



RSVpreF Older Adults

Clinical Development Program Updates

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Presentation to ACIP
June 21, 2023

RSVpreF Older Adult

– Clinical Development Program Updates



ABRYSVO™ (Respiratory Syncytial Virus Vaccine)
FDA Approval on 5/31/2023



Indication

Active immunization for the prevention of lower respiratory tract disease (LRTD) caused by respiratory syncytial virus (RSV) in individuals 60 years of age and older



Additional Clinical Trial Data

- RENOIR – End-of-Season 1 and Mid-Season 2 analyses
- RSVpreF/influenza vaccine coadministration study

RENOIR

– Phase 3 safety and efficacy study in adults ≥ 60 years of age



38,863 participants enrolled
Healthy or with stable chronic conditions



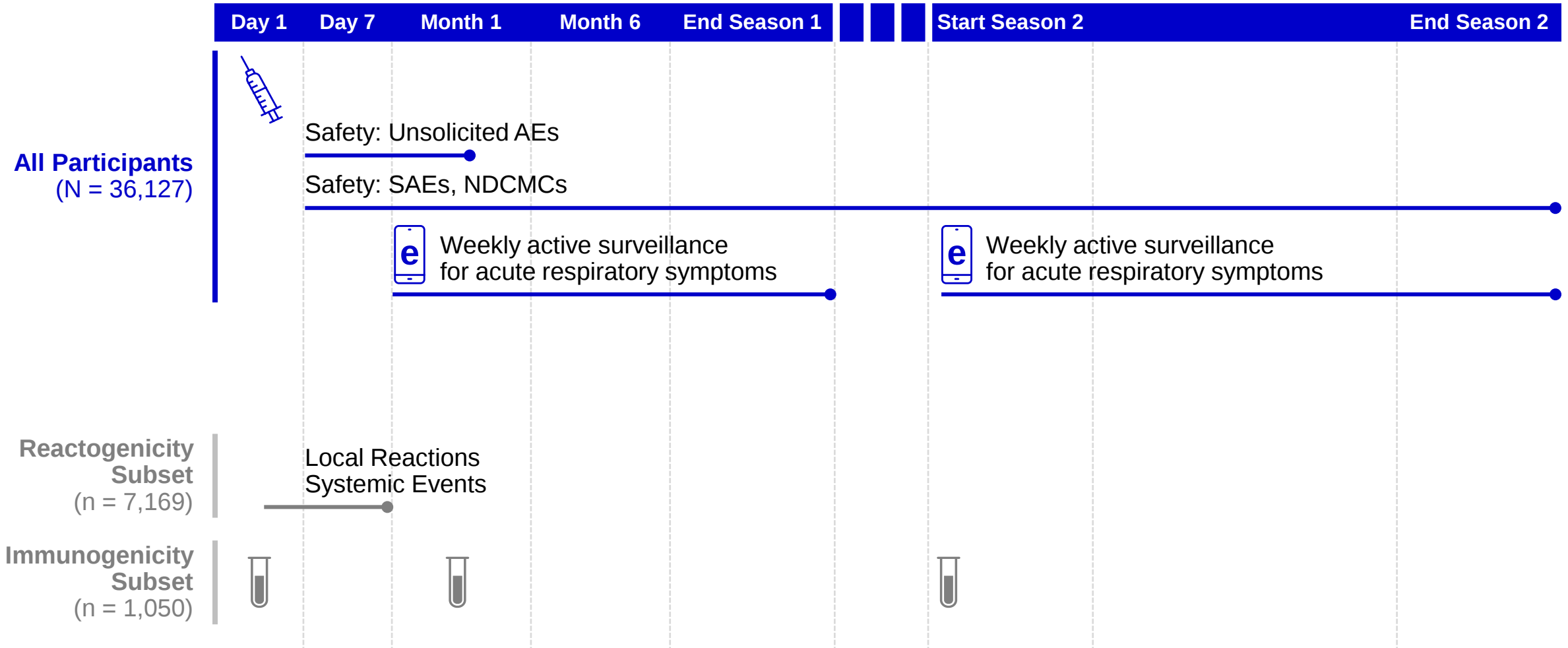
Randomized 1:1 to receive
RSVpreF 120 μg or placebo



