

nocita[®]
(bupivacaine liposome injectable suspension)

REIMAGINE THE **LOOK** OF RECOVERY



Control your patient's post-operative pain with **the only FDA-approved long-acting local anesthetic** to keep them comfortable even after going home

See Prescribing Information for
full Product Information.

ALL SURGICAL PROCEDURES RESULT IN SOME DEGREE OF TISSUE TRAUMA AND ASSOCIATED PAIN

There are 3 main reasons to minimize acute, post-surgical pain¹:

- Ethical obligation to minimize pain and suffering
- Pain delays healing and return to function
- Unmanaged, acute pain can lead to chronic, maladaptive pain

Most patients are discharged from the hospital within 24-48 hours after surgery

- Need to provide analgesia for pain relief through the post-operative period at home

THE MOST EFFECTIVE CLASS OF ANALGESICS FOR PERI-OPERATIVE PAIN CONTROL

Local anesthetics (LAs) are one of the most effective means of preventing transduction and transmission of pain signals

- Block sodium channels on the nerve cell membrane
- Prevent propagation of action potentials (pain signals)
- Considered safe, with side effects generally limited to very high doses, and do not appear to delay tissue healing¹

Previous formulations have some limitations:

- Short duration of action (<8 hours) of available LAs
- Technical difficulty associated with some nerve and epidural blocks
- Complications of indwelling soaker catheters

Current guidelines advocate use of LAs for post-operative pain.^{1,2}

“Effective pain management generally involves a balanced or multimodal strategy... Local Anesthetics (LAs) are the only class of drug that renders complete analgesia.

– 2015 AAHA/AAFP Pain Management Guidelines¹

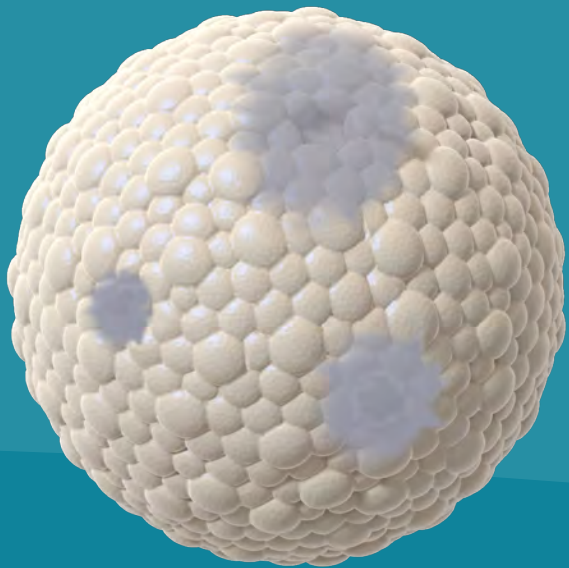


RAISING THE STANDARD OF CARE

Recovery care begins with NOCITA® (bupivacaine liposome injectable suspension)

NOCITA is a long-acting local anesthetic providing up to **72 hours of post-operative pain relief** with one dose for cranial cruciate ligament surgery in dogs and onychectomy in cats.

- Extended duration of action assists in preventing analgesic gaps in the first 72 hours post-surgery
- Provides consistent control after patient is discharged



Multivesicular liposomes

WHAT MAKES NOCITA DIFFERENT?

The extended-release bupivacaine technology used in **NOCITA** consists of multivesicular liposomes composed of hundreds to thousands of chambers encapsulating aqueous bupivacaine. The liposomes are microscopic structures designed such that bupivacaine is gradually released from vesicles over a period of time.

- Liposomes do not diffuse readily from where they are deposited
- Bupivacaine diffuses locally into surrounding tissues when it is gradually released from individual liposome vesicles

Please see Prescribing Information for full Product Information.

UP TO 72 HOURS POST-OPERATIVE PAIN CONTROL IN A SINGLE DOSE

NOCITA® (bupivacaine liposome injectable suspension) is the only long-acting, local anesthetic that controls post-op pain for up to 72 hours to help dogs undergoing cranial cruciate ligament (CCL) surgery recover comfortably, even after going home.

“In our hospital, we use fewer narcotics and plan our day to utilize the full vial, which allows for less ICU technician workload and improved hospital stay of the pet.

– Andrew Jackson, DVM, DACVS



DOG INDICATION: For single-dose infiltration into the surgical site to provide local postoperative analgesia for cranial cruciate ligament surgery in dogs.

IMPORTANT SAFETY INFORMATION FOR DOGS: NOCITA® (bupivacaine liposome injectable suspension) is for local infiltration injection in dogs only. Do not use in dogs younger than 5 months of age, dogs that are pregnant, lactating or intended for breeding. Do not administer by intravenous or intra-arterial injection. Adverse reactions in dogs may include discharge from incision, incisional inflammation and vomiting. Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics. Please see the full Prescribing Information for more detail.

