

COVID-19 Living Evidence Synthesis #6

(Version 32:16 March 2022)

Question

What is the effectiveness of available COVID-19 vaccines for adults, including variants of concern and over time frames up to 120 days?

Findings

For vaccine effectiveness in variants of concern (VOC), we present a visual summary of evidence in Table 1 and Table 2 and details in Table 3.

Methods are presented in Box 1 and in the following appendices:

- 1) reference list
- 2) glossary
- 3) data-extraction template
- 4) process for assigning variant of concern to studies
- 5) research question and critical appraisal process
- 6) <u>detailed description of the narrative</u> <u>summary statement</u>.

Overall, 444 studies were appraised and 148 used to complete this summary. The reasons for excluding the remaining 296 studies are reported in the second section of Appendix 2.

Seven new studies have been added since the previous edition of this living evidence synthesis, all of which are signaled by a lastupdated date of 16 March 2022 (highlighted in yellow). The new studies included results for VOC Omicron (4), VOC Delta (4).

Synthesis 10. Similarly, studies examining effectiveness of vaccines in children and adolescents, including those covering periods beyond 120 days, are now captured in a third synthesis, COVID-END living evidence synthesis 8. The most recent version of all three syntheses (6,8,10) can always be found on the COVID-END website.

Box 1: Our approach

We retrieved candidate studies and updates to living evidence syntheses on vaccine effectiveness using the following mechanisms: 1) PubMed via COVID-19+ Evidence Alerts; 2) systematic scanning of pre-print servers; 3) updates to the COVID-END inventory of best evidence syntheses; and 4) cross-check with updates from the VESPa team. We included studies and updates to living evidence syntheses identified up to two days before the version release date. We did not include press releases unless a preprint was available. A full list of included and excluded studies is provided in **Appendix 1**. A glossary is provided in **Appendix 2**.

Prioritized outcome measures: Infection, severe disease (as defined by the study investigators), death, and transmission.

Data extraction: We prioritized variant-confirmed and vaccine-specific data over total study population data (variant assumed and/or vaccine unspecified). We extracted data from each study in duplicate using the template provided in **Appendix 3**. Relevance to VOC is determined directly, when reported by study authors, or indirectly where reasonable assumptions can be made about the variant prevalent in the jurisdiction at the time of the study as described in **Appendix 4**.

Critical appraisal: We assessed risk of bias, direction of effect, and certainty of evidence. Risk of bias: assessed in duplicate for individual studies using an adapted version of ROBINS-I. Direction of vaccine effect: "prevented" or "protects" was applied to mean estimates or range of mean estimates of effect that are greater than or equal to 50% (the lowest acceptable limit for vaccine effectiveness as determined by WHO). Certainty of evidence: assessed for the collection of studies for each vaccine according to variant of concern using a modified version of GRADE. Details of the research question for this synopsis and the critical appraisal process are provided in Appendix 5.

Summaries: We summarized the evidence by presenting narrative evidence profiles across studies, with or without pooling, as appropriate. A template for the summary statements used on page 1 under "Findings" and in Table 1 under each VOC is provided in **Appendix 6**.

We update this document every Wednesday and post it on the COVID-END website.

Highlights of changes this week

- New data at 60 and 90 days for 2 doses and 3 doses of BNT162b2 and mRNA-1273.
- New data for ChAdOx1 (2 doses) and Ad26.COV2.S (1 dose; 1 dose + an mRNA vaccine)
- One new study has a moderate risk of bias: reports risk of infection up to 90 days after 3rd dose of BNT162b2: ref <u>168</u>)

VOC Omicron

We have low certainty evidence that **2 doses** of **BNT162b2 [Pfizer]** prevented infection from VOC **Omicron** (26 to 55% - range of means) up to 44 days and limited protection against infection (6 to 49% - range of means) up to 60 days after 2nd dose.

We have low certainty evidence that **2 doses** of **BNT162b2 [Pfizer]** prevented symptomatic infection from VOC **Omicron** (46 to 88% – range of means) at up to 60 days and limited protection (36.3% [95% CI, 25.1 to 45.8] – 1 Obs) up to 90 days after 2nd dose.

We have low certainty evidence that **2 doses** of **mRNA-1273 [Moderna]** provided limited protection from infection from VOC **Omicron** (36% [95% CI, -70 to 76.4] – 1 Obs) up to 44 days; from infection (48% [95% CI, 44 to 52] – 1 Obs) up to 60 days and from infection (24 to 30% - range of means) up to 90 days after 2nd dose.

We have low certainty evidence that **2 doses** of **mRNA-1273 [Moderna]** prevented symptomatic infection from VOC **Omicron** (44.8% [95% CI, 16 to 63.8] – 1 Obs) up to 30 days after 2nd dose.

We have low certainty evidence that **2 doses** of **ChAdOx1** prevented infection from VOC **Omicron** (51% [95% CI, 23 to 69] – 1 Obs) up to 60 days after 2nd dose.

We have low certainty evidence that **one dose of Ad26.COV2.S** provided limited protection from infection from VOC **Omicron** (47% [95% CI, 45 to 49] – 1 Obs) up to 60 days after dose and low certainty evidence that **one dose of Ad26.COV2.S followed by one dose of an mRNA vaccine** prevented infection from VOC **Omicron** (48% [95% CI, 42.5 to 53.7] – 1 Obs) at least 7 days after 2nd dose.

