

ASPERGILLUS MIX- aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

MONILIA MIX- candida albicans and neurospora intermedia solution

ASPERGILLUS FUMIGATUS- aspergillus fumigatus solution

ASPERGILLUS FLAVUS- aspergillus flavus solution

AUREOBASIDIUM PULLULANS- aureobasidium pullulans solution

MOLD MIX 3- alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum solution

MOLD MIX 2- aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

DRECHSLERA SPICIFERA- drechslera spicifera solution

EPIDERMOPHYTON FLOCCOSUM- epidermophyton floccosum solution

MUCOR CIRCINELLOIDES F. CIRCINELLOIDES- mucor circinelloides f. circinelloides solution

RHIZOPUS ORYZAE- rhizopus oryzae solution

NEUROSPORA INTERMEDIA- neurospora intermedia solution

PAECILOMYCES VARIOTII- paecilomyces variotii solution

PENICILLIUM CHRYSOGENUM (NOTATUM)- penicillium chrysogenum (notatum) solution

FUSARIUM MONILIFORME- fusarium moniliforme solution

GEOTRICHUM CANDIDUM- geotrichum candidum solution

RHODOTORULA MUCILAGINOSA- rhodotorula mucilaginosa solution

SACCHAROMYCES CEREVISIAE- saccharomyces cerevisiae solution

ALTERNARIA ALTERNATA- alternaria alternata solution

GLIOCLADIUM VIRIDE- gliocladium viride solution

HELMINTHOSPORIUM SOLANI- helminthosporium solani solution

MUCOR PLUMBEUS- mucor plumbeus solution

TRICHODERMA HARZIANUM- trichoderma harzianum solution

TRICHOPHYTON MENTAGROPHYTES- trichophyton mentagrophytes solution

TRICHOPHYTON RUBRUM- trichophyton rubrum solution

LOOSE SMUT, WHEAT- ustilago tritici solution

BARLEY LOOSE SMUT- ustilago nuda solution

LOOSE KERNEL SMUT- sporisorium cruentum solution

RHIZOPUS STOLONIFER- rhizopus stolonifer solution

TRICHOHECIUM ROSEUM- trichothecium roseum solution

ASPERGILLUS NIGER- aspergillus niger solution

CHAETOMIUM GLOBOSUM- chaetomium globosum solution

ASPERGILLUS MIX- aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

DEMATIACEAE MIX- alternaria alternata, aureobasidium pullulans, bipolaris sorokiniana, cladosporium herbarum, curvularia spicifera and helminthosporium solani solution

NEW STOCK FUNGI MIX- acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

NEW STOCK FUNGI MIX- acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis

cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

GRASS SMUT MIX- ustilago cynodontis and sporisorium cruentum solution

FUSARIUM MIX- gibberella fujikuroi and fusarium solani solution

MUCOR MIX- mucor circinelloides f. lusitanicus and mucor plumbeus solution

NEW STOCK FUNGI MIX- acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

MOLD MIX 1- alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

RHIZOPUS MIX- rhizopus stolonifer and rhizopus oryzae solution

MOLD MIX 2- aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

AHH MOLD MIX- alternaria alternata, bipolaris sorokiniana and cladosporium sphaerospermum solution

ALTERNARIA/HORMODENDRUM MIX- alternaria alternata and aspergillus fumigatus solution

ALTERNARIA HORMODENDRUM MIX- alternaria alternata and aspergillus fumigatus solution

ASPERGILLUS NIDULANS- aspergillus nidulans solution

BIPOLARIS SOROKINIANA- bipolaris sorokiniana solution

PENICILLIUM MIX- penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

GRASS SMUT MIX- ustilago cynodontis and sporisorium cruentum solution

GRAIN SMUT MIX- ustilago maydis, ustilago tritici, ustilago nuda and ustilago avenae solution

CANDIDA ALBICANS- candida albicans solution

CORN SMUT- ustilago maydis solution

ALTERNARIA/HORMODENDRUM MIX- alternaria alternata and aspergillus fumigatus solution

ACREMONIUM STRICTUM- acremonium strictum solution

NEW STOCK FUNGI MIX- acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

GRAIN SMUT MIX- ustilago maydis, ustilago tritici, ustilago nuda and ustilago avenae solution

PENICILLIUM DIGITATUM- penicillium digitatum solution

EPICOCCUM NIGRUM- epicoccum nigrum solution

STEMPHYLIUM SOLANI- stemphylium solani solution

MUCOR CIRCINELLOIDES F. LUSITANICUS- mucor circinelloides f. lusitanicus solution

MICROSPORUM CANIS- microsporium canis solution

BERMUDA GRASS SMUT- ustilago cynodontis solution
ASPERGILLUS AMSTELODAMI- aspergillus amstelodami solution
FUSARIUM SOLANI- fusarium solani solution
PHOMA BETAE- phoma betae solution
BOTRYTIS CINEREA- botrytis cinerea solution
CLADOSPORIUM HERBARUM- cladosporium herbarum solution
MYCOGONE PERNICIOSA- mycogone perniciosa solution
**CLADOSPORIUM SPHAEROSPERMUM- cladosporium
sphaerospermum solution**
**MOLD MIX 1- alternaria alternata, aspergillus niger, bipolaris sorokiniana,
cladosporium sphaerospermum and penicillium notatum solution**
OAT SMUT- ustilago avenae solution
**MOLD MIX 1- alternaria alternata, aspergillus niger, bipolaris sorokiniana,
cladosporium sphaerospermum and penicillium notatum solution**
**PENICILLIUM MIX- penicillium camemberti, penicillium chrysogenum,
penicillium digitatum, penicillium notatum and penicillium roqueforti solution**
**PHYCOMYCETES MIX- mucor circinelloides f. lusitanicus and rhizopus
stolonifer solution**
Greer Laboratories, Inc.

Non Standardized Allergenic Extracts
**Pollens, Molds, Epidermals, Insects, Dusts, Foods, and Miscellaneous
Inhalants**

WARNING

THIS ALLERGENIC PRODUCT IS INTENDED FOR USE BY PHYSICIANS WHO ARE EXPERIENCED IN THE ADMINISTRATION OF ALLERGENIC EXTRACTS AND THE EMERGENCY CARE OF ANAPHYLAXIS, OR FOR USE UNDER THE GUIDANCE OF AN ALLERGY SPECIALIST.

ALLERGENIC EXTRACTS MAY CAUSE SEVERE OR FATAL ANAPHYLAXIS IN EXTREMELY SENSITIVE PATIENTS. THE INITIAL DOSE MUST BE BASED ON SKIN TESTING AS DESCRIBED IN THE DOSAGE AND ADMINISTRATION SECTION OF THIS INSERT. PATIENTS SHOULD BE INSTRUCTED TO RECOGNIZE ADVERSE REACTION SYMPTOMS AND CAUTIONED TO CONTACT THE PHYSICIAN'S OFFICE IF REACTION SYMPTOMS OCCUR. IN CERTAIN INDIVIDUALS, THESE REACTIONS COULD BE FATAL. PATIENTS SHOULD BE OBSERVED FOR AT LEAST 20 MINUTES FOLLOWING TREATMENT.

EMERGENCY MEASURES, AS WELL AS PERSONNEL TRAINED IN THEIR USE, SHOULD BE IMMEDIATELY AVAILABLE IN THE EVENT OF A LIFE- THREATENING REACTION. PATIENTS BEING SWITCHED FROM ONE LOT OF EXTRACT TO ANOTHER FROM THE SAME MANUFACTURER SHOULD HAVE THEIR DOSE REDUCED BY 75%.

THIS PRODUCT SHOULD NOT BE INJECTED INTRAVENOUSLY.

REFER ALSO TO THE WARNINGS, PRECAUTIONS, ADVERSE REACTIONS AND OVERDOSAGE SECTION BELOW.

Allergenic Extracts are supplied as a sterile solution for intracutaneous or subcutaneous administration. Concentrates contain the soluble extractants of the source material with 0.5% sodium chloride and 0.54% sodium bicarbonate at a pH of 6.8 to 8.4 as aqueous extracts in water for injection or in 50% glycerin. Aqueous extracts contain 0.4% phenol as a preservative and 50% glycerinated extracts contain 0.2% phenol. Diluted aqueous extracts contain Buffered Saline with 0.5% sodium chloride, 0.04% potassium phosphate, 0.11% sodium phosphate heptahydrate, and 0.4% phenol in water for injection.

Source materials for these extracts are as follows: Pollens are collected from the respective grasses, weeds, trees, shrubs, cultured plants and flowers. Mold extracts are produced from pure culture mycelial mats. Rusts and smuts are obtained from natural growths. Epidermal extracts are produced from the hide, hair, or feathers containing the natural dander, or from separated dander. Insects are the whole body insects. House dust is made from various dusts ordinarily found in the home with the extract dialyzed to remove low-molecular weight irritants and concentrated to an extraction ratio of 1:1. Food extracts are prepared from the edible portions of the respective foods, obtained fresh if possible. Certain diagnostic food extracts contain 0.1% sodium formaldehyde sulfoxylate as an antioxidant. Other miscellaneous inhalants involved in respiratory allergy are obtained in the naturally occurring form to which a patient may be exposed.

Extracts are labeled either by weight-to-volume (w/v) based on the weight of the source material to the volume of the extracting fluid, or in protein nitrogen units (PNU) based on assay with one PNU representing 0.00001 mg of protein nitrogen.

The allergic reaction is dependent upon the presence of antigen-specific immunoglobulin E (IgE) antibodies that are bound to specific receptors on mast cells and basophils. The presence of IgE antibodies on mast cells and basophils sensitizes these cells and, upon interaction with the appropriate allergen, histamine and other mediators are released. IgE antibody has been shown to correlate with atopic diseases such as allergic rhinitis and allergic asthma. ⁽¹⁻⁴⁾ In the skin these mediators are responsible for the characteristic wheal and flare (erythema) reactions upon Allergenic Extract skin testing in persons with the specific allergies. ⁽³⁻⁷⁾

Specific immunotherapy with Allergenic Extracts as employed for over 45 years 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. ⁽⁸⁾ Several mechanisms have been proposed to explain the effectiveness of immunotherapy: an increase in antigen-specific IgG antibodies is frequently associated with clinical effectiveness, although correlation is not consistent in all studies; there is a decrease in specific IgE; and IgE production is suppressed during periods of seasonal or high exposure to the antigen. ⁽⁹⁾ Other changes following immunotherapy have been noted including development of auto-anti-idiotypic antibodies; a decrease in blood basophil sensitivity to allergen; a decrease in lymphokine production and lymphocyte proliferation by cells exposed to allergen; and development of allergen-specific suppressor cells. ⁽¹⁰⁾ The complete mechanisms of immunotherapy are not known and remain the subject of investigation.

Allergenic Extracts are indicated for the diagnosis and treatment of patients with immediate hypersensitivity allergy to the respective allergens, inhaled, ingested or

otherwise introduced into contact with sensitive tissues. The diagnosis of IgE-mediated allergy may be established by the allergy history, clinical evaluation, and skin test reactivity. ^(4,7,11) Immunotherapy with Allergenic Extracts is indicated when testing and patient history have identified the offending allergens and when it is not possible or practical to avoid these allergens. ⁽¹²⁻¹⁴⁾ Food extracts have not been proven effective in immunotherapy.

The use of Allergenic Extracts for the above purposes should be made only by physicians with special familiarity and knowledge of allergy. (See DOSAGE AND ADMINISTRATION)

There are no known absolute contraindications to the use of Allergenic Extracts for immunotherapy. Immunotherapy with specific antigens should not be done in those individuals who do not exhibit skin test or clinical sensitivity to the particular antigens. (See below under WARNINGS and PRECAUTIONS)

Allergenic extract injections should not be administered in the presence of diseases characterized by a bleeding diathesis.

Children with nephrotic syndrome require careful consideration and probably should not receive injection therapy because a variety of seemingly unrelated events, such as immunization, can cause an exacerbation of their nephrotic disease.

General contraindications include:

EXTREME SENSITIVITY TO THE SPECIFIC ALLERGEN - Determined from previous anaphylaxis following exposure.

AUTOIMMUNE DISEASE - Individuals with autoimmune disease may be at risk, due to the possibility of routine immunizations exacerbating symptoms of the underlying disease.

Concentrated extracts must be diluted with a sterile diluent prior to first use on a patient for treatment or intradermal testing. Allergenic Extracts are manufactured to assure high potency and have the ability during skin testing and immunotherapy to cause serious local and systemic reactions including death in sensitive patients. Most reactions occur within 20 minutes after injection, ⁽¹⁵⁾ but may occur later. ⁽¹⁶⁾ To minimize the potential for local or systemic reactions, the relative sensitivity of the patient must be assessed from the allergic history and from clinical observations. Patients should be informed of these risks prior to skin testing and immunotherapy (see PRECAUTIONS and ADVERSE REACTIONS below).

Allergenic Extract immunotherapy doses should be lowered or temporarily withheld from patients if any of the following conditions exist:

- (1) severe symptoms of rhinitis and/or asthma
- (2) infection or flu accompanied by fever
- (3) exposure to excessive amounts of clinically relevant allergen prior to a scheduled injection
- (4) evidence of a local or systemic reaction to the preceding extract injection during a course of immunotherapy

The dosage must be reduced when modifying dosages or components in a mixture or an individual prescription, or when starting a patient on fresh extract, even though the labeled strength of the old and new vials may be the same. This reduction in dosage may

be necessary due to the older vial losing potency during storage, or due to different sensitivities to different components. The amount of new extract given should not exceed 25% of the last dose given from the old vial, assuming both extracts contain comparable amounts of allergen. Any evidence of a local or generalized reaction requires a reduction in dosage during the initial stages of immunotherapy, as well as during maintenance therapy.

GENERAL:

Not for intravenous use!

Systemic allergic reactions may occur as a result of immunotherapy. The risk can be minimized by adherence to a careful injection schedule, which starts with a low concentration of extract and is increased slowly. Because of the danger of serious reactions, caution is needed in testing exquisitely sensitive patients, particularly with potent allergens, e.g., peanut, cottonseed, and flaxseed. ⁽⁸⁾ Such extracts should be appropriately diluted before use.