CLINICAL EFFICACY IN DOGS

NOCITA® (bupivacaine liposome injectable suspension):
Proven pain control for up to 72 hours following canine CCL* surgery

Clinical efficacy study design³

- 182 client-owned dogs undergoing knee surgery
- Randomized, prospective, blinded, placebo-controlled, multicenter study
- 5.3 mg/kg by infiltration injection during surgical closure
- Time intervals for evaluating treatment success were 0-24 hours, 0-48 hours and 0-72 hours
- Success ($P < 0.05$) was defined as no pain intervention**

EFFECTIVENESS RESULTS IN DOGS

	NOCITA®	Saline	p-value
Primary endpoint 0-24 hours	68.8%	36.5%	0.0322
Secondary endpoint† 0-48 hours	64.3%	34.6%	0.0402
Secondary endpoint† 0-72 hours	61.6%	32.7%	0.0432

*Cranial cruciate ligament
**Pain intervention = rescue analgesia or score of ≥ 6 on Glasgow Composite Measure Pain Scale (Licensed from New Metrics)
†Failures carried forward from each previous interval

CONCLUSION

- Percent of treatment success for the NOCITA-treated group was statistically significantly greater than the placebo-treated group over 0-24 hours
- Greater percent successes through 48 and 72 hours support effective use of NOCITA for up to 72 hours of analgesia



SAFETY RESULTS FROM FIELD STUDY IN DOGS

NOCITA® (bupivacaine liposome injectable suspension): Demonstrated safety and was well-tolerated in dogs following cranial crucial ligament surgery³

Adverse Reaction	NOCITA® N = 123	Saline N = 59
Discharge from the Incision	4 (3.3%)	0 (0.0%)
Incisional Inflammation (erythema and/or edema)	3 (2.4%)	0 (0.0%)
Vomiting	3 (2.4%)	0 (0.0%)
Abnormalities on Urinalysis (isosthenuria ±proteinuria)	2 (1.6%)	0 (0.0%)
Increased ALP	2 (1.6%)	0 (0.0%)
Surgical Limb Edema ±Erythema	1 (0.8%)	3 (5.1%)
Soft Stool/Diarrhea	1 (0.8%)	1 (1.7%)
Inappetence	1 (0.8%)	1 (1.7%)
Fever	1 (0.8%)	0 (0.0%)



NOCITA DOSING FOR DOGS

- Administer at a dose of 5.3 mg/kg (0.4 ml/kg)
- Can be given by tissue infiltration injection during surgery for post-operative analgesia following cranial cruciate ligament surgery
- NOCITA is for single-dose administration only
- Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics



ADMINISTRATION IN DOGS

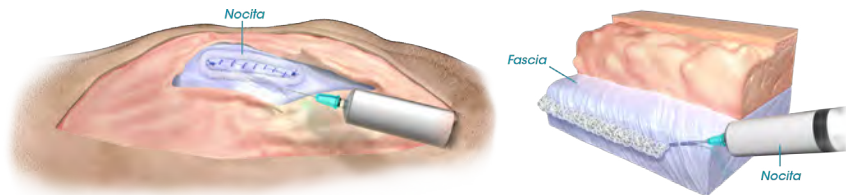
A. Incision

Medial view of the hind limb

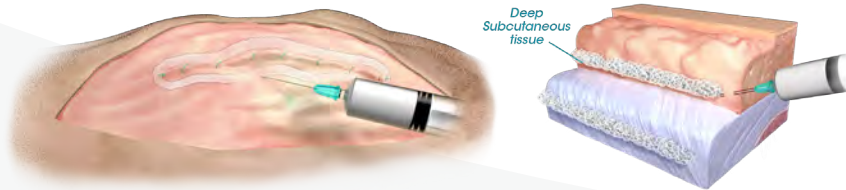
Introduce tip of the needle into the tissue

Gradually withdraw needle while injecting

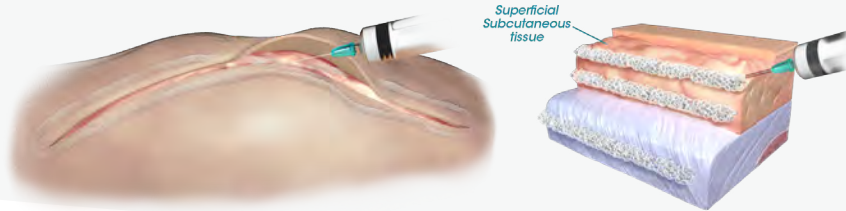
B. Fascia Layer Infiltration (post joint capsule closure)



C. Deep Subcutaneous Tissue Infiltration (post retinacular fascia closure)



D. Superficial Subcutaneous Tissue Infiltration (prior to subcuticular closure)



Administer approximately 75% of total dose volume in the surgical area (joint capsule, fascia, hardware attachment sites, osteotomies, subcutaneous layer, etc.)

Administer approximately 25% of total dose volume to ensure continuous deposition around entire incision

INDICATION: For single-dose infiltration into the surgical site to provide local postoperative analgesia for cranial cruciate ligament surgery in dogs.

IMPORTANT SAFETY INFORMATION FOR DOGS: NOCITA® (bupivacaine liposome injectable suspension) is for local infiltration injection in dogs only. Do not use in dogs younger than 5 months of age, dogs that are pregnant, lactating or intended for breeding. Do not administer by intravenous or intra-arterial injection. Adverse reactions in dogs may include discharge from incision, incisional inflammation and vomiting. Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics. Please see the full Prescribing Information for more detail.

LONG-ACTING, POST-OPERATIVE PAIN CONTROL IN A SINGLE DOSE

NOCITA® (bupivacaine liposome injectable suspension) is the only long-acting, local anesthetic that’s approved for use as a peripheral nerve block to provide up to 72 hours of regional post-operative pain control with just one dose for cats undergoing onychectomy.



CAT INDICATION: For use as a peripheral nerve block to provide regional postoperative analgesia following onychectomy in cats.

IMPORTANT SAFETY INFORMATION FOR CATS: NOCITA® (bupivacaine liposome injectable suspension) is for use as a peripheral nerve block in cats only. Do not use in cats younger than 5 months of age, that are pregnant, lactating, or intended for breeding. Do not administer by intravenous or intra arterial injection. Adverse reactions in cats may include elevated body temperature, infection or chewing/licking at the surgical site. Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics. Please see the full Prescribing Information for more detail.

