

Product Information Quick Guide

The only RSV vaccine
in a pre-filled syringe¹⁻³

RSV = respiratory syncytial virus.

INDICATION

mRESVIA® (Respiratory Syncytial Virus Vaccine) is a vaccine indicated for active immunization for the prevention of lower respiratory tract disease (LRTD) caused by respiratory syncytial virus (RSV) in adults 60 years of age and older.

IMPORTANT SAFETY INFORMATION

Contraindications

Do not administer mRESVIA to individuals with a history of severe allergic reaction (e.g., anaphylaxis) to any component of mRESVIA.

Please see Important Safety Information throughout and click for [mRESVIA Full Prescribing Information](#).

moderna®

Storage and Handling

Options for storing mRESVIA¹

Your mRESVIA pre-filled syringes can be stored in a freezer, refrigerator, or at room temperature, with varying storage times.

Refer to the content below for specifics.



In the freezer

-40°F to 5°F (-40°C to -15°C)



In the fridge

36°F to 46°F (2°C to 8°C)
for up to **30 days***



At room temp

46°F to 77°F (8°C to 25°C)
for up to **24 hours[†]**



Special precautions for storage¹

- **Do not** refreeze after thawing
- **Do not** refrigerate after thawing at room temperature
- Avoid exposure to direct sunlight and ultraviolet light
- Minimize exposure to room light

*Following frozen storage.¹

[†]Following frozen or refrigerated storage.¹

IMPORTANT SAFETY INFORMATION (CONT'D)

Warnings and Precautions

- **Management of Acute Allergic Reactions:** Appropriate medical treatment must be immediately available to manage potential anaphylactic reactions following administration of mRESVIA.

Please see Important Safety Information throughout and click for [mRESVIA Full Prescribing Information](#).

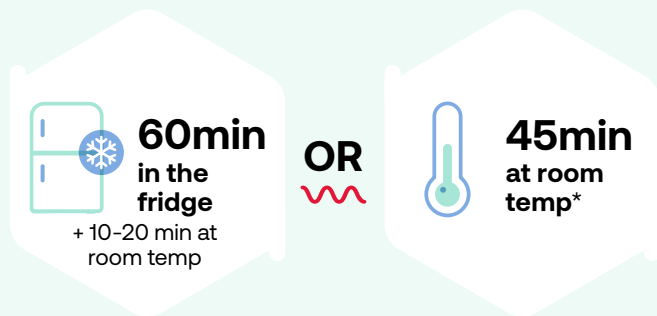
Dosage and Administration

Preparing to administer mRESVIA¹

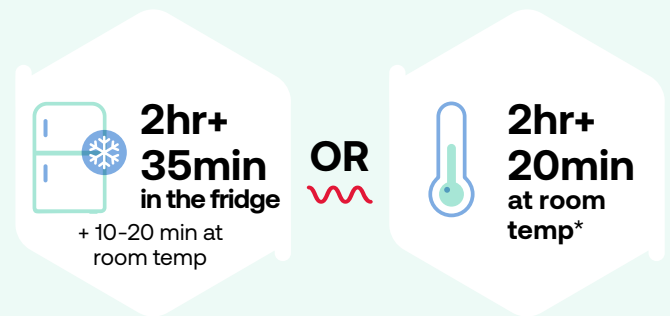
mRESVIA is a single 0.5 mL dose pre-filled syringe containing a frozen suspension that must be thawed. At room temperature, mRESVIA is ready to use.

Refer to the images below for approximate thaw times.

Thawing a syringe in a blister pack



Thawing a carton of 10 syringes



Special precautions for thawing¹

- **Do not** shake mRESVIA
- **Do not** return syringes to the refrigerator after they have been thawed to room temperature
- Discard any syringes left at room temperature for over 24 hours

*Thaw at 59°F to 77°F (15°C to 25°C), rather than at 46°F to 77°F (8°C to 25°C), as allowed for storage.¹

IMPORTANT SAFETY INFORMATION (CONT'D)

Warnings and Precautions (cont'd)

- **Syncope:** Syncope (fainting) may occur in association with administration of injectable vaccines, including mRESVIA. Procedures should be in place to avoid injury from fainting.

