

Observational Cohort Study Evaluating Real-World ABRYSCO Vaccine Effectiveness and Impact Against Medically-Attended RSV-related and All-Cause Outcomes Among Infants Born to Individuals Vaccinated During Pregnancy

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Study

Ongoing

Administrative details

EU PAS number

EUPAS1000000295

Study ID

1000000295

DARWIN EU® study

No

Study countries

Study description

This study will be conducted in collaboration with an integrated delivery health care organization in the United States using electronic medical record (EMR) data, collected during routine standard of care clinical encounters. This study will use a retrospective cohort design to study the vaccine effectiveness (VE) and impact of ABRYSV0 vaccination during pregnancy in a real-world population over multiple RSV seasons.

Study status

Ongoing

Research institutions and networks

Institutions

[Kaiser Permanente Northern California \(KPNC\)](#)

Networks

[Kaiser Permanente Northern California](#)

Contact details

Study institution contact

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Study contact

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Primary lead investigator

Nicola Klein

Primary lead investigator

Study timelines

Date when funding contract was signed

Planned: 18/06/2024

Actual: 13/11/2024

Study start date

Planned: 01/12/2025

Actual: 01/12/2025

Date of final study report

Planned: 31/03/2028

Sources of funding

- Pharmaceutical company and other private sector

More details on funding

Pfizer INC

Regulatory

Was the study required by a regulatory body?

No

Is the study required by a Risk Management Plan (RMP)?

Not applicable

Other study registration identification numbers and links

C3671048

Methodological aspects

Study type

Study type list

Study topic:

Disease /health condition
Human medicinal product

Study topic, other:

Vaccine effectiveness

Study type:

Non-interventional study

Scope of the study:

Effectiveness study (incl. comparative)

Data collection methods:

Secondary use of data

Study design:

This study will use a retrospective cohort design to assess ABRYSV0 VE and impact in a large, diverse, real-world setting. This NI study will be conducted within an integrated delivery health care organization using EMR data, collected during routine standard of care clinical encounters.

Main study objective:

The overall aim of this study is to evaluate the effectiveness and impact of maternal ABRYSV0 vaccination during pregnancy for the prevention of medically-attended (MA) respiratory syncytial virus (RSV)-associated and all-cause infant outcomes in a large, diverse, real-world population.

Study Design

Non-interventional study design

Cohort

Other

Non-interventional study design, other

The primary objective is to estimate VE of ABRYSV0 vaccination during pregnancy against RSV LRTD among infants from birth to ≤ 180 days of age. The key secondary objective is to estimate VE of ABRYSV0 vaccination during pregnancy against RSV LRTD hospitalization among infants from birth to ≤ 180 days of age. Secondary objectives include: 1) To estimate VE of ABRYSV0 vaccination during pregnancy against RSV among infants from birth to ≤ 180

days of age. 2) To estimate VE of ABRYSV0 vaccination during pregnancy against RSV hospitalization among infants from birth to ≤ 180 days of age. 3) To estimate VE of ABRYSV0 vaccination during pregnancy against all-cause LRTD among infants from birth through the RSV season (to ≤ 180 days of age). 4) To estimate VE of ABRYSV0 vaccination during pregnancy against all-cause LRTD hospitalization among infants from birth through the RSV season (to ≤ 180 days of age). 5) To estimate VE of ABRYSV0 vaccination during pregnancy against acute otitis media among infants from birth through the RSV season (to ≤ 180 days of age). 6) To estimate VE of ABRYSV0 vaccination during pregnancy against first antibiotic prescription among infants from birth through the RSV season (to ≤ 180 days of age). 7) To estimate VE and interval-specific VE of ABRYSV0 vaccination during pregnancy against RSV among infants from birth to ≤ 360 days of age. 8) To estimate VE and interval-specific VE of ABRYSV0 vaccination during pregnancy against RSV hospitalization among infants from birth to ≤ 360 days of age. 9) To estimate VE and interval-specific VE of ABRYSV0 vaccination during pregnancy against RSV LRTD among infants from birth to ≤ 360 days of age. 10) To estimate VE and interval-specific VE of ABRYSV0 vaccination during pregnancy against RSV LRTD hospitalization among infants from birth to ≤ 360 days of age.

Study drug and medical condition

Medicinal product name

ABRYSV0

Study drug International non-proprietary name (INN) or common name

RESPIRATORY SYNCYTIAL VIRUS, SUBGROUP A, STABILIZED PREFUSION F
PROTEIN 847A

Anatomical Therapeutic Chemical (ATC) code

(J07BX05) respiratory syncytial virus vaccines
respiratory syncytial virus vaccines

Population studied

Short description of the study population

The study population will comprise eligible maternal-infant pairs over a 2-year study period, identified from EMR records in the existing databases, which accrue in real-time as pregnancies/births occur. All pregnancies that reach 32 weeks of gestation during the 2-year study period, from September 22, 2023 (start of ABRYSSVO vaccination season 1 as of the date of the ACIP recommendation) to January 31, 2025 (estimated end of ABRYSSVO vaccination season 2) will be eligible for inclusion, along with all live born infants from the eligible pregnancies.

