

PANOQUELL®-CA1

Introducing the First
FDA Conditionally Approved
Treatment for Acute
Canine Pancreatitis

User Safety Warnings: Not for use in humans. Keep this medication out of reach of children. Limited data is available on the potential teratogenic effects of fuzapladib.

In case of accidental self-injection, skin contact, eye exposure, or accidental ingestion refer to the package insert.

To obtain a Safety Data Sheet, report suspected adverse drug experiences, or for technical assistance, contact Ceva Animal Health at 1-800-999-0297

For additional information about reporting adverse drug experiences for animal drugs, contact FDA at 1-888-FDA-VETS or at www.fda.gov/reportanimalae



14 mg fuzapladib sodium per vial 4 mg/mL when reconstituted

For intravenous use in dogs only. Reconstitute before using.

PANOQUELL®-CA1 is a leukocyte function-associated antigen 1 (LFA-1) activation inhibitor.

Indication: For the management of clinical signs associated with acute onset of pancreatitis in dogs.

Conditionally approved by FDA pending a full demonstration of effectiveness under application number 141-567. It is a violation of Federal law to use this product other than as directed in the labeling.

CAUTION: Federal law restricts this drug to use by or on the order of a licensed veterinarian.





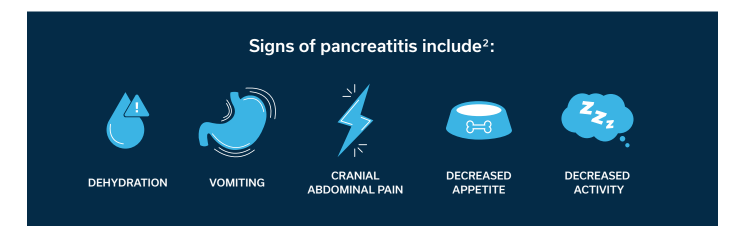


Because you never know where it could go...

Pancreatitis in dogs is a common GI disease.

Unfortunately, prevalence is unknown and causes can vary. Severity ranges from mild to severe and can change daily, which can lead to lasting damage such as recurrent pancreatitis, diabetes mellitus and exocrine pancreatic insufficiency.¹

All of this leads to unexpectedly high patient costs due to **hospitalization** and **supportive care**. This leads to added stress to the pet, veterinarian and pet owner.



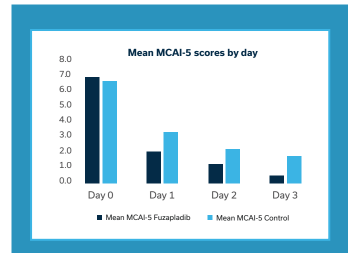
Breeds most at risk include3:



There was no current approved treatment...until now.

A fast, safe and effective treatment for the management of the inflammation associated with acute canine pancreatitis (ACP).

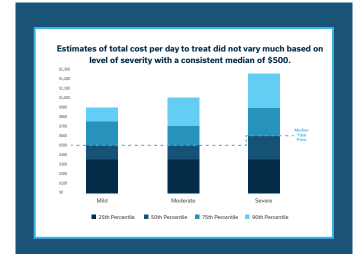
PANOQUELL®-CA1 contains fuzapladib sodium, which inhibits the infiltration of neutrophilic inflammation within the pancreas that occurs in ACP. This action is proven to reduce pancreatic inflammation and support faster recovery.



Fast & Effective

Significant reduction in clinical signs to day 3 compared to control group

PANOQUELL®-CA1 is proven to quickly reduce clinical signs and measurable values such as canine pancreas-specific lipase (Spec cpL) and C-reactive protein (CRP) associated with ACP.4



Cost Effective

PANOQUELL®-CA1 is cost effective due to quick onset and can lead to reduction in time spent in hospital.

With the multi-use vial, there is the ability to treat multiple patients and there is a reduction in waste.

Safe

PANOQUELL®-CA1 can be given in conjunction with other supportive care treatments. In the safety study, it did not produce systemic toxicity when given with other treatments and at high doses. It had an acceptable margin of safety.⁴

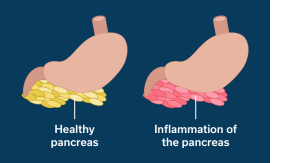
How does it work?

