarily to variations in their protein affinity. The greater the tendency for protein binding, the longer the anesthetic will prolong neuronal blockage. Bupivacaine exhibits 95% protein binding compared to 55% for mepivacaine, and this is credited for the difference in their duration of neural blockade (Error! Not a valid bookmark self-reference.) \[17\].

Time for onset
Onset of local anesthetic action depends on many factors including lipid solubility and the relative concentration of the nonionized liposoluble form (B) and the ionized water-soluble form (BH +), expressed by the pKa. The pKa represents the pH at which the proportion of ionised and nonionized medication is identical. In general, less powerful and less lipid-soluble drugs have a quicker onset than more strong and more lipid-soluble compounds \[20\].

Metabolism and elimination
The intermediate chain or linkage provides a convenient basis for classification of local anesthetics, and also determines their pattern of elimination. Amides are biotransformed in the liver but esters are hydrolyzed in the bloodstream by plasma esterases \[18\].

Drugs and Infusion Regimen for Continuous Peripheral Nerve Blocks
Local Anesthetics: Review of Pharmacological Considerations
Local anaesthetics block neural transmission by preventing sodium ions from entering neuronal membranes via channels or ionophores. In a normal, relaxed state, sodium ions are denied access to these channels. When a neuron is triggered, the channel opens or activates, allowing sodium ions to enter the cell and causing depolarization. After this abrupt shift in membrane potential, the sodium channel enters an inactive state, inhibiting future inflow, and active transport mechanisms return sodium ions to the cell's outside. This repolarization returns the channel to its usual resting condition. Understanding these sodium channel states helps explain the selective susceptibility of local anesthetics for distinct neural fiber types \[22\].

Anesthetic potency
Local anaesthetics differ in efficacy, often permitting for doses between 0.5 and 4 percent. This is
mostly due to variations in lipid solubility, which facilitates diffusion across nerve sheaths and neural membranes. The aromatic ring and its substitutions, as well as those added to the tertiary amine, determine this feature. For instance, since bupivacaine is more lipid-soluble and powerful than articaine, it may be produced at a 0.5% concentration (5mg/mL) rather than a 4% concentration (40 mg/mL) [23].

**Time for onset**

A drug's increased solubility in lipids not only increases its potency, but also facilitates its passage across cell membranes. This accelerates the onset of anesthesia for local anesthetics in isolated fibers during in vitro investigations, but it must be recognized that other variables have a role in clinical practice. Vasodilating properties may promote systemic absorption before the anesthetic reaches the neuronal membrane. A high lipid solubility may restrict dispersion in tissue fluids and increase sequestration in nearby adipose or myelin sheaths [22].

**Metabolism and elimination**

Intermediate chain or linkage facilitates the categorization of local anesthetics and modifies their clearance pattern. The liver bio-transforms amides, while plasma esterases hydrolyze esters in blood [24].

**Duration of action**

Local anesthetics' duration of action differs principally owing to changes in their attraction for protein. The higher the anesthetic's propensity for protein binding, the longer the duration of neural blockage. For instance, bupivacaine demonstrates 95 percent protein binding relative to mepivacaine's 55 percent, which accounts for the disparity in their neural blockage length [25].

Time a local anesthetic lingers near neural fibers also influences the duration of anesthesia. High lipid-soluble anesthetics Sequestration locally may allow for continued neuronal membranes release, hence extending duration, although vasoconstriction of adjacent vessels is more relevant in this respect. In order to slow absorption and extend anaesthesia, vasopressors are used in a variety of formulations [22].

**Infusates**

**Local anesthetic**

During CPNB, local anesthetic is the major analgesia administered. Ropivacaine, bupivacaine, and levobupivacaine are the most often reported medications due to their long duration of action and high sensorymotor block ratio. In human studies, while ropivacaine is less stronger than levobupivacaine and bupivacaine, all three provide comparable analgesia. Nevertheless, the concentration of ropivacaine is sometimes raised by up to 50 percent to account for diminished efficacy. Notably, both ropivacaine and bupivacaine cause tissue damage, but ropivacaine causes far less harm. Unknowns are the clinical implications of these findings [24].