The physician must be prepared to treat anaphylaxis should it occur and have the necessary drugs and equipment on hand to do so. ⁽¹⁷⁻¹⁸⁾ Extracts should not be administered by the patient or other individuals who are not prepared to treat anaphylaxis should it occur.

Patients receiving Allergenic Extracts should be kept under observation a minimum of twenty ⁽²⁰⁾ minutes so that any adverse reaction can be observed and properly handled. ⁽¹⁵⁾ This time should be extended for high-risk patients such as those with unstable asthma or those suffering an exacerbation of their symptoms.

Patients receiving beta blockers may not be responsive to beta adrenergic drugs used to treat anaphylaxis. The risks of anaphylaxis in these patients should be carefully weighed against the benefits of immunotherapy.

Check the lot number and dosage schedule of the patient to verify correctness of a prescription number, a vial number, or strength. Only after this verification has been made should an injection be given.

A separate sterile needle and syringe should be used for each patient to prevent transmission of hepatitis or other infectious agents.

INFORMATION FOR PATIENTS:

Most serious reactions following the administration of Allergenic Extracts occur within 20 minutes; the patient should remain under observation for this period of time or longer if instructed by the physician. The size of any local reaction should be recorded, because increasingly large local reactions may precede a subsequent systemic reaction with increasing dosage. The patient should be instructed to report any unusual reactions. In particular, this includes unusual swelling and/or tenderness at the injection site, or reactions such as rhinorrhea, sneezing, coughing, wheezing, shortness of breath, nausea, dizziness, or faintness. Reactions may occur some time after leaving the physician's office, in which case medical attention should be sought immediately.

DRUG INTERACTIONS: Skin test diagnosis with Allergenic Extracts is contraindicated within 24 hours after the last dose of most antihistamines, within 48 hours after the last dose of terfenadine, and within 3 weeks or longer after the last dose of astemizole. These products suppress histamine skin test reactions and could mask a positive

response.

CARCINOGENESIS, MUTAGENESIS, IMPAIRMENT OF FERTILITY: There is no evidence of carcinogenicity, mutagenesis or impairment of fertility in humans from Allergenic Extracts. No long-term studies in animals have been performed to evaluate carcinogenic potential.

PREGNANCY: PREGNANCY CATEGORY C - Animal reproduction studies have not been conducted with Allergenic Extracts. It is also not known whether Allergenic Extracts can cause fetal harm when administered to a pregnant woman or whether they can affect reproduction capacity. Allergenic Extracts should be given to a pregnant woman only if clearly needed.

There is no evidence of adverse effects of Allergenic Extracts on the fetus. ⁽⁸⁾ Studies have not been performed in animals to determine whether extracts affect fertility in males or females, have teratogenic potential, or have other adverse effects on the fetus. Caution should be exercised in testing or treating pregnant females because a systemic reaction may cause an abortion as a result of uterine muscle contractions.

LABOR AND DELIVERY: There is no known information of adverse effects during labor and delivery.

NURSING MOTHERS: It is not known whether this product is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when extracts are administered to a nursing woman.

PEDIATRIC AND GERIATRIC USE: Although most extracts have not been studied systematically in children, children and geriatric patients appear to tolerate injections of Allergenic Extracts well. Studies with pollenosis and asthma have been conducted in children (e.g. refs. 19-21). Extract usage in children should follow the same precautions as in adults.

Adverse systemic reactions may occur within minutes upon use of an Allergenic Extract to which a person has specific sensitivity. These reactions consist primarily of allergic symptoms such as generalized skin erythema, urticaria, pruritus, angioedema, rhinitis, wheezing, laryngeal edema, and hypotension. Less commonly, nausea, emesis, abdominal cramps, diarrhea and uterine contractions may occur. Severe reactions may cause shock and loss of consciousness. Fatalities have occurred rarely. ^(8,22,23) These systemic reactions occur with varying frequency in different clinics and are usually less than 1%. To some extent, the reaction rate is related to the type and dose of administered extract and to the sensitivity of the patient. In general, immunotherapy with Allergenic Extracts is considered to be safe. ⁽²⁴⁾ Despite all precautions, occasional reactions are unavoidable.

Adverse systemic reactions should be treated as follows:

A. A tourniquet should be immediately applied to the extremity above the site of injection. Release the tourniquet every few minutes for a few seconds.

B. Epinephrine 1:1000 should be injected immediately in the opposite arm in amounts of 0.3 to 0.5 mL and 0.2 mL epinephrine should be administered at the site of injection. For children below the age of 6 years, adjust the dosage of epinephrine to 0.005 mL per pound of body weight per dose. Repeat epinephrine dosage in 15 minutes if necessary and if symptoms persist.

C. Adverse reactions not responding to epinephrine therapy may require the use of parenteral bronchodilators, vasopressors, oxygen, or volume replacement therapy.

Local reactions consisting of erythema, itching, swelling, tenderness and sometimes pain may occur at the injection site. These reactions may appear within a few minutes to hours and persist for several days. Local cold applications and oral antihistamines may be effective treatment. For marked and prolonged local reactions, steroids may be helpful.

Systemic reactions are uncommon after injection, but if the patient receives more extract than can be tolerated at that particular time and begins to experience immediate hypersensitivity anaphylaxis, the procedures listed under ADVERSE REACTIONS should be instituted.

Overdosage may occur because of an error in the volume of extract injected, or an incorrect dilution injected, or because the patient may be exposed to airborne or environmental antigens simultaneously with injection of the same antigens. In the event of a systemic reaction occurring, the dosage schedule should be carefully adjusted as outlined above under WARNINGS.

1. DIAGNOSTIC TESTING

For the patient with a suspected diagnosis of allergy to more than one antigen, initial skin testing should include the individual extracts. If a screening skin test with a mixture is used, a positive response should be followed by testing with the individual extracts to determine the degree of sensitivity to each and to guide in the selection of extracts and their concentration for immunotherapy if indicated. However, because a negative skin test with a mixture may not be indicative of the absence of allergy to one or more of the components due to their dilution, testing with individual extracts is more precise. False negative responses may occur if serum levels of antihistamines remain from prior medication administration (see CONTRAINDICATIONS). The use of a positive control is especially recommended for patients on prior medications which may decrease the histamine skin test response.

Scratch or Prick-puncture Skin Testing:

Allergenic Extract concentrates may be used for scratch or prick-puncture testing or scratch tests in 50% glycerin, 1:20 w/v or strongest available strength in 5 mL vials may be used. Prick-puncture tests with concentrated extracts in patients highly sensitive to the specific antigen should yield distinctive wheals with diameters of greater than 5 mm and with much larger erythema reactions. Glycerinated histamine phosphate 5 mg/mL (1.8 mg/mL histamine base) or aqueous histamine phosphate 2.75 mg/mL (1 mg/mL histamine base; 1:1,000 W/V) may be used as a positive control.

Intradermal Skin Testing:

Extract for intradermal testing must be prepared by diluting the stock concentrate injection vials with sterile diluent (use normal or buffered saline, or normal saline with human serum albumin) or the appropriate dilutions may be purchased.

a. Patients with a negative scratch or prick-puncture test:

Patients who do not react to a scratch or prick-puncture test should be tested intradermally, using a 26 or 27 gauge 1/4 inch needle, with 0.02 to 0.05 mL of an appropriate extract dilution from 1/100 to 1/1000 of the concentrate. A negative test

should be followed by a repeat test using a higher concentration until significant wheal and flare reaction sizes are attained or until the responses remain negative. As a negative control use the diluent or, in the case of extracts in 50% glycerin, use 0.5% to 1% glycerosaline solution. As a positive control, use glycerinated histamine phosphate diluted to 0.5 mg/mL (0.18 mg/mL histamine base) or aqueous histamine phosphate 0.275 mg/mL (0.1 mg/mL histamine base).

b. Patients tested only by the intradermal method:

Since highly reactive individuals may react intracutaneously at 1:1 million or even 1:10 million dilutions, any intradermal injection should be preceded by a puncture test and the dose adjusted accordingly. Other patients suspected of being moderately allergic should be tested with 0.02 to 0.05 mL of an appropriate extract dilution on the order of 1/10,000 to 1/100,000 of the concentrate. A negative test should be followed by repeat tests using progressively stronger ten-fold concentrations until significant wheal and flare reaction sizes are attained, or until skin test responses with the higher concentrations remain negative. As a negative control, use the diluent or, in the case of extracts in 50% glycerin, use 0.5% to 1% glycerosaline solution. As a positive control, use glycerinated histamine phosphate diluted to 0.5 mg/mL (0.18 mg/mL histamine base) or aqueous histamine phosphate 0.275 mg/mL (0.1 mg/mL histamine base).

Skin tests are graded in terms of the wheal and erythema response noted at 15 to 20 minutes, and compared to the appropriate controls. Wheal and erythema sizes may be recorded by actual measurement.

2. IMMUNOTHERAPY

Immunotherapy is administered by subcutaneous injection. Dosage of Allergenic Extracts is individualized according to the patient's sensitivity, the clinical response, and tolerance to the extract administered during the phases of an injection regimen. The initial dose of the extract should be determined based on the puncture test reactivity. In patients who appear to be exquisitely sensitive by history and skin test, the initial dose of the extract should be 0.05 to 0.1 mL of a low concentration, such as dilution number 5 or 6 in below. Patients with lesser sensitivity may be started with 0.05 to 0.1 mL of the next higher concentration. The amount of Allergenic Extract is increased at each injection by no more than 50% of the previous amount, and the next increment is governed by the response to the last injection. Large local reactions which persist for longer than 24 hours are generally considered an indication for repeating the previous dose or reducing the dose at the next administration. Any evidence of systemic reaction is an indication for a reduction of 75% in the subsequent dose. The upper limits of dosage have not been established; however, doses larger than 0.2 mL of an extract in 50% glycerin may cause discomfort upon injection. The dosage of Allergenic Extract does not vary significantly with the allergic disease under treatment.

To prepare dilutions starting from a concentrate such as 1:10 W/V, 1:20 W/V, OR 20,000 PNU/mL, proceed as in Table 1 below. (Note: Add 0.5 mL of concentrate to 4.5 mL of sterile diluent and make additional dilutions in the same manner.)

TABLE 1

Ten-Fold Dilution Series *

| <u>Dilution</u> | <u>Extract</u> | <u>Diluent W/V</u> | <u>W/V</u> | <u>PNU/mL</u> |
|-----------------|----------------|--------------------|------------|---------------|
|-----------------|----------------|--------------------|------------|---------------|

| | | | | | |
|---|--------------------------------|--------|--------------|--------------|--------|
| 0 | Concentrate | | 1:10 | 1:20 | 20,000 |
| 1 | 0.5 mL in dilution concentrate | 4.5 mL | 1:100 | 1:200 | 2,000 |
| 2 | 0.5 mL in dilution 1 | 4.5 mL | 1:1,000 | 1:2,000 | 200 |
| 3 | 0.5 mL in dilution 2 | 4.5 mL | 1:10,000 | 1:20,000 | 20 |
| 4 | 0.5 mL in dilution 3 | 4.5 mL | 1:100,000 | 1:200,000 | 2 |
| 5 | 0.5 mL in dilution 4 | 4.5 mL | 1:1,000,000 | 1:2,000,000 | 0.2 |
| 6 | 0.5 mL in dilution 5 | 4.5 mL | 1:10,000,000 | 1:20,000,000 | 0.0 |

* There is no direct potency correlation across the table between PNU and W/V.

Stock concentrate extracts containing up to 40,000 PNU/mL, or 1:10 W/V or other dilutions as requested by the physician are supplied in 5, 10, 30 and 50 mL in aqueous or 50% glycerin buffered saline. House dust extract is supplied in a 1:1 W/V concentrate, or a maximum of 10,000 PNU/mL. Extracts are also supplied in dropper vials for scratch or prick testing.

Allergenic Extracts should be stored at 2-8°C and kept at this temperature range during office use. Refer to vial labels for expiration dates. Diluted extracts are inherently less stable than concentrates. Dilutions of glycerinated extracts which result in glycerin below 50% are also less stable. The more dilute extracts in aqueous diluents should be replenished daily. Potency of a particular dilution can be checked by skin test in comparison to a fresh dilution of the extract on an individual known to be allergic to the specific antigen.

