

**57955-9984-2**

**BACK, MUSCLE AND JOINT RELIEF  
National Drug Code Directory**

The Drug Listing Act of 1972 requires registered drug establishments to provide the Food and Drug Administration (FDA) with a current list of all drugs manufactured, prepared, propagated, compounded, or processed by it for commercial distribution. (See Section 510 of the Federal Food, Drug, and Cosmetic Act (Act) (21 U.S.C. § 360)). Drug products are identified and reported using a unique, three-segment number, called the National Drug Code (NDC), which serves as a universal product identifier for drugs. FDA publishes the listed NDC numbers and the information submitted as part of the listing information in the NDC Directory which is updated daily.

**57955-9984-2 Back, Muscle And Joint Relief**

<i>Labeler Name</i>	King Bio Inc.	Name of Company corresponding to the labeler code segment of the ProductNDC.
<i>NDC Package Code</i>	57955-9984-2	The labeler code, product code, and package code segments of the National Drug Code number, separated by hyphens. Asterisks are no longer used or included within the product and package code segments to indicate certain configurations of the NDC.
<i>Proprietary Name</i>	Back, Muscle And Joint Relief	Also known as the trade name. It is the name of the product chosen by the labeler.
<i>11 Digit NDC Code</i>	57955-9984-02	It should be noted that many NDCs are displayed on drug packaging in a 10-digit format. Proper billing of an NDC requires an 11-digit number in a 5-4-2 format. Converting NDCs from a 10-digit to 11-digit format requires a strategically placed zero, dependent upon the 10-digit format.
<i>Product NDC</i>	57955-9984	The labeler code and product code segments of the National Drug Code number, separated by a hyphen. Asterisks are no longer used or included within the product code segment to indicate certain configurations of the NDC.
<i>Product Type Name</i>	HUMAN OTC DRUG	Indicates the type of product, such as Human Prescription Drug or Human OTC Drug. This data element corresponds to the "Document Type" of the SPL submission for the listing. The complete list of codes and translations can be found at <a href="http://www.fda.gov/edrls">www.fda.gov/edrls</a> under Structured Product Labeling Resources.
<i>Non Proprietary Name</i>	Aesculus Hippocastanum, Arnica Montana, Bellis Perennis, Bryonia, Calcarea Carbonica, Calcarea Fluorica, Cimicifuga Racemosa, Cobaltum Metallicum, Gnaphalium Polycephalum, Hypericum Perforatum, Kali Carbonicum, Kali Phosphoricum, Magnesia Phosphorica, Oxalicum Acidum, Phosphorus, Rhus Tox, Ruta Graveolens, And Zincum Metallicum.	Sometimes called the generic name, this is usually the active ingredient(s) of the product.
<i>Package Description</i>	59 mL in 1 BOTTLE, SPRAY (57955-9984-2)	A description of the size and type of packaging in sentence form. Multilevel packages will have the descriptions concatenated together. For example: 4 BOTTLES in 1 CARTON/100 TABLETS in 1 BOTTLE.



<i>Strength Unit</i>	<p>[hp_X]/59mL;  [hp_X]/59mL;  [hp_X]/59mL;</p>	<p>These are the units to be used with the strength values above, listed in the same order as the SubstanceName and SubstanceNumber.</p>
<i>Pharmaceutical Classes</i>	<p>Allergens [CS], Calculi  Dissolution Agent [EPC],  Cell-mediated Immunity  [PE], Increased Histamine  Release [PE], Increased  Large Intestinal Motility  [PE], Inhibition Large  Intestine Fluid/Electrolyte  Absorption [PE], Inhibition  Small Intestine  Fluid/Electrolyte  Absorption [PE],  Magnesium Ion Exchange  Activity [MoA], Osmotic  Activity [MoA], Osmotic  Laxative [EPC],  Standardized Chemical  Allergen [EPC], Stimulation  Large Intestine  Fluid/Electrolyte Secretion  [PE]</p>	<p>These are the reported pharmaceutical class categories corresponding to the SubstanceNames listed above.</p>
<i>Status</i>	<p>Active</p>	<p>Possible status values:</p> <ul style="list-style-type: none"> <li>&lt;li&gt;&lt;strong&gt;Active&lt;/strong&gt; &lt;br/&gt;Active NDC Code&lt;/li&gt;</li> <li>&lt;li&gt;&lt;strong&gt;Deprecated&lt;/strong&gt; &lt;br/&gt;Deprecated NDC Code&lt;/li&gt;</li> <li>&lt;li&gt;&lt;strong&gt;Unfinished&lt;/strong&gt; (Unapproved) &lt;br/&gt;The following status describes submitted unfinished drugs, including the marketing categories of Active Pharmaceutical Ingredient (API), Drug for Further Processing, Bulk for Human Drug Compounding, and Bulk for Animal Drug Compounding. &lt;br/&gt;&lt;strong&gt;FDA does not review and approve unfinished products. Therefore, all products having "unfinished" status are considered unapproved.&lt;/strong&gt;</li> </ul>

<i>Last Update Date</i>	2026-03-10	The date that a record was last updated or changed.
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For all questions regarding this bundle please contact Support@DataLabs.Health. Also feel free to let us know about any suggestions or concerns. All additional information as well as customer support is available at <https://www.datalabs.health/>.