CLINICAL EFFICACY IN CATS

NOCITA® (bupivacaine liposome injectable suspension): Provides up to 72 hours of regional post-operative analgesia following feline onychectomy

Clinical effectiveness study design⁴

- 241 client-owned cats undergoing owner-elective onychectomy
- Randomized, prospective, blinded, placebo-controlled, multicenter study
- 5.3 mg/kg/forelimb administered once prior to surgery as a 4-point nerve block, as described in the Product Insert
- Time intervals for evaluating treatment success were 0-24 hours, 0-48 hours and 0-72 hours
- Success (*P* < 0.05) was defined as no pain intervention*

EFFECTIVENESS RESULTS IN CATS

	NOCITA®	Saline	<i>p</i> -value
Primary endpoint 0-24 hours	75.2%	40.3%	0.0252
Secondary endpoint 0-48 hours	68.7%	34.7%	0.0395
Secondary endpoint 0-72 hours	68.4%	35.3%	0.0452

*Pain intervention = rescue analgesia or score of ≥6 on Modified UNESP-Botucatu Multidimensional Composite Pain (Brondani) Scale



TREATMENT SUCCESS

- Percent of treatment success for the NOCITA-treated group was statistically significantly greater than the placebo-treated group over 0-24 hours
- Greater percent successes through 48 and 72 hours support effective use of NOCITA for up to 72 hours of analgesia

SAFETY RESULTS FROM FIELD STUDY IN CATS

NOCITA® (bupivacaine liposome injectable suspension): Demonstrated safety as a peripheral nerve block in cats undergoing onychectomy⁴

Adverse Reaction	NOCITA® N = 120	Saline N = 121
Elevated body temperature	8 (6.7%)	5 (4.1%)
Surgical site infection	4 (3.3%)	1 (0.8%)
Chewing/licking of surgical site	3 (2.5%)	2 (1.7%)
Diarrhea	2 (1.7%)	1 (0.8%)
Injection site erythema	1 (0.8%)	0 (0.0%)
Swelling of paw; erythematous digits	1 (0.8%)	0 (0.0%)

NOTE: Surgical site is NOT Injection site




NOCITA DOSING FOR CATS



FDA-approved for use as a peripheral nerve block prior to onychectomy in cats


- Administer 5.3 mg/kg/forelimb once prior to surgery as a 4-point nerve block, as described in the Product Insert
- NOCITA is for administration only once prior to surgery
- Do not dilute NOCITA prior to use as a nerve block in cats
- Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics

ADMINISTRATION IN CATS




Legend

-  Needle insertion point
-  Drug injection point
- SpU - Styloid process of the ulna
- ACb - Accessory carpal bone



Needle withdrawal + drug injection




90°
Needle redirection to a 90° angle to the palmar plane

A.

0.14 mL/kg (35%)

Superficial Branch of the Radial Nerve

At the center of the limb, on the dorsal aspect at the level of the antebrachio-carpal joint, insert the needle subcutaneously with the bevel up (●). Advance the needle subcutaneously and inject (°) adjacent to the confluence of the accessory cephalic and cephalic veins.



Dorsal

B.

0.08 mL/kg (20%)

Dorsal Branch of the Ulnar Nerve

Palpate a groove between the accessory carpal bone (ACb, in the base of the carpal pad) and the styloid process of the ulna (SpU). Distal to this groove, insert the needle subcutaneously with the bevel up and advance the needle proximally. Inject once the tip reaches the midpoint of the groove.



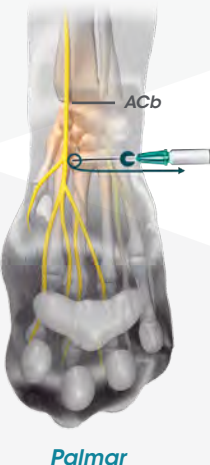
Lateral

C.

0.16 mL/kg (40%)

Median Nerve and Superficial Branch of the Palmar Branch of the Ulnar Nerve

Insert the needle subcutaneously with the bevel up lateral to the distal tip of the accessory carpal pad and advance the needle medially 2/3 the width of the limb, until the tip is located near the base of the first digit. Inject 2/3 of the volume at this point and the remaining volume while withdrawing the needle (solid teal arrow). Gently massage for 5 seconds.




Palmar

D.

0.02 mL/kg (5%)

Deep Branch of the Palmar Branch of the Ulnar Nerve

Orient the needle perpendicular to the long axis of the limb at the level of the ACb. Insert the needle subcutaneously and advance the needle laterally until it contacts the medial aspect of the ACb. Redirect the needle dorsally by rotating the needle 90°, advance it along the medial side of the ACb 2-3 mm until it penetrates the flexor retinaculum, and inject.



Palmar

INDICATION: For use as a peripheral nerve block to provide regional postoperative analgesia following onychectomy in cats.

IMPORTANT SAFETY INFORMATION FOR CATS: NOCITA® (bupivacaine liposome injectable suspension) is for use as a peripheral nerve block in cats only. Do not use in cats younger than 5 months of age, that are pregnant, lactating, or intended for breeding. Do not administer by intravenous or intra-arterial injection. Adverse reactions in cats may include elevated body temperature, infection or chewing/licking at the surgical site. Avoid concurrent use with bupivacaine HCl, lidocaine or other amide local anesthetics. Please see the full Prescribing Information for more detail.