We have low certainty evidence that **3 doses** of **BNT162b2 [Pfizer]** prevented infection from VOC **Omicron** (34 to 55% – range of means) up to 30 days; prevented infection (58% [95% CI, 57 to 58] – 1 Obs) up to 60 days and provided limited protection (35.7% [95% CI, 29.8 to 41.2) up to 90 days after 3rd dose.

We have low certainty evidence that **3 doses** of **BNT162b2 [Pfizer]** prevented symptomatic infection from VOC **Omicron** (75.5% [95% CI, 56.1 to 86.3] – 1 Obs) up to 14 days; (56.6% [95% CI, 50.8 to 61.7] – 1 Obs) up to 35 days; (43.7% [95% CI, 32.9 to 52.7] – 1 Obs) up to 77 days after 3rd dose.

We have low certainty evidence that **3 doses** of **BNT162b2 [Pfizer]** prevented severe, critical or fatal disease from VOC **Omicron** (90.8% [95% CI, 81.5 to 95.5] – 1 Obs) up to 49 days after 3rd dose.

We have low certainty evidence that **3 doses** of **mRNA-1273 [Moderna]** prevented infection by VOC **Omicron** (46 to 64% [range of means] up to 30 days and prevented infection (61% [95% CI, 60 to 62] – 1 Obs) up to 60 days after 3rd dose.

We have low certainty evidence that **3 doses** of **mRNA-1273 [Moderna]** prevented symptomatic infection by VOC **Omicron** (54.6% [95% CI, 41.1 to 65] – 1 Obs) up to 35 days and symptomatic infection (38.6% [95% CI, 19.4 to 53.1] – 1 Obs) up to 42 days after the 3rd dose.

We have low certainty evidence that **3 doses** of **mRNA-1273 [Moderna]** prevented severe, critical or fatal disease from VOC **Omicron** (80.8% [95% CI, -51.9 to 97.6] – 1 Obs) up to 42 days after 3rd dose.

We have low certainty evidence that **2 or 3 doses** of **BNT162b2 [Pfizer]** or **mRNA-1273 [Moderna]** provides little protection against transmission of VOC **Omicron** to vaccinated household or close contacts (16% [95% CI, 0 to 37] – 1 Obs) at least 7 days after 2nd dose and (47% [95% CI, 17 to 64] – 1 Obs) at least 7 days after 3rd dose.

Table 1: Visual summary of evidence for COVID-19 vaccines for variants of concern – Delta and Omicron [2 doses: 30 to 120 days since last dose; 3 doses: 1 to 90 days since last dose]

Percentages indicate <u>level of effectiveness</u> from 0% (no effect) to 100% (full protection): ranges of estimated means are provided when ≥ 1 study is available; estimated mean value is provided for single studies

Colour indicates Level of Certainty based on the evidence

High certainty evidence Moderate certainty evidence Low certainty evidence

pooling of low to moderate risk of bias RCTs or pooling of observational studies with low risk of bias and consistent findings single RCT with low to moderate risk of bias or >one observational study with low to moderate risk of bias and at least partially consistent findings

single RCT or observational study with serious risk of bias or multiple low to serious risk of bias observational studies with inconsistent findings

Outcome (vaccine)	Variant	Number of Doses	Time since Last Dose* (days)	Vaccine Effectiveness		
Infection – Omicron (2 doses: 30 to 120 days after 2 nd dose)						
Pfizer		2	44	26 to 55%		
Moderna		2	44	36.7% (-70 to 76.4)		
Pfizer		2	<mark>60</mark>	6 to 49%		
Moderna	Omicron	2	<mark>60</mark>	48% (44 to 52)		
AstraZeneca		2	<mark>60</mark>	51% (23 to 69)		
Johnson & Johnson		1	<mark>60</mark>	47% (45 to 49)		
Moderna		2	90	24 to 30%		
Infection - Omicro	n (3 doses: up to 9	00 days after	3 rd dose)			
AZ followed by		2/1	at least 7	58.6% (55.5 to 61.6)		
mRNA vaccine						
Pfizer		3	30	34 to 55%		
Moderna	Omicron	3	30	46 to 64%		
Pfizer		3	<mark>60</mark>	58% (57 to 58)		
Moderna		3	<mark>60</mark>	61% (60 to 62)		
Pfizer		3	<mark>90</mark>	35.7% (29.8 to 41.2)		
	Symptomatic Infection - Omicron (2 doses: 30 to 120 days after 2 nd dose)					
Moderna		2	30	44.8% (16 to 63.8)		
Pfizer	Omicron	2	60	46 to 88%		
Pfizer		2	90	36% (25.1 to 45.8)		
Symptomatic Infect	ion – Omicron (3	doses: up to	o 90 days after 3 rd	,		
Pfizer		3	14	75.5% (56.1 to 86.3)		
AZ followed by		2/1	14	71.4% (41.8 to 86)		
mRNA vaccine						
Pfizer	Omicron	3	28 to 35	56.6% (50.8 to 61.7)		
Moderna		3	28 to 35	54.6% (41.1 to 65)		
Moderna		3	42	38.6% (19.4 to 53.1)		
Pfizer		3	70 to 77	43.7% (32.9 to 52.7)		
Severe Disease – Or	micron (2 or 3 dos	ses)				
Pfizer	Omicron	3	7 to 42	90.6% (77.8 to 96)		
Moderna		3	7 to 42	80.5% (-51.9 to 97.6)		

