## SYNAGIS - palivizumab injection, solution Swedish Orphan Biovitrum AB (publ)

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#### HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use SYNAGIS safely and effectively. See full prescribing information for SYNAGIS.

## SYNAGIS® (palivizumab) injection, for intramuscular use Initial U.S. Approval: 1998

----- INDICATIONS AND USAGE

Synagis is a respiratory syncytial virus (RSV) F protein inhibitor monoclonal antibody indicated for the prevention of serious lower respiratory tract disease caused by respiratory syncytial virus (RSV) in pediatric patients:

- with a history of premature birth (less than or equal to 35 weeks gestational age) and who are 6 months of age or younger at the beginning of RSV season,
- with bronchopulmonary dysplasia (BPD) that required medical treatment within the previous 6 months and who are 24 months of age or younger at the beginning of RSV season,
- with hemodynamically significant congenital heart disease (CHD) and who are 24 months of age or younger at the beginning of RSV season.

<u>Limitations of Use:</u> The safety and efficacy of Synagis have not been established for treatment of RSV disease. (1)

#### ------DOSAGE AND ADMINISTRATION ------

15 mg per kg of body weight, administered intramuscularly prior to commencement of the RSV season and remaining doses administered monthly throughout the RSV season. (2.1)

Children undergoing cardio-pulmonary bypass should receive an additional dose of Synagis as soon as possible after the cardio-pulmonary bypass procedure (even if sooner than a month from the previous dose). Thereafter, doses should be administered monthly as scheduled. (2.1, 12.3)

#### ------ DOSAGE FORMS AND STRENGTHS

Single-dose liquid solution vials: 50 mg per 0.5 mL and 100 mg per 1 mL. (3)

#### ------CONTRAINDICATIONS ------

Previous significant hypersensitivity reaction to Synagis. (4)

#### ------ WARNINGS AND PRECAUTIONS

- Anaphylaxis and anaphylactic shock (including fatal cases), and other severe acute hypersensitivity reactions have been reported. Permanently discontinue Synagis and administer appropriate medications if such reactions occur. (5.1)
- As with any intramuscular injection, Synagis should be given with caution to children with thrombocytopenia or any coagulation disorder. (5.2)
- Palivizumab may interfere with immunological-based RSV diagnostic tests such as some antigen detection-based assays. (5.3, 12.4)

#### ------ ADVERSE REACTIONS ------

Adverse reactions occurring greater than or equal to 10% and at least 1% more frequently than placebo are fever and rash. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact 1-866-773-5274 or FDA at 1-800-FDA-1088 or <a href="https://www.fda.gov/medwatch">www.fda.gov/medwatch</a>.

------USE IN SPECIFIC POPULATIONS ------

Safety and effectiveness in children older than 24 months of age at the start of dosing have not been established. (8.4)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

Revised: 11/2021

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#### **FULL PRESCRIBING INFORMATION**

#### 1 INDICATIONS AND USAGE

Synagis is indicated for the prevention of serious lower respiratory tract disease caused by respiratory syncytial virus (RSV) in pediatric patients:

- with a history of premature birth (less than or equal to 35 weeks gestational age) and who are 6 months of age or younger at the beginning of RSV season,
- with bronchopulmonary dysplasia (BPD) that required medical treatment within the previous 6 months and who are 24 months of age or younger at the beginning of RSV season,
- with hemodynamically significant congenital heart disease (CHD) and who are 24

months of age or younger at the beginning of RSV season [see Clinical Studies (14)].

#### Limitations of Use:

The safety and efficacy of Synagis have not been established for treatment of RSV disease [see Warnings and Precautions (5.4)].

#### 2 DOSAGE AND ADMINISTRATION

#### 2.1 Dosing Information

The recommended dose of Synagis is 15 mg per kg of body weight given monthly by intramuscular injection. The first dose of Synagis should be administered prior to commencement of the RSV season and the remaining doses should be administered monthly throughout the RSV season. Children who develop an RSV infection should continue to receive monthly doses throughout the RSV season. In the northern hemisphere, the RSV season typically commences in November and lasts through April, but it may begin earlier or persist later in certain communities.

Synagis serum levels are decreased after cardio-pulmonary bypass [see <u>Clinical</u> <u>Pharmacology (12.3)</u>]. Children undergoing cardio-pulmonary bypass should receive an additional dose of Synagis as soon as possible after the cardio-pulmonary bypass procedure (even if sooner than a month from the previous dose). Thereafter, doses should be administered monthly as scheduled.

The efficacy of Synagis at doses less than 15 mg per kg, or of dosing less frequently than monthly throughout the RSV season, has not been established.

