

**ANIMAL ALLERGENS, AP CATTLE HAIR AND DANDER- cattle hair and dander injection, solution**

**ANIMAL ALLERGENS, AP DOG HAIR AND DANDER CANIS SPP- animal allergens, dog dander canis spp injection, solution**

**ANIMAL ALLERGENS, DOG HAIR AND DANDER CANIS SPP.- dog hair canis spp. injection, solution**

**ANIMAL ALLERGENS, AP HORSE HAIR AND DANDER- ap horse hair and dander injection, solution**

**ANIMAL ALLERGENS, FEATHER MIX- feather mix injection, solution**

**ANIMAL ALLERGENS, GUINEA PIG HAIR AND DANDER- guinea pig hair and dander injection, solution**

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, BEEF BOVINE SPP.- beef bovine spp. injection, solution**

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, CHICKEN MEAT GALLUS SP.- chicken meat gallus sp. injection, solution**

**POLLENS - TREES, OLIVE OLEA EUROPAEA- olive olea europaea injection, solution**

**POLLENS - TREES, PALM, QUEEN COCOS PLUMOSA- palm, queen cocos plumosa injection, solution**

**POLLENS - TREES, PALO VERDE CERCIDIUM FLORIDUM- palo verde cercidium floridum injection, solution**

**POLLENS - TREES, PECAN CARYA CARYA ILLINOENSIS- pecan carya carya illinoensis injection, solution**

**POLLENS - TREES, PEPPER TREE, CALIFORNIA SCHINUS MOLLE- pepper tree, califonia schinus molle injection, solution**

**POLLENS - TREES, PINE MIX- pine mix injection, solution**

**POLLENS - TREES, PRIVET LIGUSTRUM VULGARE- privet ligustrum vulgare injection, solution**

**POLLENS - TREES, RUSSIAN OLIVE ELAEAGNUS ANGUSTIFOLIA- russian olive elaeagnus angustifolia injection, solution**

**POLLENS - TREES, SYCAMORE, AMERICAN EASTERN PLATANUS OCCIDENTALIS- sycamore, american eastern platanus occidentalis injection, solution**

**POLLENS - TREES, TREE MIX 11- tree mix 11 injection, solution**

**POLLENS - TREES, TREE MIX 5- tree mix 5 injection, solution**

**POLLENS - TREES, TREE MIX 6- tree mix 6 injection, solution**

**POLLENS - TREES, TREE OF HEAVEN AILANTHUS ALTISSIMA- tree of heaven ailanthus altissima injection, solution**

**POLLENS - TREES, WALNUT, BLACK JUGLANS NIGRA- walnut, black juglans nigra injection, solution**

**POLLENS - TREES, WILLOW, BLACK SALIX NIGRA- willow, black salix nigra injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, COCKLEBUR XANTHIUM STRUMARIUM- cocklebur xanthium strumarium injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, DOG FENNEL, EASTERN EUPATORIUM CAPILLIFOLIUM- dog fennel, eastern eupatorium capillifolium injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, GOLDENROD SOLIDAGO CANADENSIS- goldenrod solidago canadensis injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, LAMBS QUARTERS CHENOPODIUM ALBUM- lambs quarters chenopodium album injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, NETTLE URTICA DIOICA- nettle urtica dioica injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, PIGWEED, ROUGH REDROOT AMARANTHUS RETROFLEXUS- pigweed, rough redroot amaranthus retroflexus injection, solution**

**POLLENS - WEEDS AND GARDEN PLANTS, PLANTAIN, ENGLISH PLANTAGO LANCEOLATA-** plantain, english plantago lanceolata injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED, GIANT AMBROSIA TRIFIDA-** ragweed, giant ambrosia trifida injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED. WESTERN AMBROSIA PSILOSTACHYA-** ragweed. western ambrosia psilostachya injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, RUSSIAN THISTLE SALSOLA KALI-** russian thistle salsola kali injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, SAGEBRUSH, MUGWORT ARTEMISIA VULGARIS-** sagebrush, mugwort artemisia vulgaris injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, SCALE, WING SHAD ATRIPLEX CANESCENS-** scale, wing shad atriplex canescens injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, SCOTCH BROOM CYTISUS SCOPARIUS-** scotch broom cytisus scoparius injection, solution

**POLLENS - WEEDS AND GARDEN PLANTS, SORREL, SHEEP RUMEX ACETOSELLA-** sorrel, sheep rumex acetosella injection, solution

**POLLENS - WEEDS, CARELESS WEED AMARANTHUS PALMERI-** careless weed amaranthus palmeri injection, solution

**POLLENS - WEEDS, CARELESS/PIGWEED MIX-** careless/pigweed mix injection, solution

**POLLENS - WEEDS, DOCK/SORREL MIX-** pollens - weeds, dock/sorrel mix injection, solution

**POLLENS - WEEDS, GIANT, SHORT, WESTERN RAGWEED MIX-** giant, short, western ragweed mix injection, solution

**POLLENS - WEEDS, KOCHIA SCOPARIA-** kochia scoparia injection, solution

**POLLENS - WEEDS, MARSHELDER/POVERTY MIX-** pollens - weeds, marshelder/poverty mix injection, solution

**POLLENS - WEEDS, WEED MIX 2630-** weed mix 2630 injection, solution

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, EGG, WHITE GALLUS SP.-** egg, white gallus sp. injection, solution

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, EGG, YOLK GALLUS SP.-** egg, yolk gallus sp. injection, solution

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, PORK SUS SP.-** pork sus sp. injection, solution

**FOOD - DAIRY PRODUCTS, CASEIN, COW MILK-** casein, cow milk injection, solution

**FOOD - DAIRY PRODUCTS, MILK, WHOLE COW-** milk, whole cow injection, solution

**FOOD - FISH AND SHELLFISH, CLAM-** clam injection, solution

**FOOD - FISH AND SHELLFISH, CODFISH GADUS CALLARIAS-** codfish gadus callarias injection, solution

**FOOD - FISH AND SHELLFISH, CRAB XIPHOSURUS SOWERBYI-** crab xiphosurus sowerbyi injection, solution

**FOOD - FISH AND SHELLFISH, LOBSTER HOMARUS AMERICANUS-** lobster homarus americanus injection, solution

**FOOD - FISH AND SHELLFISH, SALMON SALMO SALAR-** salmon salmo salar injection, solution

**FOOD - FISH AND SHELLFISH, SHRIMP CRAGO SP.-** shrimp crago sp. injection, solution

**FOOD - FISH AND SHELLFISH, TUNA THUNNUS SP.-** tuna thunnus sp. injection, solution

**FOOD - PLANT SOURCE, ALMOND PRUNUS AMYGDALUS-** almond prunus amygdalus injection, solution

**FOOD - PLANT SOURCE, APPLE MALUS SP.-** apple malus sp. injection, solution

**FOOD - PLANT SOURCE, BANANA MUSA SAPIENTUM-** banana musa

**sapientum injection, solution**

**FOOD - PLANT SOURCE, BRAZIL NUT BERTHOLLETIA EXCELSA- brazil nut**

**bertholletia excelsa injection, solution**

**FOOD - PLANT SOURCE, CARROT DAUCUS CAROTA- carrot daucus**

**carota injection, solution**

**FOOD - PLANT SOURCE, CASHEW NUT ANACARDIUM OCCIDENTALIE- cashew**

**nut anacardium occidentalis injection, solution**

**FOOD - PLANT SOURCE, CELERY APIUM GRAVEOLENS- celery apium**

**graveolens injection, solution**

**FOOD - PLANT SOURCE, CORN ZEA MAYS- corn zea mays injection, solution**

**FOOD - PLANT SOURCE, HAZELNUT FILBERT CORYLUS SPP.- hazelnut filbert**

**corylus spp. injection, solution**

**FOOD - PLANT SOURCE, MELON, CANTALOUP CUCUMIS MELO- cantaloupe**

**cucumis melo injection, solution**

**FOOD - PLANT SOURCE, ORANGE CITRUS SINENSIS- orange citrus**

**sinensis injection, solution**

**FOOD - PLANT SOURCE, PEA, GREEN OR ENGLISH PISUM SATIVUM- pea,**

**green or english pisum sativum injection, solution**

**FOOD - PLANT SOURCE, PEACH PRUNUS PERSICA- peach prunus**

**persica injection, solution**

**FOOD - PLANT SOURCE, PEANUT ARACHIS HYPOGAEA- peanut arachis**

**hypogaea injection, solution**

**FOOD - PLANT SOURCE, PECAN CARYA ILLINOENSIS- pecan carya**

**illinoensis injection, solution**

**FOOD - PLANT SOURCE, POTATO, WHITE SOLANUM TUBEROSUM- potato,**

**white solanum tuberosum injection, solution**

**FOOD - PLANT SOURCE, RICE, WHOLE GRAIN- rice, whole grain injection,**

**solution**

**FOOD - PLANT SOURCE, RYE GRAIN- rye grain injection, solution**

**FOOD - PLANT SOURCE, SOYBEAN GLYCINE SOJA- soybean glycine**

**soja injection, solution**

**FOOD - PLANT SOURCE, STRAWBERRY FRAGARIA CHILOENSIS- strawberry**

**fragaria chiloensis injection, solution**

**FOOD - PLANT SOURCE, STRING BEAN MIX- string bean mix injection,**

**solution**

**FOOD - PLANT SOURCE, TOMATO NICOTIANA SPP.- tomato nicotiana**

**spp. injection, solution**

**FOOD - PLANT SOURCE, WALNUT, BLACK JUGLANS NIGRA- walnut, black**

**juglans nigra injection, solution**

**FOOD - PLANT SOURCE, YEAST, BAKER SACCHAROMYCES CEREVISIAE- yeast,**

**baker saccharomyces cerevisiae injection, solution**

**FOOD - PLANT SOURCE, YEAST, BREWER SACCHAROMYCES CEREVISIAE-**

**yeast, brewer saccharomyces cerevisiae injection, solution**

**INSECTS WHOLE BODY COCKROACH, AMERICAN PERIPLANETA AMERICANA-**

**insects whole body cockroach, american periplaneta americana injection,**

**solution**

**INSECTS WHOLE BODY COCKROACH, GERMAN BLATELLA GERMANICA- insects**

**whole body cockroach, german blatella germanica injection, solution**

**INSECTS WHOLE BODY COCKROACH MIX- insects whole body cockroach**

**mix injection, solution**

**INSECTS WHOLE BODY, FIRE ANT MIX- insects whole body, fire ant**

**mix injection, solution**

**MOLDS - ALTERNARIA/HORMODENDRUM MIX- molds -**

**alternaria/hormodendrum mix injection, solution**

**MOLDS - MOLD MIX 10- molds - mold mix 10 injection, solution**

**MOLDS - MOLD MIX 4- molds - mold mix 4 injection, solution**

**MOLDS - TRICHOPHYTON MIX-** molds - trichophyton mix injection, solution  
**MOLDS, PENICILLIUM MIX-** molds, penicillium mix injection, solution  
**MOLDS, RUSTS AND SMUTS, ALTERNARIA TENUIS-** alternaria tenuis injection, solution  
**MOLDS, RUSTS AND SMUTS, ASPERGILLUS FUMIGATUS-** aspergillus fumigatus injection, solution  
**MOLDS, RUSTS AND SMUTS, ASPERGILLUS NIGER-** aspergillus niger injection, solution  
**MOLDS, RUSTS AND SMUTS, BOTRYTIS CINerea-** botrytis cinerea injection, solution  
**INSECTS WHOLE BODY, ANT, FIRE SOLENOPSIS INVICTA-** ant, fire solenopsis invicta injection, solution  
**MOLDS, RUSTS AND SMUTS, CANDIDA ALBICANS-** candida albicans injection, solution  
**MOLDS, RUSTS AND SMUTS, CEPHALOSPORIUM ACREMONIUM-** cephalosporium acremonium injection, solution  
**MOLDS, RUSTS AND SMUTS, CURVULARIA SPICIFERA-** curvularia spicifera injection, solution  
**MOLDS, RUSTS AND SMUTS, EPICOCCUM NIGRUM-** epicoccum nigrum injection, solution  
**MOLDS, RUSTS AND SMUTS, EPIDERMOPHYTON FLOCCOSUM-** epidermophyton floccosum injection, solution  
**MOLDS, RUSTS AND SMUTS, FUSARIUM VASINFECTUM-** fusarium vasinfectum injection, solution  
**MOLDS, RUSTS AND SMUTS, HELMINTHOSPORIUM INTERSEMINATUM-** helminthosporium interseminatum injection, solution  
**MOLDS, RUSTS AND SMUTS, HORMODENDRUM CLADOSPORIOIDES-** hormodendrum cladosporioides injection, solution  
**MOLDS, RUSTS AND SMUTS, MUCOR RACEMOSUS-** mucor racemosus injection, solution  
**MOLDS, RUSTS AND SMUTS, PENICILLIUM NOTATUM-** penicillium notatum injection, solution  
**MOLDS, RUSTS AND SMUTS, PHOMA HERBARUM-** phoma herbarum injection, solution  
**MOLDS, RUSTS AND SMUTS, PULLULARIA PULLULANS-** pullularia pullulans injection, solution  
**MOLDS, RUSTS AND SMUTS, RHIZOPUS NIGRICANS-** rhizopus nigricans injection, solution  
**MOLDS, RUSTS AND SMUTS, STEMPHYLIUM BOTRYOSUM-** stemphylium botryosum injection, solution  
**POLLENS - GRASSES, BAHIA GRASS PASPALUM NOTATUM-** bahia grass paspalum notatum injection, solution  
**POLLENS - GRASSES, BROME, SMOOTH BROMUS INERMIS-** brome, smooth bromus inermis injection, solution  
**POLLENS - GRASSES, CORN, CULTIVATED ZEA MAYS-** corn, cultivated zea mays injection, solution  
**POLLENS - GRASSES, JOHNSON GRASS SORGHUM HALEPENSE-** johnson grass sorghum halepense injection, solution  
**POLLENS - GRASSES, OATS, COMMON, CULTIVATED AVENA SATIVA-** oats, common, cultivated avena sativa injection, solution  
**POLLENS - GRASSES, GRASS MIX 8-** grass mix 8 injection, solution  
**POLLENS - GRASSES, SOUTHERN GRASS MIX-** pollens - grasses, southern grass mix injection, solution  
**POLLENS - TREES, ACACIA ACACIA LONGIFOLIA-** acacia longifolia injection, solution  
**POLLENS - TREES, ALDER, RED ALNUS RUBRA-** alder, red alnus rubra injection,

**solution**

**POLLENS - TREES, ASH, WHITE FRAXINUS AMERICANA-** ash, white fraxinus americana injection, solution  
**POLLENS - TREES, BEECH, AMERICAN FAGUS GRANDIFOLIA-** beech, american fagus grandifolia injection, solution  
**POLLENS - TREES, BIRCH MIX-** birch mix injection, solution  
**POLLENS - TREES, BOTTLE BRUSH CALLISTEMON SPP.-** bottle brush callistemon spp. injection, solution  
**POLLENS - TREES, BOXELDER/MAPLE MIX-** boxelder/maple mix injection, solution  
**POLLENS - TREES, CEDAR, MOUNTAIN JUNIPERUS ASHEI-** cedar, mountain juniperus ashei injection, solution  
**POLLENS - TREES, CEDAR, RED JUNIPERUS VIRGINIANA-** cedar, red juniperus virginiana injection, solution  
**POLLENS - TREES, COTTONWOOD, COMMON POPULUS DELTOIDES-** cottonwood, common populus deltoides injection, solution  
**POLLENS - TREES, CYPRESS, ARIZONA CUPRESSUS ARIZONICA-** cypress, arizona cupressus arizonica injection, solution  
**POLLENS - TREES, CYPRESS, BALD TAXODIUM DISTICHUM-** cypress, bald taxodium distichum injection, solution  
**POLLENS - TREES, ELM, AMERICAN ULMUS AMERICANA-** elm, american ulmus americana injection, solution  
**POLLENS - TREES, ELM, CHINESE ULMUS PARVIFOLIA-** elm, chinese ulmus parvifolia injection, solution  
**POLLENS - TREES, EUCALYPTUS, EUCALYPTUS GLOBULUS-** eucalyptus, eucalyptus globulus injection, solution  
**POLLENS - TREES, GUM, SWEET LIQUIDAMBAR STYRACIFLUA-** gum, sweet liquidambar styraciflua injection, solution  
**POLLENS - TREES, HACKBERRY CELTIS OCCIDENTALIS-** hackberry celtis occidentalis injection, solution  
**POLLENS - TREES, HICKORY, SHAGBARK CARYA OVATA-** hickory, shagbark carya ovata injection, solution  
**POLLENS - TREES, LINDEN BASSWOOD TILIA AMERICANA-** linden basswood tilia americana injection, solution  
**POLLENS - TREES, MAPLE, HARD ACER SACCHARUM-** maple, hard acer saccharum injection, solution  
**POLLENS - TREES, MELALEUCA PUNK TREE MELALEUCA QUINQUENERVIA-** melaleuca punk tree melaleuca quinquenervia injection, solution  
**POLLENS - TREES, MESQUITE, PROSOPIS JULIFLORA-** mesquite, prosopis juliflora injection, solution  
**POLLENS - TREES, MULBERRY MIX-** mulberry mix injection, solution  
**POLLENS - TREES, OAK MIX-** oak mix injection, solution  
**POLLENS - TREES, OAK, RED QUERCUS RUBRA-** oak, red quercus rubra injection, solution  
**ANIMAL ALLERGENS, UF DOG HAIR-DANDER CANIS SPP-** animal allergens, dog dander canis spp injection, solution

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**HIGHLIGHTS OF PRESCRIBING INFORMATION**

These highlights do not include all the information needed to use NON-STANDARDIZED ALLERGENIC EXTRACTS (POLLENS, MOLDS, EPIDERMALS AND INSECTS) safely and effectively. See full prescribing information for NON-STANDARDIZED ALLERGENIC EXTRACTS (POLLENS, MOLDS, EPIDERMALS AND INSECTS).