1. Lichtenstein LM, Ishizaka K, Norman PS, et al. IgE antibody measurements in ragweed hayfever: relationship to clinical severity and the results of immunotherapy. *J Clin Invest* 1973;52:474.
2. Elgefors B, Julin A, Johansson SGO. Immunoglobulin E in bronchial asthma. *Acta Allergol* 1974;29:327.
3. Norman PS. The clinical significance of IgE. *Hosp Pract* 1975;10:41-49.
4. Bryant DA, Burns MW, Lazarus L. The correlation between skin tests, bronchial provocation tests and the serum level of IgE specific for common allergens in patients with asthma. *Clin Allergy* 1975;5:145.
5. Loeffler JA, Cawley LP, Moeder M. Serum IgE levels: correlation with skin test sensitivity. *Ann Allergy* 1973;31:331.
6. Pepys J. Skin tests in diagnosis. In Gell PGH, Coombs RRA, Lachman PJ, eds. *Clinical aspects of Immunology*. Oxford: Blackwell Scientific Publications, 1975.
7. Burrows B, et al. Respiratory disorders and allergy skin test reactions. *Ann Allergy* 1976;84:134.
8. Implementation of Efficacy Review, Allergenic Extracts. *Federal Register* 1985;50:3082-3288
9. Levy DA, Lichenstein LM, Goldstein EO, Ishizaka K. Immunologic and cellular changes

- accompanying the therapy of pollen allergy. *J Clin Invest* 1973;50:360.
10. Gurka G, Rocklin R. Immunologic responses during allergen-specific immunotherapy for respiratory allergy. *Ann Allergy* 1988;61:239-43.
 11. Zeiss CR Jr. Patient evaluation. In: *Allergy and Clinical Immunology*, Locky RF, ed. Garden City, N.Y.: Medical Examination Publishing 1976:616.
 12. Frankland AW, Augustin R. Prophylaxis of summer hay-fever and asthma: a controlled trial comparing crude grass-pollen extracts with the isolated main protein component. *Lancet* 1954;1:1055.
 13. Frankland AW, Augustin R. Grass pollen antigens effective in treatment. *Clin Sci* 1962;23:95.
 14. Rohr AS, Marshall NA, Saxon A: Successful immunotherapy for *Triatoma protracta*-induced anaphylaxis. *J Allergy Clin Immunol* 1984;73:369-75.
 15. Executive Committee, American Academy of Allergy and Immunology. The waiting period after allergen skin testing and immunotherapy (Position statement). *J Allergy Clin Immunol* 1990;85:526-7.
 16. Greenberg MA, Kaufman CR, Gonzalez GE, Rosenblatt CD, Smith LJ, Summers RJ. Late and immediate systemic-allergic reactions to inhalant allergen immunotherapy. *J Allergy Clin Immunol* 1986;77:865-70.
 17. Ouellette JJ. Emergency management of the allergic reactions. *Modern Medicine* 1975;99.
 18. Anderson JA, et al. Personnel and equipment to treat systemic reactions caused by immunotherapy with allergenic extracts. *J Allergy Clin Immunol* 1986;77:271-3.
 19. Sadan N, Rhyne MB, Mellits ED, et al. Immunotherapy of pollenosis in children: investigation of the immunologic basis of clinical improvement. *N Eng J Med* 1969;280:623.
 20. Johnstone DE. Value of hyposensitization therapy for perennial bronchial asthma in children. *Pediatrics* 1961;27:39.
 21. VanAsperin PP, Kemp AS, Mellis CM. Skin test reactivity and clinical allergen sensitivity in infancy. *J Allergy Clin Immunol* 1984;73:381-6.
 22. Committee on Safety of Medicine. Desensitizing vaccines. *Br Med J* 1986;293:948.
 23. Locky RF, Benedict LM, Turkeltaub PC, Bukantz SC. Fatalities from immunotherapy (IT) and skin testing (ST). *J Allergy Clin Immunol* 1987;79:660-77.
 24. Norman PS, Van Metre TE Jr. The safety of allergenic immunotherapy. *J Allergy Clin Immunol* 1990;85:522-5.

Sterile Multiple Dose Vial U.S. Rx Only Store at 2-8°C

ALLERGENIC EXTRACT

Alternaria alternata

Item: M1A06 10 mL 20,000 PNU/mL

Preservative 0.4% Phenol.

No U.S. Standard of Potency. See Package Insert for Contents, Dose and Directions for Use.



GTIN (01) 00322840160028
S/N (21) 000000000000
LOT (10) SAMPLE
EXP (17) 01 Jan 2025

H05-1.00



NDC 22840-1600-2 DIN 02372428

GREER Laboratories, Inc. Lenoir, NC 28645 U.S. License 308

Sterile Multiple Dose Vial U.S. Rx Only Store at 2-8°C

ALLERGENIC EXTRACT

Aspergillus niger

Item: M5A08 5 mL 1:1,000 W/V

Preservative 0.4% Phenol.

No U.S. Standard of Potency. See Package Insert for Contents, Dose and Directions for Use.



GTIN (01) 00322840162817
S/N (21) 000000000000
LOT (10) SAMPLE
EXP (17) 01 Jan 2025

H04-1.00



NDC 22840-1628-1 DIN 02372428

GREER Laboratories, Inc. Lenoir, NC 28645 U.S. License 308

Sterile Multiple Dose Vial U.S. Rx Only Store at 2-8°C

ALLERGENIC EXTRACT

Candida albicans

Item: GM15A03 50 mL 1:20 W/V

Preservative 0.2% Phenol.

Contains 50% v/v Glycerin. No U.S. Standard of Potency. See Package Insert for Contents, Dose and Directions for Use.



GTIN (01) 00322840561047
S/N (21) 000000000000
LOT (10) SAMPLE
EXP (17) 01 Jan 2025

H15-1.00



NDC 22840-5610-4 DIN 02372428

GREER Laboratories, Inc. Lenoir, NC 28645 U.S. License 308

Skin Test Only Vial

U.S. Rx Only

Store at 2-8°C

ALLERGENIC EXTRACT**FUSARIUM MIX**Equal Parts *Gibberella fujikuroi*, *Fusarium solani*

Item: GMO9A01 5 mL 1:40 W/V

Preservative 0.2% Phenol.

Contains 50% v/v Glycerin. No U.S. Standard of Potency. See Package Insert for Contents, Dose and Directions for Use.



GTIN (01) 00322840964152

S/N (21) 000000000000

LOT (10) SAMPLE

EXP (17) 01 Jan 2025



H36-1.00

NDC 22840-9641-5

GREER Laboratories, Inc. Lenoir, NC 28645 U.S. License 308

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9633 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|----------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.01 g in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.01 g in 1 mL |
| EUROTIIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIIUM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIIUM AMSTELODAMI | 0.01 g in 1 mL |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.01 g in 1 mL |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.01 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9633-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9633-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9633-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MONILIA MIX

candida albicans and neurospora intermedia solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9644 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|------------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.0005 g in 1 mL |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.0005 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9644-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FUMIGATUS

aspergillus fumigatus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1616 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------|---------------------|
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1616-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FLAVUS

aspergillus flavus solution

Product Information

| | | | | |
|---|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1614 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 40000 [PNU] in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1614-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|---|---|---------------------------|----------------|
| ASPERGILLUS FUMIGATUS | | | |
| aspergillus fumigatus solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5604 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | Basis of Strength | Strength | |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.05 g in 1 mL | |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5604-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5604-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5604-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

AUREOBASIDIUM PULLULANS

aureobasidium pullulans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5607 |
| Route of Administration | INTRADERMAL, 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 | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5607-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5607-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5607-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9620 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|------------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0125 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.0125 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9620-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9620-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9620-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 2

aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9613 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|-----------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.005 g in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.005 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.005 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.005 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.005 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9613-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9613-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9613-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DRECHSLERA SPICIFERA

drechslera spicifera solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1660 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|----------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W471Q8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1660-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1660-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPIDERMOPHYTON FLOCCOSUM

epidermophyton floccosum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2666 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------------|---------------|
| EPIDERMOPHYTON FLOCCOSUM (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S) | EPIDERMOPHYTON FLOCCOSUM | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2666-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. CIRCINELLOIDES

mucor circinelloides f. circinelloides solution

| Product Information | | | | |
|--|---|--|--|---------------------------|
| Product Type | | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1692 |
| Route of Administration | | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |
| MUCOR CIRCINELLOIDES F. CIRCINELLOIDES (UNII: 48Z8OUT98B) (MUCOR CIRCINELLOIDES F. CIRCINELLOIDES - UNII:48Z8OUT98B) | | | MUCOR CIRCINELLOIDES F. CIRCINELLOIDES | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | | |
| Ingredient Name | | | | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1692-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| RHIZOPUS ORYZAE | | | | |
|--|--|---|---------------------------|---------------------|
| rhizopus oryzae solution | | | | |
| Product Information | | | | |
| Product Type | | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2620 |
| Route of Administration | | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |
| RHIZOPUS ARRHZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZUS - UNII:8476849N1Y) | | | RHIZOPUS ARRHZUS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2620-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. CIRCINELLOIDES

mucor circinelloides f. circinelloides solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2670 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|----------------|
| MUCOR CIRCINELLOIDES F. CIRCINELLOIDES (UNII: 48Z8OUT98B) (MUCOR CIRCINELLOIDES F. CIRCINELLOIDES - UNII:48Z8OUT98B) | MUCOR CIRCINELLOIDES F. CIRCINELLOIDES | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
|---|-----------|---------------------|----------------------|--------------------|

| | | | | |
|---|------------------|--|--|--|
| 1 | NDC:22840-2670-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
|---|------------------|--|--|--|

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEUROSPORA INTERMEDIA

neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1699 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------|
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1699-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1699-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEUROSPORA INTERMEDIA

neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2602 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|---|-----------------------------|---------------------------|
| 1 | NDC:22840-2602-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

PAECILOMYCES VARIOTII

paecilomyces variotii solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2673 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | | |
|--|--|--|----------------------|--------------------|
| Ingredient Name | | Basis of Strength | Strength | |
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | | PAECILOMYCES VARIOTII | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2673-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| PENICILLIUM CHRYSOGENUM (NOTATUM) | | | |
|--|---|--|---------------------|
| penicillium chrysogenum (notatum) solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2616 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2616-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MONILIFORME

fusarium moniliforme solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1672 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|----------------|
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1672-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GEOTRICHUM CANDIDUM

geotrichum candidum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5620 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|---------------------|----------------|
| GEOTRICHUM CANDIDUM (UNII: 5964J742O8) (GEOTRICHUM CANDIDUM - UNII:5964J742O8) | GEOTRICHUM CANDIDUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5620-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5620-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5620-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM CHRYSOGENUM (NOTATUM)

penicillium chrysogenum (notatum) solution

| Product Information | | | | |
|--|--|---|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2612 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.001 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2612-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| RHIZOPUS ORYZAE | | | |
|---------------------------------|---|--------------------|----------------|
| rhizopus oryzae solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2621 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | Basis of Strength | Strength | |

| | | |
|---|--------------------|--------------------|
| RHIZOPUS ARRHZIZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZIZUS - UNII:8476849N1Y) | RHIZOPUS ARRHZIZUS | 0.001 g in 1 mL |
|---|--------------------|--------------------|

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2621-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHODOTORULA MUCILAGINOSA

rhodotorula mucilaginosa solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2630 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-------------------|
| RHODOTORULA MUCILAGINOSA (UNII: 62TY3X4N9Z) (RHODOTORULA MUCILAGINOSA - UNII:62TY3X4N9Z) | RHODOTORULA MUCILAGINOSA | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2630-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2630-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

SACCHAROMYCES CEREVISIAE

saccharomyces cerevisiae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2633 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES CEREVISIAE - UNII:978D8U419H) | SACCHAROMYCES CEREVISIAE | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2633-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA ALTERNATA

alternaria alternata solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1602 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1602-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1602-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MONILIFORME

fusarium moniliforme solution

Product Information

| | | | |
|--------------------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5618 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, | | |

Route of Administration SUBCUTANEOUS

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5618-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5618-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5618-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GEOTRICHUM CANDIDUM

geotrichum candidum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2667 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------|----------|
| GEOTRICHUM CANDIDUM (UNII: 5964J742O8) (GEOTRICHUM CANDIDUM - | GEOTRICHUM | 0.1 g |

| | | |
|------------------|----------|---------|
| UNII:5964J742O8) | CANDIDUM | in 1 mL |
|------------------|----------|---------|

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2667-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1685 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|---------------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1685-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1685-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1686 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------|---------------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1686-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

HELMINTHOSPORIUM SOLANI

helminthosporium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1689 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------|------------------|
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1689-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1689-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR PLUMBEUS

mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1695 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, | | |

Route of Administration SUBCUTANEOUS

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|---------------------|
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1695-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR PLUMBEUS

mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2672 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | | Strength | | |
|---------------------------------------|--|--|----------------------|--------------------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2672-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| PAECILOMYCES VARIOTII | | | | |
|--|---|---|----------------------|--------------------|
| paecilomyces variotii solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2608 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | Basis of Strength | Strength | |
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | | PAECILOMYCES VARIOTII | 0.001 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2608-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

HELMINTHOSPORIUM SOLANI

helminthosporium solani solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1687 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------------|-----------------|
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1687-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHODOTORULA MUCILAGINOSA

rhodotorula mucilaginosa solution

| Product Information | | | | |
|---------------------------------|--|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2631 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | RHODOTORULA MUCILAGINOSA (UNII: 62TY3X4N9Z) (RHODOTORULA MUCILAGINOSA - UNII:62TY3X4N9Z) | RHODOTORULA MUCILAGINOSA | 40000 [PNU] in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | | Strength | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2631-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| RHODOTORULA MUCILAGINOSA | | | |
|-----------------------------------|--|---------------------------|-----------------|
| rhodotorula mucilaginosa solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2675 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | RHODOTORULA MUCILAGINOSA (UNII: 62TY3X4N9Z) (RHODOTORULA | RHODOTORULA | 20000 [PNU] |

MUCILAGINOSA - UNII:62TY3X4N9Z)

MUCILAGINOSA

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:22840-2675-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHODOTORULA MUCILAGINOSA

rhodotorula mucilaginosa solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5635 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|----------------|
| RHODOTORULA MUCILAGINOSA (UNII: 62TY3X4N9Z) (RHODOTORULA MUCILAGINOSA - UNII:62TY3X4N9Z) | RHODOTORULA MUCILAGINOSA | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5635-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5635-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5635-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHODERMA HARZIANUM

trichoderma harzianum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2641 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| TRICHODERMA HARZIANUM (UNII: CA33Q4013Q) (TRICHODERMA HARZIANUM - UNII:CA33Q4013Q) | TRICHODERMA HARZIANUM | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2641-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2641-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHODERMA HARZIANUM

trichoderma harzianum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2680 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------|---------------------|
| TRICHODERMA HARZIANUM (UNII: CA33Q4013Q) (TRICHODERMA HARZIANUM - UNII:CA33Q4013Q) | TRICHODERMA HARZIANUM | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2680-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON MENTAGROPHYTES

trichophyton mentagrophytes solution

Product Information

| | | | | |
|--|--|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2644 | |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.1 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | | Strength | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2644-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|--|-----------------------------|-----------------|
| TRICHOPHYTON MENTAGROPHYTES | | | |
| trichophyton mentagrophytes solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2648 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2648-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON MENTAGROPHYTES

trichophyton mentagrophytes solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5639 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|----------------|
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
| | | | | |

| | | | | |
|---|------------------|--|--|--|
| 1 | NDC:22840-5639-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5639-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5639-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON RUBRUM

trichophyton rubrum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2650 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|-----------------|
| TRICHOPHYTON RUBRUM (UNII: 2ZAU32517N) (TRICHOPHYTON RUBRUM - UNII:2ZAU32517N) | TRICHOPHYTON RUBRUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2650-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON RUBRUM

trichophyton rubrum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2681 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|---------------|
| TRICHOPHYTON RUBRUM (UNII: 2ZAU32517N) (TRICHOPHYTON RUBRUM - UNII:2ZAU32517N) | TRICHOPHYTON RUBRUM | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2681-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MONILIA MIX

candida albicans and neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9677 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|----------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.05 g in 1 mL |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9677-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MONILIA MIX

candida albicans and neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9645 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-----------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.025 g in 1 mL |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9645-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9645-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9645-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPIDERMOPHYTON FLOCCOSUM

epidermophyton floccosum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1668 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------------|-----------------|
| EPIDERMOPHYTON FLOCCOSUM (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S) | EPIDERMOPHYTON FLOCCOSUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1668-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