Stratified by age group
60–69 years | 70–79 years | ≥ 80 years

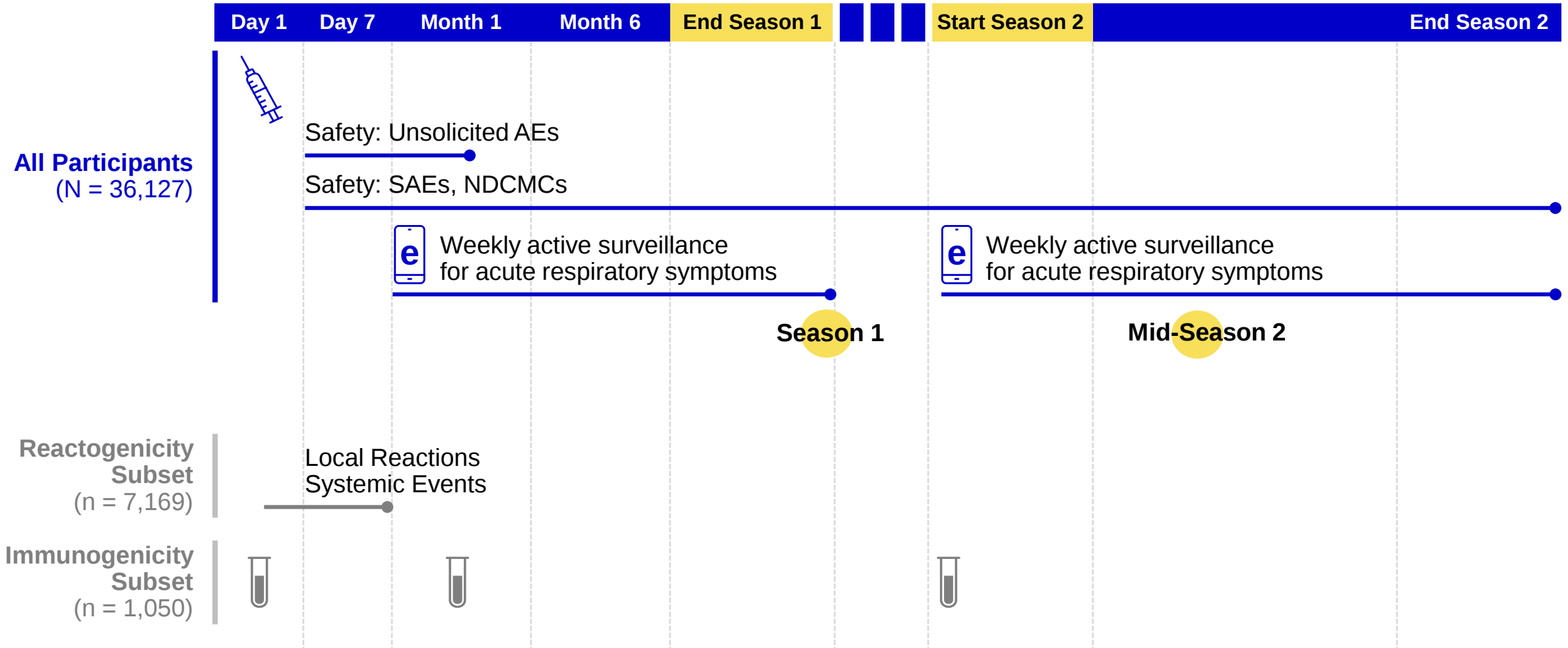


RENOIR Study Design



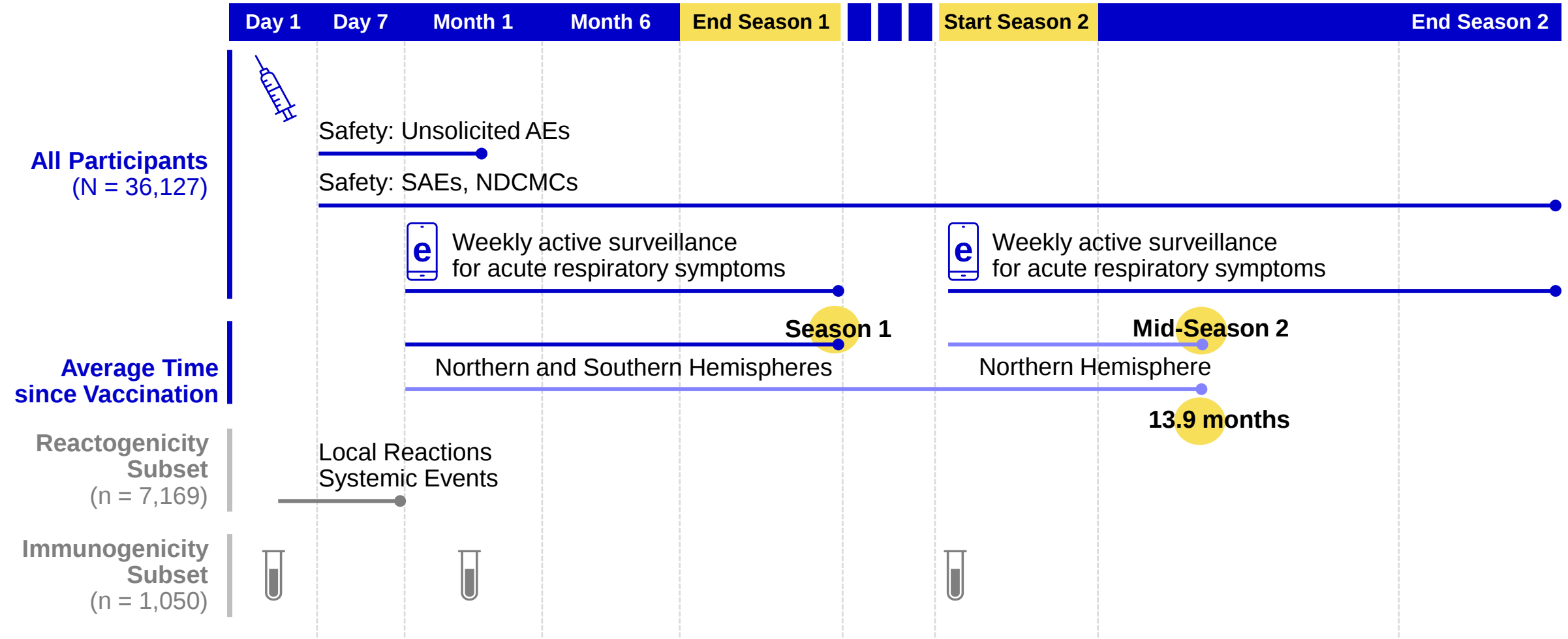
AE, adverse event; NDCMC, newly diagnosed chronic medical condition; SAE, serious adverse event