#### 2.2 Administration Instructions

- DO NOT DILUTE THE PRODUCT.
- DO NOT SHAKE OR VIGOROUSLY AGITATE THE VIAL.
- Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration. Do not use any vials exhibiting particulate matter or discoloration.
- Using aseptic techniques, attach a sterile needle to a sterile syringe. Remove the flip top from the Synagis vial and wipe the rubber stopper with a disinfectant (e.g., 70% isopropyl alcohol). Insert the needle into the vial and withdraw into the syringe an appropriate volume of solution. Administer immediately after drawing the dose into the syringe.
- Synagis should be administered in a dose of 15 mg per kg intramuscularly using aseptic technique, preferably in the anterolateral aspect of the thigh. The gluteal muscle should not be used routinely as an injection site because of the risk of damage to the sciatic nerve. The dose (volume of injection in mL) per month = patient weight (kg) x 15 mg per kg ÷ 100 mg per mL of Synagis. Injection volumes over 1 mL should be given as a divided dose.
- Synagis is supplied as a single-dose vial and does not contain preservatives. Do not re-enter the vial after withdrawal of drug; discard unused portion. Only administer one dose per vial.
- Use sterile disposable syringes and needles. To prevent the transmission of hepatitis viruses or other infectious agents from one person to another, DO NOT reuse syringes and needles.

#### **3 DOSAGE FORMS AND STRENGTHS**

Single-dose liquid solution vials: 50 mg per 0.5 mL and 100 mg per 1 mL.

#### 4 CONTRAINDICATIONS

Synagis is contraindicated in children who have had a previous significant hypersensitivity reaction to Synagis [see <u>Warnings and Precautions (5.1)</u>].

#### **5 WARNINGS AND PRECAUTIONS**

## 5.1 Hypersensitivity Reactions

Cases of anaphylaxis and anaphylactic shock, including fatal cases, have been reported following initial exposure or re-exposure to Synagis. Other acute hypersensitivity reactions, which may be severe, have also been reported on initial exposure or reexposure to Synagis. Signs and symptoms may include urticaria, pruritus, angioedema, dyspnea, respiratory failure, cyanosis, hypotonia, hypotension, and unresponsiveness. The relationship between these reactions and the development of antibodies to Synagis is unknown. If a significant hypersensitivity reaction occurs with Synagis, its use should be permanently discontinued. If anaphylaxis or other significant hypersensitivity reaction occurs, administer appropriate medications (e.g., epinephrine) and provide supportive care as required. If a mild hypersensitivity reaction occurs, clinical judgment should be used regarding cautious readministration of Synagis.

## 5.2 Coagulation Disorders

Synagis is for intramuscular use only. As with any intramuscular injection, Synagis should be given with caution to children with thrombocytopenia or any coagulation disorder.

## 5.3 RSV Diagnostic Test Interference

Palivizumab may interfere with immunological-based RSV diagnostic tests such as some antigen detection-based assays. In addition, palivizumab inhibits virus replication in cell culture, and therefore may also interfere with viral culture assays. Palivizumab does not interfere with reverse transcriptase-polymerase chain reaction based assays. Assay interference could lead to false-negative RSV diagnostic test results. Therefore, diagnostic test results, when obtained, should be used in conjunction with clinical findings to guide medical decisions [see <u>Microbiology (12.4)</u>].

#### 5.4 Treatment of RSV Disease

The safety and efficacy of Synagis have not been established for treatment of RSV disease.

## 5.5 Proper Administration

The single-dose vial of Synagis does not contain a preservative. Administration of Synagis should occur immediately after dose withdrawal from the vial. The vial should

not be re-entered. Discard any unused portion.

#### **6 ADVERSE REACTIONS**

The most serious adverse reactions occurring with Synagis are anaphylaxis and other acute hypersensitivity reactions [see <u>Warnings and Precautions</u> (5.1)].

### **6.1 Clinical Studies Experience**

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The data described below reflect exposure to Synagis (n=1639) compared with placebo (n=1143) in children 3 days to 24.1 months of age at high risk of RSV-related hospitalization in two clinical trials. Trial 1 was conducted during a single RSV season and studied a total of 1502 children less than or equal to 24 months of age with BPD or infants with premature birth (less than or equal to 35 weeks gestation) who were less than or equal to 6 months of age at study entry. Trial 2 was conducted over four consecutive seasons among a total of 1287 children less than or equal to 24 months of age with hemodynamically significant congenital heart disease.

In Trials 1 and 2 combined, fever and rash were each reported more frequently among Synagis than placebo recipients, 27% versus 25%, and 12% versus 10%, respectively. Adverse reactions observed in the 153-patient crossover study comparing the liquid and lyophilized formulations were comparable for the two formulations, and were similar to those observed with Synagis in Trials 1 and 2.

## <u>Immunogenicity</u>

In Trial 1, the incidence of anti-palivizumab antibody following the fourth injection was 1.1% in the placebo group and 0.7% in the Synagis group. In children receiving Synagis for a second season, one of the fifty-six children had transient, low titer reactivity. This reactivity was not associated with adverse events or alteration in serum concentrations. Immunogenicity was not assessed in Trial 2.