SAFETY RESULTS FROM FIELD STUDY IN CATS

NOCITA® (bupivacaine liposome injectable suspension): Demonstrated safety as a peripheral nerve block in cats undergoing onychectomy⁴

Adverse Reaction	NOCITA® N = 120	Saline N = 121
Elevated body temperature	8 (6.7%)	5 (4.1%)
Surgical site infection	4 (3.3%)	1 (0.8%)
Chewing/Licking of surgical site	3 (2.5%)	2 (1.7%)
Diarrhea	2 (1.7%)	1 (0.8%)
Injection site erythema	1 (0.8%)	0 (0.0%)
Swelling of paw; erythematous digits	1 (0.8%)	0 (0.0%)

NOTE: Surgical site is NOT Injection site

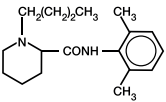
NOCITA® (bupivacaine liposome injectable suspension)

13.3 mg/mL

For use as a peripheral nerve block in cats only

Local Anesthetic

Single use vial



Caution: Federal (USA) law restricts this drug to use by or on the order of a licensed veterinarian.

Description: NOCITA® (bupivacaine liposome injectable suspension) is a sterile, non-pyrogenic white to off-white, preservative-free, aqueous suspension of multivesicular lipid-based particles containing bupivacaine. Each milliliter of NOCITA contains 13.3 mg/mL of bupivacaine. Inactive ingredients and their nominal concentrations are: cholesterol, 4.7 mg/mL; 1,2-dipalmitoyl-sn-glycero-3-phospho-rac-(1-glycerol) (DPPG), 0.9 mg/mL; triacrylin, 2.0 mg/mL; and 1,2 dieryucoylphosphatidylcholine (DEPC), 8.2 mg/mL. Bupivacaine is related chemically and pharmacologically to the amide-type local anesthetics. Chemically, bupivacaine is 1-butyl-N-(2, 6-dimethylphenyl)-2-piperidinecarboxamide with a molecular weight of 288.4. Bupivacaine chemical formula is shown in the illustration to the right.

Indication: For use as a peripheral nerve block to provide regional postoperative analgesia following onychectomy in cats.

Dosage and Administration: NOCITA is for administration only once prior to surgery. Administer 5.3 mg/kg per forelimb (0.4 mL/kg per forelimb, for a total dose of 10.6 mg/kg/cat) as a 4-point nerve block (described below) prior to onychectomy. Administration prior to surgery may provide up to 72 hours of pain control.

Prepare Dose(s):

- **Wear gloves** when handling and administering NOCITA (see **WARNINGS**).
- NOCITA should not be allowed to come into contact with topical antiseptics. When a topical antiseptic such as povidone iodine or chlorhexidine is applied, the area should be allowed to dry before NOCITA is administered.
- **Do not shake vial.** Invert the vial multiple times to re-suspend the particles immediately prior to withdrawal of the product from the vial.
- **Do not puncture the vial multiple times.** Puncture the vial stopper once with a single 25 gauge or larger needle. Use aseptic technique to sequentially attach and fill sterile syringes. Each syringe should be prepared for single patient use only. Discard the vial after all doses are withdrawn.
- Following withdrawal from the vial into a syringe, NOCITA may be stored at controlled room temperature of 68° F to 77° F (20° C to 25° C) for up to 4 hours. Because the formulation does not contain preservative, the syringe(s) must be discarded after 4 hours.
- Do not dilute NOCITA prior to use as a nerve block in cats.
- Do not mix with water or other hypotonic solutions as it will result in disruption of the liposomal particles (see **CLINICAL PHARMACOLOGY**).
- Do not mix NOCITA with other local anesthetics or other drugs prior to administration (see **PRECAUTIONS**).
- Use a 25 gauge or larger bore needle for administration.

Dose Administration:

Aspirate prior to injecting to prevent intravascular administration (see **CONTRAINDICATIONS**).

Table C-1. Dose Administration for One Forelimb.¹

Legend	Abbreviations
Needle insertion point	SpU - Styloid process of the ulna
Needle advancement	ACb - Accessory carpal bone
Drug injection point	
Needle withdrawal + drug injection	
Needle redirection to a 90° angle to the palmar plane	
Dose Volume per Injection (% of total 0.4 mL/kg/forelimb volume) and Description	
A. 0.14 mL/kg (35%) Superficial Branch of the Radial Nerve: At the center of the limb, on the dorsal aspect at the level of the antebrachio-carpal joint, insert the needle subcutaneously with the bevel up (*). Advance the needle subcutaneously as depicted by the dotted line and arrow and inject (e) adjacent to the confluence of the accessory cephalic and cephalic veins.	
B. 0.08 mL/kg (20%) Dorsal Branch of the Ulnar Nerve: Palpate a groove between the accessory carpal bone (ACb), in the base of the carpal pad) and the styloid process of the ulna (SpU). Distal to this groove, insert the needle subcutaneously with the bevel up and advance the needle proximally. Inject once the tip reaches the midpoint of the groove.	
C. 0.16 mL/kg (40%) Median Nerve and Superficial Branch of the Palmar Branch of the Ulnar Nerve: Insert the needle subcutaneously with the bevel up lateral to the distal tip of the accessory carpal pad and advance the needle medially 2/3 the width of the limb, until the tip is located near the base of the first digit. Inject 2/3 of the volume at this point and the remaining volume while withdrawing the needle (solid grey arrow). Gently massage for 5 seconds.	
D. 0.02 mL/kg (5%) Deep Branch of the Palmar Branch of the Ulnar Nerve: Orient the needle perpendicular to the long axis of the limb at the level of the ACb. Insert the needle subcutaneously and advance the needle laterally until it contacts the medial aspect of the ACb. Redirect the needle dorsally by rotating the needle 90°; advance it along the medial side of the ACb 2-3 mm until it penetrates the flexor retinaculum, and inject.	

Contraindications: Do not administer by intravenous or intra-arterial injection. If accidental intravascular administration occurs, monitor for cardiovascular (dysrhythmias, hypotension, hypertension) and neurologic (tremors, ataxia, seizures) adverse reactions.