RENOIR Study Design



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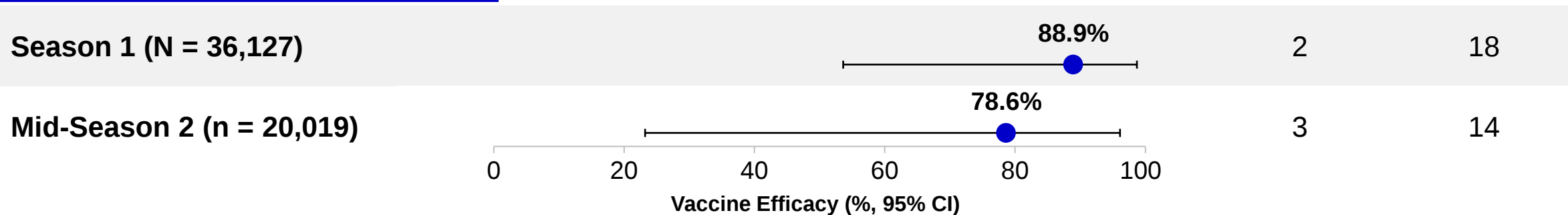


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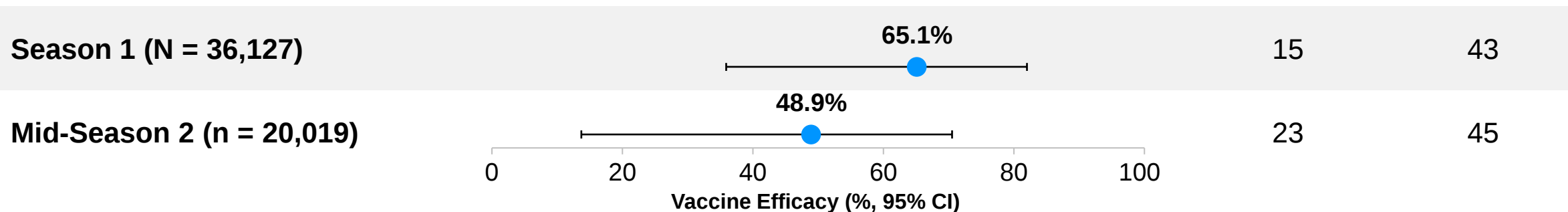
Efficacy against RSV-LRTD

– Demonstrated through Mid-Season 2 Analysis

RSV-LRTD with ≥ 3 symptoms



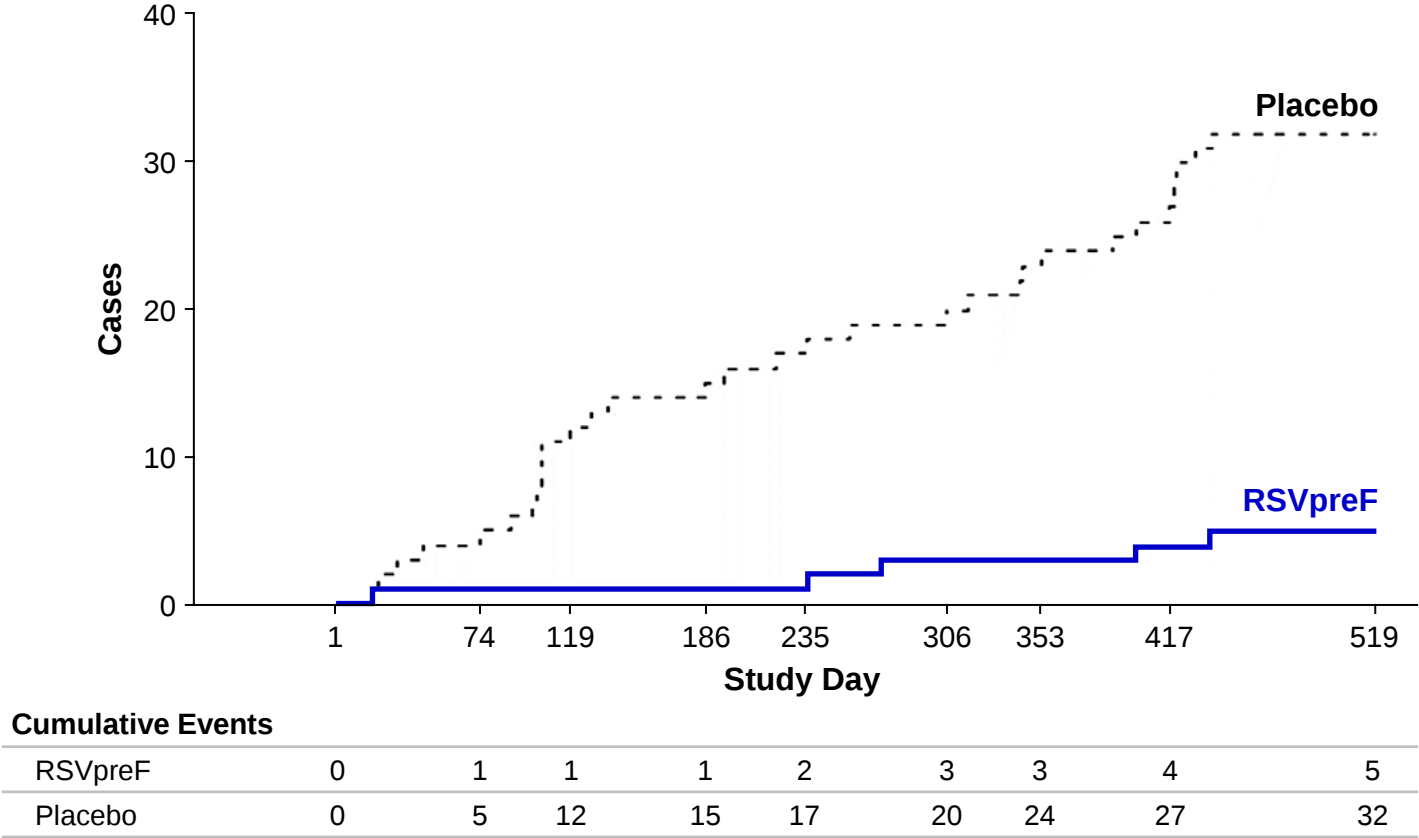
RSV-LRTD with ≥ 2 symptoms



Mid-Season 2 includes Northern Hemisphere only (US, Canada, Finland) through January 31, 2023



