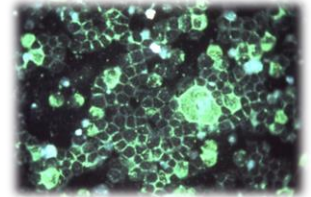


Respiratory Syncytial Virus (RSV) and Vaccination for Older Adults

WHAT IS RSV?^{1,2,3,4}

- **Respiratory syncytial virus (RSV)** is a common and contagious pulmonary pathogen that spreads through aerosolization of respiratory droplets or contact with contaminated surfaces.
- RSV produces cold-like symptoms causing significant morbidity and mortality, especially in infants and older adults.
- Adults who get infected with RSV typically have mild or no symptoms. Disease usually lasts less than 5 days. When disease is symptomatic, symptoms are usually consistent with an upper respiratory tract infection, including **rhinorrhea, pharyngitis, cough, headache, fatigue, and fever**.
- Some adults, however, may have more severe symptoms consistent with a lower respiratory tract disease (LRTD), such as pneumonia.
- The CDC estimates between 60k-160k older adults in the US are hospitalized and 6k-10k of them die due to RSV annually.
- 10-31% of older adults hospitalized with RSV are admitted into the ICU.
- For those admitted into the hospital with RSV, mortality rate is close to 8%.



WHAT RSV VACCINES ARE AVAILABLE?⁵

There are currently 2 FDA approved RSV vaccines available in the United States:

1. **AREXVY** (respiratory syncytial virus vaccine recombinant, adjuvanted; GSK; NDC: 58160-0848-11)
2. **ABRYOVO** (respiratory syncytial virus vaccine; Pfizer; NDC: 00069-0344-01)

HOW EFFECTIVE ARE THESE RSV VACCINES?⁵

In clinical trials, vaccination with a single dose of either RSV vaccine demonstrated moderate to high efficacy in preventing symptomatic RSV-associated LRTD among adults aged ≥60 years.

AREXVY

The efficacy of 1 dose of the GSK RSV vaccine in preventing symptomatic, laboratory-confirmed RSV-associated LRTD was 82.6% during the first RSV season and 56.1% during the second season. Efficacy of 1 dose over two seasons was 74.5% in preventing RSV-associated LRTD and 77.5% in preventing medically attended RSV-associated LRTD.

ABRYOVO

The efficacy of 1 dose of the Pfizer RSV vaccine in preventing symptomatic, laboratory-confirmed RSV-associated LRTD was 88.9% during the first RSV season and 78.6% during the partial second season. Efficacy of 1 dose over two seasons was 84.4% in preventing RSV-associated LRTD and 81.0% in preventing medically attended RSV-associated LRTD.

For reference, influenza vaccine efficacy is often found to be ~40-60% annually.⁶ These efficacy statements are abridged for conciseness. Consult the [MMWR](#) for more detailed findings.

HOW SAFE ARE THESE RSV VACCINES?⁵

Both vaccines were generally well-tolerated with an acceptable safety profile. Side effects after RSV vaccination are usually mild and included pain, redness, and swelling where the shot is given, fatigue, fever, headache, nausea, diarrhea, and muscle or joint pain.

In clinical trials, 6 cases of inflammatory neurologic events were reported post-vaccination (out of 38,177 study participants). Whether RSV vaccination increases the risk for inflammatory neurologic events (e.g., GBS) is currently unknown. Post-marketing surveillance is underway to clarify the existence of potential risks.

In the meantime, RSV vaccination in older adults should be targeted to those who are at highest risk for severe RSV disease and therefore, most likely to benefit from vaccination.

These safety statements are abridged for conciseness. Consult the [MMWR](#) for more detailed findings

FOR RESIDENTS

"In older adults with healthy immune systems, one dose of **Arexvy** was **83%** effective in preventing lung infections (e.g., pneumonia) due to RSV during the 1st RSV season and still **56%** effective in a 2nd season after vaccination.

Abrysvo was **89%** effective during the first RSV season after vaccination and appears to provide continued protection (2nd season is ongoing; data not finalized)."

WHAT ARE THESE VACCINES' INDICATIONS AND RECOMMENDATIONS FOR USE?⁵

INDICATION

Both RSV vaccines carry the same indication: active immunization for the prevention of LRTD caused by RSV in Individuals ≥ 60 years of age*.

CDC RECOMMENDATION

Persons aged ≥ 60 years may receive a single dose of either RSV vaccine, using shared clinical decision-making.