Do not use for intra-arterial injection. In humans, local anesthetics administered into a joint may cause chondrolysis.

Warnings: Not for use in humans. Keep out of reach of children.

NOCITA is an amide local anesthetic. In case of accidental injection or accidental topical exposure, contact a physician and seek medical attention immediately.

Wear gloves when handling vials to prevent accidental topical exposure.

Precautions: Do not administer concurrently with bupivacaine HCl, lidocaine or other amide local anesthetics. A safe interval from time of bupivacaine HCl, lidocaine or other amide local anesthetic administration to time of NOCITA administration has not been determined. The toxic effects of these drugs are additive and their administration should be used with caution including monitoring for neurologic and cardiovascular effects related to toxicity. The safe use of NOCITA in cats with cardiac disease has not been evaluated. The safe use of NOCITA in cats with hepatic or renal impairment has not been evaluated. NOCITA is metabolized by the liver and excreted by the kidneys. The ability of NOCITA to achieve effective anesthesia has not been evaluated. The safe use of NOCITA in cats for surgical procedures other than onychectomy has not been evaluated.

The safe use of NOCITA has not been evaluated in cats younger than 5 months old. The safe use of NOCITA has not been evaluated in cats that are pregnant, lactating, or intended for breeding.

Adverse Reactions: Safety was evaluated in 120 NOCITA treated cats and 121 saline (placebo) treated cats in a field study in cats undergoing onychectomy. Cats enrolled in the study were 5 months to 10 years of age, and weighed 2.0 to 9.3 kg. NOCITA was administered as a 4-point peripheral nerve block at a dose of 5.3 mg/kg per forelimb (0.4 mL/kg per forelimb).

Table C-2: Adverse Reactions Reported During the Study in the Safety Population (any cat that received treatment)

Adverse Reaction	NOCITA (n = 120)	Saline (n = 121)
Elevated body temperature*	8 (6.7%)	5 (4.1%)
Surgical site infection	4 (3.3%)	1 (0.8%)
Chewing/licking of surgical site	3 (2.5%)	2 (1.7%)
Diarrhea	2 (1.7%)	1 (0.8%)
Injection site erythema	1 (0.8%)	0 (0.0%)
Swelling of paw; erythematous digits	1 (0.8%)	0 (0.0%)

Note: If an animal experienced the same event more than once, only the first occurrence was tabulated.

*Elevated body temperature was defined as temperature $\geq 103^\circ\text{F}$ on Day 3 and normal before surgery. One of the NOCITA treated cats had an infection of one surgical site. No other cat with elevated body temperature showed evidence of infection or illness.

Eight cats, 4 in each group, had normal platelet counts before treatment on Day 0 and platelet counts below the reference range (155,000-641,000/ μL) on Day 3. The 4 cats treated with NOCITA had platelet counts of 42,000 to 100,000/ μL , and the 4 cats in the saline group had platelet counts of 114,000 to 149,000/ μL . Decreased platelet counts were not associated with clinical signs.

In a pilot study with 62 cats undergoing onychectomy (31 cats treated with NOCITA and 31 with saline), one NOCITA treated cat had a motor deficit (unilateral knuckling) which resolved by the next morning following surgery. Another NOCITA treated cat had bruising at the injection sites.

To report suspected adverse drug events and/or to obtain a copy of the Safety Data Sheet (SDS) or for technical assistance, call Aratana Therapeutics at 1-844-640-5500.

For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS or online at <http://www.fda.gov/AnimalVeterinary/SafetyHealth>

Clinical Pharmacology: Bupivacaine is an amide, non-opioid local anesthetic. It provides local analgesia by deactivating sodium channels on the nerve membrane, preventing the generation and propagation of nerve impulses. It is only present in small concentrations as uncharged molecules at tissue pH as it is a base with pKa of 8. This un-ionized form provides a lipophilicity that permits the drug to traverse across the nerve cell membrane and upon entering the cell, binds to the intracellular portion of voltage-gated sodium channels and blocks sodium influx into nerve cells, which prevents depolarization. Without depolarization, no initiation or conduction of a pain signal can occur.

Lipid Formulation

Liposomal encapsulation or incorporation in a lipid complex can substantially affect a drug's functional properties relative to those of the unencapsulated or nonlipid-associated drug. In addition, different liposomal or lipid-complexed products with a common active ingredient may vary from one another in the chemical composition and physical form of the lipid component. Such differences may affect functional properties of these drug products. Do not substitute with other bupivacaine formulations.

After injection of NOCITA, bupivacaine is released from the multivesicular liposomes over a period of time.

Pharmacokinetics

The pharmacokinetic characterization associated with bupivacaine after subcutaneous NOCITA (bupivacaine liposome injectable suspension) or bupivacaine HCl solution administered to cats evaluated for 168 hours is provided in Table C-3.

Table C-3. Plasma pharmacokinetic parameters for bupivacaine after single subcutaneous administration of NOCITA and bupivacaine HCl solution in male and female cats in a laboratory study.

PK Parameter	NOCITA® 3 mg/kg	NOCITA® 9 mg/kg	NOCITA® 15 mg/kg	bupivacaine HCl 1 mg/kg
N	6	6	6	6
T _{1/2} ^{a,b} (hr)	12.5 (1-48)	10 (1-24)	1.5 (1-24)	1 (1-4)
T _{1/2} ^b (hr)	108 (72-144)	120 (72-168)	144 (120-168)	18 (12-24)
C _{max} (ng/mL)	311.4 (82.2-565)	620.2 (374-892)	709.7 (462-1090)	263.9 (60.5-506)
AUC ₀₋₂₄ ^c (ng*hr/mL)	11347 (5176-15767)	32561 (19390-47532)	38475 (26460-48252)	1608 (314-2363)

* 5.3 mg/kg NOCITA bupivacaine base is equal to 6 mg/kg bupivacaine HCl. NOCITA doses in this table are in the bupivacaine HCl equivalent.