Outcome (vaccine)	Variant	Number of Doses	Time since Last Dose* (days)	Vaccine Effectiveness
Pfizer		3	49	90.8% (81.5 to 95.5)
Death – Omicron (2	2 or 3 doses)			
Infection – Delta (2	doses: 30 to 120 d	lavs after 2 nd	dose)	
Pfizer		2	60	82 to 87%
Moderna		2	60	71 to 94%
AstraZeneca		2	<mark>60</mark>	60% (57 to 62)
D.C.		2	0.0	Z7 - 740/
Pfizer	Delta	2	90	67 to 74%
Moderna	Delta	2	90	79 to 83%
IVIOUCIIIA			90	77 10 05/0
Pfizer		2	120	53 to 85%
Moderna				81 to 88%
THO GETTIA		2	120	01 60 0070
AstraZeneca		2	120	65 to 72%
AZ followed by		1/1	120	86% (81 to 89)
mRNA vaccine		,		,
Infection – Delta (3	doses: up to 90 da	ays after 3 rd	dose)	
AZ followed by		2/1	7	82% (68 to 90)
Pfizer				
Sinovac followed		2/1	7	93 to 98%
by Pfizer				
Sinovac followed	Delta	2/1	7	86% (74 to 93)
by AZ				
Pfizer		3	30	81 to 93%
Moderna		3	30	83 to 96%
Pfizer		3	60	90% (89 to 90)
Moderna		3	60	92% (91 to 93)
Symptomatic Infect	tion – Delta (2 dos			
Pfizer		2	60 to 90	72% (61 to 80)
AstraZeneca		1	60 to 90	65% (48 to 76)
Johnson & Johnson	Dolta	1	60 to 90	52% (33 to 66)
Moderna	Delta	2	70 to 98	90%
AstraZeneca		2	119	41 to 49%
AZ followed by mRNA vaccine		1/1	120	66% (41 to 80)
Symptomatic Infect	tion - Dolta (2 dos	200 110 to 00	dave after 2rd de	(4)
Sinovac	Dena (3 008	3 3	14	78.8% (76.8 to 80.6)
AZ followed by		2/1	14	93 to 94%
Pfizer		∠/ 1	14	75 10 7470
Sinovac followed	Delta	2/1	14	96.5% (96.2 to 96.7)
by Pfizer		<u> </u>	11	70.370 (70.2 (0 70.1)

Outcome	Variant	Number	Time since	Vaccine Effectiveness		
(vaccine)		of Doses	Last Dose* (days)			
Sinovac followed		2/1	14	93.2% (92.9 to 93.6)		
by AZ		2/1	11	73.274 (72.7 68 73.6)		
Severe Disease – Delta (2 or 3 doses)						
Pfizer		2	44 to 98	91.1% (90 to 92)		
Moderna		2	60	97.8% (83.7 to 99.7)		
Moderna		2	90	75 to 93%		
Pfizer		2	120	68 to 72%		
Moderna		2	120	91.5% (60.8 to 98.1)		
AstraZeneca	Delta		120	70.5% (67 to 73.7)		
Sinovac followed by Pfizer		2/1	14	96 to 97%		
Sinovac followed by AZ		2/1	14	98.9% (98.5 to 99.2)		
Death – Delta (2 or	3 doses)					
Johnson & Johnson		1	120	89.4% (52.3 to 97.6)		
Sinovac followed by Pfizer	Delta	2/1	14	96.8% (93.9 to 98.3)		
Sinovac followed by AZ		2/1	14	98.1% (97.3 to 98.6)		

^{*}approximate because studies did not use the same exact time frames

Table 2: Visual summary of evidence for COVID-19 vaccines for variants of concern (up to 30 days after 2 doses)

Percentages indicate <u>level of effectiveness</u> from 0% (no effect) to 100% (full protection): ranges of estimated means are provided when ≥ 1 study is available; estimated mean value is provided for single studies

Colour indicates Level of Certainty based on the evidence

High certainty evidence	Moderate certainty evidence	Low certainty evidence

pooling of low to moderate risk of bias RCTs or pooling of observational studies with low risk of bias and consistent findings single RCT with low to moderate risk of bias or >one observational study with low to moderate risk of bias and at least partially consistent findings

single RCT or observational study with serious risk of bias or multiple low to serious risk of bias observational studies with inconsistent findings