Take the "ITIS" out of pancreatitis.

PANOQUELL®-CA1 (fuzapladib sodium for injection) is a Leukocyte Function Associated Antigen-1 (LFA-1) activation inhibitor. LFA-1 plays a key role in extravasation, which is the process by which leukocytes leave the bloodstream to enter the tissues.

Role of LFA-1 in acute pancreatitis^{5,6}

Infiltration of neutrophils is a hallmark of ACP. LFA-1 is essential to this process. LFA-1 is expressed on the neutrophil surface and its ligand, ICAM-1, is expressed on the vascular endothelium. Together they mediate neutrophilic-driven inflammation and migration into pancreatic and extrapancreatic tissues.



The activation of the LFA-1 by chemokines and the upregulation of its ligand, ICAM-1, lead to "arrest" of circulating neutrophils at the site of inflammation.

Arrested neutrophils now stick to the blood vessel wall, "adhesion", and invade, "migration", into the tissues.

Once in the tissue, neutrophils release additional inflammatory mediators.

These mediators attract more neutrophils and other inflammatory cells.

Especially in severe cases, there is extrapancreatic inflammation, potentially leading to multi-organ failure, systemic inflammatory response syndrome (SIRS) and death.

How should PANOQUELL®-CA1 (fuzapladib sodium for injection) be administered?

PANOQUELL®-CA1 is indicated for the management of clinical signs associated with acute onset of pancreatitis in dogs.

PANOQUELL®-CA1 is an intravenous (IV) injection dosed once a day for 3 days. The IV injection can be given over 15 seconds to 1 minute as a bolus.

How to reconstitute and use:

- Transfer 3.5 mL of sterile diluent into the vial with PANOQUELL®-CA1 lyophilized powder.
- 2 Gently swirl the vial until the powder is fully reconstituted into solution.
- Before each use, gently swirl to ensure uniform solution.
- Draw up appropriate dose* and administer to patient.

PANOQUELL®-CA1 is a 4 mg/mL solution when reconstituted. It comes in a multi-use vial. Once reconstituted, it remains stable under refrigeration for 28 days.

^{*}See the dosage chart on the next page for more information.

Dosing Chart

BODY WEIGHT IN LBS	BODY WEIGHT IN KGS	mL/DAY
1-5*	0.5-2.3	0.05-0.23 mL
6-10*	2.7-4.5	0.27-0.45 mL
11-15*	5.0-6.8	0.50-0.68 mL
16-20*	7.3-9.1	0.73-0.91 mL
21-25*	9.5-11.4	0.95-1.1 mL
26-30	11.8-13.6	1.2-1.4 mL
31-35	14.1-15.9	1.4-1.6 mL
36-40	16.4-18.2	1.6-1.8 mL
41-45	18.6-20.5	1.9-2.0 mL
46-50	20.9-22.7	2.1-2.3 mL
51-55	23.2-25.0	2.3-2.5 mL
56-60	25.5-27.3	2.5-2.7 mL
61-65	27.7-29.5	2.8-3.0 mL
66-70	30.0-31.8	3.0-3.2 mL
71-75	32.3-34.1	3.2-3.4 mL
76-80	34.5-36.4	3.5-3.6 mL
81-85	36.8-38.6	3.7-3.9 mL
86-90	39.1-40.9	3.9-4.1 mL
91-95	41.4-43.2	4.1-4.3 mL
96-100	43.6-45.5	4.4-4.5 mL

Vial Chart

VIALS USED OVER 1 DAY	VIALS NEEDED FOR 3 DAYS
0.01-0.06	0.04-0.19
0.08-0.13	0.23-0.39
0.14-0.19	0.43-0.58
0.21-0.26	0.62-0.78
0.27-0.32	0.82-0.97
0.34-0.39	1.01-1.17
0.40-0.45	1.21-1.36
0.47-0.52	1.40-1.56
0.53-0.58	1.60-1.75
0.60-0.65	1.79-1.95
0.66-0.71	1.99-2.14
0.73-0.78	2.18-2.34
0.79-0.84	2.38-2.53
0.86-0.91	2.57-2.73
0.92-0.97	2.77-2.92
0.99-1.04	2.96-3.12
1.05-1.10	3.16-3.31
1.12-1.17	3.35-3.51
1.18-1.23	3.55-3.70
1.25-1.30	3.74-3.90

