### National Center for Immunization and Respiratory Diseases



## Summary of effectiveness of nirsevimab in infants

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### Agenda

- Real-world vaccine/product effectiveness methods
- Effectiveness of nirsevimab in the United States
  - RSV-associated emergency department encounters & hospitalization, VISION
  - RSV-associated medical encounters and hospitalization, NVSN
- Effectiveness of nirsevimab globally
- Conclusions

# Real-world vaccine/product effectiveness context and methods

## **Efficacy** ≠ **effectiveness**

- Efficacy: the degree to which an immunization prevents disease under ideal and controlled conditions (i.e., measured in clinical trials)
- Effectiveness: the degree to which an immunization prevents disease under real-world conditions (i.e., measured in post-licensure observational studies)

In this presentation, we'll discuss **product effectiveness ("PE")** from real-world data.

# Insufficient supply of nirsevimab to meet demand in 2023-2024 season

- Limited supply of nirsevimab (100mg and 50mg formulations) meant clinicians were uncertain how to ration or prioritize few available doses
- CDC issued an official Health Advisory notice via the Health Alert Network to prioritize available doses to high-risk infants and younger infants
- By January, demand had decreased and additional supply was available allowing return to original recommendations

Limited Availability of Nirsevimab in the United States—Interim CDC Recommendations to Protect Infants from Respiratory Syncytial Virus (RSV) during the 2023–2024 Respiratory Virus Season

Print





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Health Alert Network (HAN) - 00499 | Limited Availability of Nirsevimab in the United States—Interim CDC Recommendations to Protect Infants from Respiratory Syncytial Virus (RSV) during the 2023–2024 Respiratory Virus Season

Observational effectiveness measured in a test-negative design (TND) study Kev features of a TND Real-world circumstances Most heterogenous study population, Person with acute often "all comers" Reduces bias from health-care seeking behavior respiratory illness Considerations Validity dependent on test performance Residual confounding is possible Case Control **RSV** test **RSV** immunization status

Effectiveness = 1 – (odds ratio) x 100% Odds ratio =  $\frac{Odds \ of \ immunization_{cases}}{Odds \ of \ immunization_{controls}}$ 

Virtual SARS-CoV-2, Influenza, and Other respiratory viruses Network (VISION)

### **VISION Multi-Site Network of Electronic Health Records (EHRs)**

### 127 emergency rooms and 107 hospitals

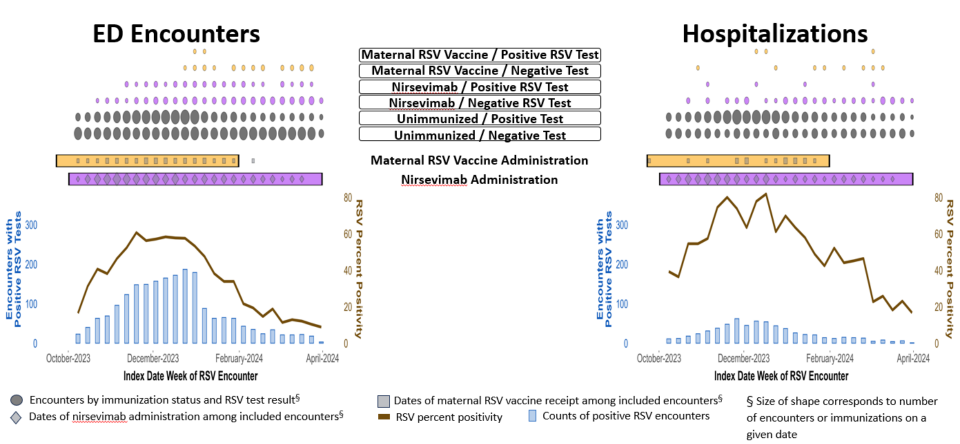
- Population: Visiting a participating ED for or hospitalized with RSV-like illness (RLI)\*
- Immunization data: Infant and maternal RSV immunization status documented by electronic health records, state and city registries, and claims data (subset of sites)
- Covariate data: Documented in electronic health records
  - Underlying medical conditions: ICD-10 discharge diagnosis codes at time of RLI encounter
  - Patient characteristics
    - Date of birth
    - Census tract of residence
    - Sex