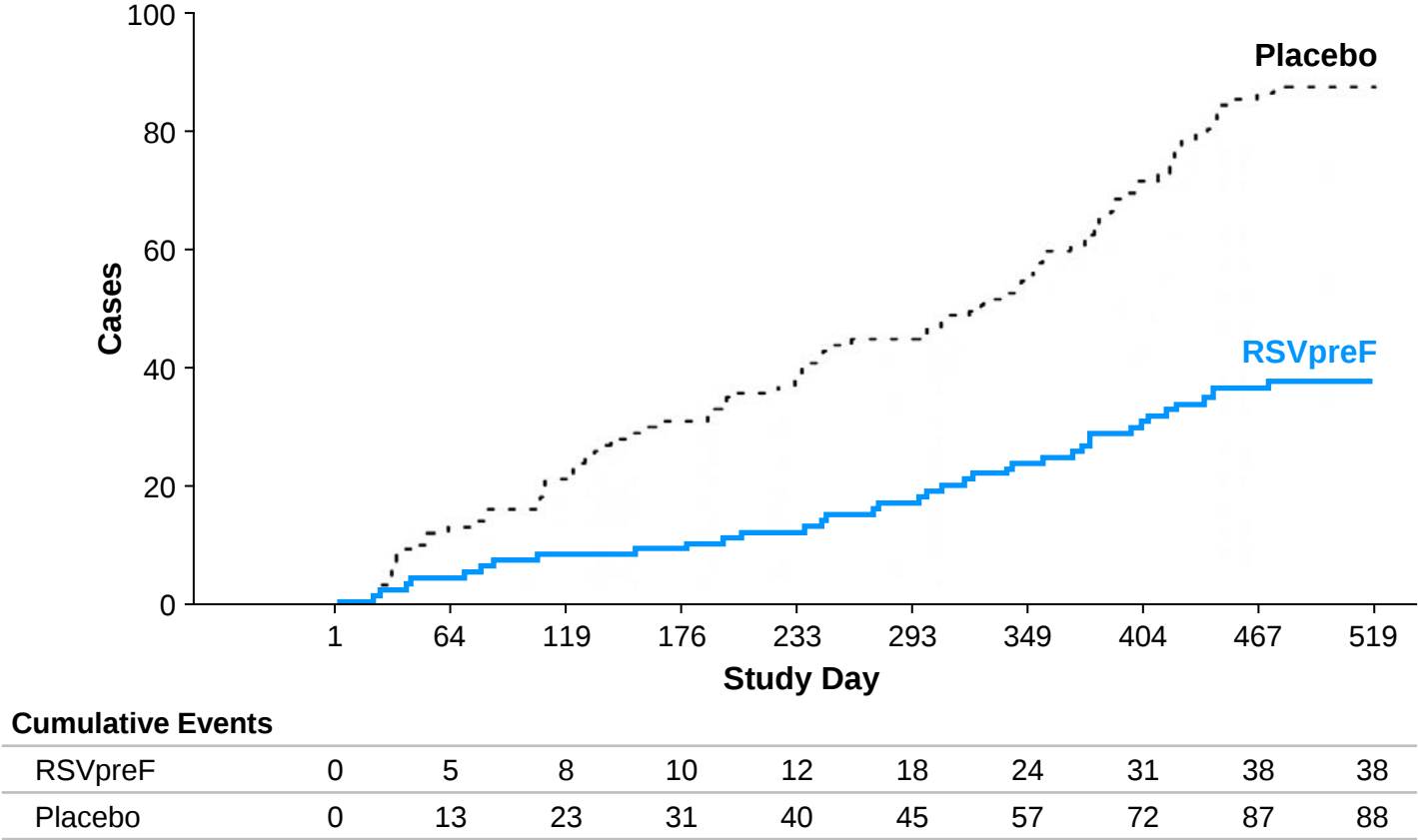
Persistent VE against RSV-LRTD with ≥ 3 Symptoms through Mid-Season 2



RSV-LRTD, lower respiratory tract disease due to RSV; RSV, respiratory syncytial virus; VE, vaccine efficacy.



Persistent VE against RSV-LRTD with ≥ 2 Symptoms through Mid-Season 2



RSV-LRTD, lower respiratory tract disease due to RSV; RSV, respiratory syncytial virus; VE, vaccine efficacy.

Adverse Events, by Category, from Vaccination through 1-Month Follow Up Visit and through Data Cutoff (31Jan2023): Safety Population

Adverse Event Category	RSVpreF N = 18,575		Placebo N = 18,288	
	n (%)	(95% CI)	n (%)	(95% CI)
From Vaccination through 1-Month Follow-Up Visit				
Any Event	1,976 (10.6)	(10.2, 11.1)	1,897 (10.4)	(9.9, 10.8)
Related	259 (1.4)	(1.2, 1.6)	178 (1.0)	(0.8, 1.1)
Immediate AE	37 (0.2)	(0.1, 0.3)	33 (0.2)	(0.1, 0.3)
Severe or life-threatening	102 (0.5)	(0.4, 0.7)	95 (0.5)	(0.4, 0.6)
From Vaccination through 31Jan2023				
NDCMC	806 (4.3)	(4.1, 4.6)	825 (4.5)	(4.2, 4.8)
SAE	790 (4.3)	(4.0, 4.6)	746 (4.1)	(3.8, 4.4)
Related SAE	3 (<0.1)	(0.0, 0.1)	0	(0.0, 0.0)
AE leading to withdrawal	12 (<0.1)	(0.0, 0.1)	11 (<0.1)	(0.0, 0.1)
AE leading to death	100 (0.5)	(0.4, 0.7)	104 (0.6)	(0.5, 0.7)

Any reactogenicity reported as adverse events (from either reactogenicity subset or non-reactogenicity subset) during the specified time period are included in this table.