Outcome (and vaccine)	Vaccine Effectiveness (2 doses unless otherwise stated) up to 30 days after last dose for each combination of vaccine, variant, and outcome					
	Alpha	Beta	Gamma	Delta	Omicron	
Any Infection						
Pfizer	78 to 95%		93%	42 to 93%		
Moderna	86 to 100%	96%	95%	52 to 91%		
AstraZeneca (AZ)	62 to 79%		90%	45 to 83%	11.4% (-18.8 to 34.6	
Johnson & Johnson				3 to 71%*		
JnJ followed by an mRNA vaccine					48% (42.5 to 53.7)	
Novavax						
Sinovac			66%	60 to 74%		
AZ followed by Pfizer or Moderna	82 to 91%		96%	88%		
Sinovac followed by AZ				74% (43 to 99)		
Symptomatic Infe	ection (reported	when data on "a	any infection" is l			
Pfizer		84 to 88%	84 to 88%	63 to 94%		
Moderna			88%	87%		
AstraZeneca		10%**	65%	61 to 92%		
Johnson & Johnson				51%*		
Novavax	86%	43%**				
Sinovac				59%		
Covaxin				50%		
AZ followed by				67 to 79%		
Pfizer or						
Moderna						
Transmission						

Outcome	Vaccine Effectiveness (2 doses unless otherwise stated) up to 30 days				
(and vaccine)	after last dose for each combination of vaccine, variant, and outcome				
Pfizer	70 to 82%			31 to 63%	
				(unvacc contact)	
				10 to 40%	
M 1	88%			(vacc contact)	
Moderna	= = :			62 to 77%	
AstraZeneca	58 to 63%			36%	
Johnson &	77%*				
Johnson					
Novavax					
Sinovac					
AZ followed by				86%	
Pfizer or					
Moderna					
Severe Disease (m	nay include dea	th for some stu	dies)		
Pfizer	92 to 100%			82 to 98%	
Moderna	96%	96%		93 to 100%	
AstraZeneca			76%		
Johnson &		82%*		93%	
Johnson					
Novavax					
Sinovac				46 to 89%	
Death					
Pfizer	91 to 97%			90%	
Moderna					
AstraZeneca				91%*	
Johnson &				90%	
Johnson					
Novavax					
Sinovac			86%	77%	

^{*}single dose

^{**}mean estimate of effect less than the lowest acceptable limit for vaccine effectiveness as determined by WHO

AZ, AstraZeneca; unvacci, unvaccinated; vacc, vaccinated; JnJ, Johnson & Johnson

Table 3: Key findings about vaccine effectiveness (revised format 13 Dec 2021)

VOC	Vaccine	Findings
Omicron	Pfizer/	BNT162b2 (2 doses) provided protection against infection by VOC Omicron
(2.1.)	BioNTech	at the following number of days after 2 nd dose:
(2 doses)	Comirnaty	• 26 to 55% up to 44 days (RME)
(any time	[BNT162b2]	• 6 to 49% up to 60 days (RME)
(any time frame)		• -76.5% (95% CI, -95.3 to -59.5) up to 164 days
iranic)		(4 Obs) [137][147][160][169]; last update 2022-03-16
		BNT162b2 (2 doses) provided protection against symptomatic infection by
		VOC Omicron at the following number of days after 2 nd dose:
		• 45.9 to 88% at up to 63 days (RME)
		• 36.3% (95% CI, 25.1 to 45.8) at 90 days
		• 34.3% (95% CI, -5 to 58.7) at 175 days
		(2 Obs) [136][162]; last update 2022-03-02
		BNT162b2 or mRNA-1273 (2 doses) provided protection against infection by
		VOC Omicron:
		 6% (95% CI, -25 to 30) 7 to 59 days after 2nd dose 13% (95% CI, -38 to 8) 60 to 119 days after 2nd dose
		• -38% (95% CI, -58 to 5) 60 to 119 days after 2 dose • -38% (95% CI, -61 to -18) 120 to 179 days after 2 nd dose
		• -16% (95% CI, -62 to 17) ≥240 days after 2 nd dose
		(1 Obs) [147]; last update 2022-01-18
Omicron	Pfizer/	BNT162b2 (3 doses) provided protection against infection by VOC Omicron
	BioNTech	at the following number of days after the 3 rd dose:
(3 doses)	Comirnaty	• 34 to 54.6% up to 30 days (RME)
	[BNT162b2]	• 58% (95% CI, 57 to 58) up to 60 days
(any time		• 35.7% (95% CI, 29.8 to 41.2) up to 90 days
frame)		(6 Obs) [137][147][160][167][168][169]; last update 2022-03-16
		BNT162b2 (3 doses) provided protection against symptomatic infection by
		VOC Omicron at the following number of days after 3 rd dose:
		• 75.5% (95% CI, 56.1 to 86.3) at 14 days
		• 56.6% (95% CI, 50.8 to 61.7) at 28 to 35 days
		• 43.7% (95% CI, 32.9 to 52.7) at 70 to 77 days
		(2 Obs) [136][162]; last update 2022-03-02
		BNT162b2 (3 doses) provided protection against severe, critical, or fatal
		disease by VOC Omicron at the following number of days after 3 rd dose:
		• 90.6% (95% CI, 77.8 to 96) at 7 to 42 days
		• 90.8% (95% CI, 81.5 to 95.5) at 49+ days (1 Obs) [162]; <i>last update 2022-03-02</i>
Omicron	Moderna	mRNA-1273 (2 doses) provided protection against infection by VOC
Officion	Spikevax	Omicron at the following number of days after 2 nd dose:
(2 doses)	[mRNA-	• 36.7% (95% CI, -69.9 to 76.4) up to 44 days
	1723]	• 48% (95% CI, 44 to 52) up to 60 days
(any time	_	• 23.7 to 30.4% up to 90 days (RME)
frame)		• -39.3% (95% CI, -61.6 to -20) up to 164 days
		• 15.2% (95% CI, 0 to 30.7) at 91 to 180 days
		■ 13.270 (95% C1, U to 30.7) at 91 to 180 days