A trial of high-risk preterm children less than or equal to 24 months of age was conducted to evaluate the immunogenicity of the lyophilized formulation of Synagis (used in Trials 1 and 2 above) and the liquid formulation of Synagis. Three hundred seventy-nine children contributed to the 4 to 6 months post-final dose analysis. The rate of anti-palivizumab antibodies at this time point was low in both formulation groups (anti-palivizumab antibodies were not detected in any subject in the liquid formulation group and were detected in one subject in the lyophilized group (0.5%), with an overall rate of 0.3% for both treatment groups combined).

These data reflect the percentage of children whose test results were considered positive for antibodies to palivizumab in an enzyme-linked immunosorbent assay (ELISA) and are highly dependent on the sensitivity and specificity of the assay.

The ELISA has substantial limitations in detecting anti-palivizumab antibodies in the presence of palivizumab. Immunogenicity samples tested with the ELISA assay likely contained palivizumab at levels that may interfere with the detection of anti-palivizumab antibodies.

An electrochemical luminescence (ECL) based immunogenicity assay, with a higher tolerance for palivizumab presence compared to the ELISA, was used to evaluate the presence of anti-palivizumab antibodies in subject samples from two additional clinical trials. The rates of anti-palivizumab antibody positive results in these trials were 1.1% and 1.5%.

## **6.2 Postmarketing Experience**

The following adverse reactions have been identified during post approval use of Synagis. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

**Blood and Lymphatic System Disorders:** severe thrombocytopenia (platelet count less than 50,000 per microliter)

General Disorders and Administration Site Conditions: injection site reactions

Limited information from post-marketing reports suggests that, within a single RSV season, adverse events after a sixth or greater dose of Synagis are similar in character and frequency to those after the initial five doses.

#### 7 DRUG INTERACTIONS

No formal drug-drug interaction studies were conducted. In Trial 1, the proportions of children in the placebo and Synagis groups who received routine childhood vaccines, influenza vaccine, bronchodilators, or corticosteroids were similar and no incremental increase in adverse reactions was observed among children receiving these agents.

#### **8 USE IN SPECIFIC POPULATIONS**

## 8.1 Pregnancy

Risk Summary

Synagis is not indicated for use in females of reproductive potential.

#### 8.2 Lactation

Risk Summary

Synagis is not indicated for use in females of reproductive potential.

#### 8.4 Pediatric Use

The safety and effectiveness of Synagis in children older than 24 months of age at the start of dosing have not been established [see <u>Clinical Studies</u> (14)].

#### **10 OVERDOSAGE**

Overdoses with doses up to 85 mg per kg have been reported in clinical studies and post-marketing experience with Synagis, and in some cases, adverse reactions were reported. In case of overdosage, it is recommended that the patient be monitored for

any signs or symptoms of adverse reactions and appropriate symptomatic treatment instituted.

#### 11 DESCRIPTION

Palivizumab is a humanized monoclonal antibody (IgG1 $\kappa$ ) produced by recombinant DNA technology, directed to an epitope in the A antigenic site of the F protein of RSV. Palivizumab is a composite of human (95%) and murine (5%) antibody sequences. The human heavy chain sequence was derived from the constant domains of human IgG1 and the variable framework regions of the  $V_H$  genes Cor and Cess. The human light chain sequence was derived from the constant domain of C $\kappa$  and the variable framework regions of the  $V_L$  gene K104 with J $\kappa$  -4. The murine sequences were derived from a murine monoclonal antibody, Mab 1129, in a process that involved the grafting of the murine complementarity determining regions into the human antibody frameworks. Palivizumab is composed of two heavy chains and two light chains and has a molecular weight of approximately 148,000 Daltons.

Synagis is supplied as a sterile, preservative-free liquid solution at 100 mg per mL to be administered by intramuscular injection. Thimerosal or other mercury-containing salts are not used in the production of Synagis. The solution has a pH of 6.0 and should appear clear or slightly opalescent.

Each 100 mg single-dose vial of Synagis liquid solution contains 100 mg of palivizumab and also contains chloride (0.5 mg), glycine (0.1 mg), and histidine (3.9 mg), in a volume of 1 mL.

Each 50 mg single-dose vial of Synagis liquid solution contains 50 mg of palivizumab and also contains chloride (0.2 mg), glycine (0.06 mg), and histidine (1.9 mg), in a volume of 0.5 mL.

#### 12 CLINICAL PHARMACOLOGY

#### 12.1 Mechanism of Action

Palivizumab is a recombinant humanized monoclonal antibody with anti-RSV activity [see Microbiology (12.4)].

#### 12.3 Pharmacokinetics

In children less than or equal to 24 months of age without congenital heart disease (CHD), the mean half-life of palivizumab was 20 days and monthly intramuscular doses of 15 mg per kg achieved mean  $\pm$  SD 30 day trough serum drug concentrations of 37  $\pm$  21 mcg per mL after the first injection, 57  $\pm$  41 mcg per mL after the second injection, 68  $\pm$  51 mcg per mL after the third injection, and 72  $\pm$  50 mcg per mL after the fourth injection. Trough concentrations following the first and fourth Synagis dose were similar in children with CHD and in non-cardiac patients. In children given Synagis for a second season, the mean  $\pm$  SD serum concentrations following the first and fourth injections were 61  $\pm$  17 mcg per mL and 86  $\pm$  31 mcg per mL, respectively.