^a Median (range)

^b Mean (range)

T_{1/2}^a = time to maximum plasma concentration

T_{1/2}^b = time to last quantifiable plasma concentration

C_{max} = maximum plasma concentration

AUC₀₋₂₄ = area under the curve from the time of dosing to the last quantifiable plasma concentration

Following a single subcutaneous dose of NOCITA, there was a less than dose proportional increase in C_{max} and AUC₀₋₂₄ across the dose range tested (3-15 mg/kg). There was a high variability in all reported parameters. Half-life is not reported for NOCITA in cats because the prolonged absorption confounds the estimation of the terminal elimination phase. Therefore, T_{1/2} is included as a more appropriate measure of the duration of quantifiable plasma concentration.

Effectiveness: Effectiveness was demonstrated in a multi-center, placebo-controlled, randomized and masked field study in client-owned cats undergoing bilateral forelimb onychectomy. In this study, 241 cats were enrolled in the study and randomized to treatment with NOCITA (n = 120) or saline (placebo, n = 121).

Cats received an opioid analgesic just prior to general anesthesia and surgery. The nerve block injection sites were shaved and a standard surgical preparation with chlorhexidine or povidone iodine was used. Prior to onychectomy, NOCITA was administered as a 4-point nerve block (see **DOSING INSTRUCTIONS**).

Pain was assessed by trained observers using a modified version of the UNESP-Botucatu Multidimensional Composite Pain Scale for up to 72 hours following extubation. Pain assessments were conducted prior to surgery, and at 0.5, 1, 2, 4, 8, 12, 24, 30, 36, 48, 56 and 72 hours post-surgery. Cats with a composite pain score ≥ 6 or that were determined to be painful by the assessor received rescue analgesic medication. Cats were classified as treatment failures. After receiving rescue analgesia, cats did not have further pain assessments performed. The primary variable for effectiveness was evaluated over the first 24-hour time interval. The percent of treatment success for NOCITA was significantly greater than saline for the 0-24 hour time interval ($p = 0.0252$). The 0-48 hour and 0-72 hour time intervals were evaluated as secondary variables and support effective use of NOCITA for up to 72 hours of analgesia.

Table C-4. Number and Percent Effectiveness for NOCITA and Saline (Placebo) Groups at each Time Interval

Time Interval for Pain Evaluation	NOCITA	Saline
0-24 hours	88/117 (75.2%)	48/119 (40.3%)
0-48 hours	79/115 (68.7%)	41/118 (34.7%)
0-72 hours	78/114 (68.4%)	42/119 (35.3%)

The per protocol populations for effectiveness varied for each pain assessment time interval because of protocol deviations affecting only one of the three time intervals for some cats.

Animal Safety: In a 22 day laboratory study, 40 healthy cats (4 cats/sex/group) aged 5-6 months were administered negative control (2.37 mL/kg saline), active control (5.3 mg/kg bupivacaine HCl), or NOCITA at 10.6, 21.2, or 31.5 mg/kg via injection using a suprainguinal approach for a femoral nerve block of the right hindlimb on Days 0, 9 and 18. These NOCITA doses correspond to 1, 2 and 3 times the maximum labeled total dose of 10.6 mg/kg/cat (representing 2, 4 and 6 times the maximum labeled dose of 5.3 mg/kg/forelimb).

Two cats died during the study. One male in the active control group died during recovery from anesthesia after the second dose and no definitive cause of death was determined. One female in the 31.5 mg/kg group was euthanized on Day 15. This cat developed a suppurative, open, necrotic wound over the region of the right stifle after the second dose administration.

For the cats who survived the study, there were no clinically relevant treatment-related effects on electrocardiograms, hematology, serum chemistry, urinalysis, coagulation, and organ weights. Right hindlimb impairment was expected because the entire dose was administered as a femoral nerve block. Right hindlimb impairment occurred in 23 of the 24 NOCITA cats which persisted for 1-5 days; 2 negative control cats which persisted for 1 day; and none of the active control cats. Left hindlimb impairment was observed the day after the first dose in one cat in the 21.2 mg/kg group. NOCITA treatment-related findings were observed on histopathology of soft tissue and the femoral nerve at the injection sites. Injection site soft tissue histopathology findings included subacute or chronic inflammation, mineralization, myofiber degeneration and myofiber necrosis. Injection site femoral nerve histopathology findings included subacute or chronic inflammation.

Sporadic clinical observations and histopathology findings throughout both negative and active control groups and NOCITA groups included: soft or watery or mucoid stool; inguinal swelling on the right hindlimb noted after only the first dose; abrasions or scabbing noted at the right abdominal and inguinal regions as well as on the right hindlimb and at the right stifle; histopathology findings at or near the injection site or right stifle included ulceration and suppurative crusts on the skin, histopathology findings at the injection site of subcutaneous foreign material and fibrosis, and myofiber regeneration.

Storage Conditions: Unopened vials should be stored refrigerated between 36° F to 46° F (2° C to 8° C). NOCITA may be held at a controlled room temperature of 68° F to 77° F (20° C to 25° C) for up to 30 days in sealed, intact (unopened) vials. Do not re-refrigerate. **Do Not Freeze.**

How Supplied: 13.3 mg/mL bupivacaine liposome injectable suspension in 10 mL or 20 mL single use vial.