LOOSE SMUT, WHEAT

ustilago tritici solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5653 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------|----------------|
| USTILAGO TRITICI (UNII: BV82OL2IZ8) (USTILAGO TRITICI - UNII:BV82OL2IZ8) | USTILAGO TRITICI | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5653-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5653-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5653-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BARLEY LOOSE SMUT

ustilago nuda solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5652 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-----------------|
| USTILAGO NUDA (UNII: 9Y53ZS6I82) (USTILAGO NUDA - UNII:9Y53ZS6I82) | USTILAGO NUDA | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5652-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5652-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5652-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. CIRCINELLOIDES

mucor circinelloides f. circinelloides solution

| Product Information | | | |
|---------------------------------|---|--|-----------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5624 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | MUCOR CIRCINELLOIDES F. CIRCINELLOIDES (UNII: 48Z8OUT98B) (MUCOR CIRCINELLOIDES F. CIRCINELLOIDES - UNII:48Z8OUT98B) | MUCOR CIRCINELLOIDES F. CIRCINELLOIDES | 0.05 g in 1 mL |
| Inactive Ingredients | | | |
| | Ingredient Name | | Strength |
| | PHENOL (UNII: 339NCG44TV) | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | |
| | GLYCERIN (UNII: PDC6A3C0OX) | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | |
| Packaging | | | |
| # | Item Code | Package Description | Marketing Start Date |
| 1 | NDC:22840-5624-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | |
| 2 | NDC:22840-5624-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | |
| Marketing Information | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| LOOSE KERNEL SMUT | | | |
|---------------------------------|---|---------------------------|-----------------|
| sporisorium cruentum solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5650 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of | Strength |

| Ingredient Name | | Strength | Strength | |
|--|--|--|----------------------|--------------------|
| SPORISORIUM CRUENTUM (UNII: GQM6LVU5V8) (SPORISORIUM CRUENTUM - UNII:GQM6LVU5V8) | | SPORISORIUM CRUENTUM | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| GLYCERIN (UNII: PDC6A3C0OX) | | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5650-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5650-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5650-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| PAECILOMYCES VARIOTII | | | |
|--|---|-----------------------|--------------------|
| paecilomyces variotii solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2607 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | | PAECILOMYCES VARIOTII | 1000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |

| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
|--|--|---|----------------------|--------------------|
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2607-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| ASPERGILLUS FUMIGATUS | | | | |
|---|---|---|----------------------|--------------------|
| aspergillus fumigatus solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1621 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.001 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1621-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS STOLONIFER

rhizopus stolonifer solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2624 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|---------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2624-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2624-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS STOLONIFER

rhizopus stolonifer solution

| Product Information | | | | |
|--|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | | Item Code (Source) | NDC:22840-2625 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | | | RHIZOPUS STOLONIFER | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | | |
| Ingredient Name | | | | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2625-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | | 09/15/1981 | |

| SACCHAROMYCES CEREVISIAE | | | | |
|--|---|--|---------------------------|-----------------|
| saccharomyces cerevisiae solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | | Item Code (Source) | NDC:22840-2634 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES | | | SACCHAROMYCES | 0.1 g |

| | | | | |
|---------------------------------------|---|--|-----------------------------|---------------------------|
| CEREVISIAE - UNII:978D8U419H) | | CEREVISIAE | in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2634-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2634-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|---|---------------------------|-----------------|
| SACCHAROMYCES CEREVISIAE | | | |
| saccharomyces cerevisiae solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2677 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES CEREVISIAE - UNII:978D8U419H) | | SACCHAROMYCES CEREVISIAE | 0.05 g in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2677-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON MENTAGROPHYTES

trichophyton mentagrophytes solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2646 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------------|----------------|
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2646-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOPHYTON RUBRUM

trichophyton rubrum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5640 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|----------------|
| TRICHOPHYTON RUBRUM (UNII: 2ZAU32517N) (TRICHOPHYTON RUBRUM - UNII:2ZAU32517N) | TRICHOPHYTON RUBRUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5640-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5640-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5640-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOTHECIUM ROSEUM

trichothecium roseum solution

Product Information

| | | | |
|---------------------|-----------------------------|------------------|------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code | NDC:22840- |
|---------------------|-----------------------------|------------------|------------|

| | | | | |
|--|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | (Source) | 2652 | |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | Basis of Strength | Strength | |
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | | TRICHOTHECIUM ROSEUM | 0.1 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2652-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2652-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|---|---------------------------|---------------------|
| TRICHOTHECIUM ROSEUM | | | |
| trichothecium roseum solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2653 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | | TRICHOTHECIUM ROSEUM | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2653-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2653-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOTHECIUM ROSEUM

trichothecium roseum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2655 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|-----------------|
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | TRICHOTHECIUM ROSEUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2655-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOTHECIUM ROSEUM

trichothecium roseum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5641 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|----------------|
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | TRICHOTHECIUM ROSEUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5641-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5641-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5641-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 4 | NDC:22840-5641-3 | 30 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA ALTERNATA

alternaria alternata solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5600 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|----------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5600-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5600-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5600-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

SACCHAROMYCES CEREVISIAE

saccharomyces cerevisiae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2676 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|---------------------|
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES CEREVISIAE - UNII:978D8U419H) | SACCHAROMYCES CEREVISIAE | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2676-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA ALTERNATA

alternaria alternata solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1600 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | | |
|--|--|--|----------------------|--------------------|
| Ingredient Name | | Basis of Strength | Strength | |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | | ALTERNARIA ALTERNATA | 20000 [PNU] in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1600-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1600-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| ALTERNARIA ALTERNATA | | | |
|--|---|----------------------|---------------------|
| alternaria alternata solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1601 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | | ALTERNARIA ALTERNATA | 40000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |

| | |
|---|--|
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1601-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1601-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FLAVUS

aspergillus flavus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1610 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|----------------|
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1610-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| | | |
|---|------------------|--|
| 2 | NDC:22840-1610-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product |
|---|------------------|--|

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FUMIGATUS

aspergillus fumigatus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1615 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------|---------------|
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1615-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1615-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIGER

aspergillus niger solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1625 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|------------------------------|------------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1625-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1625-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CHAETOMIUM GLOBOSUM

chaetomium globosum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1648 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | | |
|--|--|--|----------------------|---------------------|
| Ingredient Name | | | Basis of Strength | Strength |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | | | CHAETOMIUM GLOBOSUM | 40000 [PNU] in 1 mL |
| Inactive Ingredients | | | | |
| Ingredient Name | | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1648-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1648-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | | 09/15/1981 | |

| CHAETOMIUM GLOBOSUM | | | |
|--|---|---------------------|----------------|
| chaetomium globosum solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1647 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | | CHAETOMIUM GLOBOSUM | 0.1 g in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | | Strength |

| | |
|--|--|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

| Packaging | | | | |
|------------------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1647-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1647-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|------------------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

CHAETOMIUM GLOBOSUM

chaetomium globosum solution

| Product Information | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1650 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | |
|---|---------------------|---------------------|
| Ingredient Name | Basis of Strength | Strength |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 20000 [PNU] in 1 mL |

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

| Packaging | | | | |
|------------------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1650-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CHAETOMIUM GLOBOSUM

chaetomium globosum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1651 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|-----------------|
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1651-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum

solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9614 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|-------------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.00025 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.00025 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.00025 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.00025 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9614-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|---------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9618 |
|---------------------|-----------------------------|---------------------------|----------------|

| | | | | |
|--|---|--|-----------------------------|---------------------------|
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 10000 [PNU] in 1 mL | |
| | ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 10000 [PNU] in 1 mL | |
| | ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 10000 [PNU] in 1 mL | |
| | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 10000 [PNU] in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9618-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

| | | | |
|--|---|---------------------------|----------------|
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9631 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|------------------|
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.0002 g in 1 mL |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.0002 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.0002 g in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.0002 g in 1 mL |
| EUROTIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIUM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIUM AMSTELODAMI | 0.0002 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9631-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DEMATIACEAE MIX

alternaria alternata, aureobasidium pullulans, bipolaris sorokiniana, cladosporium herbarum, curvularia spicifera and helminthosporium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9634 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|------------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.0083 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLULANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLULANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. | 0.0083 g in 1 mL |

| | | |
|---|-------------------------|------------------|
| (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:DIAZNG09CN) | PULLUTANS | 0.0083 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0083 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0083 g in 1 mL |
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | CLADOSPORIUM HERBARUM | 0.0083 g in 1 mL |
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 0.0083 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9634-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9649 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|-------------------|
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.00625 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.00625 g in 1 mL |

| | | |
|---|--|-------------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.00625 g in 1 mL |
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 0.00625 g in 1 mL |
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.00625 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.00625 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.00625 g in 1 mL |
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.00625 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.00625 g in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.00625 g in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.00625 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.00625 g in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 19917J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:19917J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.00625 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.00625 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.00625 g in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.00625 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9649-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9649-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9652 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.003125 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.003125 g in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.003125 g in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.003125 g in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.003125 g in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.003125 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.003125 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.003125 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.003125 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.003125 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.003125 g in 1 mL |
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 0.003125 g in 1 mL |
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.003125 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.003125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.003125 g in 1 mL |
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.003125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9652-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GRASS SMUT MIX

ustilago cynodontis and sporisorium cruentum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9676 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| USTILAGO CYNODONTIS (UNII: 0V3J4YEX2W) (USTILAGO CYNODONTIS - UNII:0V3J4YEX2W) | USTILAGO CYNODONTIS | 0.05 g in 1 mL |
| SPORISORIUM CRUENTUM (UNII: GQM6LVU5V8) (SPORISORIUM CRUENTUM - UNII:GQM6LVU5V8) | SPORISORIUM CRUENTUM | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9676-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DRECHSLERA SPICIFERA

drechslera spicifera solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1664 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------|-----------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1664-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MIX

gibberella fujikuroi and fusarium solani solution

| Product Information | | | | |
|---|--|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9636 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 0.025 g in 1 mL | | |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.025 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9636-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9636-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| FUSARIUM MIX | | | |
|---|---|---------------------------|----------------|
| gibberella fujikuroi and fusarium solani solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9637 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------------------|------------------------|
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 20000 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9637-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9637-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MIX

gibberella fujikuroi and fusarium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9638 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------------------|------------------------|
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 10000 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 10000 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|-----------------|----------|
|-----------------|----------|

| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
|--|--|--|----------------------|--------------------|
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9638-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| FUSARIUM MIX | | | | |
|---|---|---|----------------------|--------------------|
| gibberella fujikuroi and fusarium solani solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9639 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 0.0005 g in 1 mL | | |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.0005 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9639-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MIX

gibberella fujikuroi and fusarium solani solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9641 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------------|------------------|
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 0.0125 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9641-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9641-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9641-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR MIX

mucor circinelloides f. lusitanicus and mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9648 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|-----------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.025 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9648-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9648-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9651 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 1250 [PNU] in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 1250 [PNU] in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 1250 [PNU] in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 1250 [PNU] in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 1250 [PNU] in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 1250 [PNU] in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 1250 [PNU] in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 1250 [PNU] in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 1250 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 1250 [PNU] in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 1250 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 1250 [PNU] in 1 mL |
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 1250 [PNU] in 1 mL |
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 1250 [PNU] in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 1250 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 1250 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9651-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9651-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 2

aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9610 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 8000 [PNU] in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 8000 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 8000 [PNU] in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 8000 [PNU] in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 8000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------|--|----------------------|--------------------|
| | NDC:22840- | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a | | |

| | | | | |
|------------------------------|---|--|-----------------------------|---------------------------|
| 1 | NDC:22840-9610-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | | 09/15/1981 | |

MOLD MIX 1
alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9603 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| | | | |
|--|---|--|-----------------|
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.01 g in 1 mL |
| | ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.01 g in 1 mL |
| | CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.01 g in 1 mL |
| | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.01 g in 1 mL |
| | ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.01 g in 1 mL |

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

| | | | | |
|------------------|------------------|--|-----------------------------|---------------------------|
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9603-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9678 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 2500 [PNU] in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 2500 [PNU] in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 2500 [PNU] in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 2500 [PNU] in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 2500 [PNU] in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 2500 [PNU] in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 2500 [PNU] in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 2500 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 2500 [PNU] in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 2500 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 2500 [PNU] in 1 mL |
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 2500 [PNU] in 1 mL |
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 2500 [PNU] in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 2500 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 2500 [PNU] in 1 mL |
| PLEOSPOA BETAE (UNII: V58BK047ES) (PLEOSPOA BETAE - UNII:V58BK047ES) | PLEOSPOA BETAE | 2500 [PNU] in 1 mL |

UNII:V58BK047ES)

FLEUSPORA DETAE

in 1 mL

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9678-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS MIX

rhizopus stolonifer and rhizopus oryzae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9680 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|---------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 10000 [PNU] in 1 mL |
| RHIZOPUS ARRHZIZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZIZUS - UNII:8476849N1Y) | RHIZOPUS ARRHZIZUS | 10000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9680-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS MIX

rhizopus stolonifer and rhizopus oryzae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9669 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0125 g in 1 mL |
| RHIZOPUS ARRHZIZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZIZUS - UNII:8476849N1Y) | RHIZOPUS ARRHZIZUS | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9669-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9669-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9669-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1684 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------|--------------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 1000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1684-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 2

aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus

stolonifer solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9612 |
| Route of Administration | INTRADERMAL, 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 | 0.0002 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.0002 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0002 g in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.0002 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.0002 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9612-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

AHH MOLD MIX

alternaria alternata, bipolaris sorokiniana and cladosporium sphaerospermum solution