In 139 children less than or equal to 24 months of age with hemodynamically significant CHD who received Synagis and underwent cardio-pulmonary bypass for open-heart surgery, the mean  $\pm$  SD serum palivizumab concentration was 98  $\pm$  52 mcg per mL

before bypass and declined to  $41 \pm 33$  mcg per mL after bypass, a reduction of 58% [see <u>Dosage and Administration (2.1)</u>]. The clinical significance of this reduction is unknown.

Specific studies were not conducted to evaluate the effects of demographic parameters on palivizumab systemic exposure. However, no effects of gender, age, body weight, or race on palivizumab serum trough concentrations were observed in a clinical study with 639 children with CHD (less than or equal to 24 months of age) receiving five monthly intramuscular injections of 15 mg per kg of Synagis.

The pharmacokinetics and safety of Synagis liquid solution and Synagis lyophilized formulation administered via intramuscular injection at 15 mg per kg were studied in a cross-over trial of 153 infants less than or equal to 6 months of age with a history of prematurity. The results of this trial indicated that the trough serum concentrations of palivizumab were comparable between the liquid solution and the lyophilized formulation, which was the formulation used in the clinical studies.

A population pharmacokinetic analysis was performed across 22 studies in 1800 patients (1684 pediatric and 116 adult patients) to characterize palivizumab pharmacokinetics and inter-subject variability in serum concentrations. Palivizumab pharmacokinetics was described by a two-compartment linear model with an elimination half-life of 24.5 days in pediatric patients. Clearance of palivizumab in a typical pediatric patient (body weight 4.5 kg) less than or equal to 24 months of age without CHD was estimated to be 11 mL per day with a bioavailability of 70% following intramuscular administration. The inter-patient variability in drug clearance was 48.7% (CV%). Covariate analysis did not identify any factors that could account for the inter-patient variability in order to predict serum concentrations a priori in an individual patient.

## 12.4 Microbiology

#### Mechanism of Action

Palivizumab, a recombinant humanized monoclonal antibody which provides passive immunity against RSV, acts by binding the RSV envelope fusion protein (RSV F) on the surface of the virus and blocking a critical step in the membrane fusion process. Palivizumab also prevents cell-to-cell fusion of RSV-infected cells.

#### **Antiviral Activity**

The antiviral activity of palivizumab was assessed in a microneutralization assay in which increasing concentrations of antibody were incubated with RSV prior to addition of the human epithelial cells HEp-2. After incubation for 4-5 days, RSV antigen was measured in an ELISA assay. The neutralization titer (50% effective concentration [EC<sub>50</sub>]) is expressed as the antibody concentration required to reduce detection of RSV antigen by 50% compared with untreated virus-infected cells. Palivizumab exhibited median EC<sub>50</sub> values of 0.65 mcg per mL (mean  $0.75 \pm 0.53$  mcg per mL; n=69, range 0.07-2.89 mcg per mL) and 0.28 mcg per mL (mean  $0.35 \pm 0.23$  mcg per mL; n=35, range 0.03-0.88 mcg per mL) against clinical RSV A and RSV B isolates, respectively. The majority of clinical RSV isolates tested (n=96) were collected from subjects across the United States (CA, CO, CT, IL, MA, NC, NY, PA, RI, TN, TX, VA), with the remainder from Japan (n=1), Australia (n=5) and Israel (n=2). These isolates encoded the most common RSV F sequence polymorphisms found among clinical isolates worldwide.

Palivizumab serum concentrations of greater than or equal to 40 mcg per mL have been

shown to reduce pulmonary RSV replication in the cotton rat model of RSV infection by 100-fold.

#### Resistance

Palivizumab binds a highly conserved region on the extracellular domain of mature RSV F, referred to as antigenic site II or site A, which encompasses amino acids 262 to 275. All RSV mutants that exhibit resistance to palivizumab have been shown to contain amino acid changes in this region on the F protein.

F protein sequence variations within antigenic site A: Amino acid substitutions in antigenic site A selected either in cell culture, in animal models, or in human subjects that resulted in palivizumab resistance were N262D, N268I, K272E/M/N/Q/T, and S275F/L. RSV variants expressing the K272N substitution in F protein showed a 5164  $\pm$  1731-fold decrease in susceptibility (i.e., fold increase in EC50 value) when compared to the wild-type RSV, while variants containing the N262D, S275F/L, or K272E/M/Q/T substitutions showed a greater than 25,000-fold decrease in susceptibility to palivizumab. The N268I substitution conferred partial resistance to palivizumab; however, fold changes in susceptibility were not quantified for this mutant. Studies carried out to investigate the mechanism of virus escape from palivizumab showed a correlation between antibody binding and virus neutralization. RSV with substitutions in antigenic site A that were resistant to neutralization by palivizumab did not bind to palivizumab.

