#### IBUPROFEN ORAL - ibuprofen oral suspension Bryant Ranch Prepack

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Ibuprofen Oral Suspension USP, 100 mg/5 mL (OTC)

## ACTIVE INGREDIENT(S)

(in each 5 mL) Ibuprofen 100 mg (NSAID)\* \*nonsteroidal anti-inflammatory drug

#### PURPOSE

Pain reliever/fever reducer

#### USE(S)

temporarily:

- relieves minor aches and pains due to the common cold, flu, sore throat, headache and toothache
- reduces fever

#### WARNINGS

Allergy alert: Ibuprofen may cause a severe allergic reaction, especially in people allergic to aspirin. Symptoms may include:

- hives
- facial swelling
- asthma (wheezing)
- shock
- skin reddening
- rash
- blisters

If an allergic reaction occurs, stop use and seek medical help right away. **Stomach bleeding warning:** This product contains an NSAID, which may cause severe stomach bleeding. The chance is higher if your child:

- has had stomach ulcers or bleeding problems
- takes a blood thinning (anticoagulant) or steroid drug
- takes other drugs containing prescription or nonprescription NSAIDs (aspirin, ibuprofen, naproxen, or others)

- takes more or for a longer time than directed
- **Heart attack and stroke warning:** NSAIDs, except aspirin, increase the risk of heart attack, heart failure, and stroke. These can be fatal. The risk is higher if you use more than directed or for longer than directed.

**Sore throat warning:** Severe or persistent sore throat or sore throat accompanied by high fever, headache, nausea, and vomiting may be serious. Consult doctor promptly. Do not use more than 2 days or administer to children under 3 years of age unless directed by doctor.

## DO NOT USE

- if the child has ever had an allergic reaction to any other pain reliever/fever reducer
- right before or after heart surgery

## ASK A DOCTOR BEFORE USE IF

- stomach bleeding warning applies to your child
- child has a history of stomach problems, such as heartburn
- child has problems or serious side effects from taking pain relievers or fever reducers
- child has not been drinking fluids
- child has lost a lot of fluid due to vomiting or diarrhea
- child has high blood pressure, heart disease, liver cirrhosis, kidney disease, or had a stroke
- child has asthma
- child is taking a diuretic

## Ask a doctor or pharmacist before use if the child is

- under a doctor's care for any serious condition
- taking any other drug

## WHEN USING THIS PRODUCT

• take with food or milk if stomach upset occurs

## Stop use and ask a doctor if

- child experiences any of the following signs of stomach bleeding:
- feels faint
- vomits blood
- has bloody or black stools
- has stomach pain that does not get better
- child has symptoms of heart problems or stroke:

- chest pain
- trouble breathing
- weakness in one part or side of body
- slurred speech
- leg swelling
- the child does not get any relief within first day (24 hours) of treatment
- fever or pain gets worse or lasts more than 3 days
- redness or swelling is present in the painful area
- any new symptoms appear

## KEEP OUT OF REACH OF CHILDREN

In case of overdose, get medical help or contact a Poison Control Center right away. (1-800-222-1222)

## DIRECTIONS

# • this product does not contain directions or complete warnings for adult use

## do not give more than directed

- shake well before using
- mL = milliliter
- find right dose on chart. If possible, use weight to dose; otherwise use age.
- use only enclosed dosing cup. Do not use any other dosing device.
- · if needed, repeat dose every 6-8 hours
- do not use more than **4 times a day**
- replace original bottle cap to maintain child resistance

## **Dosing Chart**

Weight (lb)	Age (yr)	Dose (mL)*
under 24	under 2 years	ask a doctor
24-35 lbs	2-3 years	5 mL
36-47 lbs	4-5 years	7.5 mL
48-59 lbs	6-8 years	10 mL
60-71 lbs	9-10 years	12.5 mL
72-95 lbs	11 years	15 mL

\*or as directed by a doctor

#### Other information

 $\Box$ 

- each 5 mL contains : sodium 2 mg
- store between 20-25°C (68-77°F)
- do not use if carton is opened or printed bottle neckband is broken or missing
- see bottom panel for lot number and expiration date

#### **INACTIVE INGREDIENT SECTION**

#### Berry flavor (with dye)

Acesulfame potassium, anhydrous citric acid, carboxymethylcellulose sodium, D&C Yellow #10, FD&C Red #40, flavors, glycerin, microcrystalline cellulose, polysorbate 80, propylene glycol, purified water, sodium benzoate, sucrose, and xanthan gum.