Age groups

- Neonate
 - Preterm newborn infants (0 – 27 days)
 - Term newborn infants (0 – 27 days)
- Infants and toddlers (28 days – 23 months)

Study design details

Setting

The study setting will be KPNC, which is an integrated delivery healthcare organization in Northern California with over 4.5 million members and approximately 40,000 births each year. KPNC members receive almost all their health care within KPNC clinics, hospitals, pharmacies, and laboratories. Information from encounters within these clinical settings are captured in EMRs which can be linked at an individual level across KPNC settings using a unique medical record number (MRN). The KPNC pregnancy database will be used to identify the maternal-infant study population. This database links records of newborn infants with their mother, enabling integration of maternal data from across pregnancy and delivery encounters, along with infant birth information and longitudinal follow-up for infants who remain enrolled with KPNC. Infant healthcare encounters after the birth and throughout the follow-up period will be identified in the EMR. As routinely-recommended vaccinations are provided free of charge to KPNC members and can be electronically linked with the pregnancy database and other EMR data, the receipt and gestational timing of ABRYSVO vaccination during pregnancy can be determined.

Outcomes

Medically-Attended Endpoints

- Primary: PCR-confirmed RSV LRTD occurring ≤ 180 days after birth (first episode).
- Key Secondary: PCR-confirmed RSV LRTD hospitalization occurring ≤ 180 days after birth (first episode).

Secondary:

- Follow-up from birth to ≤ 180 days of age: PCR-confirmed RSV and RSV hospitalization (first episode), as well as all-cause LRTD and LRTD hospitalization (first episode during the RSV season). Additional outcomes include acute otitis media and first antibiotic prescription (first occurrence

during the RSV season).

- Follow-up from birth to ≤ 360 days of age: PCR-confirmed RSV, RSV hospitalization, RSV LRTD, and RSV LRTD hospitalization stratified by infant age intervals, all assessed as first episodes within 360 days after birth.

Exploratory:

- Follow-up from birth to ≤ 180 days of age: PCR-confirmed RSV LRTD and RSV LRTD hospitalization (first episode) will be analyzed by stratification factors, including time from vaccination to birth, gestational age at vaccination, infant high-risk status for severe RSV, and coadministration with other vaccines during pregnancy.

- Follow-up from birth to ≤ 720 days of age: All-cause LRTD, LRTD hospitalization, and acute otitis media assessed as the total number of episodes through 720 days. Additional objectives descriptively explore PCR-confirmed RSV-positive cases: timing, severity, and sequelae.

- Preterm infants (≤ 180 days): Among infants born preterm, analyses will describe ABRYSV0 vaccination characteristics, censoring patterns, and incidence of RSV illness during the first 180 days of life.

Data analysis plan

RSV-specific infant analyses

RSV-specific from birth to ≤ 180 days of age:

Outcomes among infants born to ABRYSV0-vaccinated and ABRYSV0-unvaccinated mothers will be compared using multivariable Cox proportional hazards regression models, generating aHR and 95% CI. IPTW will be used to account for confounding bias. Calendar date will be used as the underlying time scale in the Cox models and VE will be calculated as $(1 - aHR)$ and expressed as a percentage.

RSV-specific from birth to ≤ 360 days of age:

The analyses will stratify the Cox model by infant age; the exact width of

discrete time intervals will be determined based on outcome incidence and ABRYSV0 uptake.

RSV-specific from birth ≤ 720 days of age:

Timing, severity, and sequelae, within 30 days after the index date of the first RSV illness, will be described according to maternal ABRYSV0 status. Among infants born to individuals who received ABRYSV0 vaccine during pregnancy, we will describe the gestational age at ABRYSV0 vaccination and the time interval from vaccination to birth.

Non-specific all-cause analyses

To be assessed for two overlapping follow-up periods: from birth to ≤ 180 days of age and from birth to ≤ 720 days of age.

All-cause analyses from birth to ≤ 180 days of age: Cox proportional hazards regression models will be used to estimate VE. VE will be calculated as $(1 - aHR)$ and expressed as a percentage.

All-cause analyses from birth to ≤ 720 days of age: Will be based on the total number of episodes of each outcome during follow-up. Poisson regression to generate aIRR for comparison. VE will be calculated as $(1 - aIRR)$ and expressed as a percentage.

Among preterm infants (follow-up period from birth to ≤ 180 days): An assessment of VE among preterm infants may be considered using the same methodology as for the primary endpoint, if there are sufficient data.

Data management

ENCePP Seal

The use of the ENCePP Seal has been discontinued since February 2025. The ENCePP Seal fields are retained in the display mode for transparency but are no longer maintained.

Data sources

Data sources (types)

[Electronic healthcare records \(EHR\)](#)

[Laboratory tests and analyses](#)

Use of a Common Data Model (CDM)

CDM mapping

No

Data quality specifications

Check conformance

Unknown

Check completeness

Unknown

Check stability

Unknown

Check logical consistency

Unknown

Data characterisation

Data characterisation conducted

No