Coadministration of the RSV vaccine with other adult vaccines during the same visit is acceptable, however it may increase the risk of local or systemic reactogenicity. Vaccination should be delayed for those experiencing moderate to severe acute illness with or without fever as a precaution. Until further post marketing surveillance is completed, there is no further guidance available on the necessity of revaccination.

This recommendation to use **shared clinical decision-making** is intended to provide flexibility for providers and patients to consider **patient preferences** and **individual risk factors that increase the patient's risk for severe RSV disease**.

WHO IS AT INCREASED RISK FOR SEVERE RSV DISEASE?⁵

Chronic medical conditions associated with increased risk

- Chronic lung disease (Asthma, COPD)
- Chronic cardiovascular disease (CHF, CAD)
- Moderate to severe immunocompromise
- Neurologic or neuromuscular conditions
- Kidney or liver disorders
- Hematologic disorders
- Diabetes Mellitus

Other associated risk factors

- Frailty
- Advanced age
- Residence in a long-term care facility
- Other underlying conditions or factors that a healthcare provider determines might increase risk for severe RSV disease

VACCINE PRODUCT SPECIFICATIONS STORAGE

- **Before reconstitution**
 - **AREXVY**
 - **Adjuvant suspension component vials:** Store refrigerated between 2°C and 8°C (36°F and 46°F). Store in the original package in order to protect vials from light. Do not freeze. Discard if the adjuvant suspension component has been frozen.⁷
 - **Lyophilized antigen component vials:** Store refrigerated between 2°C and 8°C (36°F and 46°F). Store in the original package in order to protect vials from light. Do not freeze. Discard if the antigen component has been frozen.⁷
 - **ABRYSVO**
 - Store **kit** refrigerated at 2°C to 8°C (36°F to 46°F) in the original carton. Do not freeze. Discard if the carton has been frozen.⁸
- **After reconstitution**
 - **AREXVY**
 - **Administer immediately or store in the refrigerator** between 2°C and 8°C (36°F to 46°F) or at room temperature [up to 25°C (77°F)] for up to 4 hours prior to use. Protect vials from light. Discard reconstituted vaccine if not used within 4 hours. Do not freeze. Discard if the vaccine has been frozen.⁷
 - **ABRYSVO**
 - After reconstitution, **administer immediately or store at room temperature** [15°C to 30°C (59°F to 86°F)] and use within 4 hours. Do **not** store reconstituted vaccine under refrigerated conditions [2°C to 8°C (36°F to 46°F)]. Do not freeze reconstituted vaccine.⁸

*ABRYSVO also carries an indication for the active immunization of pregnant individuals at 32 through 36 weeks gestational age for the prevention of LRTD and severe LRTD caused by RSV in infants from birth through 6 months of age.

PREPARATION

- **AREXVY** is supplied in 2 vials that must be combined prior to administration.
- Prepare AREXVY by reconstituting the lyophilized antigen component (a sterile white powder) with the accompanying adjuvant suspension component (an opalescent, colorless to pale brownish sterile liquid). Use only the supplied adjuvant suspension component for reconstitution. The reconstituted vaccine should be an opalescent, colorless to pale brownish liquid. Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. If either of these conditions exists, the vaccine should not be administered.⁷



Figure 1. Cleanse both vial stoppers. Using a sterile needle and sterile syringe, withdraw the entire contents of the vial containing the adjuvant suspension component (liquid) by slightly tilting the vial. Vial 1 of 2.

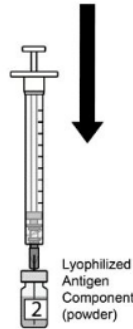


Figure 2. Slowly transfer entire contents of syringe into the lyophilized antigen component vial (powder). Vial 2 of 2.



Figure 3. Gently swirl the vial until powder is completely dissolved. **Do not shake vigorously.**

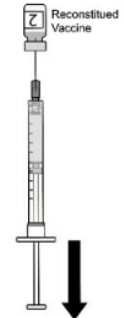
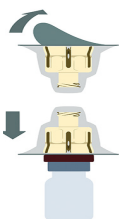


Figure 4. After reconstitution, withdraw 0.5 mL from the vial containing the reconstituted vaccine and administer intramuscularly.

