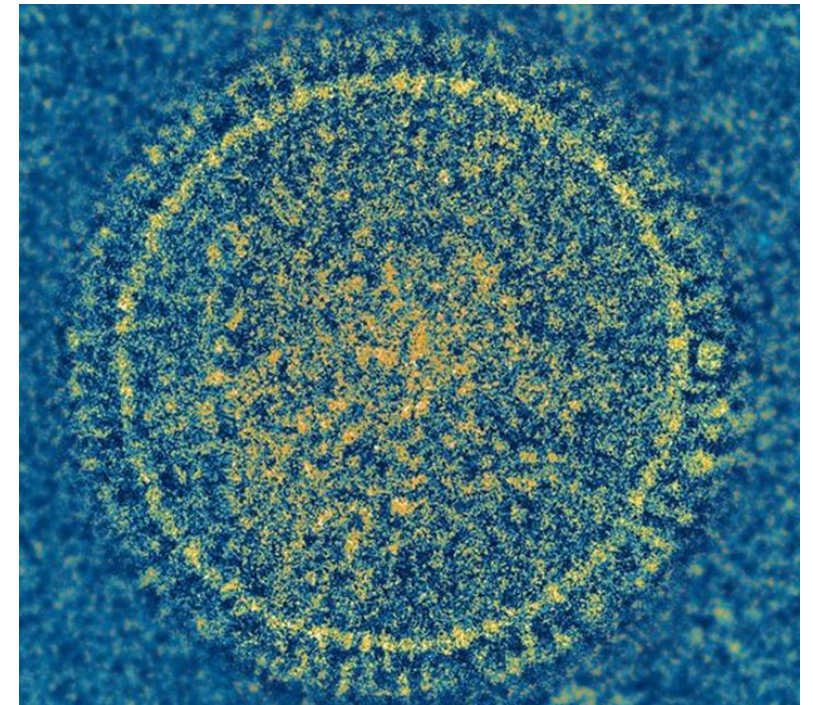


Maternal/Pediatric Respiratory Syncytial Virus (RSV) Work Group

Sarah S. Long, MD
Chair, Maternal/Pediatric RSV Work Group

ACIP General Meeting
August 3, 2023



Work group members (External)

ACIP Members

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Patsy Stinchfield (NFID)

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Anne Hause

Andrew Leidner

David Shay

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CDC ACIP Staff

Melinda Wharton

Stephanie Thomas

Jessica MacNeil

Previous maternal/pediatric RSV ACIP presentations on nirsevimab

- Epidemiology and burden of RSV in infants
- Virology and immunology of RSV
- Safety and efficacy of nirsevimab
- Cost effectiveness analysis for nirsevimab – CDC model
- Cost effectiveness analysis for nirsevimab – Comparison with manufacturer model
- Evidence to Recommendations framework for nirsevimab
- Clinical considerations for nirsevimab

Agenda: Thursday August 3, 2023

- Evidence to Recommendations Framework for nirsevimab Dr. Jefferson Jones (CDC)
- Nirsevimab implementation considerations Dr. Georgina Peacock (CDC)
- Clinical considerations for nirsevimab Dr. Jefferson Jones (CDC)
- Workgroup considerations and proposed recommendations and voting language Dr. Jefferson Jones (CDC)
- Vaccines for Children Resolution Dr. Jeanne Santoli (CDC)

For more information, contact CDC
1-800-CDC-INFO (232-4636)
TTY: 1-888-232-6348 www.cdc.gov

The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.



Proposed ACIP Voting Language

- Infants aged <8 months born during or entering their first RSV season are recommended to receive one dose of nirsevimab (50 mg for infants <5 kg and 100 mg for infants \geq 5 kg)
- Children aged 8–19 months who are at increased risk of severe RSV disease and entering their second RSV season are recommended to receive one dose of nirsevimab (200 mg)