At least one of the palivizumab resistance-associated substitutions, N262D, K272E/Q, or S275F/L was identified in 8 of 126 clinical RSV (59 RSV A and 67 RSV B) isolates from subjects who failed immunoprophylaxis, resulting in a combined resistance-associated mutation frequency of 6.3%. A review of clinical findings revealed no association between antigenic A site sequence changes and RSV disease severity among children receiving palivizumab immunoprophylaxis who develop RSV lower respiratory tract disease.

Analysis of 254 clinical RSV isolates (145 RSV A and 109 RSV B) collected from immunoprophylaxis-naïve subjects revealed palivizumab resistance-associated substitutions in 2 (1 with N262D and 1 with S275F), resulting in a resistance-associated mutation frequency of 0.79%.

F protein sequence variations outside antigenic site A: In addition to the sequence variations in antigenic site A known to confer palivizumab resistance, F protein substitutions T100A, G139S, N165D/V406I; T326A, V450A in RSV A, and T74I, A147V, I206L, S285G, V450I, T455I in RSV B were identified in viruses isolated from failures of immunoprophylaxis. These substitutions were not identified in RSV F sequences derived from 254 clinical isolates from immunoprophylaxis-naïve subjects and thus are considered treatment-associated and non-polymorphic. Recombinant RSV B encoding the S285G substitution exhibited palivizumab sensitivity (EC50 value = 0.39  $\pm$  0.02 mcg per mL) similar to recombinant wild-type RSV B (EC50 value = 0.17  $\pm$  0.02 mcg per mL).

Palivizumab susceptibility of RSV encoding common F protein sequence polymorphisms located proximal to antigenic site A was evaluated. Recombinant RSV A encoding N276S (EC $_{50}$  value = 0.72  $\pm$  0.07 mcg per mL), and recombinant RSV B with S276N (EC $_{50}$  value = 0.42  $\pm$  0.04 mcg per mL), exhibited sensitivities comparable to the corresponding recombinant wild-type RSV A (EC $_{50}$  value = 0.63  $\pm$  0.22 mcg per mL) and RSV B (EC $_{50}$  value = 0.23  $\pm$  0.07 mcg per mL). Likewise, RSV B clinical isolates containing the polymorphic variation V278A were at least as sensitive to neutralization by palivizumab

(EC $_{50}$  range 0.08-0.45 mcg per mL) as laboratory strains of wild-type RSV B (EC $_{50}$  value = 0.54  $\pm$  0.08 mcg per mL). No known polymorphic or non-polymorphic sequence variations outside the antigenic site A on RSV F have been demonstrated to render RSV resistant to neutralization by palivizumab.

## Interference of RSV Diagnostic Assays by Palivizumab

Interference with immunologically-based RSV diagnostic assays by palivizumab has been observed in laboratory studies. Rapid chromatographic/enzyme immunoassays (CIA/EIA), immunofluorescence assays (IFA), and direct immunofluorescence assays (DFA) using monoclonal antibodies targeting RSV F protein may be inhibited. Therefore, caution should be used in interpreting negative immunological assay results when clinical observations are consistent with RSV infection. A reverse transcriptase-polymerase chain reaction (RT-PCR) assay, which is not inhibited by palivizumab, may prove useful for laboratory confirmation of RSV infection [see Warnings and Precautions (5.3)].

#### 13 NONCLINICAL TOXICOLOGY

## 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis, mutagenesis, and reproductive toxicity studies have not been performed.

#### **14 CLINICAL STUDIES**

The safety and efficacy of Synagis were assessed in two randomized, double-blind, placebo-controlled trials of prophylaxis against RSV infection in children at high risk of an RSV-related hospitalization. Trial 1 was conducted during a single RSV season and studied a total of 1502 children less than or equal to 24 months of age with BPD or infants with premature birth (less than or equal to 35 weeks gestation) who were less than or equal to 6 months of age at study entry. Trial 2 was conducted over four consecutive seasons among a total of 1287 children less than or equal to 24 months of age with hemodynamically significant congenital heart disease. In both trials participants received 15 mg per kg Synagis or an equivalent volume of placebo via intramuscular injection monthly for five injections and were followed for 150 days from randomization. In Trial 1, 99% of all subjects completed the study and 93% completed all five injections. In Trial 2, 96% of all subjects completed the study and 92% completed all five injections. The incidence of RSV hospitalization is shown in Table 1. The results were shown to be statistically significant using Fisher's exact test.

Table 1: Incidence of RSV Hospitalization by Treatment Group

Trial		Placebo	Synagis	Difference Between Groups	Relative Reduction
Trial 1	N	500	1002		
Impact- RSV	Hospitalization	53 (10.6%)	48 (4.8%)	5.8%	55%
Trial 2	N	648	639		
CHD	Hospitalization	63 (9.7%)	34 (5.3%)	4.4%	45%

In Trial 1, the reduction of RSV hospitalization was observed both in children with BPD (34/266 [12.8%] placebo versus 39/496 [7.9%] Synagis) and in premature infants without BPD (19/234 [8.1%] placebo versus 9/506 [1.8%] Synagis). In Trial 2, reductions were observed in acyanotic (36/305 [11.8%] placebo versus 15/300 [5.0%] Synagis) and cyanotic children (27/343 [7.9%] placebo versus 19/339 [5.6%] Synagis).