**NON-STANDARDIZED ALLERGENIC EXTRACTS (POLLENS, MOLDS, EPIDERMALS AND INSECTS)**

**Solution for percutaneous, intradermal, or subcutaneous administration Initial U.S.  
Approval: 1925**

**WARNING: ANAPHYLAXIS**

**See full prescribing information for complete boxed warning.**

- Non-standardized allergenic extracts can cause anaphylaxis, including anaphylactic shock and death.(5.1)
- Do not administer to individuals with severe, unstable or uncontrolled asthma, history of severe systemic reaction to the allergen extract when administered for diagnosis or treatment, or with medical conditions that reduce the ability to survive anaphylaxis.(4)
- Observe individuals for at least 30 minutes following administration. Emergency measures and personnel trained in their use must be available in the event of a life-threatening reaction.(5.1)
- Individuals with extreme sensitivity to these products, on an accelerated immunotherapy build-up, switching to another lot, receiving high doses of these products, or exposed to similar allergens may be at increased risk of anaphylaxis.(5.1)
- These products may not be suitable for individuals who may be unresponsive to epinephrine or inhaled bronchodilators, such as those taking beta-blockers.(5.1)

**INDICATIONS AND USAGE**

Non-standardized allergenic extracts are indicated for:

- Skin test diagnosis of patients with a clinical history of allergy to the specific corresponding allergens.(1)
- Immunotherapy for the reduction of allergen-induced allergic symptoms confirmed by positive skin test or by *in vitro* testing for allergen-specific IgE antibodies.(1)

**DOSAGE AND ADMINISTRATION**

**For percutaneous, intradermal, or subcutaneous use only.**

Administration:

- Percutaneous for diagnostic testing.
- Intradermal for diagnostic testing.
- Subcutaneous for immunotherapy.

See full prescribing information for details on dosing and dilution preparation. (2)

**DOSAGE FORMS AND STRENGTHS**

Non-standardized allergenic extract solutions: stock concentrates, labeled in weight/volume, in a glycerin-preserved extracting fluid, supplied in 5, 10, 30, and 50 mL vials. (3, 16) Refer to the vial label for the product concentration. (11)

**CONTRAINDICATIONS**

- Severe, unstable or uncontrolled asthma.(4)
- History of any severe systemic reaction to the allergen extract when administered for diagnosis or treatment.(4)
- Medical conditions that reduce the ability to survive anaphylaxis.(4)

**WARNINGS AND PRECAUTIONS**

The risk of anaphylaxis may be increased in the following situations:

- Extreme sensitivity to non-standardized allergenic extracts.
- Concomitant environmental exposure to similar allergens.
- Receipt of high concentrations and volumes of non-standardized allergenic extracts.
- Receipt of an accelerated build-up schedule (e.g., "rush" immunotherapy).
- Changing to another lot of allergen.(5)

**ADVERSE REACTIONS**

Common adverse reactions reported for non-standardized allergenic extracts are:

- Local adverse reactions, occurring in 26 to 82% of all patients who receive subcutaneous immunotherapy (e.g., erythema, swelling, pruritus, tenderness and pain at the injection site).(6)
- Systemic adverse reactions, occurring in  $\leq$  7% of patients who receive subcutaneous immunotherapy (e.g., generalized skin erythema, urticaria, pruritus, angioedema, rhinitis, wheezing, laryngeal edema, and hypotension). Systemic reactions may be fatal.(6)

To report SUSPECTED ADVERSE REACTIONS, contact Jubilant HollisterStier at 1-800-495-7437 or Adverse.Reactions@jhs.jubl.com; or the FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

**DRUG INTERACTIONS**

- Certain medications may decrease skin test wheal and erythema responses, including antihistamines, topical corticosteroids, topical anesthetics, and tricyclic antidepressants.(7)

**See 17 for PATIENT COUNSELING INFORMATION.**

**Revised: 10/2024**

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## **FULL PRESCRIBING INFORMATION**

## **WARNING: ANAPHYLAXIS**

- Non-standardized allergenic extracts can cause anaphylaxis, including anaphylactic shock and death.(5.1)
- Do not administer to individuals with:
  - severe, unstable or uncontrolled asthma;
  - history of severe systemic reaction to the allergen extract when administered for diagnosis or treatment;
  - medical conditions that reduce the ability to survive anaphylaxis.(4)
- Observe individuals for at least 30 minutes following administration. Emergency measures and personnel trained in their use must be available in the event of a life-threatening reaction.(5.1)
- Individuals with extreme sensitivity to these products, on an accelerated immunotherapy build-up, switching to another lot, receiving high doses of these products, or exposed to similar allergens may be at increased risk of anaphylaxis.(5.1)
- These products may not be suitable for individuals who may be unresponsive to epinephrine or inhaled bronchodilators, such as those taking beta-blockers.(5.1)

## **1 INDICATIONS AND USAGE**

NON-STANDARDIZED ALLERGENIC EXTRACTS are indicated for:

- Skin test diagnosis of individuals with a clinical history of allergy to the specific corresponding allergens.

NON-STANDARDIZED ALLERGENIC EXTRACTS are indicated for:

- Immunotherapy for the reduction of allergen-induced allergic symptoms confirmed by positive skin test or by *in vitro* testing for allergen specific IgE antibodies for the specific corresponding allergens.

## **2 DOSAGE AND ADMINISTRATION**

**For percutaneous, intradermal, or subcutaneous administration only. Do not inject intravenously.**

### **2.1 Preparation for Administration**

Appearance is clear to slightly opalescent. Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Discard solution if either of these conditions exist.

Non-standardized allergenic extracts diluted with Albumin Saline with Phenol (0.4%) (stabilized diluent) may be more potent than extracts diluted with diluents that do not contain albumin. When switching from non-stabilized to stabilized diluent, consider less concentrated initial dilutions for both intradermal testing and immunotherapy.

Different formulations, preparations, or new lots of non-standardized allergenic extracts are not interchangeable. Dosing should be adjusted appropriately when formulations, preparations, or lots of non-standardized allergenic extracts are changed [see *Immunotherapy (2.3)* and *Dosage Forms and Strengths (3)*].

Allergenic extracts may be prepared for intradermal (diagnosis) or subcutaneous (immunotherapy) administration by diluting stock concentrates.

- For diluent, use sterile albumin saline with phenol or sterile normal saline with phenol.

- Dilute stock concentrates by a minimum of 100-fold for intradermal testing. Dilutions of 1,000-fold or greater are appropriate starting points for patients with a clinical history of adverse reaction.

To prepare dilutions for intradermal testing and immunotherapy, start with a stock concentrate, and prepare a ten-fold (1:10) dilution by adding 0.5 mL of concentrate to 4.5 mL of sterile aqueous diluent. Prepare subsequent dilutions in a similar manner. (see Table 1).

**Table 1: 10-fold Dilution Series**

Dilution	Extract	Milliliters of Diluent	Dilution Strength (w/v)				
0	Concentrate		1:10	1:20	1:50	1:100	1:650
1	0.5 mL Concentrate	4.5	1:100	1:200	1:500	1:1,000	1:6,500
2	0.5 mL Dilution	4.5	1:1,000	1:2,000	1:5,000	1:10,000	1:65,000
3	0.5 mL Dilution 2	4.5	1:10,000	1:20,000	1:50,000	1:100,000	1:650,000
4	0.5 mL Dilution 3	4.5	1:100,000	1:200,000	1:500,000	1:1,000,000	1:6,500,000
5	0.5 mL Dilution 4	4.5	1:1,000,000	1:2,000,000	1:5,000,000	1:10,000,000	1:65,000,000
6	0.5 mL Dilution 5	4.5	1:10,000,000	1:20,000,000	1:50,000,000	1:100,000,000	1:650,000,000

Note: A lower starting dose and/or less concentrated dilutions may be necessary for highly sensitive patients with a clinical history of sensitivity, or for those who display severe symptoms. [see *Diagnostic Testing (2.2), Percutaneous Skin Testing (2.2.1) and Intradermal (Intracutaneous) Skin Test (2.2.2)*].

## 2.2 Diagnostic Testing

Testing is performed to identify patients that exhibit an allergic response at the site of administration. False positive reactions may occur. A positive skin test reaction must be interpreted in the context of the individual's clinical history and known exposure to the allergen.

- Administer percutaneous tests prior to administration of intradermal tests to identify highly sensitive patients.
- Do not use allergen mixes for diagnostic testing because a positive reaction would not permit specific identification of the allergen(s) that elicited the reaction. In addition, a negative reaction would fail to indicate whether an individual component allergen would have elicited a positive reaction at full strength.

### 2.2.1 Percutaneous Skin Testing

#### Dose

Unless an individual is suspected to be at greater risk for anaphylaxis, the initial starting dose is 1 drop (approximately 0.05 mL) of undiluted allergenic extract. For individuals suspected to be at greater risk for anaphylaxis (for example, as indicated by a history of allergen-induced anaphylaxis), initiate percutaneous testing with a sequence of serial 10-fold dilutions of undiluted allergenic extract spaced 15-20 minutes apart [see *Preparation for Administration (2.1)*].

#### Administration

- Percutaneous Test: Place one drop (approximately 0.05 mL) of allergen on the skin and pierce through drop superficially into the skin, lifting slightly. Use a skin test device, such as a sterile needle, lancet, or bifurcated needle.
- Percutaneous Test using self-loading devices: Refer to the manufacturer's product instructions.

Concurrently, use a positive histamine skin test control to identify patients whose recent use of drugs with antihistamine activity may result in a false negative skin test. Apply a 50% glycerin solution as a negative control, to identify false positive responses to the extracting fluid used in the manufacture of allergenic extracts, or due to dermographism [see *Drug Interactions* (7)].

## **Interpreting Results**

For interpretation of percutaneous skin tests, refer to the information provided in Allergy Diagnostic Testing: An Updated Practice Parameter.<sup>1</sup> In addition, follow the directions provided with the percutaneous skin test devices. Measure wheal responses for the histamine positive control test at 15 minutes and for the allergen tests at 15 to 20 minutes.

- The negative control (50% glycerin solution) response should measure < 3 mm wheal and ≤ 10 mm flare.<sup>1</sup>
- Response to positive controls should be at least 3 millimeters larger than the response to the negative control.
- If either the response to the histamine positive control or to the negative control do not meet the criteria above for acceptable wheal size, the results for the allergenic extracts tested at the same time should be considered invalid and be repeated.
- Fire Ant: Percutaneous testing is considered positive when the response occurs at a concentration of 1:100 w/v or less.<sup>4</sup>

### **2.2.2 Intradermal (Intracutaneous) Skin Test**

Always perform percutaneous tests prior to intradermal skin tests.<sup>1, 2</sup>

#### **Dose**

Perform intradermal tests with at least 100-fold less concentrated solutions than the stock concentrates used in percutaneous tests [see *Preparation and Administration* (2.1)].

Fire Ant: Use 0.02 mL of a 1:100,000 v/v dilution of the concentrate for intradermal tests. Very sensitive individuals such as those who have had nearly fatal anaphylactic reactions may not tolerate even 1:100,000 v/v dilution of concentrate as a starting point. These patients should be tested with a 1:10,000,000 v/v dilution of concentrate [see *Preparation for Administration* (2.1)].

Use intradermal tests following a negative or equivocal percutaneous test when the patient continues to report a history of symptoms following exposure to a specific allergen.

#### **Administration**

Intradermally inject 0.02 mL of the allergen using a 1 mL intradermal testing syringe with a 26 or 27 gauge, 1/2" or 3/8" needle with intradermal bevel, graduated in 0.01 units. Insert needle at a 30° angle, bevel down.

Test concurrently with a positive histamine control at intradermal strength (0.1 mg/mL of histamine base) and an aqueous buffer negative control (Sterile Albumin Saline with Phenol, Sterile Buffered Saline with Phenol).

## **Interpreting Results**

For interpretation of intradermal skin tests, follow the information provided in Allergy Diagnostic Testing: An Updated Practice Parameter.<sup>1</sup>

- Measure wheal responses for the histamine positive control test and allergen tests at 10-15 minutes after injection
- Response to the positive control should be at least 3 millimeters larger than the response to the negative control.
- The negative control (50% glycerin solution) response should measure < 3-mm wheal and ≤ 10 mm flare (erythema).
- If either the response to the histamine positive control or to the negative control do not meet the criteria above for acceptable wheal size, the results for the allergenic extracts tested at the same time should be considered invalid and be repeated.
- Fire Ant: Intradermal testing is considered positive when the response occurs at a concentration of 1:1,000 w/v or less.<sup>4</sup>

## **2.3 Immunotherapy**

**For subcutaneous administration only.**

### **Administration of Immunotherapy**

Administer immunotherapy by subcutaneous injection in the lateral aspect of the arm or thigh. Avoid injection directly into any blood vessels. Administer injections with a sterile 1 mL allergy treatment syringe with a 26 or 27 gauge, 1/2", beveled needle, graduated in 0.01 units.

The optimal interval between doses of allergenic extract varies among individuals. Injections are usually given one or two times per week until the maintenance dose is reached, at which time the injection interval is increased to 2, 3, and finally 4 weeks.

Most adverse reactions occur within 30 minutes after injection. Therefore, observe patients for at least 30 minutes. For high risk patients, 30 minutes of observation may not be sufficient.<sup>2</sup>

Dosing of non-standardized allergenic extracts for allergen immunotherapy is highly individualized. Adjust dose according to the degree of sensitivity of the patient, tolerance to the extract administered during the early phases of an injection regimen, and the clinical response. Dosing is individualized by choice of an initial dose, the schedule of dose build-up, the target maintenance dose, the actual maintenance dose, and the duration of treatment.

The large volume of solution for immunotherapy may produce increased discomfort in the pediatric population. In order to achieve the total dose required, the volume of the dose may need to be divided into more than one injection per visit.<sup>2</sup>

#### **2.3.1 Dose Build-up**

Following the first administration of 0.03 mL of the selected initial dilution of allergenic extract, dosing is increased in 0.03 mL to 0.12 mL increments until 0.3 mL is reached, following which 0.03 mL is administered from the next most concentrated allergen extract or allergen mixture vial in the dilution series. The interval between doses is usually 3 to 7 days during dose build-up. Proceed in this manner until a maintenance dose is reached. The final maintenance dose may not be the target maintenance dose selected at the beginning of therapy.

The following adjustments may be necessary during dose build-up:

- If allergic symptoms or local reactions develop shortly after dose administration, decrease the dose volume to one-half or one-quarter of the maximum dose previously attained.

- If the patient is experiencing any seasonal allergy symptoms, decrease the dose volume to one-half or one-quarter of the maximum dose previously attained.
- Adjust the dose periodically based on the patient's tolerance and reaction.
- Decrease the dose if the previous injection resulted in a marked local reaction.
- Repeat the previous dose or reduce the dose at the next administration if local reactions persist for longer than 24 hours.
- Decrease the dose if the previous injection resulted in a systemic reaction. Any evidence of a systemic reaction is an indication for a significant (at least 75%) reduction in the subsequent dose or the cessation of immunotherapy.
- Repeated systemic reactions, however mild, are sufficient reason for the cessation of further attempts to increase the reaction-causing dose.

### **2.3.2 Maintenance Dose Selection, Adjustments, and Intervals**

The maintenance dose is the dose that provides therapeutic efficacy without severe adverse local or systemic reactions. This dose may be limited by adverse reactions and may not be the original targeted maintenance dose. Select a maintenance dose based on the patient's clinical response and tolerance.

- Suggested maintenance dose is 0.3 mL of the undiluted allergen extract. Occasionally, higher doses are necessary to relieve symptoms.
- Maintenance doses larger than 0.3 mL of undiluted allergen extract may cause patient discomfort due to the 50% glycerin content.
- After the maintenance dose is achieved, increase the injection interval to 2 weeks, then 3 weeks, and finally 4 weeks, as tolerated. Administer the maintenance dose at a given interval three or four times before further increasing the interval to assure that no reactions occur. Protection may be lost rapidly if the interval between doses is more than 4 weeks.

The following adjustments to the maintenance dose may be necessary.

***Withhold immunotherapy and/or reduce dosage, if any of the following conditions exist:***

- Severe symptoms of rhinitis and/or asthma. Decrease dose to one-half or one-quarter of the maximum dose previously attained if the patient has any seasonal symptoms.
- Allergic symptoms or a local reaction following the prior dose.
- Infection accompanied by fever.
- Exposure to excessive amounts of clinically relevant allergen prior to a scheduled injection.

In situations prompting dose reduction, a cautious increase in dosage can be attempted once the reduced dose is tolerated.

***Decrease the interval between doses*** if symptoms develop before the next injection is scheduled.

***In some patients, the dosage may be increased and/or the dosing interval shortened*** based on individual responses and dosing requirements. If the onset of symptoms is soon after the initiation of immunotherapy, decrease the interval between each dose.

***Changing to a different lot of extract:*** All extracts can lose allergenic activity over time and extracts vary in allergenic activity. Two different lots of extract could differ substantially in allergenic activity, even if they are the same formula and concentration. The volume of the first dose from the new vial should not exceed 50% of the previous dose. Do not use extracts beyond their expiry date.

**Changing to a different formulation of extract or to an extract from a different manufacturer:** Decrease the starting dose of the new extract when the extract is the same formula and dilution as the one previously used. In general, a volume dose reduction to 50% of the previous product dose is adequate, but each situation must be evaluated separately considering the patient's history of sensitivity, tolerance of previous injections, and other factors. If the patient tolerates the 50% decrease, then raise the next dose to the previous tolerated dose amount. To re-establish the maintenance dose the starting interval between doses should not be greater than one week.

**Prolonged period has elapsed since the last injection:** Patients may lose tolerance for allergen injections during prolonged intervals (> 4 weeks) between doses. The duration of tolerance is an individual characteristic and varies from patient to patient. In general, the longer the lapse in the injection schedule, the greater dose reduction required.

**Changes made in the extract concentrate formula:** Changes other than those listed above such as a difference in extracting fluid (e.g., change from non-glycerin extracts to 50% glycerin extracts), combining two or more stock concentrates, or any other change can affect a patient's tolerance of the treatment. Extra dilutions are recommended whenever starting a revised formula. The greater the change, the greater the number of dilutions required.

## **Duration of Treatment**

The duration of treatment for immunotherapy has not been established. A period of two to three years of injection therapy constitutes an average minimum course of treatment. Evaluate patients for treatment response at least every 6 to 12 months while they receive immunotherapy.

## **3 DOSAGE FORMS AND STRENGTHS**

Non-standardized allergenic extracts are solutions: stock concentrates, labeled in weight/volume, in a glycerin-preserved extracting fluid, supplied in 5, 10, 30, and 50 mL vials. (3, 16) Refer to the vial label for the product concentration. (11)

## **4 CONTRAINDICATIONS**

Non-standardized allergenic extracts are contraindicated in individuals with the following conditions:

- Severe, unstable or uncontrolled asthma.
- History of any severe systemic reaction to the allergen extract when administered for diagnosis or treatment.
- Medical conditions that reduce the ability to survive anaphylaxis.