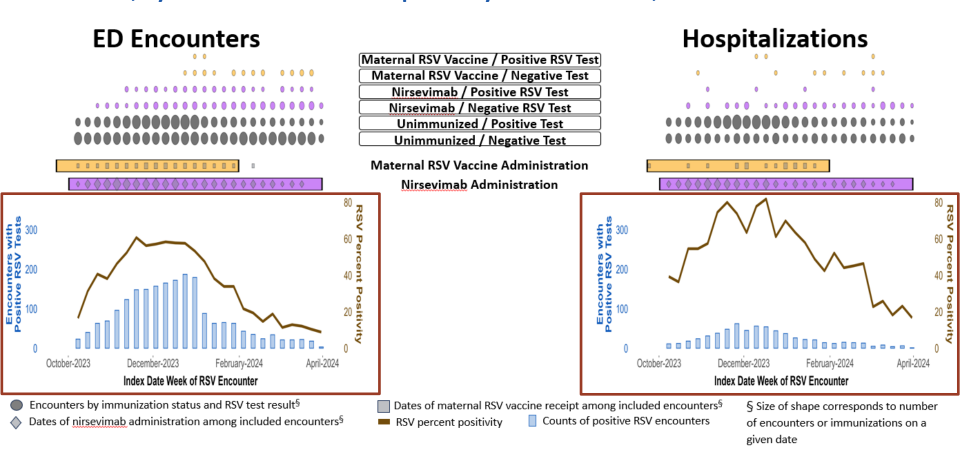
### VISION 2.0 partners included in this analysis –

**ED**: Columbia, HealthPartners Institute, Intermountain Healthcare, KPSC, KPCHR, Regenstrief

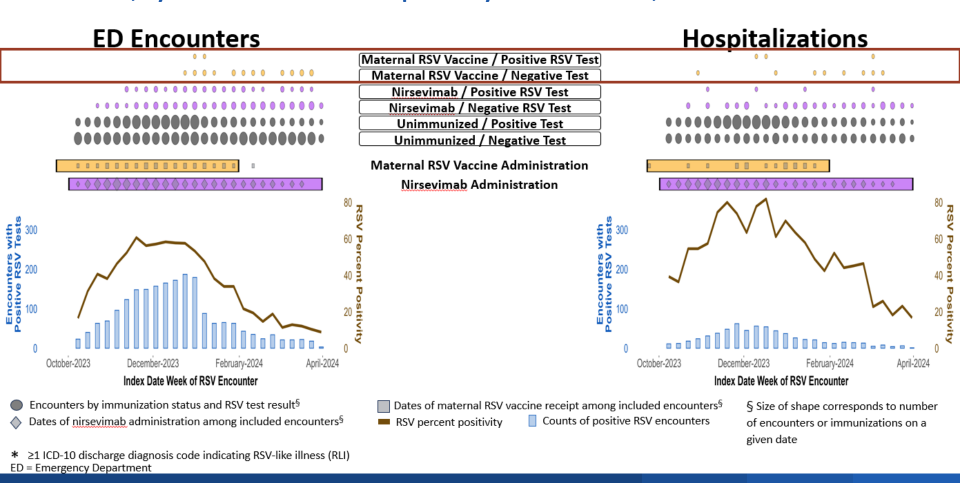
**Inpatient**: Columbia, HealthPartners Institute, Intermountain, KPSC, KPCHR, Regenstrief