Bupivacaine and ropivacaine are the most often utilized local anesthetics for CPNB, and both appear to give acceptable analgesia without severe toxicity. Compared to bupivacaine, ropivacaine is considered less cardiotoxic and more "motor sparing". Nevertheless, practically all occurrences of toxicity including bupivacaine and ropivacaine involve a massive local anaesthetic single injection. At 0.15 percent bupivacaine and 0.2 percent ropivacaine concentrations, they appear to give equal analgesia with minimal variation in preservation of hand strength. Comparing ropivacaine 0.3 percent to 0.2 percent for interscalene CPNB following open rotator cuff surgery revealed a reduction in morphine intake, no disability in strength of hand and sleep quality is improved [23].

**Additives**

During CPNB, other medicines are sometimes mixed to the local to increase analgesia with no motor block worsening. Opioid addition to perineural local anesthetics is reported, however there are presently inadequate data to determine its effectiveness. Despite clonidine was often administered to CPNB in its early years, three later RCTs failed to reveal clinically meaningful effects [26].

The safety of introducing epinephrine to perineural ropivacaine is questioned due to the possibility of extended vasoconstriction, as shown by a second randomised controlled trial. Potential additional supplements have been identified; however, none are presently licenced for perineural usage in patients, and several may have undesirable systemic side effects [27].

**Infusion rates and delivery regimen**

There are three major ways to give infusions: entirely as a basal infusion, a bolus dosage, or a mix of the two. Initial findings do not support a rigorous local anesthetic administration method for every CPNB. At this time, there is insufficient data on which to make recommendations for the ideal basal rate, bolus volume, and lockout period for the factors that may influence these parameters (catheter type, catheter location, surgical procedure, etc.) [5].

**Infusion pumps**

While perineural local anaesthetic may be supplied only utilizing human-administered bolus doses, both therapeutic variables (e.g., basal infusion advantages) as well as practical issues typically mandate the use of an infusion pump. Given the variety of clinical scenarios and practise needs, there is no one best device for all cases; thus, pump choice is often based on the required device attributes. By its power source, infusion pumps may be classified randomly. However, vacuum and spring-powered devices are usable, neither is optimal for CPNB due to a number of reasons, involving extremely variable basal infusion rates and very modest local anaesthetic reservoir capacities. Pumps of elastomeric infusion were severely constrained in comparison to electronic devices; nevertheless, this is no longer the case with the emergence of modern nonelectronic pumps [5].

Non-electronic infusion pumps are frequently preferred due to their relative ease of establishing and modifying the basal infusion rate; for their low weight
and compact size; absence of audible alerts (albeit there is no notice for an infusion stop); disposability; and for their stealthy conduct (noise generated by electronic pumps may disturb patient sleep). In contrast, reusable electronic pumps employ affordable disposable "cassettes" to provide sterile infusion to specific individuals. A limited number of electrical gadgets with a single usage are available. Although the dependability of most infusion pumps is strong independent of their power source, some electronic and non-electronic devices are more reliable than others.

**Ambulatory perineural infusion**

Patients may receive CPNB outside of the hospital using a portable infusion pump, and practically every catheter type (i.e., anatomic site) has been recorded in ambulatory patients since CPNB was first reported in 1997. As patients are seldom actively observed outside of the hospital and not all patients are willing or able to undertake the added burden of caring for the catheter and pump system, patient selection criteria for ambulatory CPNB are often more strict. Patients with renal or hepatic impairment are generally precluded from outpatient perineural infusion to minimise local anaesthetic toxicity. For infusions that may influence the phrenic nerve and weaken the ipsilateral diaphragm (e.g., interscalene and paravertebral catheters), persons with heart/lung illness and obese patients who may not be able to adjust for minor hypoxia and/or hypercarbia should exercise care. Notably, age alone is not a definite exclusion criteria, since hundreds of paediatric patients have had CPNB at home without greater complication rates or severity than adult patients.

**Indications of Continuous Peripheral Nerve Blocks**

**Intraoperative anesthesia**

There have been strong advocates of regional techniques because of the hemodynamic stability and decreased coagulability. There is a relative risk reduction in respiratory depression, pneumonia, deep venous thrombosis, pulmonary embolism, and death (by 30 %) within 30 days of surgery compared to general anesthesia. There are specific areas where regional anaesthesia provides advantages of increased mental alertness and responsiveness, and appears indicated in:

Ambulatory surgery patients benefit from the enhanced alertness, better pain control, and absence of opioid side effects, and thus may be able to reach discharge criteria sooner. There is also reduction in stay in Post anesthesia care unit (PACU), nausea and better postoperative analgesia with peripheral blocks.