**CPANOQUELL*-CA1

(fuzapladib sodium for injection)

Leukocyte function-associated antigen 1

4 mg/mL when reconstituted

(LFA-1) activation inhibitor

For intravenous use in dogs only

on the order of a licensed veterinarian. Federal law restricts this drug to use by or

under application number 141-567. a full demonstration of effectiveness Conditionally approved by FDA pending

It is a violation of Federal law to use this product other than as directed in the

structural formula: weight of 401.38 and the following cyclohexanecarboxamide monosodium. amino)-5-(trifluoromethyl)-3-pyridinyl] PANOQUELL®-CA1 (fuzapladib sodium for PANOQUELL®-CA1 has a molecular designation for N-[2-((ethylsulfonyl) Fuzapladib sodium is the non-proprietary injection) is a selective inhibitor of LFA-1.

hydroxide. mg D-mannitol, 6 mg tromethamine, contains 4 mg fuzapladib sodium, 15 milliliter of reconstituted drug product reconstituted with 3.5 mL of the provided No other diluent should be used. When sterile lyophilized powder prior to use. water for injection), containing 1.8% w/v 3.9 mL sterile diluent (bacteriostatic lyophilized powder. The second vial of and 21 mg of tromethamine as sterile adjusted with hydrochloric acid or sodium and 18 mg benzyl alcohol. The pH was Bacteriostatic Water for Injection, each benzyl alcohol, is for reconstituting the fuzapladib sodium, 52.5 mg of D-mannitol separate vials. One vial contains 14 mg of PANOQUELL®-CA1 consists of two

INDICATION:

management of clinical signs associated PANOQUELL®-CA1 is indicated for the with acute onset of pancreatitis in dogs.

DOSAGE AND ADMINISTRATION:

sterile diluent provided, resulting in a 4 mg/mL solution of PANOQUELL®-CA1. solution. before every use to ensure a uniform Once reconstituted, swirl the bottle gently powder should be reconstituted (see Prior to use, the sterile lyophilized Reconstitution Procedures) using the

injection over 15 seconds to 1 minute. at a dosage of 0.4 mg (0.1 mL) per consecutive days by intravenous (IV) bolus kg body weight once daily for three The reconstituted product is administered

Reconstitution Procedures:

The items needed for reconstitution are:

- lyophilized powder Sterile PANOQUELL®-CA1
- Sterile diluent
- diluent Sterile 5 mL syringe for transfer of
- Sterile needle

Steps for reconstitution:

- Using a sterile needle and syringe, supplied than the 3.5 mL needed for stopper. There is more sterile diluent the sterile diluent into the vial CA1 lyophilized powder through the containing the sterile PANOQUELL®trom the vial and slowly transter withdraw 3.5 mL of the sterile diluent
- Once the sterile diluent has been and needle. Discard unused sterile diluent, syringe, added to the powder vial, remove the needle and syringe from the vial.

reconstitution.

- Gently swirl the vial until the powder is no visible residue or un-dissolved fully reconstituted into solution, leaving
- a unitorm solution. Before each use, gently swirl to ensure
- <u>,</u> Draw up the appropriate dose using a Administer the dose promptly after new sterile needle and syringe.

6

7. Store any remaining reconstituted drawing into the dosing syringe. refrigeration for 28 days. product remains stable under 36° to 46°F (2° to 8°C). Reconstituted product at refrigerated conditions,

CONTRAINDICATIONS:

hypersensitivity to fuzapladib sodium. Do not use in dogs with a known

User Safety Warnings:

medication out of reach of children. Not for use in humans. Keep this

> In case of accidental self-injection: Seek medical advice immediately and the physician. show the package insert or label to

In case of accidental skin contact:

- Wash the exposed skin with water for
- If redness and swelling occur, seek the package insert or label to the medical advice immediately and show at least 15 minutes

In case of accidental eye exposure:

- Wash the eyes with water for at least 15 minutes.
- continue to rinse with water. eyes first, then remove contacts and If wearing contact lenses, rinse the
- the package insert or label to the medical advice immediately and show If redness and swelling occur, seek

In case of accidental ingestion:

- Do not induce vomiting unless Rinse the mouth out with water
- directed to do so by medical
- the physician. show the package insert or label to Seek medical advice immediately and

pregnant should avoid direct contact with PANOQUELL®-CA1. breast feeding, or planning to become teratogenic effects of fuzapladib sodium. Limited data is available on the potential Therefore, anyone who is pregnant,

is summarized in Table 1.

dogs experiencing each adverse reaction

To obtain a Safety Data Sheet, contact PANOQUELL®-CA1 excipients should avoid contact with to fuzapladib sodium or to any of the Anyone with known hypersensitivity

0297 or www.ceva.com. Ceva Animal Health, LLC, at 1-800-999.

PANOQUELL®-CA1 is highly protein

in patients requiring adjunctive therapy. Drug compatibility should be monitored medications that are highly protein bound. bound. Use with caution with other pain medications (excluding NSAIDs), anti sodium included, but were not limited to, pilot effectiveness study with fuzapladib Concurrent medications used during the diuretics, and behavioral medications. (NSAIDs), anti-emetics, antibiotics, non-steroidal anti-inflammatory drugs used protein bound drugs include not been studied in dogs. Commonly CA1 with other protein bound drugs has The concomitant use of PANOQUELL®-

> pre-existing conditions. medications used to treat well-controlled emetics, parasiticides, vaccinations, and

cardiac disease, hepatic failure, or renal has not been evaluated in dogs with The safe use of PANOQUELL®-CA1

pregnant, lactating, or intended for not been evaluated in dogs that are The safe use of PANOQUELL®-CA1 has

months of age. not been evaluated in dogs less than 6 The safe use of PANOQUELL®-CA1 has

ADVERSE REACTIONS:

In a well-controlled pilot field study

to assess the effectiveness and safety solubilized in 1 mL of Sterile Water for was excipient sterile lyophilized powder evaluated for safety. The vehicle control dogs administered vehicle control were administered fuzapladib sodium and 30 OF EFFECTIVENESS), 31 dogs (see REASONABLE EXPECTATION diagnosed with acute onset of pancreatitis formulation) in client-owned dogs of fuzapladib sodium (not commercial observed in the study and the number of Injection, USP. The adverse reactions

Pilot Field Study Table 1: Adverse Reactions During the

Hypertension	Anaphylaxis	Cerebral edema	Abrasion	Tremor/ shivering/ shaking	Subcutaneous swelling, bruising at injection site	Limb edema	Heart murmur	Hypersalivation	Pruritis, urticaria	Hyperthermia	Cardiac arrest	Arrhythmia	Diarrhea	Abnormal urine	Hepatopathy, jaundice	Respiratory tract disorders	Digestive tract disorders	Anorexia	Adverse Reaction
1 (3.2%)	1 (3.2%)	1 (3.2%)	1 (3.2%)	1 (3.2%)	1 (3.2%)	1 (3.2%)	1 (3.2%)	2 (6.5%)	2 (6.5%)	2 (6.5%)	2 (6.5%)	2 (6.5%)	3 (9.7%)	3 (9.7%)	4 (12.9%)	4 (12.9%)	5 (16.1%)	5 (16.1%)	Fuzapladib Sodium (n = 31) (%)
0	0	0	1 (3.3%)	1 (3.3%)	1 (3.3%)	2 (6.7%)	2 (6.7%)	0	0	0	0	1 (3.3%)	1 (3.3%)	2 (6.7%)	2 (6.7%)	3 (10.0%)	3 (10.0%)	2 (6.7%)	Vehicle Control (n = 30) (%)

reported with more than one abnormality. vehicle control), inspiratory crackles (1 included pneumonia (2 fuzapladib, 1 vehicle control), vomiting (1 fuzapladib, included regurgitation (1 fuzapladib, 2 (1 fuzapladib). Some of these dogs were (2 vehicle control), and malodorous urine Abnormal urine included proteinuria (2 (2 fuzapladib), and dyspnea (1 fuzapladib). fuzapladib). Respiratory tract disorders nausea (1 fuzapladib), and enteritis (1 vehicle control), flatulence (1 fuzapladib), In Table 1 above, digestive tract disorders fuzapladib, 2 vehicle control), hematuria fuzapladib, 2 vehicle control), tachypnea