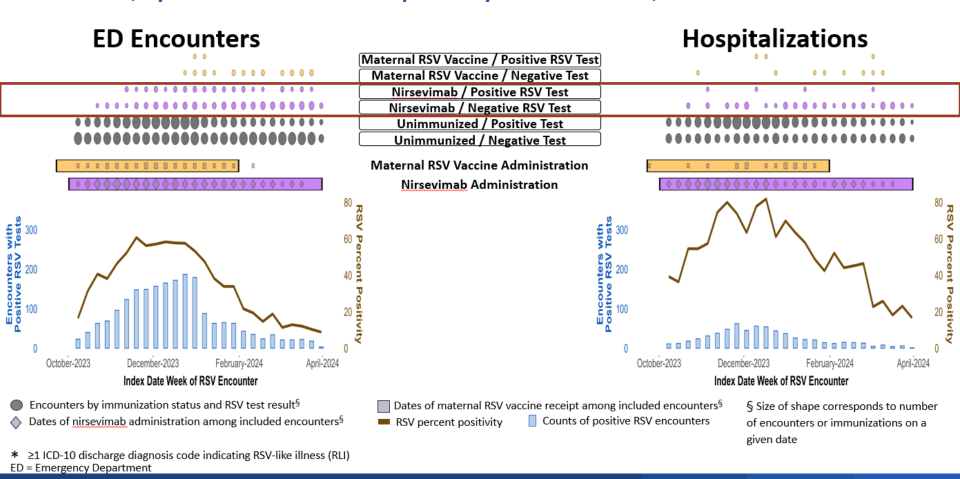


<sup>\* ≥1</sup> ICD-10 discharge diagnosis code indicating RSV-like illness (RLI) ED = Emergency Department



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## Test-negative design (TND) analysis of first season nirsevimab product effectiveness (PE) against RSV-associated ED encounters and hospitalization – VISION, October 8, 2023 – March 31, 2024

- Population:
  - Infants aged <8 months as of October 1, 2023, or born after October 1, 2023
  - Visiting a participating ED for or hospitalized with RSV-like illness (RLI)
  - With RSV test result within 10 days before or 72 hours after encounter
    - Cases: RLI with positive RSV antigen or NAAT test\*
    - Controls: RLI with negative RSV NAAT test
- Study period: October 8, 2023 March 31, 2024
- Exclusion criteria:
  - Children aged <7 days
  - Children born after September 22, 2023, without linkage to maternal records
  - Evidence of maternal RSV vaccination or palivizumab administration
  - Receipt of unrecommended nirsevimab dose(s)<sup>†</sup>
  - <7 days between nirsevimab dose and RLI encounter
  - Indeterminate RSV test result
- Statistical Analysis: Adjusted OR comparing odds of immunization<sup>‡</sup> among cases vs. controls estimated using multivariable logistic regression models, adjusting for age, race and ethnicity, sex, calendar day (days since Oct 8, 2023), and geographic region → PE = (1-aOR) X 100%

‡Immunization defined as one nirsevimab dose ≥7 days prior to encounter index date.

<sup>\*</sup>RSV-positive encounters with positive SARS-CoV-2 and/or influenza test result were (i.e., coinfections) were excluded.

<sup>†</sup> Unrecommended nirsevimab dose(s) defined as: nirsevimab doses administered on or before October 1, 2023, and receipt of >1 nirsevimab dose. Nirsevimab doses in older children may be administered as 2 injections on the same day; this was considered one 'dose'.

# First season nirsevimab product effectiveness (PE) against RSV-associated ED encounters and hospitalization – VISION, October 8, 2023 – March 31, 2024

Outcome   Nirsevimab dosage pattern	Total encounters	RSV-positive encounters N (Row %)	Median days since dose (IQR)	Adjusted PE (95% CI)*				
RSV-associated ED encoun	ter							
No nirsevimab doses	4,610	1,988 (43)	N/A	ref				
Nirsevimab, ≥7 days prior	442	63 (14)	53 (27-84)	77 (69-83)			۰	<b>0-1</b>
RSV-associated hospitaliza	tion							
No nirsevimab doses	927	601 (65)	N/A	ref				
Nirsevimab, ≥7 days prior	93	4 (4)	48 (25-84)	98 (95-99)				
					0 20	40	60	80 1

Nirsevimab was effective against RSV-associated ED encounters and hospitalization among infants in their first RSV season.

# Early Estimate of Nirsevimab Effectiveness for Prevention of Respiratory Syncytial Virus-Associated Hospitalization Among Infants Entering Their First Respiratory Syncytial Virus Season — New Vaccine Surveillance Network, October 2023-February 2024

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Update to Moline HL, Tannis A, Toepfer AP, et al. Early Estimate of Nirsevimab Effectiveness for Prevention of Respiratory Syncytial Virus—Associated Hospitalization Among Infants Entering Their First Respiratory Syncytial Virus Season — New Vaccine Surveillance Network, October 2023—February 2024. MMWR Morb Mortal Wkly Rep 2024;73:209–214. DOI: <a href="http://dx.doi.org/10.15585/mmwr.mm7309a4">http://dx.doi.org/10.15585/mmwr.mm7309a4</a>

# **New Vaccine Surveillance Network (NVSN)**

# NVSN is a prospective, population-based surveillance network for pediatric acute respiratory illness (ARI) at 7 U.S. medical centers.