**Immediate postoperative pain control**

CPNB offers greater analgesia and improved functional results with a decreased incidence of nausea, vomiting, pruritus, and drowsiness when compared to opioid analgesics. In a trial of 34 patients who had either a lumbar plexus catheter with patient-controlled analgesia (PCA) or PCA alone, the lumbar plexus catheter group had a lower need for opioids, reduced pain intensity, more patient satisfaction, and less nausea and vomiting. Single-injection peripheral nerve blockade (PNB) is straightforward to conduct and has a lower incidence of bleeding and infectious problems than continuous peripheral nerve blockade (CPNB); nevertheless, the length of the block is often shorter than 24 hours, limiting analgesia. CPNB provides for prolonged postoperative analgesia compared to PNB administered in a single injection.

In lower-limb surgery, CPNB produces comparable analgesia to neuraxial block, but with less pruritus, hypotension, urine retention, and contra lateralization of the blockade. Lumbar plexus, femoral, and sciatic CPNB may be used to give site-specific analgesia with comparable rehabilitative outcomes in comparison to epidural analgesia. However, dual catheter placement (i.e. femoral/sciatic or lumbar plexus/sciatic) might take longer than epidural catheter insertion.

Continuous peripheral nerve blocks provide the opportunity to adjust to changing demands by decreasing the amount or concentration of the local anaesthetic. This adaptability minimises the requirement for a substantial initial bolus, hence lowering the danger of systemic toxicity. In addition, the elimination of dense motor and sensory obstructions minimizes the danger of falls and positioning injuries.

**Postoperative pain control for patients at home**

Despite continuous regional blockade was originally reported more than 65 years ago, home infusion was not conceivable until 1998, when lightweight, portable infusion pumps were introduced. Patients may get CPNB outside of the hospital using a mobile infusion pump. Perineural infusion is often utilised for outpatient procedures that do not need an overnight hospital stay, but it may also be used to reduce hospitalisation and/or give advantages following home or skilled nursing facility release. Criteria of patient selection for ambulatory CPNB are often more severe. For infusions that may influence the phrenic nerve and weaken the ipsilateral diaphragm (e.g., interscalene catheter), persons with heart/lung illness and obese patients who may not be able to adjust for minor hypoxia and/or hypercarbia should exercise care.

**Treatment of intractable hiccups**

Hiccups are brought on by simultaneous contractions of the diaphragmatic and intercostal muscles, accompanied by the closure of the glottis. Persistent hiccups may be more than a harmless, short-lived annoyance; hence, they need serious thought at times. Hiccuping episodes that last only a few minutes may be obnoxious, but prolonged hiccups can lead to serious complications including dehydration, weight loss, exhaustion, insomnia, delirium, depression, cardiac arrhythmias, electrocardiogram (ECG) artefacts, wound dehiscence, severe reflux esophagitis,
Using CPNB induced sympathectomy and vasodilatation to increase blood flow

After a vascular accident, digit transfer/replantation, or limb salvage, CPNBs may be utilized to treat the vasospasm associated with Raynaud’s illness \[^{31}\].

After trauma

Continuous peripheral nerve blocks may be used to offer analgesia during transfer to a distant treatment facility or while undergoing corrective surgery. After the initial stabilization of the patient, CPNB also aids in providing instant pain relief in severe trauma cases. This may be labelled "block on arrival." In addition to providing the trauma victim with rapid pain relief, it also significantly reduces the stress reaction to tissue injury \[^{31}\].

Treatment of chronic pain

Reports mention CPNB as a treatment for chronic pain, such as complex regional pain syndrome and persistent phantom limb pain, as well as pain from terminal cancer and trigeminal neuralgia, however this has not yet been verified \[^{31}\].

Complex regional pain syndrome

CRPS, originally known as reflex sympathetic dystrophy, is a pain disease with a poorly understood mechanism and unpredictable clinical history. The condition is often resistant to treatment, and its natural course is not usually good. CRPS is diagnosed based on signs and symptoms collected from the patient's medical history and physical examination \[^{32}\]. The clinical manifestations of CRPS involve pain, sensory and vasomotor abnormalities, trophic alterations, and reduced motor function. Primarily, pharmacological pain management and physical rehabilitation of limb function should be implemented as soon as feasible. If, though, limb function does not recover and significant pain persists, interventional pain management approaches may be explored \[^{30}\]. The majority of CPNB's advantages are contingent on effectively enhancing pain management, lowering opioid intake and associated negative effects, and boosting patient satisfaction. Similar to shoulder and elbow surgery, CPNB achieves and maintains powerful analgesia owing to their total innervation by nerves impacted by the perineural injection \[^{30}\].