sodium group and one in the vehicle during the study: tour in the tuzapladib Five out of the 61 enrolled dogs died are only presented once in the table above reaction on more than one occasion but for each reported adverse reaction. Note: Some dogs experienced an adverse

tuzapladib sodium group had intestinal onset of pancreatitis: one dog in the be attributed to causes other than acute of a poor prognosis. Two deaths could control group was euthanized because cardiac arrest. One dog in the vehicle pneumonia and died after experiencing group was suspected to have aspiration group. One dog in the fuzapladib sodium group and one in the vehicle control pancreatitis: two in the fuzapladib sodium complications from severe acute onset of euthanized during or after the study, Of the seven dogs that died or were and a pheochromocytoma. dog had a cranial thromboembolic event lymphoma and one vehicle control group three deaths could be attributed to shortly after completion of the study vehicle control group were euthanized control group. Two additional dogs in the

administration. events occurred within 24 hours of collapse, and seizure. These adverse markets: facial and tongue swelling, reported voluntarily during post-approval use of the product in dogs in foreign Foreign Market Experience The tollowing adverse events were

CONTACT INFORMATION:

contact Ceva Animal Health, LLC, at experiences or for technical assistance To report suspected adverse drug Contact Ceva Animal Health, LLC, at -800-999-0297 or www.ceva.com.

at www.fda.gov/reportanimalae drugs, contact FDA at 1-888-FDA-VETS or adverse drug experiences for animal For additional information about reporting

CLINICAL PHARMACOLOGY:

Mechanism of action

such as multi-organ failure. expansion and help prevent complications into sites of tissue injury and inflammation inflammatory cell adhesion and migration of LFA-1, resulting in inhibition of PANOQUELL®-CA1 (fuzapladib sodium are thought to limit pancreatic lesion These anti-inflammatory properties through its ability to inhibit activation for injection) has anti-inflammatory effects

Pharmacokinetics

kg after the first dose and ninth dose. dose-proportional between 0.4 and 2 mg/ plasma exposure (AUC) was greater than days, minimal accumulation was observed mg/kg, and 2 mg/kg for nine consecutive of PANOQUELL®-CA1 at 0.4 mg/kg, 1.2 Following once daily IV administration with a mean accumulation ratio of 1.37 1.36, and 1.35, respectively. The extent of

doses of 0.4 mg/kg in dogs fuzapladib sodium following nine IV pharmacokinetic parameters of Table 2: Mean (± standard deviation)

C ₀ (µg/mL)	3.55 ± 1.17
AUC _{ss} (hour*μg/mL) 19.2 ± 12.7	19.2 ± 12.7
T _{1/2} (hour)	7.32 ± 4.08
V _{ss} (L/kg)	0.216 ± 0.070
Cl _{ss} (L/h/kg)	0.026 ± 0.009

V_{ss}: Volume of distribution at steady state Cl_{ss}: Clearance at steady state concentration versus time curve during AUC_{ss}: Area under the plasma administration of first two data points following IV at time zero by a log-linear regression concentration of fuzapladib sodium dosing interval at steady state $\mathsf{C}_{\scriptscriptstyle{0}}$: Back-extrapolated plasma /2: Terminal elimination half-life

EFFECTIVENESS: REASONABLE EXPECTATION OF

published literature. in the target species or studies from such as, but not limited to, pilot data may be demonstrated based on evidence A reasonable expectation of effectiveness

www.fda.gov/animalca for Conditional Approvals can be found at of effectiveness. Additional information approved pending a full demonstration PANOQUELL®-CA1 is conditionally

in dogs was based on a pilot field study. associated with acute onset of pancreatitis for the management of clinical signs effectiveness for PANOQUELL®-CA1 The reasonable expectation of

these dogs (19) began the study before from the effectiveness analysis, most of also excluded. Of the 25 dogs excluded with severe concurrent life-threatening foreign body and abdominal masses. Dogs cases of gastrointestinal obstruction/ radiographs were evaluated to exclude imaging consisting of ultrasound and/or were diagnosed with acute onset of in the effectiveness analysis. Dogs the study and 36 dogs were included and 15.9 years old were enrolled in dogs of various breeds between 1.8 (not commercial formulation) was illness other than acute pancreatitis were concentration of ≥ 400 µg/L. Abdominal pancreatic lipase immunoreactivity (cPLI) pathology results, and a Day 0 canine pancreatitis based on clinical signs, clinical field study. A total of 61 client-owned demonstrated in a well-controlled pilot The effectiveness of fuzapladib sodium