Children <18 years of age with ARI are enrolled year-round in the **outpatient**, **urgent care**, **emergency department (ED)**, **and hospital** settings.

### Surveillance Objectives:

- Determine the etiology and burden of laboratoryconfirmed acute viral respiratory diseases in children
- Characterize the clinical and epidemiologic factors of pediatric ARI and associated syndromes
- Evaluate vaccine effectiveness (VE) using a testnegative design (TND) and impact of vaccines and other immunoprophylaxis products.



### **NVSN Data Collection**

### Caregiver interview

- Race and ethnicity, preterm status, date of symptom onset, breastfeeding status

### Specimens

- Mid-turbinate nasal swab collected from all children for RSV testing by reversetranscription polymerase chain reaction; results of both clinical and surveillance testing are collected
- Sequencing of RSV-positive specimens to monitor for substitutions in the nirsevimab binding site

### Medical chart abstraction

- Age, underlying medical conditions, clinical course of illness, insurance status

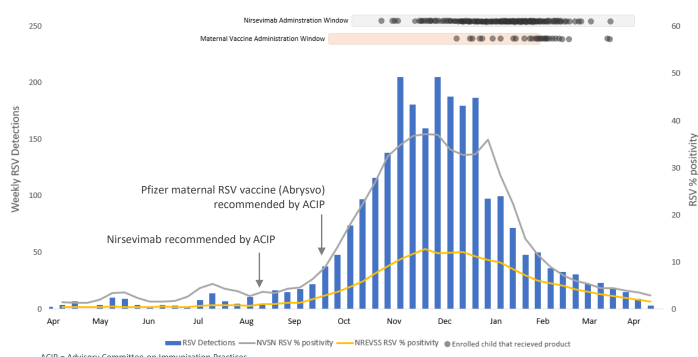
### Immunization status (nirsevimab, palivizumab, and maternal RSV vaccine)

- Ascertained by parent report and confirmed with state immunization information system, electronic health record, or birth record





RSV Detections and RSV Product Receipt in NVSN, April 2023 through April 2024



ACIP = Advisory Committee on Immunization Practices
NVSN = New Vaccine Surveillance Network
NREVSS = National Respiratory and Enteric Virus Surveillance System

All NVSN sites had some nirsevimab availability by mid-October

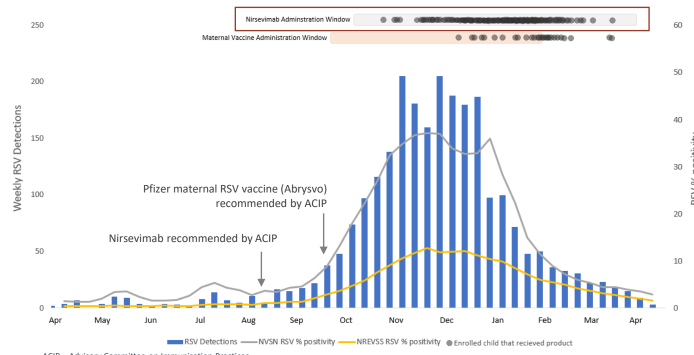
Among 2,383 infants in their first RSV season

- 11.1% received nirsevimab
- 2.2% received palivizumab
- 3.4% (of 1,542 infants <6 months of age at enrollment) had history of maternal RSV vaccination









ACIP = Advisory Committee on Immunization Practices NVSN = New Vaccine Surveillance Network NREVSS = National Respiratory and Enteric Virus Surveillance System All NVSN sites had some nirsevimab availability by mid-October