Immediate AE refers to an AE reported in the 30-minute post-vaccination observation period.

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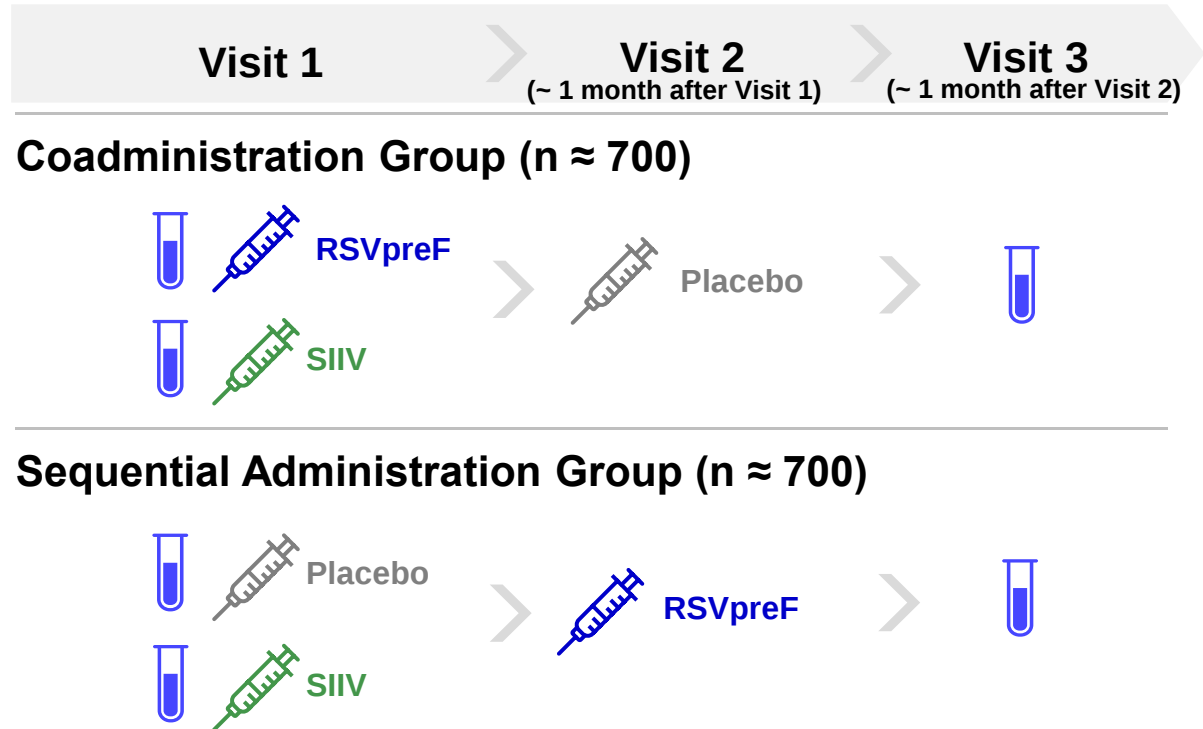


RSVpreF/influenza vaccine coadministration study

RSVpreF/SIIV Coadministration in Adults ≥ 65 Years of Age

Phase 3 Study Design and Key Procedures

- Placebo-controlled, double-blind study
- Assessing safety and immunogenicity (non-inferiority)
- Australia (31 sites)
- ~1,400 healthy participants ≥ 65 years of age
- Randomized 1:1
- SIIV: Fluad Quadrivalent
- Timeframe: April 13, 2022 – October 12, 2022