VOC	Vaccine	Findings
		• 0% (95% CI, 0 to 1.2) at 181 to 270 days
		(4 Obs) [137][148][160][169]; last update 2022-03-16
		mpDNIA 1272 (2 deces) provided protection assigns average infection by
		mRNA-1273 (2 doses) provided protection against symptomatic infection by VOC Omicron at the following number of days after 2 nd dose:
		• 44.8% (95% CI, 16 to 63.8) at 28 to 35 days
		(1 Obs) [162]; last update 2022-03-02
Omicron	Moderna	mRNA-1273 (3 doses) provided protection against infection by VOC
Officion	Spikevax	Omicron at the following number of days after 3 rd dose:
(3 doses)	[mRNA-	• 46.4 to 64% at 7 to 30 days (RME)
(0 0000)	1723]	• 61% (95% CI, 60 to 62) up to 60 days
(any time		(4 Obs) [147][148][160][167][169]; last update 2022-03-16
frame)		
,		mRNA-1273 (3 doses) provided protection against symptomatic infection by
		VOC Omicron at the following number of days after 3 rd dose:
		• 54.6% (95% CI, 41.1 to 65) at 28 to 35 days
		• 38.6% (95% CI, 19.4 to 53.1) at 42+ days
		(1 Obs) [160]; last update 2022-03-02
		mRNA-1273 (3 doses) provided protection against severe, critical, or fatal
		disease by VOC Omicron at the following number of days after 3 rd dose:
		• 80.5% (95% CI, -51.9 to 97.6) at 7 to 42 days
		(1 Obs) [160]; last update 2022-03-02
Omicron	AstraZeneca	ChAdOx1 (2 doses) provided protection against VOC Omicron for the
	[ChAd0x1]	following outcomes:
(any time	Vaxzevria	• 11.4% (95% CI, -18.8 to 34.6) from infection at 14 days after 2 nd dose
frame)	Serum	• 51% (95% CI, 23 to 69) from infection up to 60 days after 2 nd dose
	Institute of	• 5.9% (95% CI, -29.7 to 31.7) from symptomatic infection at 175 days after
	India	2 nd dose
	[Covishield]	(3 Obs) [136][160][169]; last update 2022-03-16
Omicron	AstraZeneca	ChAdOx1 (2 doses) followed by BNT162b2 provided protection against
1	[ChAd0x1]	VOC Omicron for the following outcomes:
2 doses	Vaxzevria	• 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose
followed by	Vaxzevria Serum	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd
followed by mRNA	Vaxzevria Serum Institute of	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose
followed by	Vaxzevria Serum Institute of India	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd
followed by mRNA vaccine	Vaxzevria Serum Institute of	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose
followed by mRNA vaccine (any time	Vaxzevria Serum Institute of India	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose
followed by mRNA vaccine (any time frame)	Vaxzevria Serum Institute of India [Covishield]	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16
followed by mRNA vaccine (any time	Vaxzevria Serum Institute of India [Covishield] Johnson &	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following
followed by mRNA vaccine (any time frame) Omicron	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes:
followed by mRNA vaccine (any time frame)	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson [AD26.COV	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes: 47% (95% CI, 45 to 49) from infection up to 60 days after 2nd dose
followed by mRNA vaccine (any time frame) Omicron 1 dose	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes:
followed by mRNA vaccine (any time frame) Omicron 1 dose (any time	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson [AD26.COV	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes: 47% (95% CI, 45 to 49) from infection up to 60 days after 2nd dose
followed by mRNA vaccine (any time frame) Omicron 1 dose	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson [AD26.COV	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes: 47% (95% CI, 45 to 49) from infection up to 60 days after 2nd dose
followed by mRNA vaccine (any time frame) Omicron 1 dose (any time frame)	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson [AD26.COV 2.S]	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes: 47% (95% CI, 45 to 49) from infection up to 60 days after 2nd dose (1 Obs) [169]; last update 2022-03-16
followed by mRNA vaccine (any time frame) Omicron 1 dose (any time frame)	Vaxzevria Serum Institute of India [Covishield] Johnson & Johnson [AD26.COV 2.S]	 58.6% (95% CI, 55.5 to 61.6) from infection at least 7 days after 3rd dose 71.4% (95% CI, 41.8 to 86) from symptomatic infection at 14 days after 3rd dose (2 Obs) [136][167]; last update 2022-03-16 Ad26.COV2.S provided protection against VOC Omicron for the following outcomes: 47% (95% CI, 45 to 49) from infection up to 60 days after 2nd dose (1 Obs) [169]; last update 2022-03-16 Ad26.COV2.S followed by an mRNA vaccine provided protection against