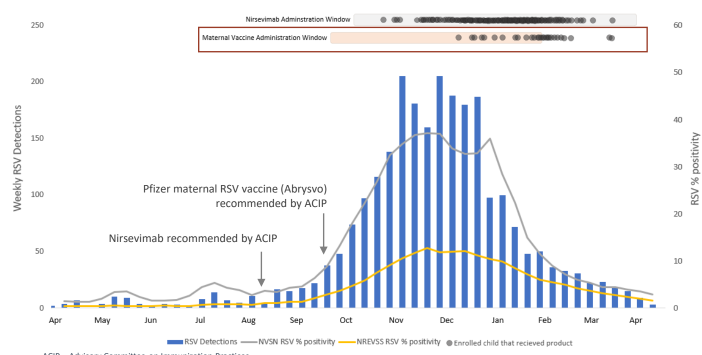
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Test-negative design (TND) analysis of first season nirsevimab product effectiveness (PE) against medically attended RSV-associated ARI episodes and RSV-associated hospitalization – NVSN, October 2023 – March 2024

### Population:

- Infants <8 months as of October 1, 2023, or born after October 1, 2023
- Enrolled from participating medical center
- ARI\*
  - Case patients children with medically attended ARI who tested positive for RSV by surveillance or clinical testing
- Control patients children with medically attended ARI who tested negative for RSV by surveillance or clinical testing
- Study period: October 2023 March 2024<sup>†</sup>

### Exclusion criteria:

- Chart review incomplete for underlying conditions, preterm status, insurance status, highest level of care, clinical course of illness
- Immunization status unverified for nirsevimab and palivizumab receipt and maternal RSV vaccination
- Receipt of palivizumab or history of maternal RSV vaccination during pregnancy
- Unknown or inconclusive RSV test result
- Receipt of nirsevimab <7 days prior to ARI symptom onset</li>
- Statistical Analysis: Adjusted OR comparing odds of immunization<sup>‡</sup> among cases vs. controls estimated using multivariable logistic regression models, adjusting for site, age in months, month of enrollment, and presence of >1 high-risk medical condition for severe RSV disease → PE = (1-aOR) x 100%

<sup>\*</sup>Acute respiratory illness (ARI) defined as >1 of the following sign/symptoms: fever, cough, earache, nasal congestion, runny nose, sore throat, vomiting after coughing, shortness of breath (rapid or shallow breathing), wheezing, apnea, or apparent life-threatening event or brief resolved unexplained event

<sup>†</sup>State-level RSV RT-PCR percent positivity thresholds of 3% were used to define the beginning and end weeks of the analysis by site

<sup>†</sup>Immunization defined as one nirsevimab dose ≥7 days prior to symptom onset. OR = odds ratio | aOR = adjusted odds ratio | PE = product effectiveness

# First season nirsevimab product effectiveness (PE) against medically attended RSV-associated ARI and RSV-associated hospitalization – NVSN, October 2023 – March 2024\*

Outcome   Nirsevimab dosage pattern	Total encounters	RSV-positive encounters N (Row %)	Median days since dose (IQR)		Adjuste E (95% (		
Medically Attended RSV-associated ARI episode <sup>‡</sup>							
No nirsevimab doses	1,575	755 (48)	N/A	ref			
Nirsevimab, ≥7 days prior <sup>§</sup>	120	9 (8)	42 (21-73)	89 (77-94)			-
RSV-associated hospitaliza	tion						
No nirsevimab doses	807	526 (65)	N/A	ref			
Nirsevimab, ≥7 days prior	63	6 (10)	38 (15-67)	91 (79-96)			
					0 20	40	50 80 10

Nirsevimab was effective against medically attended RSV-associated ARI episodes and RSV-associated hospitalization.

<sup>\*</sup>State-level RSV RT-PCR percent positivity thresholds of 3% were used to define the beginning and end weeks of the analysis by site

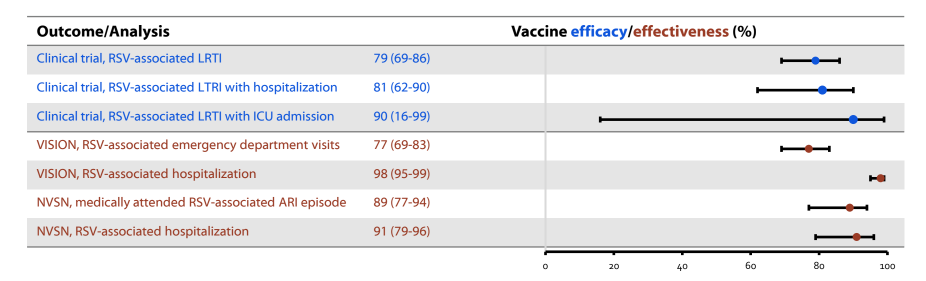
<sup>†</sup>Multivariable logistic regression models compared the odds of vaccination among RSV case and control patients while adjusting for site, age in months, month of enrollment, and presence of >1 high-risk medical condition for severe RSV disease.