Product Information

| | | | |
|---------------------|-----------------------------|------------------|------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code | NDC:22840- |
|---------------------|-----------------------------|------------------|------------|

| | | | | |
|--|---|---|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | (Source) | 9622 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.00033 g in 1 mL | |
| | COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.00033 g in 1 mL | |
| | ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.00033 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9622-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

AHH MOLD MIX

alternaria alternata, bipolaris sorokiniana and cladosporium sphaerospermum solution

| | | | |
|--|---|-----------------------------|-------------------------|
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9673 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 13333.333 [PNU] in 1 mL |

| | | |
|---|----------------------|-------------------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 13333.333 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 13333.333 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9673-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

AHH MOLD MIX

alternaria alternata, bipolaris sorokiniana and cladosporium sphaerospermum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9623 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|------------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0166 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0166 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0166 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

GLYCERIN (UNII: PDC6A3C0OX)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9623-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9623-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9623-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

AHH MOLD MIX

alternaria alternata, bipolaris sorokiniana and cladospodium sphaerospermum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9621 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|------------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0166 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0166 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0166 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9621-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9621-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA/HORMODENDRUM MIX

alternaria alternata and aspergillus fumigatus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9624 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-----------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.025 g in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.025 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:22840-9624-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DRECHSLERA SPICIFERA

drechslera spicifera solution

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1662 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | |
|--|-----------------------|---------------------|--|
| Ingredient Name | Basis of Strength | Strength | |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 20000 [PNU] in 1 mL | |

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

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1662-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA HORMODENDRUM MIX

alternaria alternata and aspergillus fumigatus solution

| Product Information | | | |
|---------------------|-----------------------------|-----------|------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code | NDC:22840- |

| | | | | |
|--|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | (Source) | 9626 | |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.025 g in 1 mL | |
| | ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.025 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | | Strength | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| | GLYCERIN (UNII: PDC6A3C0OX) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9626-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9626-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9626-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

HELMINTHOSPORIUM SOLANI

helminthosporium solani solution

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1690 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | | |
|--|--|--|----------------------|--------------------|
| Ingredient Name | | Basis of Strength | Strength | |
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | | HELMINTHOSPORIUM SOLANI | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1690-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

MUCOR PLUMBEUS

mucor plumbeus solution

| Product Information | | | |
|--|---|--------------------|---------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1696 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | | MUCOR PLUMBEUS | 40000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1696-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR PLUMBEUS

mucor plumbeus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5626 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5626-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5626-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5626-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIDULANS

aspergillus nidulans solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1622 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|---------------|
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1622-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1622-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIDULANS

aspergillus nidulans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2660 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2660-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

AUREOBASIDIUM PULLULANS

aureobasidium pullulans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1630 |
| Route of Administration | INTRADERMAL, 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 | 0.001 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1630-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| BIPOLARIS SOROKINIANA | | | |
|--|---|----------------------|-------------------|
| bipolaris sorokiniana solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1635 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | | COCHLIOBOLUS SATIVUS | 0.05 g in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1635-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1635-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1636 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.01 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1636-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5608 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|----------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5608-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5608-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5608-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FUMIGATUS

aspergillus fumigatus solution

| Product Information | | | | |
|--|--|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1617 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 40000 [PNU] in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1617-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1617-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| ASPERGILLUS FUMIGATUS | | | |
|---------------------------------|---|--------------------|----------------|
| aspergillus fumigatus solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1618 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |

| Ingredient Name | | Basis of Strength | Strength | |
|--|--|--|----------------------|--------------------|
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | | ASPERGILLUS FUMIGATUS | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1618-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1618-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9650 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | |
|--|----------------------|---------------------|
| Ingredient Name | Basis of Strength | Strength |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.0000625 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.0000625 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0000625 g in 1 mL |

| | | |
|---|--|---------------------|
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 0.0000625 g in 1 mL |
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.0000625 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0000625 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0000625 g in 1 mL |
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.0000625 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0000625 g in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.0000625 g in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.0000625 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0000625 g in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.0000625 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.0000625 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.0000625 g in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.0000625 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9650-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

| Product Information | | | | |
|---|--|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9654 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.025 g in 1 mL | | |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 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 | | |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 0.025 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9654-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9654-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| GRASS SMUT MIX | | | |
|---|-----------------------------|---------------------------|----------------|
| ustilago cynodontis and sporisorium cruentum solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9671 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, | | |

Route of Administration INTRADERMAL

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| SPORISORIUM CRUENTUM (UNII: GQM6LVU5V8) (SPORISORIUM CRUENTUM - UNII:GQM6LVU5V8) | SPORISORIUM CRUENTUM | 0.05 g in 1 mL |
| USTILAGO CYNODONTIS (UNII: 0V3J4YEX2W) (USTILAGO CYNODONTIS - UNII:0V3J4YEX2W) | USTILAGO CYNODONTIS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9671-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9671-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9671-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GRAIN SMUT MIX

ustilago maydis, ustilago tritici, ustilago nuda and ustilago avenae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9675 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of | Strength |
|-----------------|----------|----------|
|-----------------|----------|----------|

| Ingredient Name | Strength | Strength |
|---|------------------|------------------|
| USTILAGO AVENAE (UNII: YIH315U1TU) (USTILAGO AVENAE - UNII:YIH315U1TU) | USTILAGO AVENAE | 0.0125 g in 1 mL |
| USTILAGO MAYDIS (UNII: 4K7Z7K7SWG) (USTILAGO MAYDIS - UNII:4K7Z7K7SWG) | USTILAGO MAYDIS | 0.0125 g in 1 mL |
| USTILAGO TRITICI (UNII: BV82OL2IZ8) (USTILAGO TRITICI - UNII:BV82OL2IZ8) | USTILAGO TRITICI | 0.0125 g in 1 mL |
| USTILAGO NUDA (UNII: 9Y53ZS6I82) (USTILAGO NUDA - UNII:9Y53ZS6I82) | USTILAGO NUDA | 0.0125 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9675-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1641 |
| Route of Administration | INTRADERMAL, 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 | |
|---------------------------------------|--|--|----------------------|--------------------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1641-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1641-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| CORN SMUT | | | | |
|--|---|---------------------|----------------------|--------------------|
| ustilago maydis solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2664 | |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | Basis of Strength | Strength | |
| USTILAGO MAYDIS (UNII: 4K7Z7K7SWG) (USTILAGO MAYDIS - UNII:4K7Z7K7SWG) | | USTILAGO MAYDIS | 0.1 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |

| | | | | |
|----------|------------------|--|--|--|
| 1 | NDC:22840-2664-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
|----------|------------------|--|--|--|

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1643 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1643-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR MIX

mucor circinelloides f. lusitanicus and mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9646 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|------------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.0005 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.0005 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:22840-9646-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM CHRYSOGENUM (NOTATUM)

penicillium chrysogenum (notatum) solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5631 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--|-------------------|
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5631-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5631-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5631-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHODERMA HARZIANUM

trichoderma harzianum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2640 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-------------------|
| TRICHODERMA HARZIANUM (UNII: CA33Q4013Q) (TRICHODERMA HARZIANUM - UNII:CA33Q4013Q) | TRICHODERMA HARZIANUM | 0.05 g in 1 mL |

Inactive Ingredients

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

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2640-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2640-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum solution

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9616 |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | |
|--|--|--------------------|
| Ingredient Name | Basis of Strength | Strength |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 5000 [PNU] in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 5000 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 5000 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 5000 [PNU] in 1 mL |

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9616-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9616-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9617 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|-----------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 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 |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 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 |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9617-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9617-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA/HORMODENDRUM MIX

alternaria alternata and aspergillus fumigatus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9674 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 10000 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 10000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9674-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9674-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ACREMONIUM STRICTUM

acremonium strictum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5601 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|----------------|
| SAROCLADIUM STRICTUM (UNII: 3F36V0451W) (SAROCLADIUM STRICTUM - UNII:3F36V0451W) | SAROCLADIUM STRICTUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5601-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5601-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 3

alternaria alternata, aspergillus niger, cladosporium sphaerospermum and penicillium notatum

solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9619 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|------------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0125 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.0125 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0125 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9619-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MONILIA MIX

candida albicans and neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9642 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, | | |

Route of Administration SUBCUTANEOUS

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 10000 [PNU] in 1 mL |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 10000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9642-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9642-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEW STOCK FUNGI MIX

acremonium strictum, alternaria alternata, aspergillus niger, aureobasidium pullulans, bipolaris sorokiniana, botrytis cinerea, candida albicans, chaetomium globosum, cladosporium sphaerospermum, epicoccum nigrum, gibberella fujikuroi, mucor plumbeus, penicillium notatum, phoma betae, rhizopus stolonifer and trichophyton mentagrophytes solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9653 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.003125 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.003125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.003125 g in 1 mL |
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.003125 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.003125 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.003125 g in 1 mL |
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.003125 g in 1 mL |
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.003125 g in 1 mL |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 0.003125 g in 1 mL |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.003125 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.003125 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.003125 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.003125 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.003125 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.003125 g in 1 mL |
| ACREMONIUM STRICTUM (UNII: 3F36V0451W) (ACREMONIUM STRICTUM - UNII:3F36V0451W) | ACREMONIUM STRICTUM | 0.003125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9653-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9653-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9653-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GRAIN SMUT MIX

ustilago maydis, ustilago tritici, ustilago nuda and ustilago avenae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9670 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-------------------|
| USTILAGO NUDA (UNII: 9Y53ZS6I82) (USTILAGO NUDA - UNII:9Y53ZS6I82) | USTILAGO NUDA | 0.00625 g in 1 mL |
| USTILAGO AVENAE (UNII: YIH315U1TU) (USTILAGO AVENAE - UNII:YIH315U1TU) | USTILAGO AVENAE | 0.00625 g in 1 mL |
| USTILAGO MAYDIS (UNII: 4K7Z7K7SWG) (USTILAGO MAYDIS - UNII:4K7Z7K7SWG) | USTILAGO MAYDIS | 0.00625 g in 1 mL |
| USTILAGO TRITICI (UNII: BV82OL2IZ8) (USTILAGO TRITICI - UNII:BV82OL2IZ8) | USTILAGO TRITICI | 0.00625 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9670-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9670-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9670-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 4 | NDC:22840-9670-3 | 30 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEUROSPORA INTERMEDIA

neurospora intermedia solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2600 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------|---------------------|
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2600-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2600-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEUROSPORA INTERMEDIA

neurospora intermedia solution

| Product Information | | | | |
|---------------------------------|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5627 | |
| Route of Administration | SUBCUTANEOUS, PERCUTANEOUS, INTRADERMAL | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | GLYCERIN (UNII: PDC6A3C00X) | | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5627-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5627-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5627-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| PAECILOMYCES VARIOTII | | | |
|--------------------------------|---|---------------------------|----------------|
| paecilomyces variotii solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2605 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|------------------|
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | PAECILOMYCES VARIOTII | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2605-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2605-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM DIGITATUM

penicillium digitatum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5630 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

| PHENOL (UNII: 339NCG44TV) | | | | |
|---|--|--|----------------------|--------------------|
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| GLYCERIN (UNII: PDC6A3C0OX) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5630-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5630-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

| Product Information | | | |
|---|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9630 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | Basis of Strength | Strength | |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.01 g in 1 mL | |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.01 g in 1 mL | |
| EUROTIVAM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIVAM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIVAM AMSTELODAMI | 0.01 g in 1 mL | |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.01 g in 1 mL | |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.01 g in 1 mL | |
| Inactive Ingredients | | | |
| Ingredient Name | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9630-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DEMATIACEAE MIX

alternaria alternata, aureobasidium pullulans, bipolaris sorokiniana, cladosporium herbarum, curvularia spicifera and helminthosporium solani solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9635 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|------------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.0042 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 0.0042 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0042 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0042 g in 1 mL |
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | CLADOSPORIUM HERBARUM | 0.0042 g in 1 mL |
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 0.0042 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9635-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9635-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIDULANS

aspergillus nidulans solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1623 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|-----------------|
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1623-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9629 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|--------------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 4000 [PNU] in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 4000 [PNU] in 1 mL |
| EUROTIIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIIUM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIIUM AMSTELODAMI | 4000 [PNU] in 1 mL |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 4000 [PNU] in 1 mL |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 4000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9629-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9629-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPICOCCUM NIGRUM

epicoccum nigrum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2665 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-2665-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

HELMINTHOSPORIUM SOLANI

helminthosporium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2668 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------|---------------------|
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2668-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2668-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

HELMINTHOSPORIUM SOLANI

helminthosporium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5622 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------|----------------|
| HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|-----------------|----------|
|-----------------|----------|

| | |
|--|--|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5622-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5622-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5622-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR PLUMBEUS

mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1694 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|---------------|
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
|---|-----------|---------------------|----------------------|--------------------|

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1694-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1694-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS ORYZAE

rhizopus oryzae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5633 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| RHIZOPUS ARRHIUS (UNII: 8476849N1Y) (RHIZOPUS ARRHIUS - UNII:8476849N1Y) | RHIZOPUS ARRHIUS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5633-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5633-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

STEMPHYLIUM SOLANI

stemphylium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2637 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|---------------------|
| STEMPHYLIUM SOLANI (UNII: 1IEK4UDP5M) (STEMPHYLIUM SOLANI - UNII:1IEK4UDP5M) | STEMPHYLIUM SOLANI | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2637-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2637-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

STEMPHYLIUM SOLANI

stemphylium solani solution

Product Information

| | | | | |
|--|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2636 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | STEMPHYLIUM SOLANI (UNII: 1IEK4UDP5M) (STEMPHYLIUM SOLANI - UNII:1IEK4UDP5M) | STEMPHYLIUM SOLANI | 20000 [PNU] in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | | Strength | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2636-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|---|---------------------------|-----------------|
| STEMPHYLIUM SOLANI | | | |
| stemphylium solani solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5637 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | STEMPHYLIUM SOLANI (UNII: 1IEK4UDP5M) (STEMPHYLIUM SOLANI - UNII:1IEK4UDP5M) | STEMPHYLIUM SOLANI | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5637-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5637-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5637-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