VOC	Vaccine	Findings
mRNA		
vaccine		
(any time		
frame)		
Omicron	Pfizer/	BNT162b2 or mRNA-1273 (2 doses) hh contacts showed VES:
	BioNTech	• 16% (95% CI, 0 to 37) at least 7 days after 2 nd dose
Transmission	Comirnaty	BNT162b2 or mRNA-1273 (3 doses) hh contacts showed VES:
Household or close contacts	[BNT162b2]	• 47% (95% CI, 17 to 64) at least 7 days after 3 rd dose
of index case		(1 Obs) [<u>161</u>]; last update 2022-03-02
Omicron	Moderna	BNT162b2 or mRNA-1273 (2 doses) hh contacts showed VES:
	Spikevax	• 16% (95% CI, 0 to 37) at least 7 days after 2 nd dose
Transmission	[mRNA-	BNT162b2 or mRNA-1273 (3 doses) hh contacts showed VES:
Household or	1723]	• 47% (95% CI, 17 to 64) at least 7 days after 3 rd dose
close contacts		(1 Obs) [<u>161</u>]; last update 2022-03-02
of index case Delta	Pfizer/	PNTT162b2 provided protection against VOC Dalta for the following
(1-2 doses)	BioNTech	BNT162b2 provided protection against VOC Delta for the following outcome at least 14 to 21 days after 1 st dose:
(1-2 doses)	Comirnaty	• 30 to 65% from infection (RME)
(up to 30	[BNT162b2]	• 33 to 47.5% from symptomatic infection (RME)
days)		• 87 to 94% from hospitalization (RME)
		• 100% (95% CI, not reported) against severe, critical, or fatal disease
		BNT162b2 provided protection against VOC Delta for the following
		outcome at least 7 days after 2 nd dose:
		 42 to 93% from infection (RME) 63 to 94% from symptomatic infection (RME)
		82 to 98% from severe, critical, or fatal disease (RME)
		• 90% from death (RME)
		(26 Obs) [29][38][42][47][57][63][64][71][74][76][84][88][92][97][102][109][110]
		[111][118][119][121][123][133][138][156][160][163]; last update 2022-03-02
Delta	Moderna	mRNA-1273 provided protection against VOC Delta for the following
(1-2 doses)	Spikevax	outcomes at least 14 days after 1 st dose:
	[mRNA-	• 75 to 86.7% from infection (RME)
(up to 30	1723]	• 72% (95% CI, 57 to 82) from symptomatic infection
days)		• 96% (95% CI, 72 to 99) from hospitalization
		93 to 100% from severe, critical, or fatal disease (RME) mRNA-1273 provided protection against VOC Delta for the following
		outcomes 14 days after 2 nd dose:
		• 52 to 91% from infection (RME)
		87% (95% CI, 84 to 88) from symptomatic infection
		• 93 to 100% from severe, critical, or fatal disease(RME)
		(19 Obs)
		[47][57][63][64][71][74][97][101][102][109][110][111][118][121][123][133][138][
D.I.	A - 4 :: 77	140][160]; last update 2022-03-02
Delta (1-2 doses)	AstraZeneca [ChAd0x1]	ChAdOx1 provided protection against VOC Delta for the following outcome at least 21 days after 1 st dose:
(1-2 00868)	Vaxzevria	18 to 46% from infection (RME)
	1 and VIIIa	- 10 to 70 / 0 HOIR INTECTION (KIME)

VOC	Vaccine	Findings
(up to 30	Serum	• 33 to 58% from symptomatic infection (RME)
days)	Institute of India	• 71% (95% CI, 51 to 83) from hospitalization
	[Covishield]	ChAdOx1 provided protection against VOC Delta for the following outcome
		at least 7 days after 2 nd dose:
		• 44.8 to 83% from infection (RME)
		• 61 to 92% from symptomatic infection (RME)
		• 92% (95% CI, 75 to 97) from hospitalization
		• 91% (95% CI, 83 to 94) from death
		(13 Obs) [29][38][42][47][71][92][118][119][123][131][141][160][164]; last update 2022-03-02
Delta	Johnson &	Ad26.COV2.S provided protection against VOC Delta for the following
(1 dose)	Johnson	outcomes ≥ 14 days after dose:
	[AD26.COV	• 3% to 71% against infection (RME)
(up to 30	2.S]	• 50.9% (95% CI, 35.5 to 63.0) from symptomatic infection
days)		• 92.5% (95% CI, 54.9 to 99.6) from ICU admission
		• 90.5% (95% CI, 31.5 to 99.6) from death
		(6 Obs) [97][109][110][111][117][133]; last update 2021-12-15
Delta	Sinovac	CoronaVac provided protection against VOC Delta for the following
(1-2 doses)	[CoronaVac]	outcome at least 7 days after 2 nd dose:
(up to 30		• 60 to 74% from infection (RME)
days)		• 59% (95% CI, 16 to 81.6) from symptomatic infection
days		• 46 to 89% from severe disease (RME)
		• 76.5% (95% CI, 72.9 to 79.6) from death
		(3 Obs) [91][156][164]; last update 2022-03-02
		CoronaVac followed by ChAdOx1 provided protection against VOC Delta
		for the following outcomes at least 7 days after 2 nd dose:
		• 74% (95% CI, 43 to 99) from infection (1 Obs) [164]; <i>last update 2022-03-02</i>
Delta	AstraZeneca	ChAdOx1 followed by BNT162b2 at least 14 days after 2 nd dose provided
Dena	[ChAd0x1]	protection against VOC Delta for the following outcomes:
	Vaxzevria	• 67% (95% CI, 59 to 73) against symptomatic infection
	Serum	(1 Obs) [121]; last update 2021-12-01
1 dose	Institute of	
followed by an	India	ChAdOx1 followed by mRNA-1273 at least 14 days after 2 nd dose provided
mRNA	[Covishield]	protection against VOC Delta for the following outcomes:
vaccine		• 79% (95% CI, 62 to 88) against symptomatic infection
		(1 Obs) [121]; last update 2021-12-01
(up to 30		ChAdOx1 followed by either BNT162b2 or mRNA-1273 at least 14 days after
days)		2 nd dose provided protection against VOC Delta for the following outcomes:
		• 88% (95% CI, 85 to 89) against infection
		(1 Obs) [123]; last update 2021-12-01
		Chalon followed by DNIT1(2h2 and 11 1 and 11
		ChAdOx1 followed by BNT162b2 provided protection against infection by
		 VOC Delta compared to ChAdOx1 (homologous): HR 0.61 (95% CI, 0.52 to 0.71) unreported number of days after 2nd dose
		(1 Obs) [128]; last update 2021-12-01