<sup>§</sup>Immunization defined as one nirsevimab dose ≥7 days prior to symptom onset.

ARI = acute respiratory illness | N/A = not applicable | ref = reference group

# **Summary of US data**

# Observational data indicate nirsevimab is working as expected (vs. RCT results) during the first RSV season after approval among infants in their first RSV season



Results may not be comparable across studies due to differences in outcome definitions, timing, and other factors.

# Limitations of test-negative design (TND) analyses of first season nirsevimab product effectiveness (PE), October 2023 – March 2024

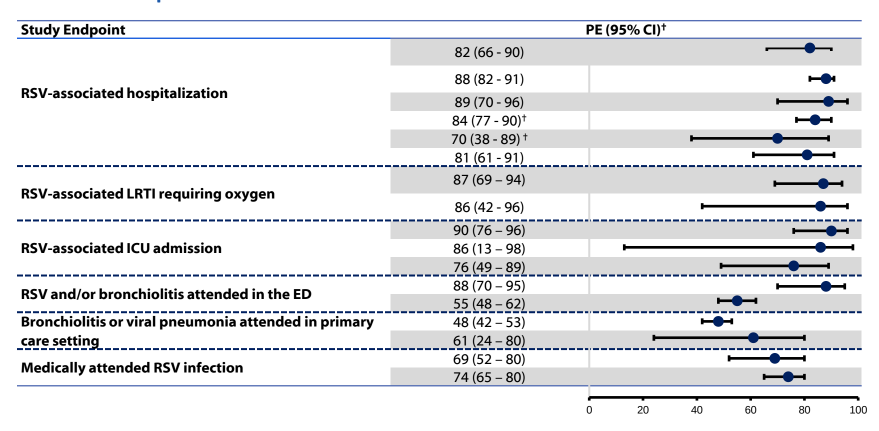
- High product effectiveness should be interpreted with caution
  - Short interval from administration to respiratory illness onset
  - Unable to assess duration of protection during the 2023-2024 RSV season
- Residual confounding was possible
- Misclassification of RSV immunization status was possible
- These results only reflect PE among infants in their first RSV season (not among children at increased risk in their second RSV season)
- VISION:
  - Cases may have sought care for something other than RSV
  - All RSV testing was clinician-directed
  - EHR data may not fully capture all underlying medical conditions, which may be associated with likelihood of immunization and risk of severe RSV disease

### NVSN:

- May not be nationally representative

# Nirsevimab effectiveness – evidence from literature

# Nirsevimab product effectiveness (PE) among infants in their first RSV season – Data\* from Spain and France



<sup>\*</sup>References provided on backup slide 32.

LRTI = lower respiratory tract infection | ICU = intensive care unit

<sup>&</sup>lt;sup>†</sup>PE estimates generated from the same study, using different methods.

# **Conclusions**

### **Conclusions**

- Nirsevimab was effective against RSV-associated ED encounters and hospitalization among infants in their first RSV season during the 2023-2024 RSV season
- Due to timing of authorization/recommendation of RSV prevention products and RSV activity during the 2023-2024 RSV season:
  - US-based analyses may be subject to residual confounding due to prioritization of nirsevimab doses
  - Short time between nirsevimab administration and outcomes, limiting ability to assess duration of protection
  - Limited ability to assess effectiveness of maternal RSV vaccines
- Ongoing monitoring of post-licensure nirsevimab and maternal RSV vaccine effectiveness will continue

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## Hospitalized population comparison for VISION and NVSN