SACCHAROMYCES CEREVISIAE

saccharomyces cerevisiae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2678 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|---------------------|
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES CEREVISIAE - UNII:978D8U419H) | SACCHAROMYCES CEREVISIAE | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2678-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

SACCHAROMYCES CEREVISIAE

saccharomyces cerevisiae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5636 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------------|----------------|
| SACCHAROMYCES CEREVISIAE (UNII: 978D8U419H) (SACCHAROMYCES CEREVISIAE - UNII:978D8U419H) | SACCHAROMYCES CEREVISIAE | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5636-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5636-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5636-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

STEMPHYLIUM SOLANI

stemphylium solani solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2635 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|----------------|
| STEMPHYLIUM SOLANI (UNII: 1IEK4UDP5M) (STEMPHYLIUM SOLANI - UNII:1IEK4UDP5M) | STEMPHYLIUM SOLANI | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2635-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2635-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

STEMPHYLIUM SOLANI

stemphylium solani solution

| Product Information | | | | |
|--|--|---|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2638 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| STEMPHYLIUM SOLANI (UNII: 1IEK4UDP5M) (STEMPHYLIUM SOLANI - UNII:1IEK4UDP5M) | STEMPHYLIUM SOLANI | 0.001 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2638-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| TRICHOPHYTON MENTAGROPHYTES | | | |
|--|---|---------------------|----------------|
| trichophyton mentagrophytes solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2649 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | Basis of Strength | Strength | |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | TRICHOPHYTON MENTAGROPHYTES | 40000 [PNU] in 1 mL | |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2649-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. LUSITANICUS

mucor circinelloides f. lusitanicus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2659 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|-----------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start | Marketing End |
|---|-----------|---------------------|-----------------|---------------|
|---|-----------|---------------------|-----------------|---------------|

| # | Item Code | Package Description | Date | Date |
|---|------------------|---|------|------|
| 1 | NDC:22840-2659-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

EPICOCCUM NIGRUM

epicoccum nigrum solution

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5616 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | |
|--|-------------------|-----------------|
| Ingredient Name | Basis of Strength | Strength |
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.025 g in 1 mL |

| Inactive Ingredients | |
|---------------------------------------|----------|
| Ingredient Name | Strength |
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5616-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5616-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5616-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|---------------------------------|-----------------|---------------|
| Marketing | Application Number or Monograph | Marketing Start | Marketing End |
| | | | |

| Category | Citation | Date | Date |
|----------|-----------|------------|------|
| BLA | BLA101833 | 09/15/1981 | |

EPIDERMOPHYTON FLOCCOSUM

epidermophyton floccosum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5617 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|----------------|
| EPIDERMOPHYTON FLOCCOSUM (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S) | EPIDERMOPHYTON FLOCCOSUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5617-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5617-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5617-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 4 | NDC:22840-5617-3 | 30 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MICROSPORUM CANIS

microsporium canis solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5623 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| MICROSPORUM CANIS (UNII: N4F4RQ7BY7) (MICROSPORUM CANIS - UNII:N4F4RQ7BY7) | MICROSPORUM CANIS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5623-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5623-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5623-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BERMUDA GRASS SMUT

ustilago cynodontis solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5648 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|-----------------|
| USTILAGO CYNODONTIS (UNII: 0V3J4YEX2W) (USTILAGO CYNODONTIS - UNII:0V3J4YEX2W) | USTILAGO CYNODONTIS | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5648-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5648-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5648-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MONILIFORME

fusarium moniliforme solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1671 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1671-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1671-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS AMSTELODAMI

aspergillus amstelodami solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1608 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-----------------|
| EUROTIVAM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIVAM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIVAM AMSTELODAMI | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1608-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIDULANS

aspergillus nidulans solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5605 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|----------------|
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5605-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5605-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIGER

aspergillus niger solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1626 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|------------------------------|----------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1626-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIGER

aspergillus niger solution

Product Information

| | | | |
|--------------|-----------------------------|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5606 |
|--------------|-----------------------------|--------------------|----------------|

| | |
|--------------------------------|---|
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS |
|--------------------------------|---|

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|----------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5606-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5606-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5606-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1644 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------|-------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - | CANDIDA ALBICANS | 40000 [PNU] |

| | | |
|------------------|------------------|---------|
| UNII:4D7G21HDBC) | CANDIDA ALBICANS | in 1 mL |
|------------------|------------------|---------|

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1644-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1644-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPICOCCUM NIGRUM

epicoccum nigrum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1666 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-----------------|
| EPICOCCUM NIGRUM (UNII: 87U156LEN7) (EPICOCCUM NIGRUM - UNII:87U156LEN7) | EPICOCCUM NIGRUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1666-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIGER

aspergillus niger solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1628 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|------------------------------|-----------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1628-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1633 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1633-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1633-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DRECHSLERA SPICIFERA

drechslera spicifera solution

Product Information

| | | | |
|--------------------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5615 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, | | |

Route of Administration SUBCUTANEOUS

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-----------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5615-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5615-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5615-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM SOLANI

fusarium solani solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1676 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|-----------------|
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 0.025 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1676-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PHOMA BETAE

phoma betae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2618 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-----------------|
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
|---|-----------|---------------------|----------------------|--------------------|

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2618-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| FUSARIUM SOLANI | | | | |
|---|---|--|----------------------|--------------------|
| fusarium solani solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5619 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | Basis of Strength | Strength | |
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII: 7TLR512M4A) | | HAEMATONECTRIA HAEMATOCOCCA | 0.025 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| GLYCERIN (UNII: PDC6A3C00X) | | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5619-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5619-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5619-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing | Application Number or Monograph | Marketing Start | Marketing End | |

| Category | Citation | Date | Date |
|----------|-----------|------------|------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1681 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|---------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1681-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1681-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

Item Code NDC:22840

| | | | | |
|--|---|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1682 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | | Strength | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1682-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|---|---------------------------|---------------------|
| HELMINTHOSPORIUM SOLANI | | | |
| helminthosporium solani solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1691 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | HELMINTHOSPORIUM SOLANI (UNII: U6Z259H815) (HELMINTHOSPORIUM SOLANI - UNII:U6Z259H815) | HELMINTHOSPORIUM SOLANI | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1691-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1691-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BOTRYTIS CINEREA

botrytis cinerea solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2662 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
|---|-----------|---------------------|----------------------|--------------------|

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2662-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| CLADOSPORIUM HERBARUM | | | | |
|--|--|--|----------------------|--------------------|
| cladosporium herbarum solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1653 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | Basis of Strength | Strength | |
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | | CLADOSPORIUM HERBARUM | 0.05 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1653-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1653-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

CLADOSPORIUM HERBARUM

cladosporium herbarum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1654 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|---------------------|
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | CLADOSPORIUM HERBARUM | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-1654-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1654-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM HERBARUM

cladosporium herbarum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1656 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-----------------|
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | CLADOSPORIUM HERBARUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1656-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA ALTERNATA

alternaria alternata solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1605 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 10000 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|-----------------|----------|
|-----------------|----------|

| | |
|--|--|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| PHENOL (UNII: 339NCG44TV) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1605-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS AMSTELODAMI

aspergillus amstelodami solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1609 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|--------------------|
| EUROTIVAM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIVAM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIVAM AMSTELODAMI | 1000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1609-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BOTRYTIS CINEREA

botrytis cinerea solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1639 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-----------------|
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1639-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MONILIFORME

fusarium moniliforme solution

| Product Information | | | | |
|---------------------------------|--|--|-----------------------------|---------------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1670 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.1 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1670-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1670-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| ASPERGILLUS FUMIGATUS | | | | |
|---------------------------------|---|---------------------------|-----------------|--|
| aspergillus fumigatus solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1619 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |

| Ingredient Name | | Strength | Strength | |
|--|--|---|----------------------|--------------------|
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | | ASPERGILLUS FUMIGATUS | 0.002 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1619-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| PENICILLIUM CHRYSOGENUM (NOTATUM) | | | |
|--|---|--|---------------------|
| penicillium chrysogenum (notatum) solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2615 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 40000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2615-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PHOMA BETAE

phoma betae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5632 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------|-----------------|
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5632-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5632-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5632-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS ORYZAE

rhizopus oryzae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2619 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|------------------|
| RHIZOPUS ARRHIZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHIZUS - UNII:8476849N1Y) | RHIZOPUS ARRHIZUS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2619-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9627 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|-----------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.02 g in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.02 g in 1 mL |
| EUROTIIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIIUM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIIUM AMSTELODAMI | 0.02 g in 1 mL |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.02 g in 1 mL |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | ASPERGILLUS NIDULANS | 0.02 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-9627-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9627-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

ALTERNARIA ALTERNATA

alternaria alternata solution

Product Information

| | | | |
|---------------------|-----------------------------|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1604 |
|---------------------|-----------------------------|---------------------------|----------------|

| | |
|--------------------------------|--|
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS |
|--------------------------------|--|

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|-----------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1604-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ACREMONIUM STRICTUM

acremonium strictum solution

Product Information

| | | | |
|--------------------------------|--|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1606 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|-----------------|
| SAROCLADIUM STRICTUM (UNII: 3F36V0451W) (SAROCLADIUM STRICTUM - UNII:3F36V0451W) | SAROCLADIUM STRICTUM | 0.001 g in 1 mL |

Inactive Ingredients

| Ingredient Name | | | Strength | |
|---------------------------------------|--|---|----------------------|--------------------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1606-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

RHIZOPUS STOLONIFER

rhizopus stolonifer solution

| Product Information | | | | |
|--|---|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2623 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.05 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2623-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| | | | | |
|---|------------------|--|--|--|
| 2 | NDC:22840-2623-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
|---|------------------|--|--|--|

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. LUSITANICUS

mucor circinelloides f. lusitanicus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2671 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|---------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2671-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOTHECIUM ROSEUM

trichothecium roseum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2654 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | TRICHOTHECIUM ROSEUM | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2654-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2654-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

TRICHOTHECIUM ROSEUM

trichothecium roseum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2656 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | | |
|--|--|----------------------|--------------------|
| Ingredient Name | | Basis of Strength | Strength |
| TRICHOTHECIUM ROSEUM (UNII: TGO054E31O) (TRICHOTHECIUM ROSEUM - UNII:TGO054E31O) | | TRICHOTHECIUM ROSEUM | 1000 [PNU] in 1 mL |

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

| Packaging | | | | |
|-----------|------------------|---|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2656-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| MYCOGONE PERNICIOSA | | | |
|------------------------------|--|--|--|
| mycogone perniciosa solution | | | |

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5642 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | |
|--|-----------------------|----------------|
| Ingredient Name | Basis of Strength | Strength |
| HYPOMYCES PERNICIOSUS (UNII: 6K41G30A6U) (HYPOMYCES PERNICIOSUS - UNII:6K41G30A6U) | HYPOMYCES PERNICIOSUS | 0.05 g in 1 mL |

| Inactive Ingredients | |
|---------------------------------------|----------|
| Ingredient Name | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |

SODIUM CHLORIDE (UNII: 451W471Q8X)

GLYCERIN (UNII: PDC6A3C0OX)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5642-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5642-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5642-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 2

aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9609 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 2000 [PNU] in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 2000 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 2000 [PNU] in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 2000 [PNU] in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 2000 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9609-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9609-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1632 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1632-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840- | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a | | |

1632-4 Combination Product

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9656 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 5000 [PNU] in 1 mL |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 5000 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 5000 [PNU] in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 5000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9656-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing | Application Number or Monograph | Marketing Start | Marketing End |
|-----------|---------------------------------|-----------------|---------------|
|-----------|---------------------------------|-----------------|---------------|

| Category | Citation | Date | Date |
|----------|-----------|------------|------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9655 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|---------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 10000 [PNU] in 1 mL |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 10000 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 10000 [PNU] in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 10000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9655-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9655-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9657 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.0125 g in 1 mL |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 0.0125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0125 g in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 0.0125 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9657-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9657-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9679 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 2500 [PNU] in 1 mL |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 2500 [PNU] in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 2500 [PNU] in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 2500 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9679-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPIDERMOPHYTON FLOCCOSUM

epidermophyton floccosum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1667 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|----------------|
| EPIDERMOPHYTON FLOCCOSUM (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S) | EPIDERMOPHYTON FLOCCOSUM | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1667-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1667-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

EPIDERMOPHYTON FLOCCOSUM

epidermophyton floccosum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1669 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|--------------------|
| EPIDERMOPHYTON FLOCCOSUM (UNII: 6JR6JTN25S) (EPIDERMOPHYTON FLOCCOSUM - UNII:6JR6JTN25S) | EPIDERMOPHYTON FLOCCOSUM | 1000 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

| | |
|---|--|
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1669-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM SPHAEROSPERMUM

cladosporium sphaerospermum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1659 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|-----------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1659-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

DRECHSLERA SPICIFERA

drechslera spicifera solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1661 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1661-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1661-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PAECILOMYCES VARIOTII

paecilomyces variotii solution

| Product Information | | | | |
|---------------------------------|--|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2606 | |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | PAECILOMYCES VARIOTII | 20000 [PNU] in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2606-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2606-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| TRICHODERMA HARZIANUM | | | |
|---------------------------------|---|--------------------|----------------|
| trichoderma harzianum solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5638 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of | Strength |

| Ingredient Name | | Strength | Strength | |
|--|--|--|----------------------|--------------------|
| TRICHODERMA HARZIANUM (UNII: CA33Q4013Q) (TRICHODERMA HARZIANUM - UNII:CA33Q4013Q) | | TRICHODERMA HARZIANUM | 0.025 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| GLYCERIN (UNII: PDC6A3C0OX) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5638-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5638-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5638-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

TRICHOPHYTON MENTAGROPHYTES

trichophyton mentagrophytes solution

| Product Information | | | |
|--|---|-----------------------------|---------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2645 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| TRICHOPHYTON MENTAGROPHYTES (UNII: 199I7J3JIV) (TRICHOPHYTON MENTAGROPHYTES - UNII:199I7J3JIV) | | TRICHOPHYTON MENTAGROPHYTES | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2645-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2645-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM MONILIFORME

fusarium moniliforme solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1673 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|----------------------|---------------------|
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1673-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