10 mL supplied in 4-vial carton. 20 mL supplied in a single vial carton and 4-vial carton.

NADA 141-461, Approved by the FDA
US Patent: 8,182,835
US Patent: 8,834,921
US Patent: 9,205,052



Manufactured for: Aratana Therapeutics, Inc., Leawood, KS 66211
Additional Information is available at www.aratana.com or by calling Aratana Therapeutics at 1-844-272-8262.

NOCITA is a registered trademark of Aratana Therapeutics, Inc.
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Reference:

1. Location and relative volumes based on: Enomoto M, Lascelles BDX and Gerard MP. Defining the local nerve blocks for feline distal thoracic limb surgery: a cadaveric study. *Journal of Feline Medicine and Surgery*; 2016 18 (10): 838-845.

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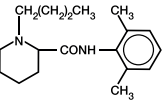
NOCITA® (bupivacaine liposome injectable suspension)

13.3 mg/mL

For local infiltration injection in dogs only

Local Anesthetic

Single use vial



Caution: Federal (USA) law restricts this drug to use by or on the order of a licensed veterinarian.

Description: NOCITA® (bupivacaine liposome injectable suspension) is a sterile, non-pyrogenic, white to off-white, preservative-free, aqueous suspension of multivesicular lipid-based particles containing bupivacaine. Each milliliter of NOCITA contains 13.3 mg of bupivacaine. Inactive ingredients and their nominal concentrations are: cholesterol, 4.7 mg/mL; 1,2-dipalmitoyl-sn-glycero-3-phospho-rac-(1-glycerol) (DPPG), 0.9 mg/mL; triacrylin, 2.0 mg/mL; and 1,2 dieryucoylphosphatidylcholine (DEPC), 8.2 mg/mL. Bupivacaine is related chemically and pharmacologically to the amide-type local anesthetics. Chemically, bupivacaine is 1-butyl-N-(2, 6-dimethylphenyl)-2-piperidinecarboxamide with a molecular weight of 288.4. Bupivacaine structural formula is shown in the illustration to the right.

Indication: For single-dose infiltration into the surgical site to provide local postoperative analgesia for cranial cruciate ligament surgery in dogs.

Dosage and Administration: NOCITA is for single dose administration only. A dose of 5.3 mg/kg (0.4 mL/kg) is administered by infiltration injection into the tissue layers at the time of incisional closure. A single dose administered during surgical closure may provide up to 72 hours of pain control.

Dosing Instructions:

- **Wear gloves** when handling and administering NOCITA (see **WARNINGS**).
- NOCITA should not be allowed to come into contact with topical antiseptics. When a topical antiseptic such as povidone iodine or chlorhexidine is applied, the area should be allowed to dry before NOCITA is administered into the surgical site.
- **Do not shake vial.** Invert the vial multiple times to re-suspend the particles immediately prior to withdrawal of the product from the vial.
- **Do not puncture the vial multiple times.** Puncture the vial stopper once with a single 25 gauge or larger needle. Use aseptic technique to sequentially attach and fill sterile syringes for dosing. Each syringe should be prepared for single patient use only. Discard the vial after all doses are withdrawn.

- Following withdrawal from the vial into a syringe, NOCITA may be stored at controlled room temperature of 68° F to 77° F (20° C to 25° C) for up to 4 hours. Because the formulation does not contain preservative, the syringe(s) must be discarded after 4 hours.
- If the dose volume of NOCITA (0.4 mL/kg) is not sufficient to cover the surgical site, add up to an equal volume of normal (0.9%) sterile saline or Lactated Ringer's solution. If saline or Lactated Ringer's is added to the NOCITA dose, administer the entire volume by tissue infiltration into the surgical site. Do not mix with water or other hypotonic solutions as it will result in disruption of the liposomal particles (see **CLINICAL PHARMACOLOGY**).

Do not mix NOCITA with other local anesthetics or other drugs prior to administration (see **PRECAUTIONS**).

- Use a 25 gauge or larger bore needle for administration.
- Administer by infiltration injection: Inject slowly into the tissues using an infiltration injection technique. To obtain adequate coverage, infiltrate all of the tissues in each surgical closure layer. Aspirate frequently to prevent intravascular administration (see **CONTRAINDICATIONS**).

Contraindications: Do not administer by intravenous or intra-arterial injection. If accidental intravascular administration occurs, monitor for cardiovascular (dysrhythmias, hypotension, hypertension) and neurologic (tremors, ataxia, seizures) adverse reactions.

Do not use for intra-arterial injection. In humans, local anesthetics administered into a joint may cause chondrolysis.

Warnings: Not for use in humans. Keep out of reach of children.

NOCITA is an amide local anesthetic. In case of accidental injection or accidental topical exposure, contact a physician and seek medical attention immediately.

Wear gloves when handling vials to prevent accidental topical exposure.

Precautions: Do not administer concurrently with bupivacaine HCl, lidocaine or other amide local anesthetics. A safe interval from time of bupivacaine HCl, lidocaine or other amide local anesthetic administration to time of NOCITA administration has not been determined. The toxic effects of these drugs are additive and their administration should be used with caution including monitoring for neurologic and cardiovascular effects related to toxicity.

The safe use of NOCITA in dogs with cardiac disease has not been evaluated. The safe use of NOCITA in dogs with hepatic or renal impairment has not been evaluated. NOCITA is metabolized by the liver and excreted by the kidneys.

The ability of NOCITA to achieve effective anesthesia has not been studied. Therefore, NOCITA is not indicated for pre-incisional or pre-procedural loco-regional anesthetic techniques that require deep and complete sensory block in the area of administration.