	VISION, no. (col %)			NVSN, no. (col %)			
Characteristic	Total no. of patients	RSV case-patients	RSV control-patients	Total no. of patients	RSV case-patients	RSV control-patients	
All hospitalizations	1,020	605	415	870	532	338	
Median age, months (IQR)	4 (1-7)	3 (1-6)	4 (1-7)	3 (1-6)	3 (1-5)	3 (1-6)	
Gestational age							
Preterm (<37 weeks)	133 (13)	59 (10)	74 (18)	180 (21)	102 (19)	78 (23)	
Term (≥37 weeks)	508 (50)	323 (53)	185 (45)	687 (79)	428 (81)	259 (77)	
Unknown	379 (37)	223 (37)	156 (38)	3 (0)	2 (0)	1 (0)	
Race/ethnicity							
Black or African American, Non-Hispanic	77 (8)	48 (8)	29 (7)	113 (13)	56 (11)	57 (17)	
White, Non-Hispanic	437 (43)	268 (44)	169 (41)	390 (45)	266 (50)	124 (37)	
Hispanic or Latino	394 (39)	234 (39)	160 (39)	248 (29)	146 (27)	102 (30)	
Other, Non-Hispanic	75 (7)	33 (6)	42 (10)	107 (12)	58 (11)	49 (14)	
Unknown	37 (4)	22 (4)	15 (4)	12 (1)	6 (1)	6 (2)	
High risk conditions for severe RSV disease							
None	796 (78)	529 (87)	267 (64)	832 (96)	520 (99)	281 (83)	
≥1	224 (22)	76 (13)	148 (36)	38 (4)	12 (2)	26 (8)	
Immunization status							
No nirsevimab	927 (91)	601 (99)	326 (79)	807 (93)	526 (99)	281 (83)	
Nirsevimab, ≥7 days earlier	93 (9)	4 (1)	89 (21)	63 (7)	6 (1)	57 (17)	

### Empirical studies\* on nirsevimab product effectiveness (PE) among infants in their first RSV season

Citation	Country	Sample Size (Number of Infants)	Study Design	PE (95% Confidence Interval)
Ares-Gomez et al., 2024	Spain	10,259	Prospective Cohort	Hospitalization for RSV-related LRTI: 82% (95% CI: 66% - 90%) Severe RSV-related LRTI requiring oxygen support: 87% (95% CI: 69% - 94%) All-cause LRTI hospitalizations: 69% (56% - 78%) All-cause hospitalizations: 66% (56% - 74%)
Coma et al., 2024	Spain	26,525	Retrospective Cohort	Hospital admission for RSV-related disease: 88% (95% CI: 82% - 91%) Hospital ER visits due to bronchiolitis: 55% (95% CI: 48% - 62%) Medically attended RSV infection: 69% (95% CI: 52% - 80%) Primary care attended bronchiolitis: 48% (95% CI: 42% - 53%) Viral pneumonia diagnosed in primary care: 61% (95% CI: 24% - 80%) ICU admission for RSV-related disease: 90% (95% CI: 76% - 96%)
Estrella-Porter et al., 2024	Spain	27,362	Retrospective Cohort	Medically attended RSV infection: 74% (95% CI: 65% - 80%)
Ezpeleta et al., 2024	Spain	1,177	Prospective Cohort	Hospitalization due to RSV: 89% (95% CI: 70% - 96%) RSV infection attended in the ER: 88% (95% CI: 70% - 95%) RSV ICU admission: 86% (95% CI: 13% - 98%)
Lopez-Lacort et al., 2024	Spain	166	Screening and Test negative case control	RSV-LRTI hospital admission (pooled data across several regions): Screening methods: 84% (95% CI: 77% - 90%) Test negative design: 70% (95% CI: 38% - 89%)
Paireau et al., 2024	France	288	Test negative case control	RSV bronchiolitis hospitalized In the pediatric ICU: 76% (95% CI: 49% - 89%)
Aguera et al., 2024	Spain	181	Test negative case control	Hospitalization for RSV-related LRTI: 81% (95% CI: 61% - 91%) Severe RSV-related LRTI requiring NIV/CMV: 86% (95% CI: 42% - 96%)

<sup>\*</sup>Published during June 20, 2023, through June 21, 2024

LRTI = lower respiratory tract infection | ER = emergency room | ICU = intensive care unit | CI = confidence interval | NIV: noninvasive ventilation | CMV: continuous mandatory ventilation