FUSARIUM SOLANI

fusarium solani solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1678 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-----------------------------|-----------------|
| HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1678-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 1

alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

| Product Information | | | | |
|---|--|--|----------------------|--------------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9601 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | Basis of Strength | Strength | | |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 4000 [PNU] in 1 mL | | |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 4000 [PNU] in 1 mL | | |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 4000 [PNU] in 1 mL | | |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 4000 [PNU] in 1 mL | | |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 4000 [PNU] in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | Strength | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9601-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9601-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| FUSARIUM MONILIFORME | | | |
|-------------------------------|-----------------------------|---------------------------|----------------|
| fusarium moniliforme solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1674 |

| | | | | |
|--|---|---|-----------------------------|---------------------------|
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 0.001 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1674-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| | | | |
|--|---|-----------------------------|---------------------|
| FUSARIUM SOLANI | | | |
| fusarium solani solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1677 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| | Ingredient Name | Basis of Strength | Strength |
| | HAEMATONECTRIA HAEMATOCOCCA (UNII: 7TLR512M4A) (HAEMATONECTRIA HAEMATOCOCCA - UNII:7TLR512M4A) | HAEMATONECTRIA HAEMATOCOCCA | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | |

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1677-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GEOTRICHUM CANDIDUM

geotrichum candidum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1680 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|-----------------|
| GEOTRICHUM CANDIDUM (UNII: 5964J742O8) (GEOTRICHUM CANDIDUM - UNII:5964J742O8) | GEOTRICHUM CANDIDUM | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1680-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5621 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------|----------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5621-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5621-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5621-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR PLUMBEUS

mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1698 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|---|-----------------------------|---------------------------|
| 1 | NDC:22840-1698-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

NEUROSPORA INTERMEDIA

neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2601 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | | | |
|--|--|--|-----------------------|---------------------|
| Ingredient Name | | | Basis of Strength | Strength |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | | | NEUROSPORA INTERMEDIA | 20000 [PNU] in 1 mL |
| Inactive Ingredients | | | | |
| Ingredient Name | | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2601-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | | 09/15/1981 | |

| PAECILOMYCES VARIOTII | | | |
|--|---|-----------------------|---------------------|
| paecilomyces variotii solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2609 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | | PAECILOMYCES VARIOTII | 40000 [PNU] in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2609-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2609-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PAECILOMYCES VARIOTII

paecilomyces variotii solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5629 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|----------------|
| PAECILOMYCES VARIOTII (UNII: KO7V58BY40) (PAECILOMYCES VARIOTII - UNII:KO7V58BY40) | PAECILOMYCES VARIOTII | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5629-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840- | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a | | |

| | | | | |
|---|------------------|--|--|--|
| 4 | 5629-2 | Combination Product | | |
| 3 | NDC:22840-5629-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM CHRYSOGENUM (NOTATUM)

penicillium chrysogenum (notatum) solution

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2614 |
| Route of Administration | SUBCUTANEOUS, INTRADERMAL, PERCUTANEOUS | | |

| Active Ingredient/Active Moiety | | |
|--|--|---------------|
| Ingredient Name | Basis of Strength | Strength |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.1 g in 1 mL |

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

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2614-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2614-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

PHOMA BETAE

phoma betae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2674 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| PLEOSPORA BETAE (UNII: V58BK047ES) (PLEOSPORA BETAE - UNII:V58BK047ES) | PLEOSPORA BETAE | 0.05 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-2674-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS NIGER

aspergillus niger solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1627 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|------------------------------|---------------------|
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1627-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CORN SMUT

ustilago maydis solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5649 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| USTILAGO MAYDIS (UNII: 4K7Z7K7SWG) (USTILAGO MAYDIS - UNII:4K7Z7K7SWG) | USTILAGO MAYDIS | 0.05 g in 1 mL |

Inactive Ingredients

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

GLYCERIN (UNII: PDC6A3C0OX)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5649-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5649-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5649-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

OAT SMUT

ustilago avenae solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5651 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|-------------------|-----------------|
| USTILAGO AVENAE (UNII: YIH315U1TU) (USTILAGO AVENAE - UNII:YIH315U1TU) | USTILAGO AVENAE | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
|---|-----------|---------------------|----------------------|--------------------|

| | | | | |
|---|------------------|--|--|--|
| 1 | NDC:22840-5651-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5651-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5651-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 1

alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9602 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 8000 [PNU] in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 8000 [PNU] in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 8000 [PNU] in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 8000 [PNU] in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 8000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|-----------|---------------------|----------------------|--------------------|
| | | | | |

| | | | | |
|---|------------------|--|--|--|
| 1 | NDC:22840-9602-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9602-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 1

alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9604 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|------------------|
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0002 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.0002 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.0002 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0002 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.0002 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9604-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 1

alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9606 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|----------------|
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.01 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.01 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.01 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | ASPERGILLUS NIGER VAR. NIGER | 0.01 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 0.01 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9606-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9606-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9606-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MOLD MIX 2

aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9608 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|--------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 4000 [PNU] in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | COCHLIOBOLUS SPICIFER | 4000 [PNU] in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | GIBBERELLA FUJIKUROI | 4000 [PNU] in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | AUREOBASIDIUM PULLULANS VAR. PULLUTANS | 4000 [PNU] in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 4000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9608-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9608-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ACREMONIUM STRICTUM

acremonium strictum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2679 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|----------------------|---------------|
| SAROCLADIUM STRICTUM (UNII: 3F36V0451W) (SAROCLADIUM STRICTUM - UNII:3F36V0451W) | SAROCLADIUM STRICTUM | 0.1 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-2679-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BOTRYTIS CINEREA

botrytis cinerea solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5609 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|-----------------|
| BOTRYTIS CINEREA (UNII: TBW53313S7) (BOTRYTIS CINEREA - UNII:TBW53313S7) | BOTRYTIS CINEREA | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5609-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5609-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5609-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1646 |
| Route of Administration | INTRADERMAL, 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 | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-1646-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

| RHIZOPUS STOLONIFER | | | |
|--|---|---------------------|--------------------|
| rhizopus stolonifer solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2627 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | | RHIZOPUS STOLONIFER | 0.001 g in 1 mL |
| Inactive Ingredients | | | |
| Ingredient Name | | Strength | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| PHENOL (UNII: 339NCG44TV) | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2627-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS STOLONIFER

rhizopus stolonifer solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5634 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|-----------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.025 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5634-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5634-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5634-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHODOTORULA MUCILAGINOSA

rhodotorula mucilaginosa solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2628 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| RHODOTORULA MUCILAGINOSA (UNII: 62TY3X4N9Z) (RHODOTORULA MUCILAGINOSA - UNII:62TY3X4N9Z) | RHODOTORULA MUCILAGINOSA | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-2628-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5610 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5610-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5610-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5610-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM HERBARUM

cladosporium herbarum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5613 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | | Basis of Strength | Strength | |
|--|--|--|----------------------|--------------------|
| CLADOSPORIUM HERBARUM (UNII: O64JF1198) (CLADOSPORIUM HERBARUM - UNII:O64JF1198) | | CLADOSPORIUM HERBARUM | 0.025 g in 1 mL | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| GLYCERIN (UNII: PDC6A3C0OX) | | | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-5613-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5613-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5613-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

MOLD MIX 1

alternaria alternata, aspergillus niger, bipolaris sorokiniana, cladosporium sphaerospermum and penicillium notatum solution

| Product Information | | | |
|--|---|--|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9600 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.02 g in 1 mL |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | | ASPERGILLUS NIGER VAR. NIGER | 0.02 g in 1 mL |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - | | COCHLIOBOLUS SATIVUS | 0.02 g |

| | | |
|---|-----------------------------|-------------------|
| UNII:3LN5B70U4W) | COCCIDIUM BURNII | in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.02 g in 1 mL |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.02 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9600-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9600-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

GLIOCLADIUM VIRIDE

gliocladium viride solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1683 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------|--------------------|
| GLIOCLADIUM VIRIDE (UNII: 8Z8C642TPL) (GLIOCLADIUM VIRIDE - UNII:8Z8C642TPL) | GLIOCLADIUM VIRIDE | 0.001 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1683-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9660 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--|------------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.0125 g in 1 mL |
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 0.0125 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.0125 g in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9660-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9660-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9660-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM MIX

penicillium camemberti, penicillium chrysogenum, penicillium digitatum, penicillium notatum and penicillium roqueforti solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9659 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--|-------------------|
| PENICILLIUM CAMEMBERTI (UNII: T069R9OPVG) (PENICILLIUM CAMEMBERTI - UNII:T069R9OPVG) | PENICILLIUM CAMEMBERTI | 0.00025 g in 1 mL |
| PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM (UNII: 3Y1PE1GCIG) (PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM - UNII:3Y1PE1GCIG) | PENICILLIUM CHRYSOGENUM VAR. CHRYSOGENUM | 0.00025 g in 1 mL |
| PENICILLIUM ROQUEFORTI (UNII: 7ORP6R724L) (PENICILLIUM ROQUEFORTI - UNII:7ORP6R724L) | PENICILLIUM ROQUEFORTI | 0.00025 g in 1 mL |
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.00025 g in 1 mL |

Inactive Ingredients

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

Packaging

| Marketing Start | Marketing End |
|-----------------|---------------|
|-----------------|---------------|

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9659-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| PHYCOMYCETES MIX | | | | |
|--|---|--|----------------------|--------------------|
| mucor circinelloides f. lusitanicus and rhizopus stolonifer solution | | | | |
| Product Information | | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9661 | |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | | |
| Active Ingredient/Active Moiety | | | | |
| | Ingredient Name | Basis of Strength | Strength | |
| | MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.025 g in 1 mL | |
| | RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.025 g in 1 mL | |
| Inactive Ingredients | | | | |
| | Ingredient Name | Strength | | |
| | SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | |
| | PHENOL (UNII: 339NCG44TV) | | | |
| | SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9661-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

PHYCOMYCETES MIX

mucor circinelloides f. lusitanicus and rhizopus stolonifer solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9662 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|------------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.0005 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0005 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9662-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PHYCOMYCETES MIX

mucor circinelloides f. lusitanicus and rhizopus stolonifer solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9664 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------------------------|------------------|
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.0125 g in 1 mL |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0125 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9664-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9664-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-9664-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 4 | NDC:22840-9664-3 | 30 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS MIX

rhizopus stolonifer and rhizopus oryzae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9665 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|-----------------|-------------------|----------|
|-----------------|-------------------|----------|

| | | |
|---|---------------------|-----------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.025 g in 1 mL |
| RHIZOPUS ARRHZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZUS - UNII:8476849N1Y) | RHIZOPUS ARRHZUS | 0.025 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9665-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9665-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS MIX

rhizopus stolonifer and rhizopus oryzae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9666 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|---------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 20000 [PNU] in 1 mL |
| RHIZOPUS ARRHZUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZUS - UNII:8476849N1Y) | RHIZOPUS ARRHZUS | 20000 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|-----------------|----------|
|-----------------|----------|

| | |
|--|--|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9666-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

RHIZOPUS MIX

rhizopus stolonifer and rhizopus oryzae solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9668 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|---------------------|------------------|
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | RHIZOPUS STOLONIFER | 0.0005 g in 1 mL |
| RHIZOPUS ARRHZISUS (UNII: 8476849N1Y) (RHIZOPUS ARRHZISUS - UNII:8476849N1Y) | RHIZOPUS ARRHZISUS | 0.0005 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9668-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FLAVUS

aspergillus flavus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1612 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------|-----------------|
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.001 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-1612-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

| Product Information | | | | |
|--|------------------|--|-----------------------------|---------------------------|
| Product Type | | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2661 |
| Route of Administration | | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | | | COCHLIOBOLUS SATIVUS | 10000 [PNU] in 1 mL |
| Inactive Ingredients | | | | |
| Ingredient Name | | | | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2661-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | | BLA101833 | 09/15/1981 | |

| TRICHODERMA HARZIANUM | | | | |
|---------------------------------|--|---|---------------------------|-----------------|
| trichoderma harzianum solution | | | | |
| Product Information | | | | |
| Product Type | | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2642 |
| Route of Administration | | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | | |
| Ingredient Name | | | Basis of Strength | Strength |

| | | |
|---|-----------------------|-----------------|
| TRICHODERMA HARZIANUM (UNII: CA33Q4013Q) (TRICHODERMA HARZIANUM - UNII:CA33Q4013Q) | TRICHODERMA HARZIANUM | 0.001 g in 1 mL |
|---|-----------------------|-----------------|

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

| Packaging | | | | |
|-----------|------------------|---|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2642-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. LUSITANICUS

mucor circinelloides f. lusitanicus solution

| Product Information | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5625 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | |
|---|-------------------------------------|----------------|
| Ingredient Name | Basis of Strength | Strength |
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.05 g in 1 mL |

| Inactive Ingredients | |
|--|----------|
| Ingredient Name | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5625-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5625-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5625-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CHAETOMIUM GLOBOSUM

chaetomium globosum solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5612 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|---------------------|----------------|
| CHAETOMIUM GLOBOSUM (UNII: 5016WB8B8A) (CHAETOMIUM GLOBOSUM - UNII:5016WB8B8A) | CHAETOMIUM GLOBOSUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5612-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5612-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |
| 3 | NDC:22840- | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a | | |

| 5612-2 | Combination Product | | |
|------------------------------|--|----------------------|--------------------|
| Marketing Information | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM SPHAEROSPERMUM

cladosporium sphaerospermum solution

| Product Information | | | |
|----------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2663 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | |
|--|-----------------------------|---------------|--|
| Ingredient Name | Basis of Strength | Strength | |
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.1 g in 1 mL | |

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

| Packaging | | | | |
|------------------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2663-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|------------------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM SPHAEROSPERMUM

cladosporium sphaerospermum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5614 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------------|-----------------|
| CLADOSPORIUM SPHAEROSPERMUM (UNII: P87YCA1U8R) (CLADOSPORIUM SPHAEROSPERMUM - UNII:P87YCA1U8R) | CLADOSPORIUM SPHAEROSPERMUM | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|-----------------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-5614-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5614-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5614-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR CIRCINELLOIDES F. CIRCINELLOIDES

mucor circinelloides f. circinelloides solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2669 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | |
|--|--|--|------------------|
| Ingredient Name | | Basis of Strength | Strength |
| MUCOR CIRCINELLOIDES F. CIRCINELLOIDES (UNII: 48Z8OUT98B) (MUCOR CIRCINELLOIDES F. CIRCINELLOIDES - UNII:48Z8OUT98B) | | MUCOR CIRCINELLOIDES F. CIRCINELLOIDES | 0.1 g in 1 mL |