## **5 WARNINGS AND PRECAUTIONS**

### **5.1 Anaphylaxis**

Anaphylaxis, which may lead to death, can occur in individuals following the administration of non-standardized allergenic extracts, particularly in the following situations:

- Extreme sensitivity to the non-standardized allergenic extract.
- Concomitant environmental exposure to allergens.
- Receipt of high doses of the non-standardized allergenic extract.
- Receipt of an accelerated build-up schedule ("rush" immunotherapy).

- Change from one lot of a particular non-standardized allergenic extract to another lot of the same non-standardized allergenic extract.

Administer non-standardized allergenic extracts in a healthcare setting under the supervision of a physician prepared to manage anaphylaxis; management may include use of inhaled bronchodilators and use of epinephrine. Non-standardized allergenic extracts may not be suitable for individuals who may be unresponsive to epinephrine or inhaled bronchodilators, such as those taking beta-blockers. See prescribing information for epinephrine for complete information, particularly on medications that blunt or potentiate epinephrine activity. Individuals should remain in the physician's office for a minimum of 30 minutes after receiving an injection of non-standardized allergenic extracts, so that any adverse reaction can be observed and properly handled.

## **5.2 Cross-reactions and Dose Sensitivity**

When determining the final dose of an allergen mixture for immunotherapy, consider cross-reactivity among component extracts.

Determine the initial dilution of allergenic extract, starting dose, and progression of dosage based on the patient's history and results of skin tests [see *Dosage and Administration (2)*]. Strongly positive skin tests can be indicators for potential adverse reactions.

## **6 ADVERSE REACTIONS**

Common adverse reactions reported for non-standardized allergenic extracts are:

- Local reactions occurring in 26 to 82% of all patients who receive subcutaneous immunotherapy, at the injection site (e.g., erythema, swelling, pruritus, tenderness and pain).<sup>2</sup>
- Systemic adverse reactions, occurring in ≤ 7% of patients who receive subcutaneous immunotherapy (e.g., generalized skin erythema, urticaria, pruritus, angioedema, rhinitis, wheezing, laryngeal edema, hypotension, and shock).<sup>3</sup> Systemic reactions may be fatal.<sup>2</sup>

No clinical trials of non-standardized allergenic extracts have been conducted.

Published studies of non-standardized allergenic extracts report systemic reactions occurring in fewer than 1% in patients receiving conventional immunotherapy and greater than 36% in patients receiving rush immunotherapy. Most systemic reactions occurred within 30 minutes of injection. However, systemic reactions have been reported to occur up to 2 hours after the final injection with rush schedules. Some reactions have occurred up to 6 hours after skin tests or immunotherapy.<sup>2, 3</sup>

## **7 DRUG INTERACTIONS**

### **7.1 Antihistamines**

Do not perform skin testing with non-standardized allergenic extracts within 3 to 10 days of first-generation H1-histamine receptor blockers (e.g., clemastine, diphenhydramine) and second-generation antihistamines (e.g., loratadine, fexofenadine) being used. These products suppress histamine skin test reactions and could mask a positive response.<sup>1, 2</sup>

### **7.2 Topical Corticosteroids and Topical Anesthetics**

Topical corticosteroids may suppress skin reactivity; therefore, discontinue use at the

skin test site for at least 2 to 3 weeks before skin testing. Avoid use of topical local anesthetics at skin test sites because they can suppress flare responses.<sup>1, 2</sup>

### **7.3 Tricyclic Antidepressants**

Tricyclic antidepressants, such as doxepin, can have potent antihistamine effects and may alter skin test results. Allow 7 to 14 days after discontinuation of tricyclic medication prior to skin testing.<sup>1, 2</sup>

## **8 USE IN SPECIFIC POPULATIONS**

### **8.1 Pregnancy**

#### **Risk Summary**

All pregnancies have a risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively. There are no human or animal data to establish the presence or absence of non-standardized allergenic extracts-associated risks during pregnancy.

### **8.2 Lactation**

#### **Risk Summary**

It is not known whether non-standardized allergenic extracts are present in human milk. Data are not available to assess the effects of these extracts on the breastfed child or on milk production/excretion. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for non-standardized allergenic extracts and any potential adverse effects on the breastfed child from the extracts or from the underlying maternal condition.

### **8.4 Pediatric Use**

For use of these products in children younger than 5 years of age, consideration should be given to the patient's ability to comply and cooperate with receipt of the product and the potential for difficulty in communicating with the child regarding systemic reactions.<sup>2</sup>

The volume of a dose for immunotherapy may need to be divided for pediatric patients [see *Dosage and Administration (2.3)*]

### **8.5 Geriatric Use**

Data are not available to determine if subjects 65 years of age and older respond differently to allergen immunotherapy than younger subjects.

## **11 DESCRIPTION**

Non-standardized allergenic extracts are labeled "No U.S. Standard of Potency".

Non-standardized allergenic extracts are supplied in a Glycero Cocos extraction solution, which consists of 0.5% sodium chloride for isotonicity, 0.275% sodium bicarbonate as a buffer, and 50% glycerin (volume/volume) as preservative.

Non-standardized allergenic extracts are supplied as a weight to volume (w/v) solution of allergen in extraction solution. Product concentrations vary based on the source. Refer to the vial label for the product concentration.

Source material mold mycelia and *Candida albicans* cells are cultivated on liquid medium which may contain one or more of the following constituents: casein hydrolysate; malt

extract; yeast extract; maltose; dextrose; ammonium nitrate, calcium carbonate, calcium chloride, ammonium citrate, potassium phosphate, sodium citrate, citric acid; magnesium sulfate; or trace elements. Acetone and ether may be used as drying and de-fattening agents. *Candida albicans* cells are treated with phenol, which is removed by dialysis.

Dog Hair and Dander extracts are manufactured in 3 product forms:

- Dog Hair and Dander (Regular Process) is derived from extraction of the source material without additional processing, and is prepared at 1:10 w/v in Glycero-Cocas.
- Acetone Precipitated (AP) Dog Hair and Dander is derived from the acetone precipitated aqueous extract and is prepared at 1:100 w/v in Glycero-Cocas.
- Ultrafiltered (UF) Dog Hair and Dander is derived from the UF aqueous extract and is prepared at 1:650 w/v in Glycero-Cocas.

## **12 CLINICAL PHARMACOLOGY**

### **12.1 Mechanism of Action**

The skin test reaction results from interaction of the introduced allergen and allergen-specific IgE antibodies bound to mast cells, leading to mast cell degranulation and release of histamine, tryptase and other mediators, which results in the formation of the wheal and flare.

The precise mechanisms of action of allergen immunotherapy are not known.

Immunologic responses to immunotherapy include changes in allergen-specific IgE levels, allergen-specific IgG levels, and regulatory T cell responses.<sup>2</sup>

## **14 CLINICAL STUDIES**

Specific immunotherapy with allergenic extracts is helpful in reducing symptoms associated with exposure to the offending allergens. A summary of effectiveness by the Panel on Review of Allergenic Extracts, an advisory committee to the U.S. Food and Drug Administration, has been published.<sup>5</sup>

## **15 REFERENCES**

1. Bernstein IL, Li JT, Bernstein DI, et al. Allergy diagnostic testing: and updated practice parameter. Ann Allergy Asthma Immunol. 2008 Mar;100:S1-148.
2. Cox L, Nelson H, Lockey R, Calabria C, Chacko T, Finegold I, et al. Allergen immunotherapy: A practice parameter third update. J Allergy Clin Immunol. 2011 Jan;127:S1-55.
3. Greineder DK. Risk management in allergen immunotherapy. J Allergy Clin Immunol. 1996 Dec;98(6 Pt 3):S330-4
4. Golden D B K, Demain J, Freeman T, Graft D, et al. Stinging insect hypersensitivity: A practice parameter update 2016. Ann Allergy Asthma Immunol 118 (2017) 28-54.
5. Federal Register Proposed Rule: Biological Products: Implementation of Efficacy Review, Allergenic Extracts, Federal Register 1985;50:3082-3288.

## **16 HOW SUPPLIED**

Non-standardized allergenic extracts and mixes are supplied as 50% glycerin stock concentrates labeled in weight/volume and provided in 10 milliliter, 30 milliliter and 50 milliliter vials for use in percutaneous skin testing and subcutaneous

immunotherapy. These extracts may also be supplied in 5 milliliter dropper vials for percutaneous testing only.

These products are supplied as listed in Table 2.

**TABLE 2: AVAILABLE PRODUCTS****POLLEN - GRASS ALLERGENS**

Bahia Grass,*Paspalum notatum*

Brome, Smooth*Bromus inermis*

Corn, Cultivated*Zea mays*

Grass Mix 8-100,000 BAU/mL each of *P. pratensis*; *A. gigantean*; *P. pretense*; 10,000 BAU/mL of *C. dactylon*; 1:20w/v of *S. halepense*

Johnson Grass, *Sorghum halepense*

Oats, Common Cultivated, *Avena sativa*

**POLLEN - TREE ALLERGENS**

Acacia, Golden, *Acacia longifolia*

Alder, Red, *Alnus rubra*

Ash, White, *Fraxinus americana*

Beech, American, *Fagus grandifolia*

Birch Mix (PRW)-*B. papyrifera*, *B. pendula*, *B. nigra*

Bottlebrush, *Melaleuca citrina*

Boxelder/Maple Mix (BHR)-*A. negundo*, *A. saccharum*, *A. rubrum*

Cedar, Mountain, *Juniperus ashei*

Cedar, Red, *Juniperus virginiana*

Cottonwood, Common, *Populus deltoides*

Cypress, Arizona, *Cupressus arizonica*

Cypress, Bald, *Taxodium distichum*

Elm, American, *Ulmus americana*

Elm, Chinese, *Ulmus parvifolia*

Gum, Sweet, *Liquidambar styraciflua*

Hackberry, *Celtis occidentalis*

Hickory, Shagbark, *Carya ovata*

Maple, Hard/Sugar, *Acer saccharum*

Melaleuca, *Melaleuca quinquenervia*

Mesquite, *Prosopis glandulosa*

Mulberry Mix (RW)-*M. rubra*, *M. alba*

Oak Mix (RVW)-*Q. rubra*, *Q. virginiana*, *Q. alba*

Oak, Red, *Quercus Rubra*

Olive Tree, *Olea europaea*

Palm, Queen, *Syagrus romanzoffiana*

Pecan Tree, *Carya illinoinensis*

Pine Mix (LY)-*P. contorta*, *P. ponderosa*

Privet, Common, *Ligustrum vulgare*

Russian Olive, *Elaeagnus angustifolia*

Sycamore, American, *Platanus occidentalis*

Tree Mix 5-20% each of *F. Americana*; *J. nigra*; *P. deltoides*; *U. Americana*; 6.7% each of *B. papyrifera*; *B. nigra*; *B. pendula*

Tree Mix 6- Tree Mix 6-20% each of *F. Americana*; *J. nigra*; *P. deltoides*; *U. Americana*; 6.7% each of *B. papyrifera*; *B. nigra*; *B. pendula*

Tree Mix 11-10% each of *F. americana*; *B. nigra*; *J. nigra*; *P. deltoides*; *U. americana*; *C. ovata*; *A. saccharum*; *Q. rubra*; *P. occidentalis*; *S. nigra*

Walnut, Black, *Juglans nigra*

Willow, Black, *Salix nigra*

**POLLEN - WEED AND GARDEN PLANT ALLERGENS**

Careless Weed, <i>Amaranthus palmeri</i>
Careless/Pigweed Mix (CR)- <i>A. palmeri</i> , <i>A. retroflexus</i>
Cocklebur, Common, <i>Xanthium strumarium</i>
Dock/Sorrel Mix (DS)- <i>R. crispus</i> , <i>R. acetosella</i>
Dog Fennel, Eastern, <i>Eupatorium capillifolium</i>
Goldenrod, <i>Solidago canadensis</i>
Kochia, <i>Kochia scoparia</i>
Lamb's Quarters, <i>Chenopodium album</i>
Marshelder/Poverty Mix (BPT)- <i>C. xanthifolia</i> , <i>I. annua</i> , <i>I. axillaris</i>
Nettle, <i>Urtica dioica</i>
Pigweed, Rough Redroot, <i>Amaranthus retroflexus</i>
Plantain, English, <i>Plantago lanceolata</i>
Ragweed, Giant, <i>Ambrosia trifida</i>
Ragweed Mix (GSW)- <i>A. trifida</i> , <i>A. artemisiifolia</i> , <i>A. psilostachya</i>
Ragweed, Western, <i>Ambrosia psilostachya</i>
Russian Thistle, <i>Salcoloma kali</i>
Sagebrush, Mugwort, <i>Artemisia vulgaris</i>
Scale, Wing, <i>Atriplex canescens</i>
Sorrel, Sheep, <i>Rumex acetosella</i>
Weed Mix 2630-25% each of <i>X. strumarium</i> ; <i>C. album</i> ; <i>A. retroflexus</i> ; 12.5% each of <i>R. crispus</i> ; <i>R. acetosella</i>
MOLDS
<i>Alternaria/Hormodendrum</i> Mix- <i>A. tenuis</i> , <i>H. cladosporioides</i>
<i>Alternaria tenuis</i> ( <i>Alternaria alternata</i> )
<i>Aspergillus fumigatus</i>
<i>Aspergillus niger</i> var. <i>niger</i>
<i>Botrytis cinerea</i>
<i>Candida albicans</i>
<i>Cephalosporium acremonium</i> ( <i>Sarocladium strictum</i> )
<i>Curvularia spicifera</i> ( <i>Cochliobolus spicifer</i> )
<i>Epicoccum nigrum</i>
<i>Epidermophyton floccosum</i>
<i>Fusarium vasinfectum</i> ( <i>Fusarium oxysporum</i> <i>vasinfectum</i> )
<i>Helminthosporium interseminatum</i> ( <i>Dendryphiella vinosa</i> )
<i>Hormodendrum cladosporioides</i> ( <i>Cladosporium cladosporioides</i> )
Mold Mix 4-25% each of <i>A. alternata</i> ; <i>C. cladosporioides</i> ; 6.2% each of <i>A. fumigatus</i> ; <i>A. nidulans</i> ; <i>A. nigervar. niger</i> ; <i>A. terreus</i> ; <i>P. digitatum</i> ; <i>P. expansum</i> ; <i>P. chrysogenum</i> var. <i>chrysogenum</i> ; <i>C. rosea</i> f. <i>rosea</i>
Mold Mix 10-2.5% each of <i>A. fumigatus</i> ; <i>A. nidulans</i> ; <i>A. nigervar. niger</i> ; <i>A. terreus</i> ; <i>P. digitatum</i> ; <i>P. expansum</i> ; <i>P. chrysogenum</i> var. <i>chrysogenum</i> ; <i>C. roseaf. rosea</i> ; 10% each of <i>A. alternata</i> ; <i>F. oxysporum</i> <i>vasinfectum</i> ; <i>D. vinosa</i> ; <i>C. cladosporioides</i> ; <i>M. racemosus</i> ; <i>P. exigua</i> var. <i>exigua</i> ; <i>A. pullulans</i> var. <i>pullutans</i> ; <i>R. stolonifer</i>
<i>Mucor racemosus</i>
<i>Penicillium</i> Mix- <i>P. expansum</i> , <i>P. digitatum</i> , <i>P. chrysogenum</i> , <i>C. rosea</i>
<i>Penicillium notatum</i> ( <i>Penicillium chrysogenum</i> var. <i>chrysogenum</i> )
<i>Phoma herbarum</i> ( <i>Phoma exigua</i> var. <i>exigua</i> )
<i>Pullularia pullulans</i> ( <i>Aerobasidium</i> <i>pullulans</i> var. <i>pullulans</i> )
<i>Rhizopus nigricans</i> ( <i>Rhizopus stolonifer</i> )
<i>Stemphylium botryosum</i> ( <i>Pleosporatarda</i> )
Trichophyton Mix- <i>T. tonsurans</i> , <i>T. rubrum</i> , <i>T. mentagrophytes</i>
EPIDERMALS
AP Horse Hair and Dander, <i>Equuscaballus</i>
AP Cattle Hair and Dander, <i>Bostaurus</i>
AP Dog Hair and Dander, <i>Canislupusfamiliaris</i>

Dog Hair and Dander, <i>Canis lupus familiaris</i>
UF Dog Hair and Dander, <i>Canis lupus familiaris</i>
Feather Mix- <i>G. gallus, A. platyrhynchos, A. anser</i>
Guinea Pig Hair and Dander, <i>Cavia porcellus</i>
<b>INSECTS</b>
Cockroach, American, <i>Periplaneta americana</i>
Cockroach, German, <i>Blattella germanica</i>
Cockroach Mix- <i>P. americana, B. germanica</i>
Fire Ant, <i>Solenopsis invicta</i>

## **16.2 Storage and Handling**

Store extracts at 2°C to 8°C (36°F to 46°F).

## **17 PATIENT COUNSELING INFORMATION**

Instruct patients to remain in the office under observation for a minimum of 30 minutes after an injection or longer, if deemed necessary for the individual.

Inform patients that reactions may occur more than 30 minutes after skin testing or an injection.

Instruct patient to recognize the following symptoms as systemic adverse reactions and seek emergency medical care right away if any of these symptoms occur:

- Unusual swelling and/or tenderness at the injection site.
- Hives or itching of the skin.
- Swelling of face and/or mouth.
- Sneezing, coughing, or wheezing.
- Shortness of breath.
- Nausea.
- Dizziness or faintness.

Manufacturer:

**Jubilant HollisterStier LLC**

Spokane, WA 99207 U.S.A.