SIIV, seasonal inactivated influenza vaccine

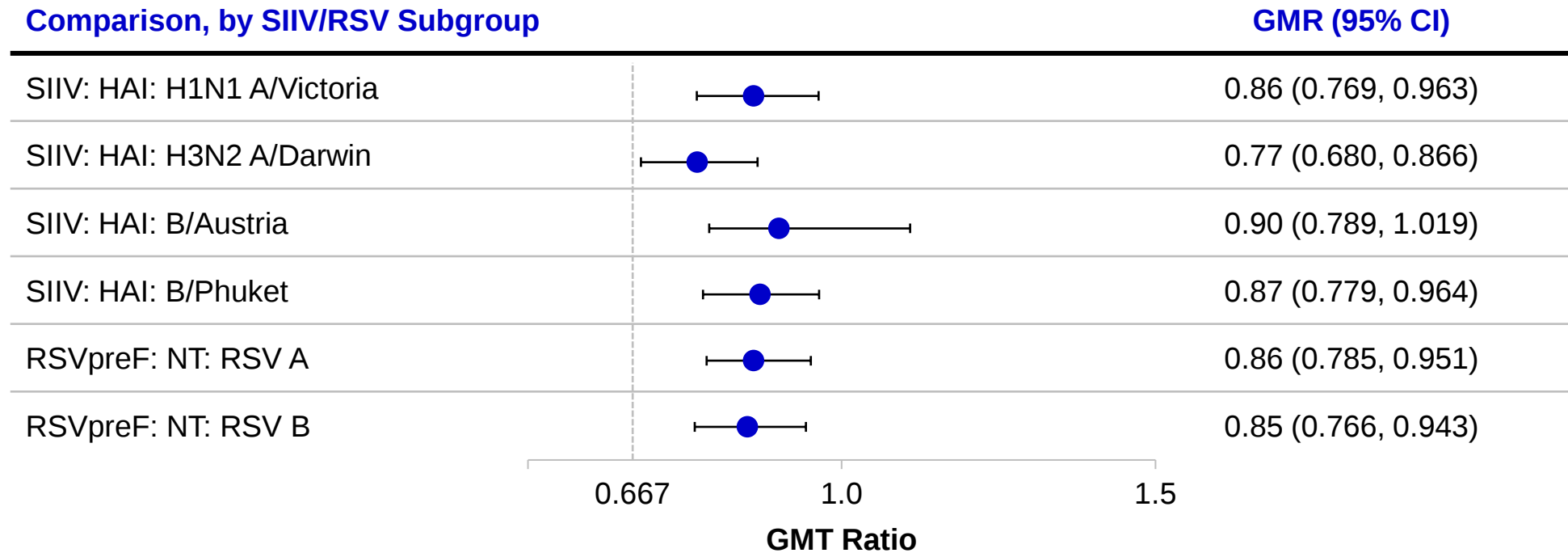
Demographics

	Coadministration (RSVpreF + SIIV) / Placebo (N = 703) n (%)	Sequential Administration (Placebo + SIIV) / RSVpreF (N = 696) n (%)	Total (N = 1,399) n (%)
Sex			
Female	398 (56.6)	372 (53.4)	770 (55.0)
Age at Visit 1 (years)			
Mean (SD)	70.7 (4.7)	70.7 (4.7)	70.7 (4.7)
Median	70.0	70.0	70.0
Min, max	(65, 91)	(65, 88)	(65, 91)
Age group at Visit 1			
65-74 years	567 (80.7)	559 (80.3)	1126 (80.5)
≥ 75 years	136 (19.3)	137 (19.7)	273 (19.5)
Race			
White	669 (95.2)	665 (95.5)	1,334 (95.4)
Asian	22 (3.1)	21 (3.0)	43 (3.1)
Multiracial	4 (0.6)	1 (0.1)	5 (0.4)
Other	4 (0.6)	4 (0.5)	8 (0.6)
Not reported or unknown	4 (0.6)	5 (0.7)	9 (0.6)

SIIV, seasonal inactivated influenza vaccine

Non-inferiority Demonstrated by SIIV HAI and RSV Neutralizing Titer GMRs

Geometric Mean Ratios with 95% CIs – Evaluable RSV Immunogenicity Population and Evaluable SIIV Immunogenicity Population



GMRs and 2-sided confidence intervals (CIs) calculated by exponentiating the mean difference of the logarithms of the titers (coadministration minus sequential-administration) and corresponding confidence intervals (CIs) (based on Student's t distribution).

GMR, geometric mean ratio; GMT, geometric mean titer; HAI, hemagglutination inhibition assay; NT, neutralizing titer; RSV, respiratory syncytial virus

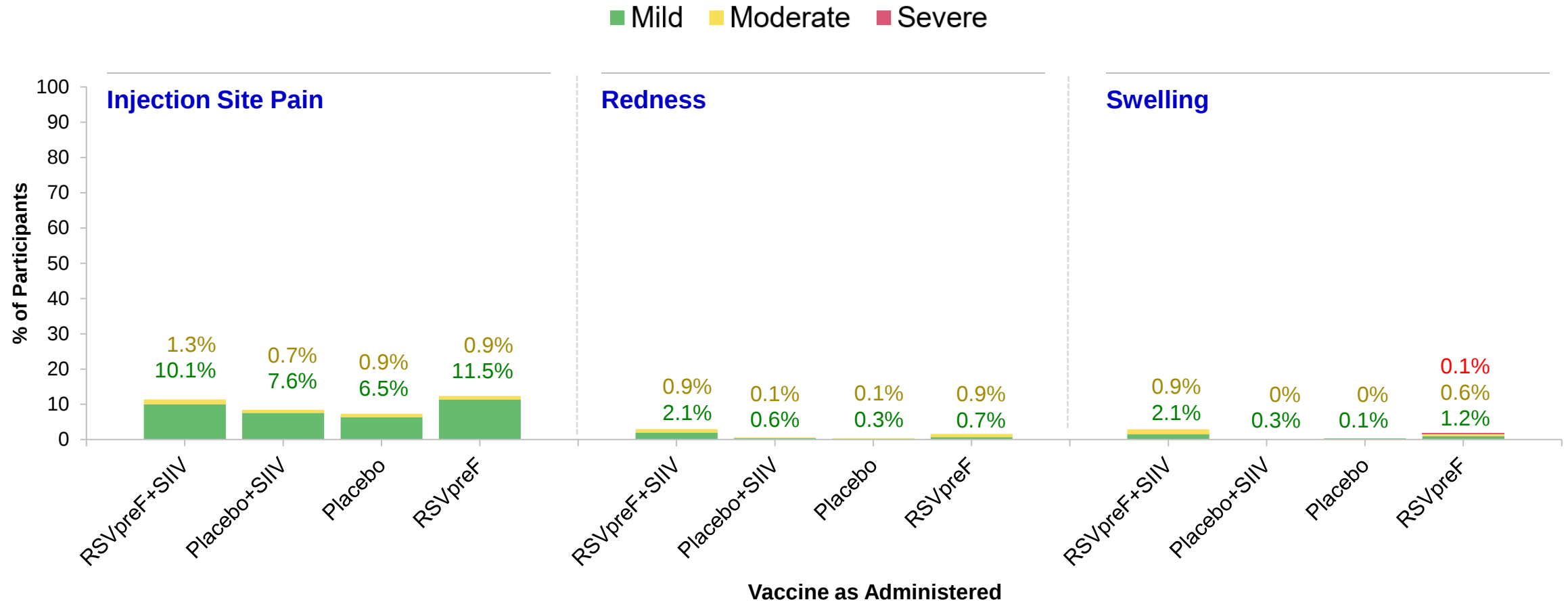
Similar HAI Titer Seroprotection and Seroconversion at 1 Month in Coadministration and Sequential Administration Groups