VOC	Vaccine	Findings
Delta	Pfizer/	BNT162b2 showed a higher risk of infection by VOC Delta in participants
(2 doses)	BioNTech	fully vaccinated (≥14 days after 2 nd dose) longer than or equal to 146 days ago
	Comirnaty	vs fully vaccinated less than 146 days ago [OR 2.06 (95% CI, 1.69 to 2.51)]
(>30 days)	[BNT162b2]	(1 Obs) [69]; last update 2021-08-25
		BNT162b2 provided protection against infection by VOC Delta for the following number of days after 2 nd dose: 82 to 87% up to 60 days (RME) 67 to 74% from 21 to 98 days (RME) 53 to 85% up to 120 days (RME) 57 to 84% up to 150 days (RME) 90bs) [76][84][123][137][152][156] [158][163][169]; last update 2022-03-16 BNT162b2 provided protection against symptomatic infection by VOC Delta for the following number of days after 2 nd dose: 76% (95% CI, 72 to 81) – at 30 to 59 days (age 30-59) 72% (95% CI, 61 to 80) – at 60 to 89 days (age 30-59) 47% (95% CI, 39 to 55) – at 121 to 180 days 70.1% (95% CI, 68.9 to 71.2) – at 7 months (210 days) (4 Obs) [92][114][124][141]; last update 2022-01-05 BNT162b2 provided protection against severe, critical, or fatal disease by VOC Delta for the following number of days after 2 nd dose: 91.1% (95% CI, 90 to 92) at 44 to 98 days 68 to 72% up to 120 days 92 to 94% - age 40 to 59 up to 150 days (RME) 57 to 86% - age 60+ up to 150 days (RME) 57 to 86% - age 60+ up to 150 days (RME) (5 Obs) [76][125][156] [158][163]; last update 2022-03-02 BNT162b2 provided protection against death by VOC Delta for the following number of days after 2 nd dose: 81 to 89% up to 150 days (RME) (3 Obs) [124][125][156]; last update 2022-02-02 BNT162b2 provided protection against infection by VOC Delta at the following intervals between doses: 92% (95% CI, 91 to 93) at 14 to 27 days after 2 nd dose (interval 7+ weeks) 90% (95% CI, 88 to 91) at 4 months after 2 nd dose (interval 7+ weeks)
		(1 Obs) [<u>123</u>]; last update 2021-11-17
Delta	Moderna	mRNA-1273 provided protection against infection by VOC Delta the
(2 doses)	Spikevax	following number of days after 2 nd dose:
4.20.1	[mRNA-	• 71 to 94% up to 60 days (RME)
(>30 days)	1723]	• 79 to 83% up to 90 days (RME)
		• 81 to 88% at 120 days (RME)
		• 63.6% (95% CI, 51.8 to 72.5) at 91 to 180 days
		• 65 to 88% at 151 to 180 days (RME)
		 61.4% (95% CI, 56.8 to 65.5) at 181 to 270 days 52.9% (95% CI, 43.7 to 60.5) at >270 days
		(8 Obs) [101][123][137][143][152][157][158][169]; last update 2022-03-16
		(0 000) [101][122][121][122][121][120][102], mst upunt 2022-02-10