- **ABRYSVO** is supplied in a kit that includes a vial of Lyophilized Antigen Component (a sterile white powder), a prefilled syringe containing Sterile Water Diluent Component and a vial adapter.⁸



CLICK

STEP 1. Open and attach vial adapter

- Open the vial adapter packaging by peeling off the top cover, but do not remove the adapter
- Align the adapter spike over the center of the vial's rubber stopper
- Connect the vial adapter to the vial with a straight downward push, locking it into place. **Do not push at an angle**



STEP 2. Remove cap and connect syringe

- Remove the syringe cap while holding the Luer lock adapter
- Continue holding the syringe by the Luer lock and connect it to the vial adapter by turning clockwise. **Do not overtighten**



MIX

STEP 3. Inject diluent and gently swirl

- Inject the entire contents of the syringe containing the sterile water diluent into the vial. **Do not remove the empty syringe**
- While holding the plunger rod down, gently swirl the vial until the powder is completely dissolved. **Do not shake**



PREP

STEP 4. Withdraw the contents

- Slowly withdraw the entire contents of the vial into the syringe to ensure an approximately 0.5 mL dose for administration

STEP 5. Disconnect syringe and attach needle

- Disconnect the syringe from the vial by holding the Luer lock adapter and turning counter-clockwise
- Attach a sterile needle suitable for intramuscular injection to the syringe
- Inspect prepared vaccine for particulate matter or discoloration prior to administration. **Do not use if either is present**

STORAGE AND HANDLING

Storage after reconstitution: ABRYSVO should be administered immediately or stored at room temperature [15°C to 30°C (59°F to 86°F)] and used within 4 hours. Do not store reconstituted vaccine under refrigerated conditions [2°C to 8°C (36°F to 46°F)]. Do not freeze reconstituted vaccine.



Scan the code or visit
[ABRYSVOPrep.com](https://www.abrysvoprep.com) to watch
a video on how to prepare ABRYSVO

ADMINISTRATION

- **AREXVY** is administered as a 0.5mL intramuscular injection only.⁷
 - **ABRYSVO** is administered as a 0.5mL intramuscular injection only.⁸
- Consult the [ACIP Vaccine Administration General Best Practice Guidelines for Immunization](#) for guidance on administering IM injections.

ADVERSE EFFECTS

- **AREXVY**
 - The most commonly reported solicited local adverse reaction ($\geq 10\%$) was injection site pain (60.9%). The most commonly reported solicited systemic adverse reactions ($\geq 10\%$) were fatigue (33.6%), myalgia (28.9%), headache (27.2%), and arthralgia (18.1%).⁷
- **ABRYVO**
 - The most commonly reported solicited local and systemic adverse reactions in individuals 60 years of age and older ($\geq 10\%$) were fatigue (15.5%), headache (12.8%), pain at the injection site (10.5%), and muscle pain (10.1%).⁸ The most commonly reported solicited local and systemic adverse reactions in pregnant individuals ($\geq 10\%$) were pain at the injection site (40.6%), headache (31.0%), muscle pain (26.5%), and nausea (20.0%).

HOW ARE RSV VACCINES CODED FOR AND DOCUMENTED IN EHRs?

RSV vaccines may be identified by sale proprietary (brand) name, generic name, description, and/or NDC.

Respiratory Syncytial Virus (RSV) Vaccine Codes Fall 2023								
CVX Code	CVX Description	Sale Proprietary Name	Sale Labeler	Unit of Sale NDC11	Unit of Use NDC11	Presentation	CPT Code	CPT Description
303	Respiratory syncytial virus (RSV), vaccine, recombinant, protein subunit RSV prefusion F, adjuvant reconstituted, 0.5 mL, preservative free	Arexvy	GlaxoSmithKline Biologicals SA	58160-0848-11	58160-0723-03	VIAL, 0.5 mL, reconstituted	90679	Respiratory syncytial virus vaccine, preF, recombinant, subunit, adjuvanted, for intramuscular use
305	Respiratory syncytial virus (RSV), vaccine, bivalent, protein subunit RSV prefusion F, diluent reconstituted, 0.5 mL, preservative free	Abrysvo	Pfizer Laboratories DivPfizer Inc	00069-0344-01 00069-0344-05 00069-0344-10	00069-0207-01	VIAL, 0.5 mL, reconstituted	90678	Respiratory syncytial virus vaccine, preF, subunit, bivalent, for intramuscular use

HOW ARE RSV VACCINES BILLED?

- RSV vaccines are covered by Medicare Part D, with no patient copay.
- They can be billed to Part D regardless of SNF Part A or long-stay status and they are not subject to consolidated billing.
- RSV vaccines have been covered by commercial plans for eligible recipients (i.e., those who meet the FDA-approved indication and CDC recommendations) but may carry copay liabilities on the part of the patient.

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