The clinical studies do not suggest that RSV infection was less severe among children hospitalized with RSV infection who received Synagis for RSV prophylaxis compared to those who received placebo.

#### 16 HOW SUPPLIED/STORAGE AND HANDLING

Synagis is supplied in single-dose vials as a preservative-free, sterile liquid solution at 100 mg per mL for intramuscular injection.

50 mg vial NDC 66658-230-01

The 50 mg vial contains 50 mg Synagis in 0.5 mL.

100 mg vial NDC 66658-231-01

The 100 mg vial contains 100 mg Synagis in 1 mL.

The rubber stopper used for sealing vials of Synagis is not made with natural rubber latex.

#### Storage

Upon receipt and until use, Synagis should be stored between 2°C and 8°C (36°F and 46°F) in its original container. DO NOT freeze. DO NOT use beyond the expiration date.

#### 17 PATIENT COUNSELING INFORMATION

# Advise the patient's caregiver to read the FDA-approved patient labeling (Patient Information)

## **Hypersensitivity Reactions**

Inform the patient's caregiver of the signs and symptoms of potential hypersensitivity reactions, and advise the caregiver to seek medical attention immediately if the child experiences a severe hypersensitivity reaction to Synagis [see <u>Contraindications (4) and Warnings and Precautions (5.1)</u>].

#### <u>Administration</u>

Advise the patient's caregiver that Synagis should be administered by a healthcare provider once a month during the RSV season by intramuscular injection and the importance of compliance with the full course of therapy [see <u>Dosage and Administration</u> (2)].

#### sobi

#### Manufactured by:

Swedish Orphan Biovitrum AB (publ) Stockholm, Sweden

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# PATIENT INFORMATION SYNAGIS® (Si-na-jis)

(palivizumab) injection

#### What is SYNAGIS?

SYNAGIS is a prescription medication that is used to help prevent a serious lung disease caused by Respiratory Syncytial Virus (RSV) in children:

- born prematurely (at or before 35 weeks) **and** who are 6 months of age or less at the beginning of RSV season,
- who have a chronic lung condition called bronchopulmonary dysplasia (BPD), that needed medical treatment within the last 6 months, **and** who are 24 months of age or less at the beginning of RSV season,
- born with certain types of heart disease **and** who are 24 months of age or less at the beginning of RSV season.

SYNAGIS contains man-made, disease-fighting proteins called antibodies.

It is not known if SYNAGIS is safe and effective to treat the symptoms of RSV in a child who already has RSV. Synagis is used to help prevent RSV disease.

It is not known if SYNAGIS is safe and effective in children who are older than 24 months of age at the start of dosing.

#### Who should not receive SYNAGIS?

Your child should not receive SYNAGIS if they have ever had a severe allergic reaction to it. See the end of this leaflet for a complete list of ingredients in SYNAGIS. Signs and symptoms of a severe allergic reaction could include:

- severe rash, hives, or itching skin
- swelling of the lips, tongue, or face
- swelling of the throat, difficulty swallowing
- difficult, rapid, or irregular breathing
- bluish color of skin, lips, or under fingernails
- muscle weakness or floppiness
- unresponsiveness

# Before your child receives SYNAGIS, tell your healthcare provider about all of your child:

- has ever had a reaction to SYNAGIS.
- has bleeding or bruising problems. SYNAGIS is given by injection. If your child has a problem with bleeding or bruises easily, an injection could cause a problem.

Tell your child's healthcare provider about all the medicines your child takes, including prescription and over-the-counter medicines, vitamins, and herbal supplements.

## How is SYNAGIS given?

- SYNAGIS is given as a monthly injection, usually in the thigh (leg) muscle, by your child's healthcare provider.
- Your child's healthcare provider will give you detailed instructions on when SYNAGIS will be given.
  - "RSV season" is the time of year when RSV infections most commonly happen, usually fall through spring, but it may begin earlier or last longer in certain areas. During this time, when RSV is most active, your child will need to receive SYNAGIS injections. Your healthcare provider can tell you when the RSV season starts in your area.
  - Your child should receive the first SYNAGIS injection before the RSV season starts to help prevent RSV infection. If the season has already started, your child should receive their first SYNAGIS injection as soon as possible to help protect them when exposure to the virus is more likely.
  - SYNAGIS is needed every 28-30 days during the RSV season. Each injection of SYNAGIS helps protect your child from severe RSV disease for about 1 month. Keep all of your child's appointments with your healthcare provider.
- If your child misses an injection, talk to your healthcare provider and schedule another injection as soon as possible.
- Your child may still get severe RSV disease after receiving SYNAGIS. Talk to your healthcare provider about what symptoms to look for. If your child gets a RSV infection, they should continue to receive their scheduled SYNAGIS injections to help prevent severe disease from new RSV infections.
- If your child has certain types of heart disease and has corrective surgery, your healthcare provider may need to give your child an additional SYNAGIS injection soon after surgery.