Multiple studies have established the advantages of continuous interscalene block (cISNB) over single-shot interscalene block (sISNB) for moderately uncomfortable and highly painful surgery. The advantages of continuous infrACLavicular perineural block for elbow surgery are well-established, but effective analgesia requires a large dosage of local anaesthetic, resulting in insensible extremities. Nevertheless, for surgical operations distal to the elbow, continuous brachial blocks seem to be less effective \[^{3}\].

For continuous regional anaesthesia after shoulder and knee arthroplasty, a rapid increase in passive joint range of motion has been observed, which may result in a shorter hospital stay. There have been reports of ambulatory shoulder arthroplasty and 23-hour-stay knee and hip arthroplasty employing ambulatory continuous interscalene, femoral, and psoas compartment nerve blocks \[^{33}\].

**Contraindications of Continuous Peripheral Nerve Blocks**

**Absolute contraindications**

**Patient refusal**

It is an absolute contraindication. If regional techniques offer significant advantages in risk reduction in a specific situation, these need to be discussed with the patient and the surgeon. If the patient still refuses, other alternatives should be pursued \[^{34}\].

**Infection**

Active infection at needle insertion site is a contraindication for regional techniques because of the risk of spreading or seeding infection. Good aseptic technique remains important. Although data supporting the notion that perineural catheters are often infected, abscess at the site of a perineural catheter and septicemia are very uncommon with continuous nerve blocks. Although it is assumed that the occurrence of perineural catheter infection is very low, its actual occurrence has not been determined \[^{30}\].

This idea is explained by the short duration of perineural infusions and the frequent use of antibiotics in major orthopaedic operation. This discovery may explain reports of infections associated with the use of continuous nerve block in ambulatory surgical patients. Additionally, there is evidence that local anesthetic solutions are effective against bacteria \[^{35}\].

**An allergy to local anesthetic**

Ensure that the allergy is "real." During dental anaesthetic, some patients may report experiencing symptoms such as dizziness, nausea, etc. Inquire whether the patient had difficulty breathing, a rash, or other symptoms that might suggest a "real" allergy. If the patient has a genuine allergic response to a local anaesthetic, indicate which local anaesthetic caused the reaction. Due to their conversion to PABA, there is a greater prevalence of allergic responses with ester local anaesthetics. Local anaesthetics derived from amides have a very low frequency of allergic responses. No cross-reactions exist between amides and esters. A real allergy is an unequivocal contraindication for a peripheral nerve block using the allergenic local anaesthetic or others in the same family \[^{34}\]. Absolute contraindication is the inability to ensure sterile equipment for the block. This might lead to the introduction of infectious pathogens in healthy cells \[^{36}\].

**Relative contraindications**

1. **Coagulopathy**

Surgeons have become more liberal with chemoprophylaxis against deep venous thrombosis, increasing the population at risk. In general, it has been
regarded safe to execute peripheral nerve blocks on anticoagulated patients. Nevertheless, the possibility of perineural hematomas in these individuals with compressive neuropathy should be considered [37]. Blocks in more superficial anatomic locations, which permit direct compression of a bleeding vessel, are at lower risk of significant sequelae. It may be reasonable to proceed with a CPNB in a patient with abnormal coagulation status when the benefit of the block is felt to be greater than the risk of bleeding and any significant bleeding would be both apparent and manageable [37].

2. Pre-existent neurological deficit

May be a reason to avoid regional techniques if the deficit is unstable, as in progressive multiple sclerosis, but that risk may not be as great as once thought. Stable deficits should be carefully documented before a block is performed. Neuropathy may make elicitation of a paresthesia more difficult (as in diabetics), and a nerve stimulator or ultrasound guidance may be more appropriate in this situation [38].

3. Lack of cooperation

The agitated uncooperative patient is a poor candidate for a regional block, as in the performance of any regional technique on an awake child. Likewise, the inability to communicate with the patient because of mental status or language problems is a relative contraindication [34].