U.S. Lic. No. 1272

Version Date: February 24, 2022

## **PRINCIPAL DISPLAY PANEL**

## ALLERGENIC EXTRACT

Preservative:  
50% Glycerin v/v

Inactive Ingredients:  
0.5% Sodium chloride  
0.275% Sodium bicarbonate

UF DOG  
HAIR-DANDER  
(ULTRAFILTERED)  
*Canis lupus familiaris*  
hair & dander



(01)00000000000000  
(17)190702  
(10)S1234567  
(21)00000000000001

1:650 w/v

Item: XXXXX  
Lot: S1234567  
Exp: 2019Jul02

No U.S. standard of potency  
Dose/Route: See Package Insert

NDC: 65044-4850-2  
U.S. License No. 1272

10 mL      Item: XXXXX

5000000XXXX-H01

Non-Returnable

Store at 2-8°C

5000000XXXX-H01

Rx Only - Sterile

### UF Dog Hair-Dander, 10 mL 1.650wv Carton Label

## ALLERGENIC EXTRACT

UF DOG HAIR-DANDER  
(ULTRAFILTERED)  
*Canis lupus familiaris* hair & dander

10 mL      **1:650 w/v**  
Dose/Route: See Package Insert

Rx Only - Sterile  
Store at 2-8°C

5000000XXXX-H01

Preservative 50% Glycerin v/v  
U.S. License No. 1272

Item: XXXXX  
Lot: S1234567  
Exp: 2019Jul01

5000000XXXX-H01

Jubilant HollisterStier LLC Spokane, WA 99207

### UF Dog Hair-Dander, 10 mL 1.650wv Vial Label

## ANIMAL ALLERGENS, AP CATTLE HAIR AND DANDER

cattle hair and dander injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4811
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BOS TAURUS HAIR</b> (UNII: TOQ97Z8644) (BOS TAURUS HAIR - UNII:TOQ97Z8644)	BOS TAURUS HAIR	0.01 g in 1 mL
<b>BOS TAURUS DANDER</b> (UNII: C8VYS72608) (BOS TAURUS DANDER - UNII:C8VYS72608)	BOS TAURUS DANDER	0.01 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4811-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4811-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-4811-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	01/30/1978	

## ANIMAL ALLERGENS, AP DOG HAIR AND DANDER CANIS SPP

animal allergens, dog dander canis spp injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4824
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>CANIS LUPUS FAMILIARIS HAIR</b> (UNII: 05S7L91ZTR) (CANIS LUPUS FAMILIARIS HAIR - UNII:05S7L91ZTR)	CANIS LUPUS FAMILIARIS HAIR	0.005 g in 1 mL
<b>CANIS LUPUS FAMILIARIS DANDER</b> (UNII: 11JCK302I4) (CANIS LUPUS FAMILIARIS DANDER - UNII:11JCK302I4)	CANIS LUPUS FAMILIARIS DANDER	0.005 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4824-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4824-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-4824-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	08/24/1976	

## ANIMAL ALLERGENS, DOG HAIR AND DANDER CANIS SPP.

dog hair canis spp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4083
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CANIS LUPUS FAMILIARIS HAIR (UNII: 05S7L91ZTR) (CANIS LUPUS FAMILIARIS HAIR - UNII:05S7L91ZTR)	CANIS LUPUS FAMILIARIS HAIR	0.05 g in 1 mL
CANIS LUPUS FAMILIARIS DANDER (UNII: 11JCK302I4) (CANIS LUPUS FAMILIARIS DANDER - UNII:11JCK302I4)	CANIS LUPUS FAMILIARIS DANDER	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4083-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4083-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
	NDC:65044-4083-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

3	NDC:65044-4083-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		
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## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## ANIMAL ALLERGENS, DOG HAIR AND DANDER CANIS SPP.

dog hair canis spp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4085
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CANIS LUPUS FAMILIARIS HAIR (UNII: 05S7L91ZTR) (CANIS LUPUS FAMILIARIS HAIR - UNII:05S7L91ZTR)	CANIS LUPUS FAMILIARIS HAIR	0.05 g in 1 mL
CANIS LUPUS FAMILIARIS DANDER (UNII: 11JCK302I4) (CANIS LUPUS FAMILIARIS DANDER - UNII:11JCK302I4)	CANIS LUPUS FAMILIARIS DANDER	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4085-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4085-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-4085-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## ANIMAL ALLERGENS, AP HORSE HAIR AND DANDER

ap horse hair and dander injection, solution

## Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-4855
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>EQUUS CABALLUS HAIR</b> (UNII: 4F35XG0149) (EQUUS CABALLUS HAIR - UNII:4F35XG0149)	EQUUS CABALLUS HAIR	0.01 g in 1 mL
<b>EQUUS CABALLUS DANDER</b> (UNII: J81SZ18495) (EQUUS CABALLUS DANDER - UNII:J81SZ18495)	EQUUS CABALLUS DANDER	0.01 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4855-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4855-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-4855-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	01/30/1978	

## ANIMAL ALLERGENS, FEATHER MIX

feather mix injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-4349
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>GALLUS GALLUS FEATHER</b> (UNII: 1FCM16V0FV) (GALLUS GALLUS FEATHER - UNII:1FCM16V0FV)	GALLUS GALLUS FEATHER	0.1 g in 1 mL
<b>ANAS PLATYRHYNCHOS FEATHER</b> (UNII: 83B65P4796) (ANAS PLATYRHYNCHOS FEATHER - UNII:83B65P4796)	ANAS PLATYRHYNCHOS FEATHER	0.1 g in 1 mL
<b>ANSER ANSER FEATHER</b> (UNII: 15XI414745) (ANSER ANSER FEATHER - UNII:15XI414745)	ANSER ANSER FEATHER	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4349-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4349-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-4349-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## ANIMAL ALLERGENS, FEATHER MIX

feather mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4352
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
GALLUS GALLUS FEATHER (UNII: 1FCM16V0FV) (GALLUS GALLUS FEATHER - UNII:1FCM16V0FV)	GALLUS GALLUS FEATHER	0.1 g in 1 mL
ANAS PLATYRHYNCHOS FEATHER (UNII: 83B65P4796) (ANAS PLATYRHYNCHOS FEATHER - UNII:83B65P4796)	ANAS PLATYRHYNCHOS FEATHER	0.1 g in 1 mL
ANSER ANSER FEATHER (UNII: 15XI414745) (ANSER ANSER FEATHER - UNII:15XI414745)	ANSER ANSER FEATHER	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
PHENOL (UNII: 339NCG44TV)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4352-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>2</b>	NDC:65044-4352-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-4352-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## ANIMAL ALLERGENS, GUINEA PIG HAIR AND DANDER

guinea pig hair and dander injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4401
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>CAVIA PORCELLUS HAIR</b> (UNII: KBA5Y6X57N) (CAVIA PORCELLUS HAIR - UNII:KBA5Y6X57N)	CAVIA PORCELLUS HAIR	0.05 g in 1 mL
<b>CAVIA PORCELLUS DANDER</b> (UNII: 84Q71TU5SU) (CAVIA PORCELLUS DANDER - UNII:84Q71TU5SU)	CAVIA PORCELLUS DANDER	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-4401-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, BEEF BOVINE SPP.

beef bovine spp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3077
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BEEF (UNII: 4PIB2155QP) (BEEF - UNII:4PIB2155QP)	BEEF	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3077-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, CHICKEN MEAT GALLUS SP.

chicken meat gallus sp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3173
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
POULTRY, UNSPECIFIED (UNII: L7WXO2P5HM) (POULTRY, UNSPECIFIED - UNII:L7WXO2P5HM)	POULTRY, UNSPECIFIED	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3173-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OLIVE OLEA EUROPaea

olive olea europaea injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2053
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
OLEA EUROPaea POLLEN (UNII: 43R41XZ627) (OLEA EUROPaea POLLEN - UNII:43R41XZ627)	OLEA EUROPaea POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2053-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2053-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2053-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PALM, QUEEN COCOS PLUMOSA

palm, queen cocos plumosa injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2074
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SYAGRUS ROMANZOFFIANA POLLEN (UNII: 84ZOM591BB) (SYAGRUS ROMANZOFFIANA POLLEN - UNII:84ZOM591BB)	SYAGRUS ROMANZOFFIANA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2074-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2074-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2074-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PALO VERDE CERCIDIUM FLORIDUM

palo verde cercidium floridum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2018
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PARKINSONIA FLORIDA POLLEN (UNII: 57586C96ZL) (PARKINSONIA FLORIDA POLLEN - UNII:57586C96ZL)	PARKINSONIA FLORIDA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C00X)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2098-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2098-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2098-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>POLLENS - TREES, PECAN CARYA CARYA ILLINOENSIS</b>				
pecan carya carya illinoensis injection, solution				

Product Information				
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2098	
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
CARYA ILLINOINENSIS POLLEN (UNII: PYO4JR720Y) (CARYA ILLINOINENSIS POLLEN - UNII:PYO4JR720Y)	CARYA ILLINOINENSIS POLLEN	0.05 g in 1 mL		

Inactive Ingredients				
Ingredient Name	Strength			
GLYCERIN (UNII: PDC6A3C00X)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
SODIUM BICARBONATE (UNII: 8MDF5V39QO)				

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2098-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2098-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2098-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PECAN CARYA CARYA ILLINOENSIS

pecan carya carya illinoensis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2101
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CARYA ILLINOINENSIS POLLEN (UNII: PYO4JR720Y) (CARYA ILLINOINENSIS POLLEN - UNII:PYO4JR720Y)	CARYA ILLINOINENSIS POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2101-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2101-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2101-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PEPPER TREE, CALIFORNIA SCHINUS MOLLE

pepper tree, california schinus molle injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2107
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>SCHINUS MOLLE POLLEN</b> (UNII: M0G28FH9K1) (SCHINUS MOLLE POLLEN - UNII:M0G28FH9K1)	SCHINUS MOLLE POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2107-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2107-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2107-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PINE MIX

pine mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2203
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>PINUS CONTORTA POLLEN</b> (UNII: FB7IP650ET) (PINUS CONTORTA POLLEN - UNII:FB7IP650ET)	PINUS CONTORTA POLLEN	0.05 g in 1 mL
<b>PINUS PONDEROSA POLLEN</b> (UNII: 042SUA2DS9) (PINUS PONDEROSA POLLEN - UNII:042SUA2DS9)	PINUS PONDEROSA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2203-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2203-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2203-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, PRIVET LIGUSTRUM VULGARE

privet ligustrum vulgare injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2251
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
LIGUSTRUM VULGARE POLLEN (UNII: Y3FRX92Z0E) (LIGUSTRUM VULGARE POLLEN - UNII:Y3FRX92Z0E)	LIGUSTRUM VULGARE POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2251-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2251-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2251-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, RUSSIAN OLIVE ELAEAGNUS ANGUSTIFOLIA

russian olive elaeagnus angustifolia injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2359
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ELAEAGNUS ANGUSTIFOLIA POLLEN (UNII: 68P4F4M6VD) (ELAEAGNUS ANGUSTIFOLIA POLLEN - UNII:68P4F4M6VD)	ELAEAGNUS ANGUSTIFOLIA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2359-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2359-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2359-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, SYCAMORE, AMERICAN EASTERN PLATANUS OCCIDENTALIS

sycamore, american eastern platanus occidentalis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2563
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PLATANUS OCCIDENTALIS POLLEN (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2563-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2563-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2563-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - TREES, TREE MIX 11**

tree mix 11 injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2619
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	0.05 g in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.05 g in 1 mL
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	0.05 g in 1 mL
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	0.05 g in 1 mL
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.05 g in 1 mL
<b>CARYA OVATA POLLEN</b> (UNII: 54UN9R2798) (CARYA OVATA POLLEN - UNII:54UN9R2798)	CARYA OVATA POLLEN	0.05 g in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.05 g in 1 mL
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.05 g in 1 mL
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	0.05 g in 1 mL
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C00X)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2619-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2619-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2619-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 11

tree mix 11 injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2622
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
FRAXINUS AMERICANA POLLEN (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	0.1 g in 1 mL
BETULA NIGRA POLLEN (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.1 g in 1 mL
JUGLANS NIGRA POLLEN (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	0.1 g in 1 mL
POPULUS DELTOIDES POLLEN (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	0.1 g in 1 mL
ULMUS AMERICANA POLLEN (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.1 g in 1 mL
CARYA OVATA POLLEN (UNII: 54UN9R2798) (CARYA OVATA POLLEN - UNII:54UN9R2798)	CARYA OVATA POLLEN	0.1 g in 1 mL
ACER SACCHARUM POLLEN (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.1 g in 1 mL
QUERCUS RUBRA POLLEN (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.1 g in 1 mL
PLATANUS OCCIDENTALIS POLLEN (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	0.1 g in 1 mL
SALIX NIGRA POLLEN (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2622-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2622-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2622-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>POLLENS - TREES, TREE MIX 11</b>	
tree mix 11 injection, solution	

Product Information			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2624
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	20000 [PNU] in 1 mL	
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	20000 [PNU] in 1 mL	
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	20000 [PNU] in 1 mL	
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DV63WP)	POPULUS DELTOIDES POLLEN	20000 [PNU] in 1 mL	
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	20000 [PNU] in 1 mL	
<b>CARYA OVATA POLLEN</b> (UNII: 54UN9R2798) (CARYA OVATA POLLEN - UNII:54UN9R2798)	CARYA OVATA POLLEN	20000 [PNU] in 1 mL	
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	20000 [PNU] in 1 mL	
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	20000 [PNU] in 1 mL	
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	20000 [PNU] in 1 mL	
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	20000 [PNU] in 1 mL	

Inactive Ingredients	
Ingredient Name	Strength

<b>PHENOL</b> (UNII: 339NCG44TV)
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2624-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2624-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2624-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 11

tree mix 11 injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2623
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	40000 [PNU] in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	40000 [PNU] in 1 mL
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	40000 [PNU] in 1 mL
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	40000 [PNU] in 1 mL
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	40000 [PNU] in 1 mL
<b>CARYA OVATA POLLEN</b> (UNII: 54UN9R2798) (CARYA OVATA POLLEN - UNII:54UN9R2798)	CARYA OVATA POLLEN	40000 [PNU] in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	40000 [PNU] in 1 mL
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	40000 [PNU] in 1 mL
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	40000 [PNU] in 1 mL
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	40000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	

**SODIUM CHLORIDE** (UNII: 451W47IQ8X)

**SODIUM BICARBONATE** (UNII: 8MDF5V39QO)

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2623-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2623-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2623-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 5

tree mix 5 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2854
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>CARYA ILLINOINENSIS POLLEN</b> (UNII: PYO4JR720Y) (CARYA ILLINOINENSIS POLLEN - UNII:PYO4JR720Y)	CARYA ILLINOINENSIS POLLEN	0.05 g in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.017 g in 1 mL
<b>ACER NEGUNDO POLLEN</b> (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII:P6K070AR8V)	ACER NEGUNDO POLLEN	0.017 g in 1 mL
<b>ACER RUBRUM POLLEN</b> (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII:700NK45C76)	ACER RUBRUM POLLEN	0.017 g in 1 mL
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.017 g in 1 mL
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	0.017 g in 1 mL
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	0.017 g in 1 mL
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	0.05 g in 1 mL
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2854-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2854-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2854-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## **POLLENS - TREES, TREE MIX 5**

tree mix 5 injection, solution

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2856
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
Ingredient Name	Basis of Strength	Strength	
<b>CARYA ILLINOINENSIS POLLEN</b> (UNII: PYO4JR720Y) (CARYA ILLINOINENSIS POLLEN - UNII:PYO4JR720Y)	CARYA ILLINOINENSIS POLLEN	0.1 g in 1 mL	
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.033 g in 1 mL	
<b>ACER NEGUNDO POLLEN</b> (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII:P6K070AR8V)	ACER NEGUNDO POLLEN	0.033 g in 1 mL	
<b>ACER RUBRUM POLLEN</b> (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII:700NK45C76)	ACER RUBRUM POLLEN	0.033 g in 1 mL	
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.033 g in 1 mL	
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	0.033 g in 1 mL	
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	0.033 g in 1 mL	
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	0.1 g in 1 mL	
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	0.1 g in 1 mL	

<b>Inactive Ingredients</b>		
Ingredient Name	Strength	
<b>PHENOL</b> (UNII: 339NCG44TV)		
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)		
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)		

<b>Packaging</b>		
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#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2856-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2856-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2856-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 5

tree mix 5 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2855
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>CARYA ILLINOINENSIS POLLEN</b> (UNII: PYO4JR720Y) (CARYA ILLINOINENSIS POLLEN - UNII:PYO4JR720Y)	CARYA ILLINOINENSIS POLLEN	20000 [PNU] in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	20000 [PNU] in 1 mL
<b>ACER NEGUNDO POLLEN</b> (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII:P6K070AR8V)	ACER NEGUNDO POLLEN	20000 [PNU] in 1 mL
<b>ACER RUBRUM POLLEN</b> (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII:700NK45C76)	ACER RUBRUM POLLEN	20000 [PNU] in 1 mL
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	20000 [PNU] in 1 mL
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	20000 [PNU] in 1 mL
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	20000 [PNU] in 1 mL
<b>PLATANUS OCCIDENTALIS POLLEN</b> (UNII: E03U1K03LK) (PLATANUS OCCIDENTALIS POLLEN - UNII:E03U1K03LK)	PLATANUS OCCIDENTALIS POLLEN	20000 [PNU] in 1 mL
<b>SALIX NIGRA POLLEN</b> (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	20000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-	10 mL in 1 VIAL; Type 0: Not a Combination		

1	2855-2	Product		
2	NDC:65044-2855-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2855-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 6

tree mix 6 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2863
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	0.05 g in 1 mL
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	0.017 g in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.017 g in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	0.017 g in 1 mL
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	0.05 g in 1 mL
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	0.05 g in 1 mL
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2863-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2863-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2863-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 6

tree mix 6 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2861
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	0.1 g in 1 mL
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	0.033 g in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.033 g in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	0.033 g in 1 mL
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	0.1 g in 1 mL
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	0.1 g in 1 mL
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2861-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2861-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2861-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE MIX 6

tree mix 6 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2862
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	20000 [PNU] in 1 mL
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	20000 [PNU] in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	20000 [PNU] in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	20000 [PNU] in 1 mL
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	20000 [PNU] in 1 mL
<b>POPULUS DELTOIDES POLLEN</b> (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	20000 [PNU] in 1 mL
<b>ULMUS AMERICANA POLLEN</b> (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	20000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2862-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2862-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2862-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, TREE OF HEAVEN AILANTHUS ALTISSIMA

tree of heaven ailanthus altissima injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2599
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AILANTHUS ALTISSIMA POLLEN</b> (UNII: 2A64U81OQ3) (AILANTHUS ALTISSIMA POLLEN - UNII:2A64U81OQ3)	AILANTHUS ALTISSIMA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2599-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2599-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2599-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, WALNUT, BLACK JUGLANS NIGRA

walnut, black juglans nigra injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2626
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>JUGLANS NIGRA POLLEN</b> (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)	JUGLANS NIGRA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2626-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2626-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2626-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## **POLLENS - TREES, WALNUT, BLACK JUGLANS NIGRA**

walnut, black juglans nigra injection, solution

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2629
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
Ingredient Name		Basis of Strength	Strength
JUGLANS NIGRA POLLEN (UNII: 1BV28146ZR) (JUGLANS NIGRA POLLEN - UNII:1BV28146ZR)		JUGLANS NIGRA POLLEN	0.1 g in 1 mL

<b>Inactive Ingredients</b>		
Ingredient Name		Strength
PHENOL (UNII: 339NCG44TV)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2629-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2629-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2629-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>			
Marketing	Application Number or Monograph	Marketing Start	Marketing End