Serostatus, Strain	Coadministration (RSVpreF + SIIV)/Placebo		Sequential Administration (Placebo + SIIV)/RSVpreF	
	%	(95% CI)	%	(95% CI)
Seroprotection¹				
1 month after vaccination				
H1N1 A/Victoria	91.6	(89.3, 93.6)	94.3	(92.3, 95.9)
H3N2 A/Darwin	87.9	(85.2, 90.3)	91.0	(88.6, 93.0)
B/Austria	85.3	(82.4, 87.9)	89.5	(87.0, 91.7)
B/Phuket	88.4	(85.7, 90.7)	92.6	(90.4, 94.4)
Seroconversion²				
H1N1 A/Victoria	36.6	(32.9, 40.3)	43.9	(40.1, 47.7)
H3N2 A/Darwin	58.8	(55.0, 62.6)	62.6	(58.9, 66.3)
B/Austria	39.5	(35.8, 43.3)	47.3	(43.5, 51.1)
B/Phuket	25.3	(22.1, 28.8)	28.0	(24.6, 31.5)

¹Seroprotection: HAI titer $\geq$ 1:40

²Seroconversion: $\geq$ 4-fold rise from before to after receipt of SIIV if the HAI titer is $\geq$ 1:10 before SIIV or if the after SIIV HAI titer is $\geq$ 1:40 where the before SIIV is $<$ 1:10

Local Reactions Mostly Mild or Moderate



Participants reporting local reactions by maximum severity within 7 days after each vaccination

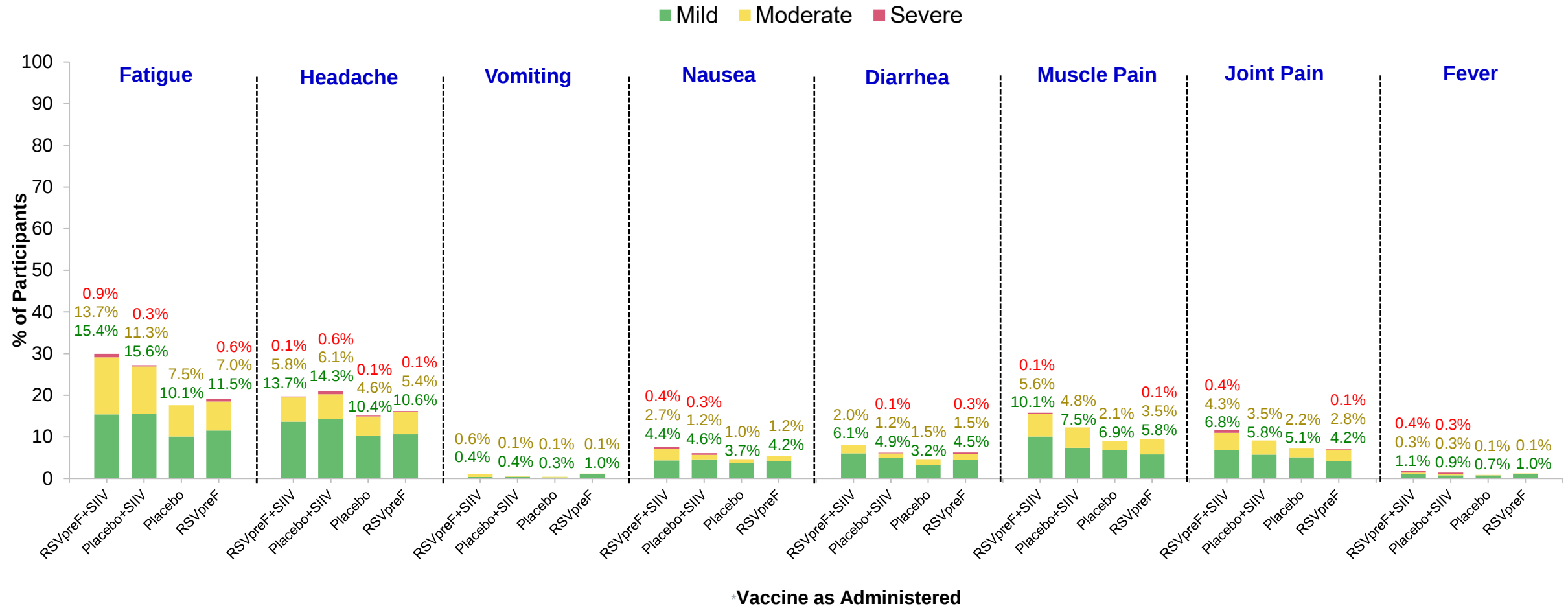
Local reactions evaluated on the arm receiving RSVpreF/placebo; no assessment of local reactogenicity at SIIV injection site.