The safe use of NOCITA for surgical procedures other than cranial cruciate ligament surgery has not been evaluated (see **ANIMAL SAFETY** and **ADVERSE REACTIONS**).

The safe use of NOCITA has not been evaluated in dogs younger than 5 months old.

The safe use of NOCITA has not been evaluated in dogs that are pregnant, lactating, or intended for breeding.

Adverse Reactions: Safety was evaluated in 123 NOCITA treated dogs and 59 saline (placebo) treated dogs in a field study in dogs that underwent cranial cruciate ligament stabilization surgery. Dogs enrolled in the study were 1-13 years of age, and weighed 3.4 to 61.3 kg. NOCITA was administered by infiltrative injection at the surgical site at a dose of 5.3 mg/kg (0.4 mL/kg).

Table D-1. Adverse Reactions Reported During the Study in the Safety Population (any dog that received treatment)

Adverse Reaction	NOCITA (n = 123)	Saline (n = 59)
Discharge from the Incision	4 (3.3%)	0 (0.0%)
Incisional Inflammation (erythema and/or edema)	3 (2.4%)	0 (0.0%)
Vomiting	3 (2.4%)	0 (0.0%)
Abnormalities on Urinalysis (isosthenuria \pm proteinuria)	2 (1.6%)	0 (0.0%)
Increased ALP	2 (1.6%)	0 (0.0%)
Surgical Limb Edema \pm Erythema	1 (0.8%)	3 (5.1%)
Soft Stool/Diarrhea	1 (0.8%)	1 (1.7%)
Inappetence	1 (0.8%)	1 (1.7%)
Fever	1 (0.8%)	0 (0.0%)

Note: If an animal experienced the same event more than once, only the first occurrence was tabulated.

To report suspected adverse drug events and/or to obtain a copy of the Safety Data Sheet (SDS) or for technical assistance, call Aratana Therapeutics at 1-844-640-5500.

For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS or online at <http://www.fda.gov/AnimalVeterinary/SafetyHealth>

Clinical Pharmacology: Bupivacaine is an amide, non-opioid local anesthetic. It provides local analgesia by deactivating sodium channels on the nerve membrane, preventing the generation and propagation of nerve impulses. It is only present in small concentrations as uncharged molecules at tissue pH as it is a base with pKa of 8. This un-ionized form provides a lipophilicity that permits the drug to traverse across the nerve cell membrane and upon entering the cell, binds to the intracellular portion of voltage-gated sodium channels and blocks sodium influx into nerve cells, which prevents depolarization. Without depolarization, no initiation or conduction of a pain signal can occur.

Lipid Formulation

Liposomal encapsulation or incorporation in a lipid complex can substantially affect a drug's functional properties relative to those of the unencapsulated or nonlipid-associated drug. In addition, different liposomal or lipid-complexed products with a common active ingredient may vary from one another in the chemical composition and physical form of the lipid component. Such differences may affect functional properties of these drug products. Do not substitute with other bupivacaine formulations.

After injection of NOCITA into the soft tissue, bupivacaine is released from the multivesicular liposomes over a period of time.

Pharmacokinetics

The pharmacokinetic characterization associated with bupivacaine after subcutaneous NOCITA (bupivacaine liposome injectable suspension) or bupivacaine HCl solution administered to Beagle dogs is provided in Table D-2.

Table D-2. Mean (\pm SD) Plasma Pharmacokinetic Parameters for bupivacaine after single subcutaneous administration of NOCITA and bupivacaine HCl solution in male and female Beagle dogs in a laboratory study

pharmacokinetics

pharmacokinetic characterization associated with bupivacaine after octaneure NOCITA (bupivacaine liposome injectable suspension) bupivacaine HCl solution administered to Beagle dogs is provided in table D-2. Mean (±SD) Plasma Pharmacokinetic Parameters for bupivacaine after single subcutaneous administration of NOCITA and bupivacaine HCl solution in male and female Beagle dogs in a laboratory study

PK Parameter	NOCITA® 9 mg/kg	NOCITA® 18 mg/kg	NOCITA® 30 mg/kg	bupivacaine HCl 9 mg/kg
N, sex	6, (3M/3F)	6, (3M/3F)	6, (3M/3F)	6, (3M/3F)
T _{1/2} ^{a,b} (hr)	0.5 (0.5-0.5)	0.5 (0.5-0.5)	60.0 (0.5-72)	0.5 (0.5-0.5)
C _{max} (ng/mL)	488 (335)	560 (299)	633 (280)	1420 (355)
AUC ₀₋₇₂ (ng·hr/mL)	9100 (4460)	12800 (2020)	25600 (8160)	9720 (1860)
T _{1/2} ^c (hr)	36.2 (12.4)	25.7 (8.15)	43.9 (12.5)	10.1 (8.54)

EXTENDED, POST-OPERATIVE PAIN CONTROL WITH JUST ONE DOSE

Recovery care begins with NOCITA® (bupivacaine liposome injectable suspension)

- All surgical procedures result in some degree of tissue trauma and associated pain¹
- Local anesthetics are one of the most effective means of preventing pain, however, previous options have limitations
- NOCITA is the only long-acting local anesthetic that **controls post-op pain with one dose for up to 72 hours** following canine CCL surgery or feline onychectomy



AVAILABLE IN TWO SIZES

NOCITA is available in convenient 10 mL and 20 mL vials.

To learn more, visit nocita.aratana.com
or call Aratana Customer Care at
1-844-ARATANA (272-8262).

nocita
(bupivacaine liposome injectable suspension)

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3. NOCITA Freedom of Information Summary, NADA 141-461, 12 AUG 2016.
4. NOCITA Freedom on Information Summary, Supplemental NADA 141-461, 03 AUG 2018.