Category	Citation	Date	Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, WILLOW, BLACK SALIX NIGRA

willow, black salix nigra injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2677
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALIX NIGRA POLLEN (UNII: 6M2JIH93ZN) (SALIX NIGRA POLLEN - UNII:6M2JIH93ZN)	SALIX NIGRA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2677-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2677-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2677-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, COCKLEBUR XANTHIUM STRUMARIUM

cocklebur xanthium strumarium injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1405
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
XANTHIUM STRUMARIUM POLLEN (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1405-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1405-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1405-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, COCKLEBUR XANTHIUM STRUMARIUM

cocklebur xanthium strumarium injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1408
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
XANTHIUM STRUMARIUM POLLEN (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

Marketing Start Date	Marketing End Date

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1408-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1408-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1408-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, COCKLEBUR XANTHIUM STRUMARIUM

cocklebur xanthium strumarium injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1409
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
XANTHIUM STRUMARIUM POLLEN (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	40000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1409-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1409-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1409-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - WEEDS AND GARDEN PLANTS, DOG FENNEL, EASTERN EUPATORIUM CAPILLIFOLIUM

dog fennel, eastern eupatorium capillifolium injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2057
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
EUPATORIUM CAPILLIFOLIUM POLLEN (UNII: B67NF86HF0) (EUPATORIUM CAPILLIFOLIUM POLLEN - UNII:B67NF86HF0)	EUPATORIUM CAPILLIFOLIUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2057-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2057-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2057-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - WEEDS AND GARDEN PLANTS, GOLDENROD SOLIDAGO CANADENSIS

goldenrod solidago canadensis injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1630
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SOLIDAGO CANADENSIS POLLEN (UNII: 644CZ16IR5) (SOLIDAGO CANADENSIS POLLEN - UNII:644CZ16IR5)	SOLIDAGO CANADENSIS POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1630-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1630-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1630-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - WEEDS AND GARDEN PLANTS, LAMBS QUARTERS CHENOPODIUM ALBUM

lambs quarters chenopodium album injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1786
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CHENOPODIUM ALBUM POLLEN (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1786-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1786-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		

3	NDC:65044-1786-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		
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## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, LAMBS QUARTERS CHENOPODIUM ALBUM

lambs quarters chenopodium album injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1789
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CHENOPODIUM ALBUM POLLEN (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1789-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1789-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1789-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, LAMBS QUARTERS CHENOPODIUM ALBUM

lambs quarters chenopodium album injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1790
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CHENOPODIUM ALBUM POLLEN (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	40000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1790-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1790-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1790-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, LAMBS QUARTERS CHENOPODIUM ALBUM

lambs quarters chenopodium album injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1791
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CHENOPODIUM ALBUM POLLEN (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	20000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
<b>PHENOL</b> (UNII: 339NCG44TV)		
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)		
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1791-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1791-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1791-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>POLLENS - WEEDS AND GARDEN PLANTS, NETTLE URTICA DIOICA</b>				
nettles urtica dioica injection, solution				

Product Information				
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1945	
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS			

Active Ingredient/Active Moiety				
Ingredient Name			Basis of Strength	Strength
<b>URTICA DIOICA POLLEN</b> (UNII: DNB59M1NVU) (URTICA DIOICA POLLEN - UNII:DNB59M1NVU)			URTICA DIOICA POLLEN	0.05 g in 1 mL

Inactive Ingredients				
Ingredient Name				Strength
GLYCERIN (UNII: PDC6A3C0OX)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
SODIUM BICARBONATE (UNII: 8MDF5V39QO)				

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1945-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1945-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1945-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, NETTLE URTICA DIOICA

nettles urtica dioica injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1947
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
URTICA DIOICA POLLEN (UNII: DNB59M1NVU) (URTICA DIOICA POLLEN - UNII:DNB59M1NVU)	URTICA DIOICA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1947-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1947-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1947-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, PIGWEED, ROUGH REDROOT AMARANTHUS RETROFLEXUS

pigweed, rough redroot amaranthus retroflexus injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2125
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2125-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2125-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2125-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - WEEDS AND GARDEN PLANTS, PLANTAIN, ENGLISH PLANTAGO LANCEOLATA**

plantain, english plantago lanceolata injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2212
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>PLANTAGO LANCEOLATA POLLEN</b> (UNII: D087T1U2CI) (PLANTAGO LANCEOLATA POLLEN - UNII:D087T1U2CI)	PLANTAGO LANCEOLATA POLLEN	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2212-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2212-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2212-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

  

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## **POLLENS - WEEDS AND GARDEN PLANTS, PLANTAIN, ENGLISH PLANTAGO LANCEOLATA**

plantain, english plantago lanceolata injection, solution

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2215
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
Ingredient Name	Basis of Strength	Strength	
PLANTAGO LANCEOLATA POLLEN (UNII: DO87T1U2CI) (PLANTAGO LANCEOLATA POLLEN - UNII:DO87T1U2CI)	PLANTAGO LANCEOLATA POLLEN	0.1 g in 1 mL	

<b>Inactive Ingredients</b>		
Ingredient Name	Strength	
PHENOL (UNII: 339NCG44TV)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2215-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2215-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2215-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	

BLA

BLA103888

04/19/1941

## POLLENS - WEEDS AND GARDEN PLANTS, PLANTAIN, ENGLISH PLANTAGO LANCEOLATA

plantain, english plantago lanceolata injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2217
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PLANTAGO LANCEOLATA POLLEN (UNII: DO87T1U2CI) (PLANTAGO LANCEOLATA POLLEN - UNII:DO87T1U2CI)	PLANTAGO LANCEOLATA POLLEN	20000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2217-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2217-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2217-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, PLANTAIN, ENGLISH PLANTAGO LANCEOLATA

plantain, english plantago lanceolata injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2216
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>PLANTAGO LANCEOLATA POLLEN</b> (UNII: DO87T1U2CI) (PLANTAGO LANCEOLATA POLLEN - UNII:DO87T1U2CI)	PLANTAGO LANCEOLATA POLLEN	40000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2216-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2216-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2216-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED, GIANT AMBROSIA TRIFIDA

ragweed, giant ambrosia trifida injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2293
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AMBROSIA TRIFIDA POLLEN</b> (UNII: KU1V1898XX) (AMBROSIA TRIFIDA POLLEN - UNII:KU1V1898XX)	AMBROSIA TRIFIDA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2293-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2293-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2293-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED, GIANT AMBROSIA TRIFIDA

ragweed, giant ambrosia trifida injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2296
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AMBROSIA TRIFIDA POLLEN (UNII: KU1V1898XX) (AMBROSIA TRIFIDA POLLEN - UNII:KU1V1898XX)	AMBROSIA TRIFIDA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2296-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2296-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2296-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED. WESTERN AMBROSIA PSILOSTACHYA

ragweed. western ambrosia psilostachya injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2308
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AMBROSIA PSILOSTACHYA POLLEN (UNII: RX18M46K8L) (AMBROSIA PSILOSTACHYA POLLEN - UNII:RX18M46K8L)	AMBROSIA PSILOSTACHYA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2308-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2308-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2308-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - WEEDS AND GARDEN PLANTS, RAGWEED. WESTERN AMBROSIA PSILOSTACHYA

ragweed. western ambrosia psilostachya injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2311
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AMBROSIA PSILOSTACHYA POLLEN (UNII: RX18M46K8L) (AMBROSIA PSILOSTACHYA POLLEN - UNII:RX18M46K8L)	AMBROSIA PSILOSTACHYA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2311-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2311-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2311-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, RUSSIAN THISTLE SALSO LA KALI

russian thistle salsola kali injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2362
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALSOLA KALI POLLEN (UNII: 2MH135KC6G) (SALSOLA KALI POLLEN - UNII:2MH135KC6G)	SALSOLA KALI POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2362-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>2</b>	NDC:65044-2362-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2362-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SAGEBRUSH, MUGWORT ARTEMISIA VULGARIS

sagebrush, mugwort artemisia vulgaris injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2413
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ARTEMISIA VULGARIS POLLEN (UNII: ANT994T71D) (ARTEMISIA VULGARIS POLLEN - UNII:ANT994T71D)	ARTEMISIA VULGARIS POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2413-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2413-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2413-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SAGEBRUSH, MUGWORT ARTEMISIA VULGARIS

sagebrush, mugwort artemisia vulgaris injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2416
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ARTEMISIA VULGARIS POLLEN (UNII: ANT994T71D) (ARTEMISIA VULGARIS POLLEN - UNII:ANT994T71D)	ARTEMISIA VULGARIS POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2416-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2416-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2416-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SAGEBRUSH, MUGWORT ARTEMISIA VULGARIS

sagebrush, mugwort artemisia vulgaris injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2417
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ARTEMISIA VULGARIS POLLEN (UNII: ANT994T71D) (ARTEMISIA VULGARIS POLLEN - UNII:ANT994T71D)	ARTEMISIA VULGARIS POLLEN	40000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
PHENOL (UNII: 339NCG44TV)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2417-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2417-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2417-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SCALE, WING SHAD ATRIPLEX CANESCENS

scale, wing shad atriplex canescens injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2482
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ATRIPLEX CANESCENS POLLEN (UNII: 26U0BU8G83) (ATRIPLEX CANESCENS POLLEN - UNII:26U0BU8G83)	ATRIPLEX CANESCENS POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2482-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2482-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-	50 mL in 1 VIAL; Type 0: Not a Combination		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SCALE, WING SHAD ATRIPLEX CANESCENS

scale, wing shad atriplex canescens injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2485
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ATRIPLEX CANESCENS POLLEN (UNII: 26U0BU8G83) (ATRIPLEX CANESCENS POLLEN - UNII:26U0BU8G83)	ATRIPLEX CANESCENS POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2485-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2485-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2485-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SCOTCH BROOM CYTISUS SCOPARIUS

scotch broom cytisus scoparius injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2487
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

#### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>CYTISUS SCOPARIUS FLOWERING TOP (UNII: XZC6H8R666) (CYTISUS SCOPARIUS FLOWERING TOP - UNII:XZC6H8R666)</b>	CYTISUS SCOPARIUS FLOWERING TOP	0.05 g in 1 mL

#### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN (UNII: PDC6A3C0OX)</b>	
<b>SODIUM CHLORIDE (UNII: 451W47IQ8X)</b>	
<b>SODIUM BICARBONATE (UNII: 8MDF5V39QO)</b>	

#### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2487-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2487-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2487-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

#### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS AND GARDEN PLANTS, SORREL, SHEEP RUMEX ACETOSELLA

sorrel, sheep rumex acetosella injection, solution

#### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2506
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

#### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>RUMEX ACETOSELLA POLLEN (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)</b>	RUMEX ACETOSELLA POLLEN	0.05 g in 1 mL

#### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN (UNII: PDC6A3C0OX)</b>	
<b>SODIUM CHLORIDE (UNII: 451W47IQ8X)</b>	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2506-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2506-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2506-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - WEEDS AND GARDEN PLANTS, SORREL, SHEEP RUMEX ACETOSELLA**

sorrel, sheep rumex acetosella injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2508
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
RUMEX ACETOSELLA POLLEN (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2508-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2508-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2508-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, CARELESS WEED AMARANTHUS PALMERI

careless weed amaranthus palmeri injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1297
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AMARANTHUS PALMERI POLLEN (UNII: 1GH3W23KH) (AMARANTHUS PALMERI POLLEN - UNII:1GH3W23KH)	AMARANTHUS PALMERI POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1297-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1297-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1297-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, CARELESS/PIGWEED MIX

careless/pigweed mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1300
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AMARANTHUS PALMERI POLLEN</b> (UNII: 1GH3W23KH) (AMARANTHUS PALMERI POLLEN - UNII:1GH3W23KH)	AMARANTHUS PALMERI POLLEN	0.05 g in 1 mL
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1300-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1300-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1300-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, CARELESS/PIGWEED MIX

careless/pigweed mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1303
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AMARANTHUS PALMERI POLLEN</b> (UNII: 1GH3W23KH) (AMARANTHUS PALMERI POLLEN - UNII:1GH3W23KH)	AMARANTHUS PALMERI POLLEN	0.1 g in 1 mL
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1303-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1303-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1303-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## **POLLENS - WEEDS, DOCK/SORREL MIX**

pollens - weeds, dock/sorrel mix injection, solution

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1516
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>		
Ingredient Name	Basis of Strength	Strength
<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	0.05 g in 1 mL
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	0.05 g in 1 mL

<b>Inactive Ingredients</b>		
Ingredient Name	Strength	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)		
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)		
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)		

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1516-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1516-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1516-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## POLLENS - WEEDS, DOCK/SORREL MIX

pollens - weeds, dock/sorrel mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1519
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
RUMEX CRISPUS POLLEN (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	0.1 g in 1 mL
RUMEX ACETOSELLA POLLEN (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1519-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1519-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1519-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, DOCK/SORREL MIX

pollens - weeds, dock/sorrel mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1520
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength

<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	20000 [PNU] in 1 mL
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	20000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1520-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1520-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1520-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, GIANT, SHORT, WESTERN RAGWEED MIX

giant, short, western ragweed mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2320
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AMBROSIA TRIFIDA POLLEN</b> (UNII: KU1V1898XX) (AMBROSIA TRIFIDA POLLEN - UNII:KU1V1898XX)	AMBROSIA TRIFIDA POLLEN	0.05 g in 1 mL
<b>AMBROSIA ARTEMISIIFOLIA POLLEN</b> (UNII: K20Y81AC03) (AMBROSIA ARTEMISIIFOLIA POLLEN - UNII:K20Y81AC03)	AMBROSIA ARTEMISIIFOLIA POLLEN	0.05 g in 1 mL
<b>AMBROSIA PSILOSTACHYA POLLEN</b> (UNII: RX18M46K8L) (AMBROSIA PSILOSTACHYA POLLEN - UNII:RX18M46K8L)	AMBROSIA PSILOSTACHYA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2320-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2320-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2320-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>POLLENS - WEEDS, KOCHIA SCOPARIA</b>				
kochia scoparia injection, solution				

<b>Product Information</b>				
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1780	
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS			

<b>Active Ingredient/Active Moiety</b>				
Ingredient Name			Basis of Strength	Strength
<b>BASSIA SCOPARIA POLLEN</b> (UNII: 07A108ZKW5) (BASSIA SCOPARIA POLLEN - UNII:07A108ZKW5)			BASSIA SCOPARIA POLLEN	0.05 g in 1 mL

<b>Inactive Ingredients</b>				
Ingredient Name			Strength	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)				
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)				
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)				

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1780-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1780-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1780-5	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## POLLENS - WEEDS, KOCHIA SCOPARIA

kochia scoparia injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1783
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BASSIA SCOPARIA POLLEN (UNII: 07A108ZKW5) (BASSIA SCOPARIA POLLEN - UNII:07A108ZKW5)	BASSIA SCOPARIA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1783-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1783-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1783-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, MARSHELDER/POVERTY MIX

pollens - weeds, marshelder/poverty mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1858
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IVA AXILLARIS POLLEN (UNII: 13KFG30UBR) (IVA AXILLARIS POLLEN - UNII:13KFG30UBR)	IVA AXILLARIS POLLEN	0.05 g in 1 mL
IVA ANNUA POLLEN (UNII: Y2U5S5PF22) (IVA ANNUA POLLEN -	IVA ANNUA POLLEN	0.05 g

UNII:Y2U5S5PF22)	IVANNUA POLLEN	in 1 mL
<b>CYCLACHAENA XANTHIFOLIA POLLEN</b> (UNII: V80TPZ0T6J) (CYCLACHAENA XANTHIFOLIA POLLEN - UNII:V80TPZ0T6J)	CYCLACHAENA XANTHIFOLIA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1858-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1858-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1858-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, MARSHELDER/POVERTY MIX

pollens - weeds, marshelder/poverty mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1861
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>IVA AXILLARIS POLLEN</b> (UNII: 13KFG30UBR) (IVA AXILLARIS POLLEN - UNII:13KFG30UBR)	IVA AXILLARIS POLLEN	0.1 g in 1 mL
<b>IVA ANNUA POLLEN</b> (UNII: Y2U5S5PF22) (IVA ANNUA POLLEN - UNII:Y2U5S5PF22)	IVA ANNUA POLLEN	0.1 g in 1 mL
<b>CYCLACHAENA XANTHIFOLIA POLLEN</b> (UNII: V80TPZ0T6J) (CYCLACHAENA XANTHIFOLIA POLLEN - UNII:V80TPZ0T6J)	CYCLACHAENA XANTHIFOLIA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1861-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1861-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1861-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, WEED MIX 2630

weed mix 2630 injection, solution

Product Information			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2634
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>XANTHIUM STRUMARIUM POLLEN</b> (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	0.05 g in 1 mL	
<b>CHENOPODIUM ALBUM POLLEN</b> (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	0.05 g in 1 mL	
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	0.05 g in 1 mL	
<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	0.025 g in 1 mL	
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	0.025 g in 1 mL	

Inactive Ingredients		
Ingredient Name	Strength	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)		
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)		
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2634-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2634-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2634-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, WEED MIX 2630

weed mix 2630 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2632
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>XANTHIUM STRUMARIUM POLLEN</b> (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	0.1 g in 1 mL
<b>CHENOPODIUM ALBUM POLLEN</b> (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	0.1 g in 1 mL
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	0.1 g in 1 mL
<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	0.05 g in 1 mL
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2632-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2632-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2632-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, WEED MIX 2630

weed mix 2630 injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2635
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>XANTHIUM STRUMARIUM POLLEN</b> (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	20000 [PNU] in 1 mL
<b>CHENOPODIUM ALBUM POLLEN</b> (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	20000 [PNU] in 1 mL
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	20000 [PNU] in 1 mL
<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	20000 [PNU] in 1 mL
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	20000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2635-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2635-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2635-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - WEEDS, WEED MIX 2630

weed mix 2630 injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2633
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength

<b>XANTHIUM STRUMARIUM POLLEN</b> (UNII: 2QOF601J1M) (XANTHIUM STRUMARIUM POLLEN - UNII:2QOF601J1M)	XANTHIUM STRUMARIUM POLLEN	40000 [PNU] in 1 mL
<b>CHENOPODIUM ALBUM POLLEN</b> (UNII: 098LKX5NCN) (CHENOPODIUM ALBUM POLLEN - UNII:098LKX5NCN)	CHENOPODIUM ALBUM POLLEN	40000 [PNU] in 1 mL
<b>AMARANTHUS RETROFLEXUS POLLEN</b> (UNII: 73B14PX5FW) (AMARANTHUS RETROFLEXUS POLLEN - UNII:73B14PX5FW)	AMARANTHUS RETROFLEXUS POLLEN	40000 [PNU] in 1 mL
<b>RUMEX CRISPUS POLLEN</b> (UNII: V825XJG64G) (RUMEX CRISPUS POLLEN - UNII:V825XJG64G)	RUMEX CRISPUS POLLEN	40000 [PNU] in 1 mL
<b>RUMEX ACETOSELLA POLLEN</b> (UNII: N52MIQ81ZW) (RUMEX ACETOSELLA POLLEN - UNII:N52MIQ81ZW)	RUMEX ACETOSELLA POLLEN	40000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-2633-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2633-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2633-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, EGG, WHITE GALLUS SP.

egg, white gallus sp. injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3248
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>EGG WHITE</b> (UNII: 3E0I92Z2GR) (EGG WHITE - UNII:3E0I92Z2GR)	EGG WHITE	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3248-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, EGG, YOLK GALLUS SP.**

egg, yolk gallus sp. injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3254
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
EGG YOLK (UNII: 4IPS17B70T) (EGG YOLK - UNII:4IPS17B70T)	EGG YOLK	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3254-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - ANIMAL PRODUCTS AND POULTRY PRODUCTS, PORK SUS SP.**

pork sus sp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3509
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PORK (UNII: O138UB266J) (PORK - UNII:O138UB266J)	PORK	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3509-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - DAIRY PRODUCTS, CASEIN, COW MILK

casein, cow milk injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3380
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CASEIN (UNII: 48268V50D5) (CASEIN - UNII:48268V50D5)	CASEIN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3380-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

FOOD - DAIRY PRODUCTS, MILK, WHOLE COW				
milk, whole cow injection, solution				

Product Information				
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3389	
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
COW MILK (UNII: 917J3173FT) (COW MILK - UNII:917J3173FT)	COW MILK	0.05 g in 1 mL		

Inactive Ingredients				
Ingredient Name	Strength			
GLYCERIN (UNII: PDC6A3C0OX)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
SODIUM BICARBONATE (UNII: 8MDF5V39QO)				

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3389-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

FOOD - FISH AND SHELLFISH, CLAM				
clam injection, solution				
Product Information				
Product Type				
NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3191		

**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>QUAHOG, UNSPECIFIED</b> (UNII: 226LY0AFR9) (QUAHOG, UNSPECIFIED - UNII:226LY0AFR9)	QUAHOG, UNSPECIFIED	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-3191-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - FISH AND SHELLFISH, CODFISH GADUS CALLARIAS**

codfish gadus callarias injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3203
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>COD, UNSPECIFIED</b> (UNII: 8D6Q5LNG3D) (COD, UNSPECIFIED - UNII:8D6Q5LNG3D)	COD, UNSPECIFIED	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3203-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - FISH AND SHELLFISH, CRAB XIPHOSURUS SOWERBYI

crab xiphosurus sowerbyi injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3215
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CRAB LEG, UNSPECIFIED (UNII: S1VF61QL09) (CRAB LEG, UNSPECIFIED - UNII:S1VF61QL09)	CRAB LEG, UNSPECIFIED	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3215-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - FISH AND SHELLFISH, LOBSTER HOMARUS AMERICANUS

lobster homarus americanus injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3362
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>LOBSTER, UNSPECIFIED</b> (UNII: ZQ6LG2C39M) (LOBSTER, UNSPECIFIED - UNII:ZQ6LG2C39M)	LOBSTER, UNSPECIFIED	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3362-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - FISH AND SHELLFISH, SALMON SALMO SALAR**

salmon salmo salar injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3565
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>SALMON, UNSPECIFIED</b> (UNII: 6122W2M0GB) (SALMON, UNSPECIFIED - UNII:6122W2M0GB)	SALMON, UNSPECIFIED	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3565-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - FISH AND SHELLFISH, SHRIMP CRAGO SP.

shrimp crago sp. injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3584
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SHRIMP, UNSPECIFIED (UNII: 1891LE191T) (SHRIMP, UNSPECIFIED - UNII:1891LE191T)	SHRIMP, UNSPECIFIED	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3584-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - FISH AND SHELLFISH, TUNA THUNNUS SP.

tuna thunnus sp. injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3674
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>TUNA, UNSPECIFIED</b> (UNII: V2T3IHT3E2) (TUNA, UNSPECIFIED - UNII:V2T3IHT3E2)	TUNA, UNSPECIFIED	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3674-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, ALMOND PRUNUS AMYGDALUS**

almond prunus amygdalus injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3014
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>ALMOND</b> (UNII: 3Z252A2K9G) (ALMOND - UNII:3Z252A2K9G)	ALMOND	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
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<b>1</b>	NDC:65044-3014-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
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## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, APPLE MALUS SP.

apple malus sp. injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3020
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
APPLE (UNII: B423VGH5S9) (APPLE - UNII:B423VGH5S9)	APPLE	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-3020-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, BANANA MUSA SAPIENTUM

banana musa sapientum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3041
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>BANANA</b> (UNII: 4AJZ4765R9) (BANANA - UNII:4AJZ4765R9)	BANANA	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3041-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, BRAZIL NUT BERTHOLLETIA EXCELSA**

brazil nut bertholletia excelsa injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3107
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>BRAZIL NUT</b> (UNII: XKR79OET1K) (BRAZIL NUT - UNII:XKR79OET1K)	BRAZIL NUT	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3107-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, CARROT DAUCUS CAROTA

carrot daucus carota injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3125
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CARROT (UNII: L56Z1JK48B) (CARROT - UNII:L56Z1JK48B)	CARROT	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3125-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, CASHEW NUT ANACARDIUM OCCIDENTALIE

cashew nut anacardium occidentalis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3134
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CASHEW (UNII: 3H5U5CX7KO) (CASHEW - UNII:3H5U5CX7KO)	CASHEW	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3134-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, CELERY APIUM GRAVEOLENS

celery apium graveolens injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3140
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CELERY (UNII: 44IDY6DTKX) (CELERY - UNII:44IDY6DTKX)	CELERY	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3140-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, CORN ZEA MAYS

corn zea mays injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3212
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CORN (UNII: 0N8672707O) (CORN - UNII:0N8672707O)	CORN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3212-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, HAZELNUT FILBERT CORYLUS SPP.

hazelnut filbert corylus spp. injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3305
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
HAZELNUT, UNSPECIFIED (UNII: IW0OM96F6O) (HAZELNUT, UNSPECIFIED - UNII:IW0OM96F6O)	HAZELNUT, UNSPECIFIED	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name		Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)		
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)		
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3305-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>FOOD - PLANT SOURCE, MELON, CANTALOUPE CUCUMIS MELO</b>			
cantaloupe cucumis melo injection, solution			

Product Information			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3116
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
CANTALOUPE (UNII: 8QF5D5H6UH) (CANTALOUPE - UNII:8QF5D5H6UH)	CANTALOUPE	0.1 g in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
GLYCERIN (UNII: PDC6A3C00X)			
SODIUM CHLORIDE (UNII: 451W47IQ8X)			
SODIUM BICARBONATE (UNII: 8MDF5V39QO)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3116-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## FOOD - PLANT SOURCE, ORANGE CITRUS SINENSIS

orange citrus sinensis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3428
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ORANGE (UNII: 5EVU04N5QU) (ORANGE - UNII:5EVU04N5QU)	ORANGE	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3428-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, PEA, GREEN OR ENGLISH PISUM SATIVUM

pea, green or english pisum sativum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3449
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PEA (UNII: W4X7H8GYFM) (PEA - UNII:W4X7H8GYFM)	PEA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3449-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, PEACH PRUNUS PERSICA**

peach prunus persica injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3452
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
PEACH (UNII: 30KE88I3QG) (PEACH - UNII:30KE88I3QG)	PEACH	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3452-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, PEANUT ARACHIS HYPOGAEA**

peanut arachis hypogaea injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-3455
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PEANUT (UNII: QE1QX6B99R) (PEANUT - UNII:QE1QX6B99R)	PEANUT	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3455-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, PECAN CARYA ILLINOENSIS

pecan carya illinoensis injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-3461
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PECAN (UNII: F14P91GB5F) (PECAN - UNII:F14P91GB5F)	PECAN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date

1	NDC:65044-3461-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	

<b>FOOD - PLANT SOURCE, POTATO, WHITE SOLANUM TUBEROSUM</b>	
potato, white solanum tuberosum injection, solution	

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3518
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>		
Ingredient Name	Basis of Strength	Strength
POTATO (UNII: CFE1S8DYWD) (POTATO - UNII:CFE1S8DYWD)	POTATO	0.1 g in 1 mL

<b>Inactive Ingredients</b>		
Ingredient Name	Strength	
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3518-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>FOOD - PLANT SOURCE, RICE, WHOLE GRAIN</b>				
rice, whole grain injection, solution				
<b>Product Information</b>				
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3548	
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS			

**Active Ingredient/Active Moiety**

Ingredient Name		Basis of Strength	Strength
<b>BROWN RICE</b> (UNII: 659G217HPG) (BROWN RICE - UNII:659G217HPG)		BROWN RICE	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3548-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, RYE GRAIN**

rye grain injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3554
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>RYE</b> (UNII: 0R4AQI398X) (RYE - UNII:0R4AQI398X)	RYE	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3554-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, SOYBEAN GLYCINE SOJA

soybean glycine soja injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3596
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SOYBEAN (UNII: L7HT8F1ZOD) (SOYBEAN - UNII:L7HT8F1ZOD)	SOYBEAN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3596-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## FOOD - PLANT SOURCE, STRAWBERRY FRAGARIA CHILOENSIS

strawberry fragaria chiloensis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3626
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
STRAWBERRY (UNII: 4J2TY8Y81V) (STRAWBERRY - UNII:4J2TY8Y81V)	STRAWBERRY	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3626-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# FOOD - PLANT SOURCE, STRING BEAN MIX

string bean mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3074
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
STRING BEAN (UNII: N9D69B2Q7Y) (STRING BEAN - UNII:N9D69B2Q7Y)	STRING BEAN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3074-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## **FOOD - PLANT SOURCE, TOMATO NICOTIANA SPP.**

tomato nicotiana spp. injection, solution

### **Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-3656
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### **Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>TOMATO</b> (UNII: Z4KHF2C175) (TOMATO - UNII:Z4KHF2C175)	TOMATO	0.1 g in 1 mL

### **Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### **Packaging**

#	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:65044-3656-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### **Marketing Information**

<b>Marketing Category</b>	<b>Application Number or Monograph Citation</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
BLA	BLA103888	04/19/1941	

## **FOOD - PLANT SOURCE, WALNUT, BLACK JUGLANS NIGRA**

walnut, black juglans nigra injection, solution

### **Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-3695
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### **Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>BLACK WALNUT</b> (UNII: 02WM57RXZJ) (BLACK WALNUT - UNII:02WM57RXZJ)	BLACK WALNUT	0.1 g in 1 mL

### **Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3695-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, YEAST, BAKER SACCHAROMYCES CEREVISIAE**

yeast, baker saccharomyces cerevisiae injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3713
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
YEAST (UNII: 3NY3SM6B8U) (YEAST - UNII:3NY3SM6B8U)	YEAST	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3713-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**FOOD - PLANT SOURCE, YEAST, BREWER SACCHAROMYCES CEREVISIAE**

yeast, brewer saccharomyces cerevisiae injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-3716
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
YEAST (UNII: 3NY3SM6B8U) (YEAST - UNII:3NY3SM6B8U)	YEAST	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-3716-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY COCKROACH, AMERICAN PERiplaneta americana

insects whole body cockroach, american periplaneta americana injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6580
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PERiplaneta americana (UNII: 2RQ1L9N089) (PERiplaneta americana - UNII:2RQ1L9N089)	PERiplaneta americana	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-6580-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-6580-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-6580-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**INSECTS WHOLE BODY COCKROACH, GERMAN BLATELLA GERMANICA**

insects whole body cockroach, german blatella germanica injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6581
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
BLATTELLA GERMANICA (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-6581-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-6581-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-6581-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY COCKROACH MIX

insects whole body cockroach mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6584
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PERIPLANETA AMERICANA (UNII: 2RQ1L9N089) (PERIPLANETA AMERICANA - UNII:2RQ1L9N089)	PERIPLANETA AMERICANA	0.1 g in 1 mL
BLATTELLA GERMANICA (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-6584-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-6584-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-6584-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY COCKROACH MIX

insects whole body cockroach mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6587
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>PERIPLANETA AMERICANA</b> (UNII: 2RQ1L9N089) (PERIPLANETA AMERICANA - UNII:2RQ1L9N089)	PERIPLANETA AMERICANA	0.1 g in 1 mL
<b>BLATTELLA GERMANICA</b> (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-6587-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-6587-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-6587-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**INSECTS WHOLE BODY COCKROACH MIX**

insects whole body cockroach mix injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6588
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>PERIPLANETA AMERICANA</b> (UNII: 2RQ1L9N089) (PERIPLANETA AMERICANA - UNII:2RQ1L9N089)	PERIPLANETA AMERICANA	20000 [PNU] in 1 mL
<b>BLATTELLA GERMANICA</b> (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	20000 [PNU] in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	

**SODIUM CHLORIDE** (UNII: 451W47IQ8X)

**SODIUM BICARBONATE** (UNII: 8MDF5V39QO)

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-6588-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-6588-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-6588-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY COCKROACH MIX

insects whole body cockroach mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6589
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>PERIPLANETA AMERICANA</b> (UNII: 2RQ1L9N089) (PERIPLANETA AMERICANA - UNII:2RQ1L9N089)	PERIPLANETA AMERICANA	20000 [PNU] in 1 mL
<b>BLATTELLA GERMANICA</b> (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	20000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-6589-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-6589-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-6589-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY COCKROACH MIX

insects whole body cockroach mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6590
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PERIPLANETA AMERICANA (UNII: 2RQ1L9N089) (PERIPLANETA AMERICANA - UNII:2RQ1L9N089)	PERIPLANETA AMERICANA	5000 [PNU] in 1 mL
BLATTELLA GERMANICA (UNII: G9O67I0A8Q) (BLATTELLA GERMANICA - UNII:G9O67I0A8Q)	BLATTELLA GERMANICA	5000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-6590-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-6590-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-6590-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY, FIRE ANT MIX

insects whole body, fire ant mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-6518
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>SOLENOPSIS RICHTERI</b> (UNII: 739684T11W) (SOLENOPSIS RICHTERI - UNII:739684T11W)	SOLENOPSIS RICHTERI	0.1 g in 1 mL
<b>SOLENOPSIS INVICTA</b> (UNII: 507CR4P444) (SOLENOPSIS INVICTA - UNII:507CR4P444)	SOLENOPSIS INVICTA	0.1 g in 1 mL

**Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:65044-6518-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-6518-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-6518-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

<b>Marketing Category</b>	<b>Application Number or Monograph Citation</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
BLA	BLA103888	04/19/1941	

**INSECTS WHOLE BODY, FIRE ANT MIX**

insects whole body, fire ant mix injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-6517
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>SOLENOPSIS RICHTERI</b> (UNII: 739684T11W) (SOLENOPSIS RICHTERI - UNII:739684T11W)	SOLENOPSIS RICHTERI	0.1 g in 1 mL
<b>SOLENOPSIS INVICTA</b> (UNII: 507CR4P444) (SOLENOPSIS INVICTA - UNII:507CR4P444)	SOLENOPSIS INVICTA	0.1 g in 1 mL

**Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>

<b>PHENOL</b> (UNII: 339NCG44TV)
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-6517-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-6517-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-6517-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS - ALTERNARIA/HORMODENDRUM MIX

molds - alternaria/hormodendrum mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5004
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ALTERNARIA ALTERNATA</b> (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H)	ALTERNARIA ALTERNATA	0.1 g in 1 mL
<b>CLADOSPORIUM CLADOSPORIOIDES</b> (UNII: 4ZWY20GTGO) (CLADOSPORIUM CLADOSPORIOIDES - UNII:4ZWY20GTGO)	CLADOSPORIUM CLADOSPORIOIDES	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5004-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5004-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5004-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS - MOLD MIX 10

molds - mold mix 10 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5136
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ALTERNARIA ALTERNATA</b> (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H)	ALTERNARIA ALTERNATA	0.1 g in 1 mL
<b>ASPERGILLUS FUMIGATUS</b> (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48)	ASPERGILLUS FUMIGATUS	0.025 g in 1 mL
<b>ASPERGILLUS NIDULANS</b> (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80)	ASPERGILLUS NIDULANS	0.025 g in 1 mL
<b>ASPERGILLUS NIGER VAR. NIGER</b> (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6)	ASPERGILLUS NIGER VAR. NIGER	0.025 g in 1 mL
<b>ASPERGILLUS TERREUS</b> (UNII: QBN8K7055X) (ASPERGILLUS TERREUS - UNII:QBN8K7055X)	ASPERGILLUS TERREUS	0.025 g in 1 mL
<b>FUSARIUM OXYSPORUM VASINFECTUM</b> (UNII: 6M98DC08TZ) (FUSARIUM OXYSPORUM VASINFECTUM - UNII:6M98DC08TZ)	FUSARIUM OXYSPORUM VASINFECTUM	0.1 g in 1 mL
<b>DENDRYPHIELLA VINOSA</b> (UNII: 7S6NW5FH8X) (DENDRYPHIELLA VINOSA - UNII:7S6NW5FH8X)	DENDRYPHIELLA VINOSA	0.1 g in 1 mL
<b>CLADOSPORIUM CLADOSPORIOIDES</b> (UNII: 4ZWY20GTGO) (CLADOSPORIUM CLADOSPORIOIDES - UNII:4ZWY20GTGO)	CLADOSPORIUM CLADOSPORIOIDES	0.1 g in 1 mL
<b>MUCOR RACEMOSUS</b> (UNII: 17RH99LQ7G) (MUCOR RACEMOSUS - UNII:17RH99LQ7G)	MUCOR RACEMOSUS	0.1 g in 1 mL
<b>PENICILLIUM DIGITATUM</b> (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239)	PENICILLIUM DIGITATUM	0.02 g in 1 mL
<b>PENICILLIUM EXPANSUM</b> (UNII: 1XSC3BB35Z) (PENICILLIUM EXPANSUM - UNII:1XSC3BB35Z)	PENICILLIUM EXPANSUM	0.02 g in 1 mL
<b>PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM</b> (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG)	PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM	0.02 g in 1 mL
<b>CLONOSTACHYS ROSEA F. ROSEA</b> (UNII: I5F729WZ2H) (CLONOSTACHYS ROSEA F. ROSEA - UNII:I5F729WZ2H)	CLONOSTACHYS ROSEA F. ROSEA	0.02 g in 1 mL
<b>PHOMA EXIGUA VAR. EXIGUA</b> (UNII: 8JAG41IE4M) (PHOMA EXIGUA VAR. EXIGUA - UNII:8JAG41IE4M)	PHOMA EXIGUA VAR. EXIGUA	0.1 g in 1 mL
<b>AUREOBASIDIUM PULLULANS VAR. PULLUTANS</b> (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK)	AUREOBASIDIUM PULLULANS VAR. PULLUTANS	0.1 g in 1 mL
<b>RHIZOPUS STOLONIFER</b> (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q)	RHIZOPUS STOLONIFER	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5136-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5136-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5136-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS - MOLD MIX 4