Local reactions after RSVpreF had median onset 2 to 3 days after vaccination and median duration of 1 to 2 days

Only 1 severe local reaction reported (swelling), in the sequential administration group at Visit 2

SIIV, seasonal inactivated influenza vaccine

Systemic Events Mostly Mild or Moderate



Participants reporting systemic events by maximum severity within 7 days after each vaccination

Systemic events after RSVpreF+SIIV had median onset 2 to 4 days after vaccination and median duration of 1 to 2 days

SIIV, seasonal inactivated influenza vaccine

Adverse Events, by Category, within One Month after Vaccination: Safety Population

	RSVpreF+SIIV (N = 703)	Placebo+SIIV (N = 695)	Placebo (N = 689)	RSVpreF (N = 691)
Adverse Event Category	n (%)	n (%)	n (%)	n (%)
Any event	154 (21.9)	134 (19.3)	117 (17.0)	115 (16.6)
Related	9 (1.3)	3 (0.4)	3 (0.4)	5 (0.7)
Serious	8 (1.1)	6 (0.9)	2 (0.3)	5 (0.7)
Related	0 (0.0)	0 (0.0)	0 (0.0)	0 (0.0)
Death	0 (0.0)	1 (0.1)	0 (0.0)	0 (0.0)
Immediate	3 (0.4)	1 (0.1)	1 (0.1)	0 (0.0)

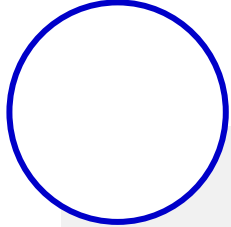
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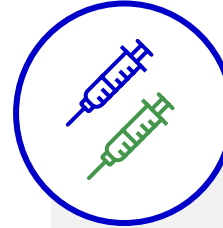
RSVpreF Older Adult Clinical Development Program Updates

– Conclusions



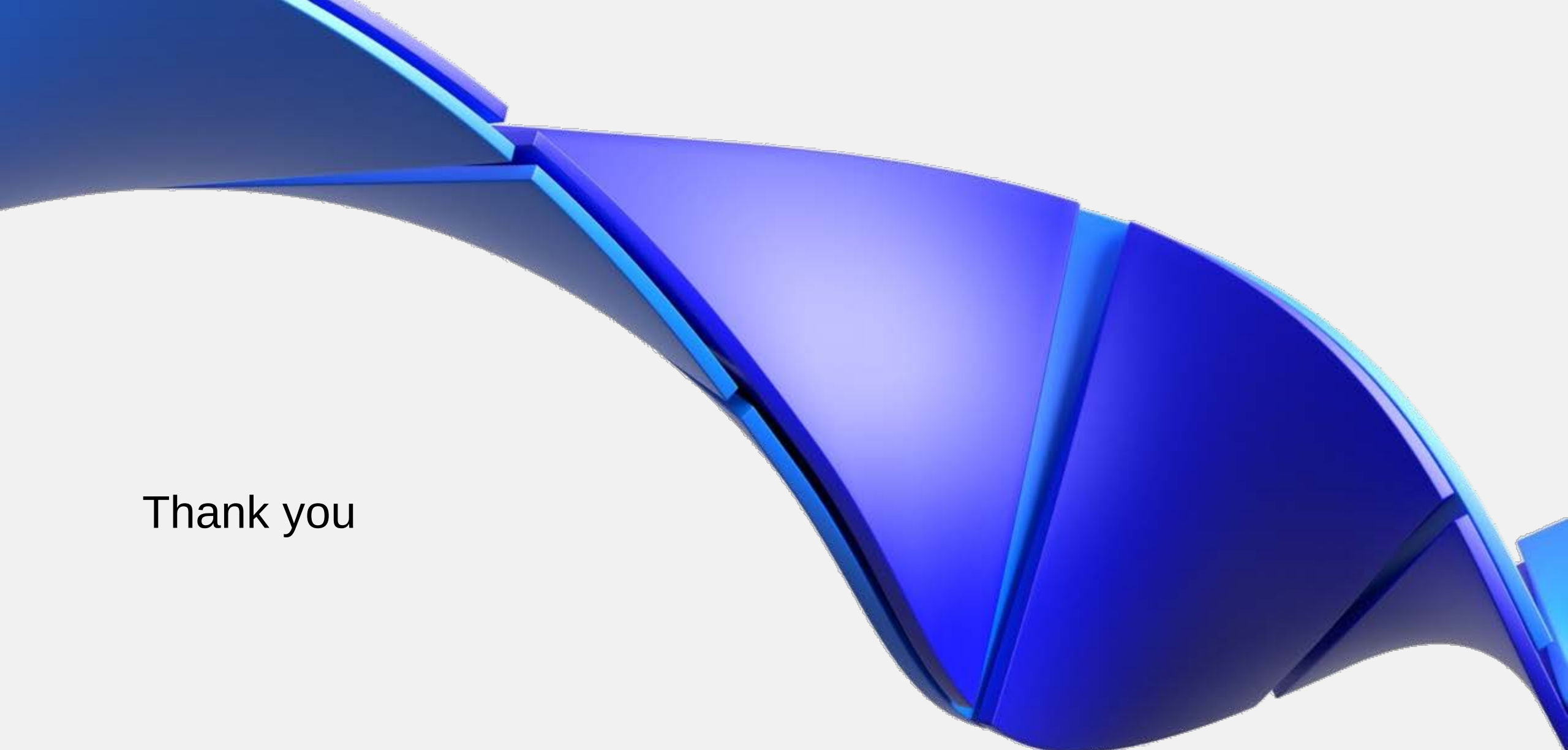
RENOIR Phase 3 Pivotal Efficacy – Season 1 and Mid-Season 2

- Favorable overall safety profile of RSVpreF
- RSVpreF remained efficacious in prevention of RSV-LRTD
 - Through end of season 1
 - In mid-season 2
 - Average 13.9 months of follow up since vaccination



RSVpreF Coadministration with Influenza Vaccine

- RSVpreF safe and well tolerated when coadministered with influenza vaccine
- Non-inferior immune responses when RSVpreF coadministered with influenza vaccine

A large, abstract graphic composed of several overlapping, curved, and faceted planes in various shades of blue and purple. The shapes create a sense of depth and movement, resembling a stylized, modern architectural structure or a complex molecular model. The lighting is soft, highlighting the edges and surfaces of the planes.

Thank you