Strategies of Local Anesthetics Application in CPNB

When determining the infusion technique for a PNB, there are a great number of possible alternatives and factors to be considered. The optimal infusion protocol for the majority of PNBs would offer dependable and powerful analgesia, little motor blockage, and minimum toxicity and adverse reactions. Local anaesthetic is the centrepiece of all PNB infusion treatment modalities; however, when determining a particular plan, one must take into account the different local anaesthetic chosen, concentration of the agent, the addition of analgesic adjuncts to the infusion, continuous infusion rates, and intermittent or patient-controlled boluses, as well as PNB-specific, surgical, and patient-specific factors. Due to the intricacy of these factors, it is challenging to determine the optimal infusion approach for all PNBs [39].

Infusates

During CPNB, local anaesthetic is the major analgesia used. Although intermediate-length medications may be utilised, ropivacaine, bupivacaine, and levobupivacaine are the most often described medicines due to their extended duration of action. Despite the fact that bupivacaine and levobupivacaine are more potent than ropivacaine, all three give comparable analgesia [40]. Unknown is whether local anaesthetic dosage (mass) is the only driver of CPNB actions, or whether volume (rate) and/or concentration exhibit additional impact [28].

Infusions of diluted, long-acting local anaesthetics (such as bupivacaine, levobupivacaine, or ropivacaine) are often used. Each of these substances causes dependable analgesia and varied but limited motor block at diluted quantities. Generally, a low concentration is not only sufficient, but may even be preferable in comparison to greater concentrations [41].

Infusion strategies for PNBs

Comparing the various local anaesthetics is also challenging. In general, longer-acting local anaesthetics are preferred for PNB infusion, but it is impossible to compare individual agents because of the difficulty in identifying the relative strength of the different agents. Generally, bupivacaine and levobupivacaine are likely more strong than ropivacaine; hence, normal infusion doses for ropivacaine are greater than for the former in order to provide equivalent analgesia [42].

The best infusion strategy for PNBs also keeps unknown. Patient-controlled intermittent bolus, automated intermittent bolus, continuous infusion, or a mix of continuous infusion and intermittent bolus are all viable choices. Inherently, the mixture of continuous infusion with occasional patient-controlled bolus delivery would give the optimal balance of baseline analgesia and the capacity to react to breakthrough pain as required. Moreover, with a reduced continuous infusion rate, this method may assist reduce the total local anaesthetic provided to patients, thereby prolonging the duration of the block for patients using fixed-volume, prefilled infusion systems [43].

Infusion rates and delivery regimen

There are three major ways to give infusions: entirely as a basal infusion, a bolus dosage, or a mix of the two. Initial findings does not support a rigorous local anaesthetic administration method for every CPNB. At this time, there is insufficient data on which to make recommendations for the ideal basal rate, bolus volume, and lockout period for the factors that may influence these values (catheter type, catheter location, surgical procedure, etc.) [44].

Infusion pumps

Although perineural local anaesthetic may be supplied using just human-administered bolus doses, clinical reasons (e.g., the advantages of basal infusion) and practical concerns often mandate the use of an infusion pump. Given the variety of clinical scenarios and practise needs, there is no one best device for all cases; thus, pump choice is often based on the required device attributes [44].

By its power source, infusion pumps may be classified randomly. Although spring-powered and vacuum-powered devices are available, neither is optimal for CPNB due to a number of reasons, including extremely variable basal infusion rates and very modest local anaesthetic storage capacities, correspondingly. Till past, elastomeric infusion pumps were significantly constrained in comparison to electronic equipment. [42].
Ambulatory perineural infusion

Patients may receive CPNB outside of the hospital using a mobile infusion pump, and practically every catheter type (i.e., anatomic site) has been recorded in ambulatory patients since CPNB was first published in 1997 [45].

As patients are seldom actively supervised outside of the hospital and not all patients are ready or willing to undertake the added burden of caring for the catheter and pump system, criteria of patient selection for ambulatory CPNB are often more strict. To reduce local anaesthetic toxicity, patients with renal or hepatic impairment are often excluded from outpatient perineural infusion [46].

For infusions that may influence the phrenic nerve and weaken the ipsilateral diaphragm (e.g., interscalene and paravertebral catheters), persons with heart/lung illness and obese patients who may not be able to adjust for minor hypoxia and/or hypercarbia should exercise care. Notably, age alone is not a definite exclusion criteria, since hundreds of paediatric patients have had CPNB at home without greater complication rates or severity than adult patients [45].

References

Continuous Nerve Block