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

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-2669-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| ALTERNARIA/HORMODENDRUM MIX | | | |
|---|---|--------------------|----------------|
| alternaria alternata and aspergillus fumigatus solution | | | |
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9625 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | |
|--|-----------------------|---------------------|
| Ingredient Name | Basis of Strength | Strength |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | ASPERGILLUS FUMIGATUS | 0.0005 g in 1 mL |
| ALTERNARIA ALTERNATA (UNII: 52B29REC7H) (ALTERNARIA ALTERNATA - UNII:52B29REC7H) | ALTERNARIA ALTERNATA | 0.0005 g in 1 mL |

| Inactive Ingredients | |
|---------------------------------------|----------|
| Ingredient Name | Strength |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9625-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

MONILIA MIX

candida albicans and neurospora intermedia solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9643 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|-------------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 500 [PNU] in 1 mL |
| NEUROSPORA INTERMEDIA (UNII: 2072U60DUI) (NEUROSPORA INTERMEDIA - UNII:2072U60DUI) | NEUROSPORA INTERMEDIA | 500 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:22840-9643-1 | 5 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS FLAVUS

aspergillus flavus solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5603 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|--|--------------------|---------------------|
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | ASPERGILLUS FLAVUS | 0.025 [PNU] in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|---------------------------------------|----------|
| GLYCERIN (UNII: PDC6A3C0OX) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5603-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5603-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5603-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

BIPOLARIS SOROKINIANA

bipolaris sorokiniana solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1634 |
| Route of Administration | PERCUTANEOUS, INTRADERMAL, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|---------------------|
| COCHLIOBOLUS SATIVUS (UNII: 3LN5B70U4W) (COCHLIOBOLUS SATIVUS - UNII:3LN5B70U4W) | COCHLIOBOLUS SATIVUS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-1634-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-1634-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

CLADOSPORIUM HERBARUM

cladosporium herbarum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1657 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|---------------------|
| CLADOSPORIUM HERBARUM (UNII: O64JF11198) (CLADOSPORIUM HERBARUM - UNII:O64JF11198) | CLADOSPORIUM HERBARUM | 40000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1657-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1640 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|----------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 0.01 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |

PHENOL (UNII: 339NCG44TV)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1640-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

CANDIDA ALBICANS

candida albicans solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-1642 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-------------------|---------------------|
| CANDIDA ALBICANS (UNII: 4D7G21HDBC) (CANDIDA ALBICANS - UNII:4D7G21HDBC) | CANDIDA ALBICANS | 20000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-1642-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

ASPERGILLUS AMSTELODAMI

aspergillus amstelodami solution

Product Information

| | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-5602 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|-----------------------|----------------|
| EUROTIIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIIUM AMSTELODAMI - UNII:D932NLL87Z) | EUROTIIUM AMSTELODAMI | 0.05 g in 1 mL |

Inactive Ingredients

| Ingredient Name | Strength |
|--|----------|
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | |
| PHENOL (UNII: 339NCG44TV) | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | |
| GLYCERIN (UNII: PDC6A3C0OX) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-5602-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-5602-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 3 | NDC:22840-5602-5 | 5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

PENICILLIUM DIGITATUM

penicillium digitatum solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-2610 |
| Route of Administration | INTRADERMAL, SUBCUTANEOUS, PERCUTANEOUS | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| PENICILLIUM DIGITATUM (UNII: 1SB49SV239) (PENICILLIUM DIGITATUM - UNII:1SB49SV239) | PENICILLIUM DIGITATUM | 0.025 g in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|----------|------------------|--|-----------------------------|---------------------------|
| 1 | NDC:22840-2610-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-2610-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|---------------------------|---|-----------------------------|---------------------------|
| BLA | BLA101833 | 09/15/1981 | |

MUCOR MIX

mucor circinelloides f. lusitanicus and mucor plumbeus solution

Product Information

| | | | |
|--------------------------------|---|---------------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9672 |
| Route of Administration | PERCUTANEOUS, SUBCUTANEOUS, INTRADERMAL | | |

| Active Ingredient/Active Moiety | | | |
|--|--|-------------------------------------|-------------------|
| Ingredient Name | | Basis of Strength | Strength |
| MUCOR CIRCINELLOIDES F. LUSITANICUS (UNII: 0J0X819B3C) (MUCOR CIRCINELLOIDES F. LUSITANICUS - UNII:0J0X819B3C) | | MUCOR CIRCINELLOIDES F. LUSITANICUS | 0.05 g in 1 mL |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | | MUCOR PLUMBEUS | 0.05 g in 1 mL |

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

| Packaging | | | | |
|-----------|------------------|--|----------------------|--------------------|
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9672-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

| Marketing Information | | | |
|-----------------------|--|----------------------|--------------------|
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
| BLA | BLA101833 | 09/15/1981 | |

| MOLD MIX 2 | | | |
|--|--|--|--|
| aureobasidium pullulans, curvularia spicifera, gibberella fujikuroi, mucor plumbeus and rhizopus stolonifer solution | | | |

| Product Information | | | |
|-------------------------|---|--------------------|----------------|
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9607 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |

| Active Ingredient/Active Moiety | | | |
|--|--|------------------------------|-------------------|
| Ingredient Name | | Basis of Strength | Strength |
| RHIZOPUS STOLONIFER (UNII: FEE198DK4Q) (RHIZOPUS STOLONIFER - UNII:FEE198DK4Q) | | RHIZOPUS STOLONIFER | 0.01 g in 1 mL |
| COCHLIOBOLUS SPICIFER (UNII: 91M9RWP3TD) (COCHLIOBOLUS SPICIFER - UNII:91M9RWP3TD) | | COCHLIOBOLUS SPICIFER | 0.01 g in 1 mL |
| GIBBERELLA FUJIKUROI (UNII: 815V716OR2) (GIBBERELLA FUJIKUROI - UNII:815V716OR2) | | GIBBERELLA FUJIKUROI | 0.01 g in 1 mL |
| AUREOBASIDIUM PULLULANS VAR. PULLUTANS (UNII: D1A2NG69CK) | | AUREOBASIDIUM PULLULANS VAR. | 0.01 g |

| | | | | |
|---|---|--|-----------------------------|---------------------------|
| (AUREOBASIDIUM PULLULANS VAR. PULLUTANS - UNII:D1A2NG69CK) | PULLULANS VAR. PULLUTANS | in 1 mL | | |
| MUCOR PLUMBEUS (UNII: D7401PWY6E) (MUCOR PLUMBEUS - UNII:D7401PWY6E) | MUCOR PLUMBEUS | 0.01 g in 1 mL | | |
| Inactive Ingredients | | | | |
| Ingredient Name | | Strength | | |
| SODIUM BICARBONATE (UNII: 8MDF5V39QO) | | | | |
| PHENOL (UNII: 339NCG44TV) | | | | |
| SODIUM CHLORIDE (UNII: 451W47IQ8X) | | | | |
| Packaging | | | | |
| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
| 1 | NDC:22840-9607-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9607-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| Marketing Information | | | | |
| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date | |
| BLA | BLA101833 | 09/15/1981 | | |

ASPERGILLUS MIX

aspergillus amstelodami, aspergillus flavus, aspergillus fumigatus, aspergillus nidulans and aspergillus niger solution

| | | | |
|--|---|------------------------------|--------------------|
| Product Information | | | |
| Product Type | NON-STANDARDIZED ALLERGENIC | Item Code (Source) | NDC:22840-9628 |
| Route of Administration | INTRADERMAL, PERCUTANEOUS, SUBCUTANEOUS | | |
| Active Ingredient/Active Moiety | | | |
| Ingredient Name | | Basis of Strength | Strength |
| ASPERGILLUS NIGER VAR. NIGER (UNII: 9IOA40ANG6) (ASPERGILLUS NIGER VAR. NIGER - UNII:9IOA40ANG6) | | ASPERGILLUS NIGER VAR. NIGER | 8000 [PNU] in 1 mL |
| ASPERGILLUS FUMIGATUS (UNII: X88DF51T48) (ASPERGILLUS FUMIGATUS - UNII:X88DF51T48) | | ASPERGILLUS FUMIGATUS | 8000 [PNU] in 1 mL |
| EUROTIIUM AMSTELODAMI (UNII: D932NLL87Z) (EUROTIIUM AMSTELODAMI - UNII:D932NLL87Z) | | EUROTIIUM AMSTELODAMI | 8000 [PNU] in 1 mL |
| ASPERGILLUS FLAVUS (UNII: 3J888Y9L13) (ASPERGILLUS FLAVUS - UNII:3J888Y9L13) | | ASPERGILLUS FLAVUS | 8000 [PNU] in 1 mL |
| ASPERGILLUS NIDULANS (UNII: 242A53RB80) (ASPERGILLUS NIDULANS - UNII:242A53RB80) | | ASPERGILLUS NIDULANS | 8000 [PNU] in 1 mL |

Inactive Ingredients

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

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22840-9628-2 | 10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |
| 2 | NDC:22840-9628-4 | 50 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| BLA | BLA101833 | 09/15/1981 | |

Labeler - Greer Laboratories, Inc. (024671414)

Registrant - Greer Laboratories, Inc. (024671414)

Establishment

| Name | Address | ID/FEI | Business Operations |
|--------------------------|---------|-----------|---|
| Greer Laboratories, Inc. | | 024671414 | manufacture(22840-9643, 22840-1684, 22840-5633, 22840-1609, 22840-2656, 22840-1669, 22840-9631, 22840-9633, 22840-1622, 22840-1623, 22840-2660, 22840-5605, 22840-1625, 22840-1626, 22840-1627, 22840-1628, 22840-5606, 22840-1630, 22840-5607, 22840-5648, 22840-1632, 22840-1633, 22840-1634, 22840-1635, 22840-1636, 22840-2661, 22840-5608, 22840-1639, 22840-2662, 22840-5609, 22840-1640, 22840-1641, 22840-1642, 22840-1643, 22840-1644, 22840-1646, 22840-5610, 22840-1647, 22840-1648, 22840-1650, 22840-1651, 22840-5612, 22840-1653, 22840-1654, 22840-1656, 22840-1657, 22840-5613, 22840-1659, 22840-2663, 22840-5614, 22840-2664, 22840-1660, 22840-1661, 22840-1662, 22840-1664, 22840-5615, 22840-9634, 22840-9635, 22840-1666, 22840-2665, 22840-5616, 22840-1667, 22840-2620, 22840-1668, 22840-2666, 22840-5617, 22840-9636, 22840-9637, 22840-9638, 22840-9639, 22840-9641, 22840-1676, 22840-1677, 22840-1678, 22840-5619, 22840-1680, 22840-2667, 22840-5620, 22840-1670, 22840-1671, 22840-1672, 22840-1673, 22840-1674, 22840-5618, 22840-1681, 22840-1682, 22840-1683, 22840-1685, 22840-1686, 22840-5621, 22840-9670, 22840-9675, 22840-9671, 22840-9676, 22840-1687, 22840-1689, 22840-1690, 22840-1691, 22840-2668, 22840-5622, 22840-5642, 22840-5623, 22840-9600, 22840-9601, 22840-9602, 22840-9603, 22840-9604, 22840-9606, 22840-9607, 22840-9608, 22840-9609, 22840-9610, 22840-9612, 22840-9613, 22840-9614, 22840-9616, 22840-9617, 22840-9618, 22840-9619, 22840-9620, 22840-9642, 22840-9644, 22840-9645, 22840-9677, 22840-1692, 22840-2669, 22840-2670, 22840-5624, 22840-2659, 22840-2671, 22840-5625, 22840-9646, 22840-9648, 22840-9672, 22840-1694, 22840-1695, 22840-1696, 22840-1698, 22840-2672, 22840-5626, 22840-1699, 22840-2600, 22840-2601, 22840-2602, 22840-5627, 22840-9649, 22840-9650, 22840-9651, 22840-9652, 22840-9653, 22840-9678, 22840-2605, 22840-2606, 22840-2608, 22840-2609, |

22840-2673, 22840-5629, 22840-2612, 22840-2614, 22840-2615, 22840-2616,
22840-5631, 22840-2610, 22840-5630, 22840-9654, 22840-9655, 22840-9656,
22840-9657, 22840-9659, 22840-9660, 22840-9679, 22840-2618, 22840-2674,
22840-5632, 22840-9661, 22840-9662, 22840-9664, 22840-2619, 22840-2621,
22840-9665, 22840-9666, 22840-9668, 22840-9669, 22840-9680, 22840-2623,
22840-2624, 22840-2625, 22840-2627, 22840-5634, 22840-2628, 22840-2630,
22840-2631, 22840-2675, 22840-5635, 22840-2633, 22840-2634, 22840-2676,
22840-2677, 22840-2678, 22840-5636, 22840-1606, 22840-2679, 22840-5601,
22840-2635, 22840-2636, 22840-2637, 22840-2638, 22840-5637, 22840-2640,
22840-2641, 22840-2642, 22840-2680, 22840-5638, 22840-2644, 22840-2645,
22840-2646, 22840-2648, 22840-2649, 22840-5639, 22840-2650, 22840-2681,
22840-5640, 22840-2652, 22840-2653, 22840-2654, 22840-2655, 22840-5641,
22840-9621, 22840-9622, 22840-9623, 22840-9673, 22840-1600, 22840-1601,
22840-1602, 22840-1604, 22840-9628, 22840-9629, 22840-9630, 22840-1605,
22840-5600, 22840-9624, 22840-9625, 22840-9626, 22840-9674, 22840-1608,
22840-5602, 22840-1610, 22840-1612, 22840-1614, 22840-5603, 22840-1615,
22840-1616, 22840-1617, 22840-1618, 22840-1619, 22840-1621, 22840-5604,
22840-9627, 22840-2607)

Revised: 6/2025

Greer Laboratories, Inc.