molds - mold mix 4 injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5000
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H)	ALTERNARIA ALTERNATA	0.1 g in 1 mL
ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48)	ASPERGILLUS FUMIGATUS	0.025 g in 1 mL
ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80)	ASPERGILLUS NIDULANS	0.025 g in 1 mL
ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6)	ASPERGILLUS NIGER VAR. NIGER	0.025 g in 1 mL
ASPERGILLUS TERREUS (UNII: QBN8K7055X) (ASPERGILLUS TERREUS - UNII:QBN8K7055X)	ASPERGILLUS TERREUS	0.025 g in 1 mL
CLADOSPORIUM CLADOSPORIOIDES (UNII: 4ZWY20GTGO) (CLADOSPORIUM CLADOSPORIOIDES - UNII:4ZWY20GTGO)	CLADOSPORIUM CLADOSPORIOIDES	0.1 g in 1 mL
PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239)	PENICILLIUM DIGITATUM	0.025 g in 1 mL
PENICILLIUM EXPANSUM (UNII: 1XSC3BB35Z) (PENICILLIUM EXPANSUM - UNII:1XSC3BB35Z)	PENICILLIUM EXPANSUM	0.025 g in 1 mL
PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG)	PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM	0.025 g in 1 mL
CLONOSTACHYS ROSEA F. ROSEA (UNII: I5F729WZ2H) (CLONOSTACHYS ROSEA F. ROSEA - UNII:I5F729WZ2H)	CLONOSTACHYS ROSEA F. ROSEA	0.025 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5000-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5000-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5000-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>MOLDS - TRICHOPHYTON MIX</b>			
molds - trichophyton mix injection, solution			
<b>Product Information</b>			
<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5284
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
Ingredient Name	Basis of Strength	Strength	
<b>TRICHOPHYTON TONSURANS</b> (UNII: JY1BE33I3Y) (TRICHOPHYTON TONSURANS - UNII:JY1BE33I3Y)	TRICHOPHYTON TONSURANS	0.1 g in 1 mL	
<b>TRICHOPHYTON RUBRUM</b> (UNII: 2ZAU32517N) (TRICHOPHYTON RUBRUM - UNII:2ZAU32517N)	TRICHOPHYTON RUBRUM	0.1 g in 1 mL	
<b>TRICHOPHYTON MENTAGROPHYTES</b> (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV)	TRICHOPHYTON MENTAGROPHYTES	0.1 g in 1 mL	

<b>Inactive Ingredients</b>			
Ingredient Name	Strength		
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)			
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)			
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)			

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5284-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5284-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5284-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>			
Marketing	Application Number or Monograph	Marketing Start	Marketing End

Category	Citation	Date	Date
BLA	BLA103888	04/19/1941	

## MOLDS, PENICILLIUM MIX

molds, penicillium mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5168
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239)	PENICILLIUM DIGITATUM	0.1 g in 1 mL
PENICILLIUM EXPANSUM (UNII: 1XSC3BB35Z) (PENICILLIUM EXPANSUM - UNII:1XSC3BB35Z)	PENICILLIUM EXPANSUM	0.1 g in 1 mL
PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG)	PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM	0.1 g in 1 mL
CLONOSTACHYS ROSEA F. ROSEA (UNII: I5F729WZ2H) (CLONOSTACHYS ROSEA F. ROSEA - UNII:I5F729WZ2H)	CLONOSTACHYS ROSEA F. ROSEA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5168-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5168-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5168-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, ALTERNARIA TENUIS

alternaria tenuis injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5008
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>ALTERNARIA ALTERNATA</b> (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H)	ALTERNARIA ALTERNATA	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5008-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5008-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5008-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**MOLDS, RUSTS AND SMUTS, ASPERGILLUS FUMIGATUS**

aspergillus fumigatus injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5020
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>ASPERGILLUS FUMIGATUS</b> (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48)	ASPERGILLUS FUMIGATUS	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5020-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5020-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5020-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>MOLDS, RUSTS AND SMUTS, ASPERGILLUS NIGER</b>			
aspergillus niger injection, solution			
<b>Product Information</b>			
<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5032
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		
<b>Active Ingredient/Active Moiety</b>			
<b>Ingredient Name</b>			<b>Basis of Strength</b>
<b>ASPERGILLUS NIGER VAR. NIGER</b> (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6)			ASPERGILLUS NIGER VAR. NIGER 0.1 g in 1 mL
<b>Inactive Ingredients</b>			
<b>Ingredient Name</b>			<b>Strength</b>
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)			
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)			
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)			

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5032-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5032-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5032-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

  

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

## MOLDS, RUSTS AND SMUTS, BOTRYTIS CINEREA

botrytis cinerea injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5048
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BOTRYTIS CINEREA</b> (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7)	BOTRYTIS CINEREA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5048-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5048-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5048-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## INSECTS WHOLE BODY, ANT, FIRE SOLENOPSIS INVICTA

ant, fire solenopsis invicta injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-6513
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>SOLENOPODIS INVICTA</b> (UNII: E07CB4B111) (SOLENOPODIS INVICTA		0.1 g in 1 mL

**SOLENOPSIS INVICTA** (UNII: SU/CR4P444) (SOLENOPSIS INVICTA - UNII:507CR4P444)

**SOLENOPSIS INVICTA** 0.1 g  
in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-6513-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, CANDIDA ALBICANS

candida albicans injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5052
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC)	CANDIDA ALBICANS	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5052-1	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5052-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5052-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, CANDIDA ALBICANS

candida albicans injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5055
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC)	CANDIDA ALBICANS	0.001 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5055-1	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5055-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5055-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, CEPHALOSPORIUM ACREMONIUM

cephalosporium acremonium injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5056
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>SAROCLADIUM STRICTUM</b> (UNII: 3F36V0451W) (SAROCLADIUM STRICTUM - UNII:3F36V0451W)	SAROCLADIUM STRICTUM	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5056-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5056-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5056-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**MOLDS, RUSTS AND SMUTS, CURVULARIA SPICIFERA**

curvularia spicifera injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5076
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>COCHLIOBOLUS SPICIFER</b> (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD)	COCHLIOBOLUS SPICIFER	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5076-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5076-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5076-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>MOLDS, RUSTS AND SMUTS, EPICOCCUM NIGRUM</b>	
epicoccum nigrum injection, solution	

<b>Product Information</b>			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5100
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
Ingredient Name	Basis of Strength	Strength	
EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7)	EPICOCCUM NIGRUM	0.1 g in 1 mL	

<b>Inactive Ingredients</b>		
Ingredient Name	Strength	
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5100-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5100-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5100-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	

BLA

BLA103888

04/19/1941

**MOLDS, RUSTS AND SMUTS, EPIDERMOPHYTON FLOCCOSUM**

epidermophyton floccosum injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5104
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>EPIDERMOPHYTON FLOCCOSUM</b> (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S)	EPIDERMOPHYTON FLOCCOSUM	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5104-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5104-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5104-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**MOLDS, RUSTS AND SMUTS, FUSARIUM VASINFECTUM**

fusarium vasinfectum injection, solution

**Product Information**

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-5112
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>FUSARIUM OXYSPORUM VASINFECTUM</b> (UNII: 6M98DC08TZ) (FUSARIUM OXYSPORUM VASINFECTUM - UNII:6M98DC08TZ)	FUSARIUM OXYSPORUM VASINFECTUM	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5112-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5112-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5112-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# MOLDS, RUSTS AND SMUTS, HELMINTHOSPORIUM INTERSEMINATUM

helminthosporium interseminatum injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5124
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DENDRYPHIELLA VINOSA (UNII: 7S6NW5FH8X) (DENDRYPHIELLA VINOSA - UNII:7S6NW5FH8X)	DENDRYPHIELLA VINOSA	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5124-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>2</b>	NDC:65044-5124-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5124-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, HORMODENDRUM CLADOSPORIOIDES

hormodendrum cladosporioides injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5128
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CLADOSPORIUM CLADOSPORIOIDES (UNII: 4ZWY20GTGO) (CLADOSPORIUM CLADOSPORIOIDES - UNII:4ZWY20GTGO)	CLADOSPORIUM CLADOSPORIOIDES	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5128-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5128-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5128-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, MUCOR RACEMOSUS

mucor racemosus injection, solution

Product Information			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5144
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>MUCOR RACEMOSUS</b> (UNII: 17RH99LQ7G) (MUCOR RACEMOSUS - UNII:17RH99LQ7G)	MUCOR RACEMOSUS	0.1 g in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)			
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)			
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5144-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5144-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5144-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA103888	04/19/1941		

<b>MOLDS, RUSTS AND SMUTS, PENICILLIUM NOTATUM</b>			
penicillium notatum injection, solution			

Product Information			
Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5208
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM</b> (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG)	PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM	0.1 g in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)			

**SODIUM CHLORIDE** (UNII: 451W47IQ8X)

**SODIUM BICARBONATE** (UNII: 8MDF5V39QO)

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5208-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5208-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5208-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, PHOMA HERBARUM

phoma herbarum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5220
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>PHOMA EXIGUA VAR. EXIGUA</b> (UNII: 8JAG41IE4M) (PHOMA EXIGUA VAR. EXIGUA - UNII:8JAG41IE4M)	PHOMA EXIGUA VAR. EXIGUA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5220-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5220-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5220-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, PULLULARIA PULLULANS

pullularia pullulans injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5235
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK)	AUREOBASIDIUM PULLULANS VAR. PULLUTANS	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-5235-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-5235-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-5235-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, RHIZOPUS NIGRICANS

rhizopus nigricans injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5230
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>RHIZOPUS STOLONIFER</b> (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q)	RHIZOPUS STOLONIFER	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-5230-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5230-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5230-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## MOLDS, RUSTS AND SMUTS, STEMPHYLIUM BOTRYOSUM

stemphylium botryosum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-5264
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>PLEOSPORA TARDA</b> (UNII: TPL549N9R8) (PLEOSPORA TARDA - UNII:TPL549N9R8)	PLEOSPORA TARDA	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date

<b>1</b>	NDC:65044-5264-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-5264-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-5264-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, BAHIA GRASS PASPALUM NOTATUM

bahia grass paspalum notatum injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1081
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PASPALUM NOTATUM POLLEN (UNII: V003SHB7VK) (PASPALUM NOTATUM POLLEN - UNII:V003SHB7VK)	PASPALUM NOTATUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1081-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1081-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1081-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, BAHIA GRASS PASPALUM NOTATUM

bahia grass paspalum notatum injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1084
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PASPALUM NOTATUM POLLEN (UNII: V003SHB7VK) (PASPALUM NOTATUM POLLEN - UNII:V003SHB7VK)	PASPALUM NOTATUM POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1084-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1084-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1084-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, BROME, SMOOTH BROMUS INERMIS

brome, smooth bromus inermis injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1237
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BROMUS INERMIS POLLEN (UNII: 766QT72BK6) (BROMUS INERMIS POLLEN - UNII:766QT72BK6)	BROMUS INERMIS POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
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**GLYCERIN** (UNII: PDC6A3C0OX)**SODIUM CHLORIDE** (UNII: 451W47IQ8X)**SODIUM BICARBONATE** (UNII: 8MDF5V39QO)**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1237-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1237-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1237-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - GRASSES, CORN, CULTIVATED ZEA MAYS**

corn, cultivated zea mays injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1414
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
ZEA MAYS POLLEN (UNII: 74PD8J616H) (ZEA MAYS POLLEN - UNII:74PD8J616H)	ZEA MAYS POLLEN	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1414-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1414-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1414-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, JOHNSON GRASS SORGHUM HALEPENSE

johnson grass sorghum halepense injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1744
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SORGHUM HALEPENSE POLLEN (UNII: 577VA5B4HP) (SORGHUM HALEPENSE POLLEN - UNII:577VA5B4HP)	SORGHUM HALEPENSE POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1744-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1744-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1744-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, JOHNSON GRASS SORGHUM HALEPENSE

johnson grass sorghum halepense injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1747
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>SORGHUM HALEPENSE POLLEN</b> (UNII: 577VA5B4HP) (SORGHUM HALEPENSE POLLEN - UNII:577VA5B4HP)	SORGHUM HALEPENSE POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1747-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1747-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1747-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, OATS, COMMON, CULTIVATED AVENA SATIVA

oats, common, cultivated avena sativa injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2041
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>AVENA SATIVA POLLEN</b> (UNII: A7IKY24TR7) (AVENA SATIVA POLLEN - UNII:A7IKY24TR7)	AVENA SATIVA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date

<b>1</b>	NDC:65044-2041-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-2041-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-2041-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, GRASS MIX 8

grass mix 8 injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-0879
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>POA PRATENSIS POLLEN</b> (UNII: SCB8J7LS3T) (POA PRATENSIS POLLEN - UNII:SCB8J7LS3T)	POA PRATENSIS POLLEN	100000 [BAU] in 1 mL
<b>CYNODON DACTYLON POLLEN</b> (UNII: 175F461W10) (CYNODON DACTYLON POLLEN - UNII:175F461W10)	CYNODON DACTYLON POLLEN	10000 [BAU] in 1 mL
<b>SORGHUM HALEPENSE POLLEN</b> (UNII: 577VA5B4HP) (SORGHUM HALEPENSE POLLEN - UNII:577VA5B4HP)	SORGHUM HALEPENSE POLLEN	0.05 g in 1 mL
<b>AGROSTIS GIGANTEA POLLEN</b> (UNII: HU8V6E7HOA) (AGROSTIS GIGANTEA POLLEN - UNII:HU8V6E7HOA)	AGROSTIS GIGANTEA POLLEN	100000 [BAU] in 1 mL
<b>PHLEUM PRATENSE POLLEN</b> (UNII: 65M88RW2EG) (PHLEUM PRATENSE POLLEN - UNII:65M88RW2EG)	PHLEUM PRATENSE POLLEN	100000 [BAU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-0879-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, SOUTHERN GRASS MIX

pollens - grasses, southern grass mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-0854
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>POA PRATENSIS POLLEN</b> (UNII: SCB8J7LS3T) (POA PRATENSIS POLLEN - UNII:SCB8J7LS3T)	POA PRATENSIS POLLEN	100000 [BAU] in 1 mL
<b>DACTYLIS GLOMERATA POLLEN</b> (UNII: 83N78IDA7P) (DACTYLIS GLOMERATA POLLEN - UNII:83N78IDA7P)	DACTYLIS GLOMERATA POLLEN	100000 [BAU] in 1 mL
<b>AGROSTIS GIGANTEA POLLEN</b> (UNII: HU8V6E7HOA) (AGROSTIS GIGANTEA POLLEN - UNII:HU8V6E7HOA)	AGROSTIS GIGANTEA POLLEN	100000 [BAU] in 1 mL
<b>PHLEUM PRATENSE POLLEN</b> (UNII: 65M88RW2EG) (PHLEUM PRATENSE POLLEN - UNII:65M88RW2EG)	PHLEUM PRATENSE POLLEN	100000 [BAU] in 1 mL
<b>ANTHOXANTHUM ODORATUM POLLEN</b> (UNII: 2KIK19R45Y) (ANTHOXANTHUM ODORATUM POLLEN - UNII:2KIK19R45Y)	ANTHOXANTHUM ODORATUM POLLEN	100000 [BAU] in 1 mL
<b>SORGHUM HALEPENSE POLLEN</b> (UNII: 577VA5B4HP) (SORGHUM HALEPENSE POLLEN - UNII:577VA5B4HP)	SORGHUM HALEPENSE POLLEN	0.05 g in 1 mL
<b>CYNODON DACTYLON POLLEN</b> (UNII: 175F461W10) (CYNODON DACTYLON POLLEN - UNII:175F461W10)	CYNODON DACTYLON POLLEN	10000 [BAU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-0854-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-0854-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-0854-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - GRASSES, SOUTHERN GRASS MIX

pollens - grasses, southern grass mix injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-0856
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>POA PRATENSIS POLLEN</b> (UNII: SCB8J7LS3T) (POA PRATENSIS POLLEN - UNII:SCB8J7LS3T)	POA PRATENSIS POLLEN	10000 [BAU] in 1 mL
<b>DACTYLIS GLOMERATA POLLEN</b> (UNII: 83N78IDA7P) (DACTYLIS GLOMERATA POLLEN - UNII:83N78IDA7P)	DACTYLIS GLOMERATA POLLEN	10000 [BAU] in 1 mL
<b>AGROSTIS GIGANTEA POLLEN</b> (UNII: HU8V6E7HOA) (AGROSTIS GIGANTEA POLLEN - UNII:HU8V6E7HOA)	AGROSTIS GIGANTEA POLLEN	10000 [BAU] in 1 mL
<b>PHLEUM PRATENSE POLLEN</b> (UNII: 65M88RW2EG) (PHLEUM PRATENSE POLLEN - UNII:65M88RW2EG)	PHLEUM PRATENSE POLLEN	10000 [BAU] in 1 mL
<b>ANTHOXANTHUM ODORATUM POLLEN</b> (UNII: 2KIK19R45Y) (ANTHOXANTHUM ODORATUM POLLEN - UNII:2KIK19R45Y)	ANTHOXANTHUM ODORATUM POLLEN	10000 [BAU] in 1 mL
<b>SORGHUM HALEPENSE POLLEN</b> (UNII: 577VA5B4HP) (SORGHUM HALEPENSE POLLEN - UNII:577VA5B4HP)	SORGHUM HALEPENSE POLLEN	0.005 g in 1 mL
<b>CYNODON DACTYLON POLLEN</b> (UNII: 175F461W10) (CYNODON DACTYLON POLLEN - UNII:175F461W10)	CYNODON DACTYLON POLLEN	1000 [BAU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-0856-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-0856-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-0856-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ACACIA ACACIA LONGIFOLIA

acacia longifolia injection, solution

### Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-1006
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ACACIA LONGIFOLIA POLLEN</b> (UNII: 24SO2J296O) (ACACIA LONGIFOLIA POLLEN - UNII:24SO2J296O)	ACACIA LONGIFOLIA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1006-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1006-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1006-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ALDER, RED ALNUS RUBRA

alder, red alnus rubra injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1018
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ALNUS RUBRA POLLEN</b> (UNII: Z0F2YK1B7H) (ALNUS RUBRA POLLEN - UNII:Z0F2YK1B7H)	ALNUS RUBRA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date