# What are the possible side effects of SYNAGIS? SYNAGIS may cause serious side effects including:

Severe allergic reactions. Severe allergic reactions may happen after any
injection of SYNAGIS, and may be life-threatening or cause death. Call your
healthcare provider or get medical help right away if your child has any of the signs
or symptoms of a serious allergic reaction. See "Who should not receive
SYNAGIS?".

The most common side effects of SYNAGIS include fever and rash.

These are not all the possible side effects of SYNAGIS.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

You may also report side effects at 1866-773-5274.

#### General information about the safe and effective use of SYNAGIS.

Medicines are sometimes prescribed for purposes other than those listed in Patient Information leaflets.

You can ask your pharmacist or healthcare provider for information about SYNAGIS that is written for health professionals.

## What are the ingredients in SYNAGIS?

Active ingredient: palivizumab

**Inactive ingredients**: chloride, glycine, and histidine Manufactured by: Swedish Orphan **sobi** 

Biovitrum AB (publ) , Stockholm, Sweden Synagis® is a registered trademark of Arexis AB c/o Swedish Orphan Biovitrum AB (publ).

For more information, go to www.synagis.com or call 1866-773-5274.

This Patient Information has been approved by the U.S. Food and Drug Revised: 11/2021 Administration

## Principal Display Panel - 50 mg/0.5 mL Carton Label

NDC 66658-230-01

**SYNAGIS®** 

**PALIVIZUMAB** 

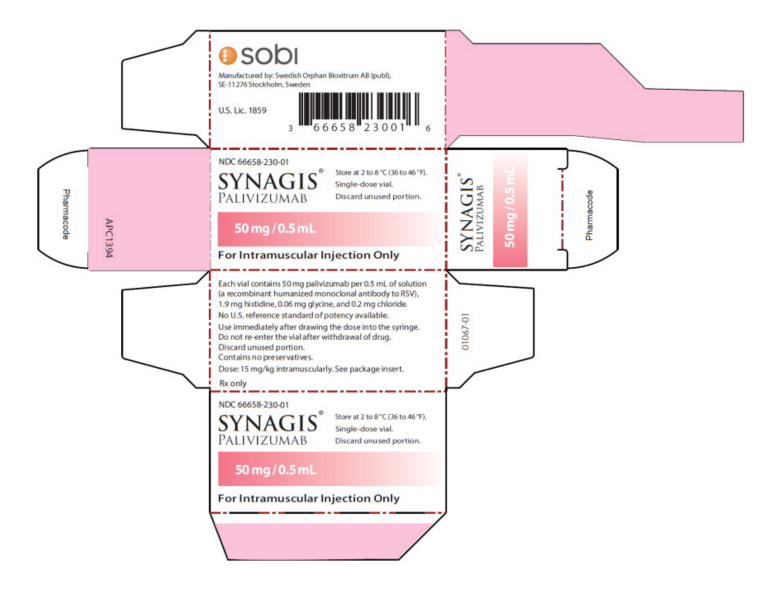
50 mg/0.5 mL

## For Intramuscular Injection Only

Store at 2 to 8°C (36 to 46°F).

Single-dose vial.

Discard unused portion.



## Principal Display Panel - 50 mg/0.5 mL Vial Label

NDC 66658-230-01

50 mg/0.5 mL

**SYNAGIS®** 

**PALIVIZUMAB** 

For Intramuscular Injection Only

US Lic. 1859



## Principal Display Panel - 100 mg/mL Carton Label

NDC 66658-231-01

**SYNAGIS®** 

**PALIVIZUMAB** 

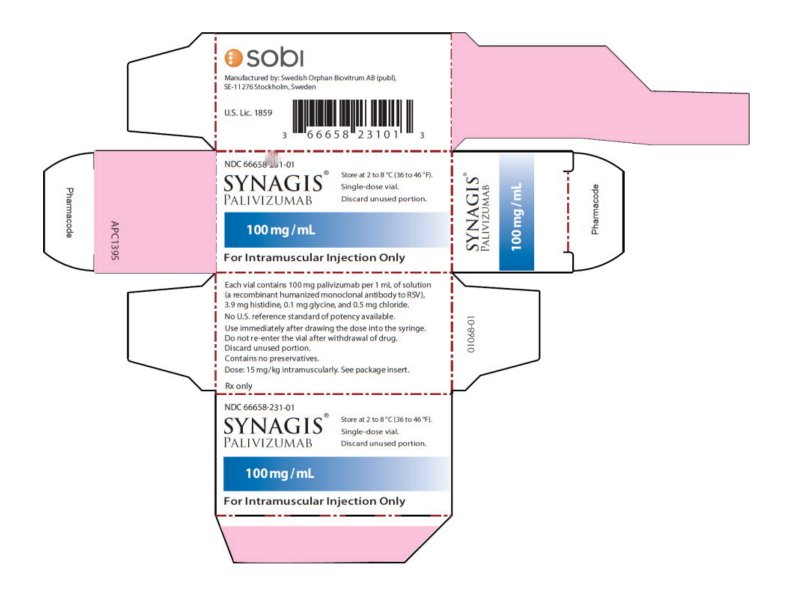
**100 mg/mL** 

## For Intramuscular Injection Only

Store at 2 to 8°C (36 to 46°F).