#### Berry flavor (dye free)

Acesulfame potassium, anhydrous citric acid, carboxymethylcellulose sodium, flavors, glycerin, microcrystalline cellulose, polysorbate 80, propylene glycol, purified water, sodium benzoate, sucrose, and xanthan gum.

#### Grape flavor

Acesulfame potassium, anhydrous citric acid, carboxymethylcellulose sodium, D&C Red #33, FD&C Blue #1, flavors, glycerin, microcrystalline cellulose, polysorbate 80, propylene glycol, purified water, sodium benzoate, sucrose, and xanthan gum.

#### <u>Bubble Gum flavor</u>

Acesulfame potassium, anhydrous citric acid, carboxymethylcellulose sodium, D&C Red #33, FD&C Red #40, flavors, glycerin, microcrystalline cellulose, polysorbate 80, propylene glycol, purified water, sodium benzoate, sucrose, and xanthan gum.

#### **QUESTIONS SECTION**

Call 1-888-588-1418 from 9 AM to 5 PM EST,



**Distributed by:** Camber Consumer Care, Inc. Piscataway, NJ 08854, USA

#### HOW SUPPLIED

Ibuprofen Oral Suspension Berry Dye Free

• NDC 72162-1912-2: 118 mL in a BOTTLE

Repackaged/Relabeled by: Bryant Ranch Prepack, Inc. Burbank, CA 91504

## **Ibuprofen Oral Suspension Berry Dye Free**



Insert



Product Info	mation							
Product Type		HUMAN OTC DRUG	ltem Code (	Source)	NDC:72162-192	12(NDC:69230-311		
Route of Admin	istration	ORAL						
Active Ingred	lient/Active	Moietv						
<b>J</b>	Basis of Strength Stre							
BUPROFEN (UNII:	-	e <b>nt Name</b> BUPROFEN - UNII:WK2	2XYI10QM)	IBUPROF	-	100 mg in 5 m		
Inactive Ingro	edients							
		Ingredient N	Name			Strengtl		
ACESULFAME PO	TASSIUM (UNII:	230V73Q5G9)						
ANHYDROUS CITI	RIC ACID (UNII: >	(F417D3PSL)						
CARBOXYMETHY	LCELLULOSE SO	DDIUM, UNSPECIFI	ED (UNII: K6790	DBS311)				
GLYCERIN (UNII: PDC6A3C0OX)								
	MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)							
POLYSORBATE 80 (UNII: 60ZP39ZG8H)								
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)								
WATER (UNII: 059QF0KO0R)								
SODIUM BENZOA	TE (UNII: OJ245F	E5EU)						
SODIUM BENZOA SUCROSE (UNII: C	<b>TE</b> (UNII: OJ245F 151H8M554)							
SODIUM BENZOA SUCROSE (UNII: C XANTHAN GUM (U	<b>TE</b> (UNII: OJ245F 151H8M554) INII: TTV12P4NEE							
SODIUM BENZOA SUCROSE (UNII: C XANTHAN GUM (U Product Char	<b>TE</b> (UNII: OJ245F 151H8M554) INII: TTV12P4NEE		<b>6</b>					
SODIUM BENZOA SUCROSE (UNII: C XANTHAN GUM (U Product Char Color	<b>TE</b> (UNII: OJ245F 151H8M554) INII: TTV12P4NEE		Score					
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## Registrant - Bryant Ranch Prepack (171714327)

Establishment					
Name	Address	ID/FEI	<b>Business Operations</b>		
Bryant Ranch Prepack		171714327	REPACK(72162-1912), RELABEL(72162-1912)		

Revised: 1/2025

Bryant Ranch Prepack