Please see Important Safety Information throughout and click for [mRESVIA Full Prescribing Information](#).

Administering mRESVIA with a pre-filled syringe⁴

Your mRESVIA pre-filled syringe may look and feel slightly different than a standard pre-filled syringe, as it is designed with a Luer lock.

Refer to the steps below for guidance.



< Tip cap

Luer lock >

< Plunger

Syringe barrel >

Plunger rod >

- 1 Hold the syringe vertically, with the tip cap facing upright
- 2 Grip the tip cap using one hand
- 3 Twist the tip cap counter-clockwise. **You may feel resistance.** Avoid pulling or applying pressure to the tip cap as you twist
- 4 Remove the tip cap in a slow, steady motion once released
- 5 Select a sterile Luer lock needle of appropriate length and gauge for intramuscular injection
- 6 Administer the entire dose, then discard the syringe

Always inspect your mRESVIA syringe prior to administration.¹

mRESVIA is a white to off-white suspension that may contain visible white or translucent product-related particulates. **Do not** administer the vaccine if it is discolored or contains other particulate matter.

IMPORTANT SAFETY INFORMATION (CONT'D)

Warnings and Precautions (cont'd)

- **Altered Immunocompetence:** Immunocompromised individuals, including those receiving immunosuppressive therapy, may have a diminished immune response to mRESVIA.

Please see Important Safety Information throughout and click for [mRESVIA Full Prescribing Information](#).

Reimbursement, Ordering, and Reporting Codes


Most payers, including Medicare Part D, process claims for mRESVIA under the pharmacy benefit.

VACCINE TYPE	NDC 10/NDC 11 LABELER PRODUCT ID	CVX CODE
mRESVIA (Carton of 10 Pre-Filled Syringes)	NDC 80777-345-96	326

Some payers, including Medicaid and commercial, may allow access in an office setting and process under the medical benefit.

VACCINE CPT CODE (Carton)	90683 Respiratory syncytial virus vaccine, mRNA lipid nanoparticles, for intramuscular use	VACCINE ADMINISTRATION CPT CODES (Carton)	90471 Immunization administration (1 vaccine) 90472 Immunization administration (each additional vaccine)
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Order mRESVIA today.
It's easy on [modernadirect.com](https://www.modernadirect.com)

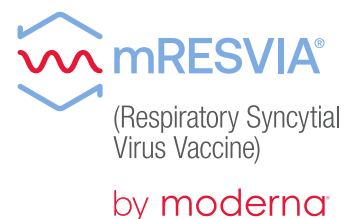
For any questions related to billing, contact Moderna Customer Care at: [1-866-MODERNA \(1-866-663-3762\)](tel:1-866-MODERNA)
8:00 AM - 8:00 PM ET

IMPORTANT SAFETY INFORMATION (CONT'D)

Adverse Reactions

In a clinical trial, the most commonly reported (≥10%) adverse reactions were injection-site pain (55.9%), fatigue (30.8%), headache (26.7%), myalgia (25.6%), arthralgia (21.7%), axillary (underarm) swelling or tenderness (15.2%) and chills (11.6%).

Please see Important Safety Information throughout and click for [mRESVIA Full Prescribing Information](#).



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- **Altered Immunocompetence:** Immunocompromised individuals, including those receiving immunosuppressive therapy, may have a diminished immune response to mRESVIA.

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To report suspected adverse reactions, contact ModernaTX, Inc. at 1-866-663-3762 or VAERS at 1-800-822-7967 or www.vaers.hhs.gov.

Please click for [mRESVIA Full Prescribing Information](#).

Colorado and Connecticut prescribers and pharmacists may view the price disclosure at <https://modernadirect.com/wac-disclosure>

References: 1. mRESVIA Prescribing Information. ModernaTX, Inc. 2. AREXVY Prescribing Information. GlaxoSmithKline Biologics SA. 3. ABRYSVO Product Information. Pfizer Inc. 4. Data on file. Moderna, Inc; 2024.