Single-dose vial.

Discard unused portion.



## Principal Display Panel - 100 mg/mL Vial Label

NDC 66658-231-01

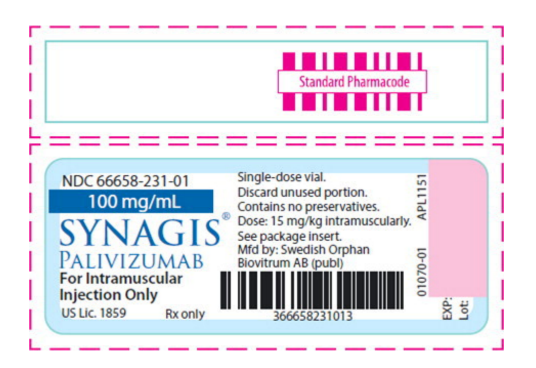
100 mg/mL

**SYNAGIS®** 

**PALIVIZUMAB** 

For Intramuscular Injection Only

US Lic. 1859



## **SYNAGIS**

palivizumab injection, solution

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Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:66658-231
Route of Administration	INTRAMUSCULAR		

<b>Active Ingredient/Active M</b>	loiety
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Ingredient Name	<b>Basis of Strength</b>	Strength
PALIVIZUMAB (UNII: DQ448MW7KS) (PALIVIZUMAB - UNII:DQ448MW7KS)	PALIVIZ UMAB	100 mg in 1 mL

Inactive Ingredients				
Ingredient Name	Strength			
CHLORIDE ION (UNII: Q32ZN48698)	0.5 mg in 1 mL			
GLYCINE (UNII: TE7660XO1C)	0.1 mg in 1 mL			
HISTIDINE (UNII: 4QD397987E)	3.9 mg in 1 mL			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:66658- 231-01	1 mL in 1 VIAL, SINGLE-DOSE; Type 0: Not a Combination Product	09/28/2005	

Marketing Information			
Marketing	Application Number or Monograph	Marketing Start	Marketing End

Category	Citation	Date	Date
BLA	BLA103770	06/19/1998	

## **SYNAGIS**

palivizumab injection, solution

#### **Product Information**

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:66658-230
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Route of Administration INTRAMUSCULAR

## **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength

PALIVIZUMAB (UNII: DQ448MW7KS) (PALIVIZUMAB - UNII:DQ448MW7KS) PALIVIZUMAB 50 mg in 0.5 mL

### **Inactive Ingredients**

Ingredient Name	Strength
CHLORIDE ION (UNII: Q32ZN48698)	0.2 mg in 0.5 mL
GLYCINE (UNII: TE7660XO1C)	0.06 mg in 0.5 mL
HISTIDINE (UNII: 4QD397987E)	1.9 mg in 0.5 mL

## **Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:66658- 230-01	0.5 mL in 1 VIAL, SINGLE-DOSE; Type 0: Not a Combination Product	09/28/2005	

## **Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA103770	12/09/1999	

## Labeler - Swedish Orphan Biovitrum AB (publ) (354010589)

## **Establishment**

Name	Address	ID/FEI	Business Operations
Almac Pharma Services LLC		078607239	PACK(66658-231, 66658-230), LABEL(66658-231, 66658-230)

Name	Address	ID/FEI	Business Operations
AstraZ eneca			MANUFACTURE(66658-231, 66658-230), PACK(66658-231, 66658-230).

Establishment						
Name	Address	ID/FEI	Business Operations			
AstraZeneca Pharmaceuticals LP		968459953	API MANUFACTURE(66658-231, 66658-230) , ANALYSIS(66658-231, 66658-230)			

Establishment							
Name	Address	ID/FEI	Business Operations				
BioReliance Corporation		147227730	ANALYSIS (66658-231, 66658-230)				

Establishment							
Name	Address	ID/FEI	Business Operations				
Boehringer Ingelheim Pharma GmbH and Co.			MANUFACTURE(66658-231, 66658-230), API MANUFACTURE(66658-231, 66658-230), ANALYSIS(66658-231, 66658-230), PACK(66658-231, 66658-230), LABEL(66658-231, 66658-230)				

Establishment						
Name	Address	ID/FEI	<b>Business Operations</b>			
Charles River Laboratories, Inc.		078495006	ANALYSIS (66658-231, 66658-230)			

Establishment						
Name	Address	ID/FEI	<b>Business Operations</b>			
Eurofins Lancaster Laboratories, Inc.		069777290	ANALYSIS (66658-231, 66658-230)			

Revised: 11/2021 Swedish Orphan Biovitrum AB (publ)