> were excluded for other reasons. cPLI results were ≤ 400 µg/L. Six dogs and were later excluded because their the cPLI results from Day 0 were finalized

dogs received 0.1 mL/kg vehicle control parasiticides and vaccinations. conditions. Some dogs also received used to treat well-controlled pre-existing NSAIDs), anti-emetics, and medications support, pain medications (excluding pancreatitis, including fluids, nutritional the standard of care for acute onset of dogs enrolled in the study received USP) IV once daily for three days. All in 1 mL of Sterile Water for Injection, (excipient lyophilized powder solubilized 0.4 mg/kg fuzapladib sodium and 19 effectiveness analysis, 17 dogs received Of the 36 dogs included in the

Stool consistency, and Blood in the stool Cranial abdominal pain, Dehydration, Appetite (voluntary food intake), Vomiting dogs with acute pancreatitis: Activity, following seven clinical signs relevant in was used to evaluate and score the A Modified Canine Activity Index (MCAI)

0 mean MCAI scores for the fuzapladib control (p = 0.0193). reduction in MCAI scores compared to sodium had a statistically significant respectively. Dogs treated with fuzapladib vehicle control groups were -7.7 and -5.7, 0 to 3 for the fuzapladib sodium and in the mean total MCAI scores from Day 8.53 and 7.68, respectively. The changes sodium and vehicle control groups were 3, as assessed by the Investigator. Day score from Day 0 (pre-treatment) to Day the change in the group mean total MCA The primary effectiveness variable was

TARGET ANIMAL SAFETY:

dog on one day each. One dog in the kg group dogs and one 2 mg/kg group x 10³/L) was observed in two 0.4 mg/ 169 x 10³/L (reference range: 171- 361 study. Mild thrombocytopenia of 121-CA1 and occurred only at the end of the only in dogs administered PANOQUELL®. values of \geq 160 mmHg) was observed Hypertension (systolic blood pressure frequency in the higher dose groups. dose dependent manner with increased injection site swelling and bruising in a CA1 resulted in hypertension and All dogs survived to study termination. by IV injection once daily for 9 days. PANOQUELL®-CA1, or saline control, 0.4 (1X), 1.2 (3X), or 2 (5X) mg/kg aged 6 to 7 months were administered intact Beagle dogs (4 dogs/sex/group) In a 9-day laboratory study, 32 healthy The administration of PANOQUELL®-

> all groups, including control, but increased administered PANOQUELL®-CA1. injection sites were found only in dogs and subcutaneous hemorrhage of the fibroplasia, subcutaneous inflammation, increased incidence and severity of dermal On histopathology, observations of in severity in a dose dependent manner. sites was observed on gross necropsy in subcutaneous hemorrhage of the injection injection on the last day of dosing. Focal kg group had pain associated with the thrombocytopenia. One dog in the 2 mg/ the injection site that coincided with the 0.4 mg/kg group also had bruising of

HOW SUPPLIED:

diluent should be used. lyophilized powder prior to use. No other alcohol, is for reconstituting the sterile mL sterile diluent (bacteriostatic water lyophilized powder. The second vial of 3.9 and 21 mg of tromethamine as sterile separate vials. One vial contains 14 mg of PANOQUELL®-CA1 consists of two for injection), containing 1.8% w/v benzyl fuzapladib sodium, 52.5 mg of D-mannitol,

STORAGE, HANDLING, AND DISPOSAL:

8°C). Use within 28 days of first puncture. refrigerated conditions, 36° to 46°F (2° to Store the reconstituted product at temperature, 59° to 77°F (15° to 25°C). Store unopened vials at room

MANUFACTURED FOR:

Ishihara Sangyo Kaisha, Ltd., Osaka, Japan

DISTRIBUTED BY:

Lenexa, KS 66215 Ceva Animal Health, LLC

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Revision date: 10/22 ISK/PAN/P1/1