<b>1</b>	NDC:65044-1018-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1018-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1018-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ALDER, RED ALNUS RUBRA

alder, red alnus rubra injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1021
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALNUS RUBRA POLLEN (UNII: Z0F2YK1B7H) (ALNUS RUBRA POLLEN - UNII:Z0F2YK1B7H)	ALNUS RUBRA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1021-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1021-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1021-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ASH, WHITE FRAXINUS AMERICANA

ash, white fraxinus americana injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1060
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FRAXINUS AMERICANA POLLEN</b> (UNII: G684LX721Q) (FRAXINUS AMERICANA POLLEN - UNII:G684LX721Q)	FRAXINUS AMERICANA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1060-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1060-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1060-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BEECH, AMERICAN FAGUS GRANDIFOLIA

beech, american fagus grandifolia injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1120
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>FAGUS GRANDIFOLIA POLLEN</b> (UNII: 34X886W1H4) (FAGUS GRANDIFOLIA POLLEN - UNII:34X886W1H4)	FAGUS GRANDIFOLIA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
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**GLYCERIN** (UNII: PDC6A3C0OX)

**SODIUM CHLORIDE** (UNII: 451W47IQ8X)

**SODIUM BICARBONATE** (UNII: 8MDF5V39QO)

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1120-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1120-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1120-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BIRCH MIX

birch mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1168
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	0.05 g in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	0.05 g in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1168-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1168-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-	50 mL in 1 VIAL; Type 0: Not a Combination		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BIRCH MIX

birch mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1171
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	0.1 g in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	0.1 g in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1171-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1171-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1171-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BIRCH MIX

birch mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1172
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BETULA PAPYRIFERA POLLEN</b> (UNII: 3538FNV8AY) (BETULA PAPYRIFERA POLLEN - UNII:3538FNV8AY)	BETULA PAPYRIFERA POLLEN	20000 [PNU] in 1 mL
<b>BETULA PENDULA POLLEN</b> (UNII: ZL5TV40C5Y) (BETULA PENDULA POLLEN - UNII:ZL5TV40C5Y)	BETULA PENDULA POLLEN	40000 [PNU] in 1 mL
<b>BETULA NIGRA POLLEN</b> (UNII: 93963RFO1P) (BETULA NIGRA POLLEN - UNII:93963RFO1P)	BETULA NIGRA POLLEN	40000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1172-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1172-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1172-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## **POLLENS - TREES, BOTTLE BRUSH CALLISTEMON SPP.**

bottle brush callistemon spp. injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1207
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>MELALEUCA CITRINA POLLEN</b> (UNII: 620II98F1T) (MELALEUCA CITRINA POLLEN - UNII:620II98F1T)	MELALEUCA CITRINA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C00X)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1207-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1207-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1207-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BOXELDER/MAPLE MIX

boxelder/maple mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1213
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ACER NEGUNDO POLLEN (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII: P6K070AR8V)	ACER NEGUNDO POLLEN	0.05 g in 1 mL
ACER SACCHARUM POLLEN (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII: V38QUQ7861)	ACER SACCHARUM POLLEN	0.05 g in 1 mL
ACER RUBRUM POLLEN (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII: 700NK45C76)	ACER RUBRUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C00X)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1213-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>2</b>	NDC:65044-1213-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1213-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BOXELDER/MAPLE MIX

boxelder/maple mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1216
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ACER NEGUNDO POLLEN</b> (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII:P6K070AR8V)	ACER NEGUNDO POLLEN	0.1 g in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.1 g in 1 mL
<b>ACER RUBRUM POLLEN</b> (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII:700NK45C76)	ACER RUBRUM POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1216-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1216-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1216-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, BOXELDER/MAPLE MIX

boxelder/maple mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1217
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ACER NEGUNDO POLLEN</b> (UNII: P6K070AR8V) (ACER NEGUNDO POLLEN - UNII:P6K070AR8V)	ACER NEGUNDO POLLEN	40000 [PNU] in 1 mL
<b>ACER SACCHARUM POLLEN</b> (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	40000 [PNU] in 1 mL
<b>ACER RUBRUM POLLEN</b> (UNII: 700NK45C76) (ACER RUBRUM POLLEN - UNII:700NK45C76)	ACER RUBRUM POLLEN	40000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1217-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1217-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1217-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CEDAR, MOUNTAIN JUNIPERUS ASHEI

cedar, mountain juniperus ashei injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1336
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
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<b>JUNIPERUS ASHEI POLLEN</b> (UNII: 544F8MEY0Y) (JUNIPERUS ASHEI POLLEN - UNII:544F8MEY0Y)	JUNIPERUS ASHEI POLLEN	0.05 g in 1 mL
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### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1336-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1336-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1336-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CEDAR, MOUNTAIN JUNIPERUS ASHEI

cedar, mountain juniperus ashei injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1339
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>JUNIPERUS ASHEI POLLEN</b> (UNII: 544F8MEY0Y) (JUNIPERUS ASHEI POLLEN - UNII:544F8MEY0Y)	JUNIPERUS ASHEI POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-	10 mL in 1 VIAL; Type 0: Not a Combination		

<b>1</b>	1339-2	Product		
<b>2</b>	NDC:65044-1339-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1339-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CEDAR, RED JUNIPERUS VIRGINIANA

cedar, red juniperus virginiana injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1343
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
JUNIPERUS VIRGINIANA POLLEN (UNII: PY0JA16R2G) (JUNIPERUS VIRGINIANA POLLEN - UNII:PY0JA16R2G)	JUNIPERUS VIRGINIANA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1343-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1343-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1343-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CEDAR, RED JUNIPERUS VIRGINIANA

cedar, red juniperus virginiana injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1342
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
JUNIPERUS VIRGINIANA POLLEN (UNII: PY0JA16R2G) (JUNIPERUS VIRGINIANA POLLEN - UNII:PY0JA16R2G)	JUNIPERUS VIRGINIANA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1342-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1342-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1342-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, COTTONWOOD, COMMON POPULUS DELTOIDES

cottonwood, common populus deltoides injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1435
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
POPULUS DELTOIDES POLLEN (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1435-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1435-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1435-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - TREES, COTTONWOOD, COMMON POPULUS DELTOIDES**

cottonwood, common populus deltoides injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1438
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
POPULUS DELTOIDES POLLEN (UNII: 476DVV63WP) (POPULUS DELTOIDES POLLEN - UNII:476DVV63WP)	POPULUS DELTOIDES POLLEN	20000 [PNU] in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1438-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1438-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1438-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CYPRESS, ARIZONA CUPRESSUS ARIZONICA

cypress, arizona cupressus arizonica injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1450
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CUPRESSUS ARIZONICA POLLEN (UNII: 232DMH0XVF) (CUPRESSUS ARIZONICA POLLEN - UNII:232DMH0XVF)	CUPRESSUS ARIZONICA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1450-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1450-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1450-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, CYPRESS, BALD TAXODIUM DISTICHUM

cypress, bald taxodium distichum injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1453
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TAXODIUM DISTICHUM POLLEN (UNII: O12H03B41R) (TAXODIUM DISTICHUM POLLEN - UNII:O12H03B41R)	TAXODIUM DISTICHUM POLLEN	0.02 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1453-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1453-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1453-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ELM, AMERICAN ULMUS AMERICANA

elm, american ulmus americana injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1540
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ULMUS AMERICANA POLLEN (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date

<b>1</b>	NDC:65044-1540-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1540-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1540-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ELM, AMERICAN ULMUS AMERICANA

elm, american ulmus americana injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1543
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ULMUS AMERICANA POLLEN (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	0.1 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1543-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1543-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1543-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ELM, AMERICAN ULMUS AMERICANA

elm, american ulmus americana injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1544
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ULMUS AMERICANA POLLEN (UNII: 89BAT511BD) (ULMUS AMERICANA POLLEN - UNII:89BAT511BD)	ULMUS AMERICANA POLLEN	40000 [PNU] in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
PHENOL (UNII: 339NCG44TV)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1544-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1544-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1544-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, ELM, CHINESE ULMUS PARVIFOLIA

elm, chinese ulmus parvifolia injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1546
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ULMUS PARVIFOLIA POLLEN (UNII: IU0Z41653U) (ULMUS PARVIFOLIA POLLEN - UNII:IU0Z41653U)	ULMUS PARVIFOLIA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1564-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1564-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1564-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, EUCALYPTUS, EUCALYPTUS GLOBULUS

eucalyptus, eucalyptus globulus injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1564
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
EUCALYPTUS GLOBULUS POLLEN (UNII: 7XW7TB10X9) (EUCALYPTUS GLOBULUS POLLEN - UNII:7XW7TB10X9)	EUCALYPTUS GLOBULUS POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1564-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1564-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1564-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, GUM, SWEET LIQUIDAMBAR STYRACIFLUA

gum, sweet liquidambar styraciflua injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1660
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
LIQUIDAMBAR STYRACIFLUA POLLEN (UNII: 5Q246DS5BS) (LIQUIDAMBAR STYRACIFLUA POLLEN - UNII:5Q246DS5BS)	LIQUIDAMBAR STYRACIFLUA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1660-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1660-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1660-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, GUM, SWEET LIQUIDAMBAR STYRACIFLUA

gum, sweet liquidambar styraciflua injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1662
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
<b>Ingredient Name</b>		<b>Basis of Strength</b>	<b>Strength</b>
LIQUIDAMBAR STYRACIFLUA POLLEN (UNII: 5Q246DS5BS) (LIQUIDAMBAR STYRACIFLUA POLLEN - UNII:5Q246DS5BS)		LIQUIDAMBAR STYRACIFLUA POLLEN	0.1 g in 1 mL

<b>Inactive Ingredients</b>			
<b>Ingredient Name</b>			<b>Strength</b>
PHENOL (UNII: 339NCG44TV)			
SODIUM CHLORIDE (UNII: 451W47IQ8X)			
SODIUM BICARBONATE (UNII: 8MDF5V39QO)			

<b>Packaging</b>				
#	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:65044-1662-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1662-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1662-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>Marketing Information</b>				
<b>Marketing Category</b>	<b>Application Number or Monograph Citation</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>	
BLA	BLA103888	04/19/1941		

<b>POLLENS - TREES, HACKBERRY CELTIS OCCIDENTALIS</b>			
hackberry celtis occidentalis injection, solution			

<b>Product Information</b>			
<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-1663
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

<b>Active Ingredient/Active Moiety</b>			
<b>Ingredient Name</b>		<b>Basis of Strength</b>	<b>Strength</b>
CELTIS OCCIDENTALIS POLLEN (UNII: 68R9X9Y96X) (CELTIS OCCIDENTALIS POLLEN - UNII:68R9X9Y96X)		CELTIS OCCIDENTALIS POLLEN	0.05 g in 1 mL

<b>Inactive Ingredients</b>			
<b>Ingredient Name</b>			<b>Strength</b>
GLYCERIN (UNII: PDC6A3C0OX)			
SODIUM CHLORIDE (UNII: 451W47IQ8X)			
SODIUM BICARBONATE (UNII: 8MDF5V39QO)			

<b>Packaging</b>			

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1663-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1663-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1663-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, HICKORY, SHAGBARK CARYA OVATA

hickory, shagbark carya ovata injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1702
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CARYA OVATA POLLEN (UNII: 54UN9R2798) (CARYA OVATA POLLEN - UNII:54UN9R2798)	CARYA OVATA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1702-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1702-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1702-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - TREES, LINDEN BASSWOOD TILIA AMERICANA

linden basswood tilia americana injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1801
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TILIA AMERICANA POLLEN (UNII: E2B4Q4BXJG) (TILIA AMERICANA POLLEN - UNII:E2B4Q4BXJG)	TILIA AMERICANA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C00X)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1801-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1801-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1801-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

# POLLENS - TREES, MAPLE, HARD ACER SACCHARUM

maple, hard acer saccharum injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1831
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ACER SACCHARUM POLLEN (UNII: V38QUQ7861) (ACER SACCHARUM POLLEN - UNII:V38QUQ7861)	ACER SACCHARUM POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1831-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1831-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1831-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, MELALEUCA PUNK TREE MELALEUCA QUINQUENERVIA

melaleuca punk tree melaleuca quinquenervia injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1873
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
MELALEUCA QUINQUENERVIA POLLEN (UNII: NX974IRT8E) (MELALEUCA QUINQUENERVIA POLLEN - UNII:NX974IRT8E)	MELALEUCA QUINQUENERVIA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1873-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1873-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		

<b>3</b>	NDC:65044-1873-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		
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## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, MESQUITE, PROSOPIS JULIFLORA

mesquite, prosopis juliflora injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1876
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PROSOPIS JULIFLORA POLLEN (UNII: 6EIJ3D04MR) (PROSOPIS JULIFLORA POLLEN - UNII:6EIJ3D04MR)	PROSOPIS JULIFLORA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1876-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1876-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1876-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, MULBERRY MIX

mulberry mix injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1909
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**Route of Administration**

PERCUTANEOUS, SUBCUTANEOUS

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>MORUS ALBA POLLEN</b> (UNII: 3I9T68187H) (MORUS ALBA POLLEN - UNII:3I9T68187H)	MORUS ALBA POLLEN	0.05 g in 1 mL
<b>MORUS RUBRA POLLEN</b> (UNII: 9LYI4RTZ52) (MORUS RUBRA POLLEN - UNII:9LYI4RTZ52)	MORUS RUBRA POLLEN	0.05 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b>	NDC:65044-1909-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>2</b>	NDC:65044-1909-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
<b>3</b>	NDC:65044-1909-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

**POLLENS - TREES, MULBERRY MIX**

mulberry mix injection, solution

**Product Information**

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-1912
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>MORUS ALBA POLLEN</b> (UNII: 3I9T68187H) (MORUS ALBA POLLEN - UNII:3I9T68187H)	MORUS ALBA POLLEN	0.1 g in 1 mL
<b>MORUS RUBRA POLLEN</b> (UNII: 9LYI4RTZ52) (MORUS RUBRA POLLEN - UNII:9LYI4RTZ52)	MORUS RUBRA POLLEN	0.1 g in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
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<b>PHENOL</b> (UNII: 339NCG44TV)
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-1912-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-1912-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-1912-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OAK MIX

oak mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2035
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.05 g in 1 mL
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	0.05 g in 1 mL
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	0.05 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>GLYCERIN</b> (UNII: PDC6A3C00X)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2035-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2035-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2035-4	50 mL in 1 VIAL; Type 0: Not a Combination		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OAK MIX

oak mix injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2038
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.1 g in 1 mL
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	0.1 g in 1 mL
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	0.1 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2038-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2038-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2038-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OAK MIX

oak mix injection, solution

## Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2039
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	20000 [PNU] in 1 mL
<b>QUERCUS VIRGINIANA POLLEN</b> (UNII: 8KDG09A4GO) (QUERCUS VIRGINIANA POLLEN - UNII:8KDG09A4GO)	QUERCUS VIRGINIANA POLLEN	20000 [PNU] in 1 mL
<b>QUERCUS ALBA POLLEN</b> (UNII: Z4Y9ZSV4KK) (QUERCUS ALBA POLLEN - UNII:Z4Y9ZSV4KK)	QUERCUS ALBA POLLEN	20000 [PNU] in 1 mL

## Inactive Ingredients

Ingredient Name	Strength
<b>PHENOL</b> (UNII: 339NCG44TV)	
<b>SODIUM CHLORIDE</b> (UNII: 451W47IQ8X)	
<b>SODIUM BICARBONATE</b> (UNII: 8MDF5V39QO)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2039-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2039-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2039-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OAK, RED QUERCUS RUBRA

oak, red quercus rubra injection, solution

## Product Information

<b>Product Type</b>	NON-STANDARDIZED ALLERGENIC	<b>Item Code (Source)</b>	NDC:65044-2014
<b>Route of Administration</b>	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>QUERCUS RUBRA POLLEN</b> (UNII: SVW19ET93C) (QUERCUS RUBRA POLLEN - UNII:SVW19ET93C)	QUERCUS RUBRA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2014-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2014-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
3	NDC:65044-2014-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## POLLENS - TREES, OLIVE OLEA EUROPAEA

olive olea europaea injection, solution

## Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-2050
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
OLEA EUROPAEA POLLEN (UNII: 43R41XZ627) (OLEA EUROPAEA POLLEN - UNII:43R41XZ627)	OLEA EUROPAEA POLLEN	0.05 g in 1 mL

## Inactive Ingredients

Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM CHLORIDE (UNII: 451W47IQ8X)		
SODIUM BICARBONATE (UNII: 8MDF5V39QO)		

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-2050-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-2050-3	30 mL in 1 VIAL; Type 0: Not a Combination Product		
-	NDC:65044-	50 mL in 1 VIAL; Type 0: Not a Combination		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	04/19/1941	

## ANIMAL ALLERGENS, UF DOG HAIR-DANDER CANIS SPP

animal allergens, dog dander canis spp injection, solution

### Product Information

Product Type	NON-STANDARDIZED ALLERGENIC	Item Code (Source)	NDC:65044-4850
Route of Administration	PERCUTANEOUS, SUBCUTANEOUS		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CANIS LUPUS FAMILIARIS HAIR (UNII: 05S7L91ZTR) (CANIS LUPUS FAMILIARIS HAIR - UNII:05S7L91ZTR)	CANIS LUPUS FAMILIARIS HAIR	0.0008 g in 1 mL
CANIS LUPUS FAMILIARIS DANDER (UNII: 11JCK302I4) (CANIS LUPUS FAMILIARIS DANDER - UNII:11JCK302I4)	CANIS LUPUS FAMILIARIS DANDER	0.0008 g in 1 mL

### Inactive Ingredients

Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65044-4850-2	10 mL in 1 VIAL; Type 0: Not a Combination Product		
2	NDC:65044-4850-4	50 mL in 1 VIAL; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103888	07/01/2022	

**Labeler** - Jubilant HollisterStier LLC (069263643)

**Registrant** - Jubilant HollisterStier LLC - HollisterStier Allergy (119241112)