VOC	Vaccine	Findings
		mRNA-1273 provided protection against symptomatic infection by VOC
		Delta the following number of days after 2 nd dose:
		• 91% (95% CI, 85 to 95) – at 30 to 59 days (age 30-59)
		• 90% – at 70 to 98 days (RME)
		• 71% (95% CI, 56 to 81) – at 121 to 180 days
		• 81.9% (95% CI, 81 to 82.7) – at 7 months (210 days)
		(4 Obs) [92][114][124][141]; last update 2022-01-05
		mRNA-1273 provided protection against severe disease by VOC Delta the
		following number of days after 2 nd dose:
		• 97.8% (95% CI, 83.7 to 99.7) at 60 days
		• 74.5 to 93.4% up to 90 days (RME)
		• 91.5% (95% CI, 60.8 to 98.1) up to 120 days (RME)
		• 85.2% (95% CI, 82.7 to 87.7) at 150 days
		(3 Obs)[<u>143</u>][<u>157</u>][<u>158</u>]; last update 2022-02-16
		mRNA-1273 provided protection against death by VOC Delta the following
		number of days after 2 nd dose:
		• 96% (95% CI, 91.9 to 98) at 60 days
		• 93.7% (95% CI, 90.2 to 95.9) at 210 days
		(1 Obs) [124]; last update 2022-02-02
		mRNA-1273 provided protection against infection by VOC Delta at the
		following intervals between doses:
		• 92% (95% CI, 90 to 94) at 14 to 27 days after 2 nd dose (interval 7+ weeks)
		• 91% (95% CI, 87 to 94) at 4 months after 2 nd dose (interval 7+ weeks) (1 Obs) [123]; <i>last update 2021-11-17</i>
Delta	AstraZeneca	ChAdOx1 provided protection against infection by VOC Delta the following
	[ChAd0x1]	number of days after 2 nd dose:
(2 doses)	Vaxzevria	• 65 to 72% (95% CI, 66 to 77) at 120 days (RME)
	Serum	(2 Obs) [123][169]; last update 2022-03-16
(>30 days)	Institute of	
	India	ChAdOx1 provided protection against symptomatic infection by VOC Delta
	[Covishield]	the following number of days after 2 nd dose:
		• 63 to 67% – at 30 to 59 days (RME)
		• 65% (95% CI, 48 to 76) – at 60 to 89 days
		• 41 to 49% – at 120 days (17 weeks) (RME)
		• 69.7% (95% CI, 68.7 to 70.5) – at 140 days
		(4 Obs) [92][114][141][142]; last update 2022-01-05
		ChAdOx1 provided protection against severe disease by VOC Delta the
		following number of days after 2 nd dose:
		• 79.0% (95% CI, 75.9 to 81.7) at 56 to 63 days
		• 70.5% (95% CI, 67 to 73.7) at 112 to 119
		(1 Obs)[<u>142</u>]; last update 2022-01-05
		ChAdOx1 provided protection against infection by VOC Delta at the
		following intervals between doses:
		• 85% (95% CI, 60 to 94) at 14 to 27 days after 2 nd dose (interval 7+ weeks)
		• 72% (95% CI, 66 to 77) at 84+ days after 2 nd dose (interval 7+ weeks)

(1 Obs) [123]; last update 2021-11-17 Delta Johnson & Johnson [AD26.COV Delta the following number of days after dose: • 60% (95% CI, 57 to 62) from infection up to 60 days • 74% (95% CI, 70 to 76) from infection at ≥150 days • 89.4% (95% CI, 52.3 to 97.6) from death at 120 days (3 Obs) [124][152][169]; last update 2022-03-16	oy VOC
Johnson [AD26.COV Delta the following number of days after dose: (>30 days) Delta the following number of days after dose: 60% (95% CI, 57 to 62) from infection up to 60 days 74% (95% CI, 70 to 76) from infection at ≥150 days 89.4% (95% CI, 52.3 to 97.6) from death at 120 days	by VOC
(>30 days) [AD26.COV 2.S] • 60% (95% CI, 57 to 62) from infection up to 60 days • 74% (95% CI, 70 to 76) from infection at ≥150 days • 89.4% (95% CI, 52.3 to 97.6) from death at 120 days	
(>30 days) • 74% (95% CI, 70 to 76) from infection at ≥150 days • 89.4% (95% CI, 52.3 to 97.6) from death at 120 days	
• 89.4% (95% CI, 52.3 to 97.6) from death at 120 days	
$ (3 \text{ ODS}) 124 132 109 $, tast update $\frac{2022-09-10}{2000}$	
Ad26.COV2.S provided protection against symptomatic infection by	y VOC
Delta the following number of days after dose:	
• 50% (95% CI, 36 to 62) – at 30 to 59 days	
• 52% (95% CI, 33 to 66) – at 60 to 89 days	
• 64.3% (95% CI, 62.3 to 66.1) – at 150 days	
(2 Obs) [124][141]; last update 2022-01-05	
Delta Sinovac CoronaVac provided protection against the following outcomes by	VOC
[CoronaVac] Delta the following number of days after the 2 nd dose:	
(2 doses) • 30% (95% CI, 18.4 to 39.9) from infection up to 150 days	
• 30.2% (95% CI, 7.6 to 47.3) from ICU admission up to 150 days	
(>30 days) • 75.7% (95% CI, 67.0 to 82.1) from death up to 150 days	
(1 Obs) [156]; last update 2022-02-02	
Delta AstraZeneca ChAdOx1 followed by an mRNA provided protection against infect	ion by
[ChAd0x1] VOC Delta the following number of days after 2 nd dose:	
ChAdOx1 (1 Vaxzevria • 86% (95% CI, 81 to 89) at 120 days	
dose) followed Serum (1 Obs) [123]; last update 2021-11-17	
by mRNA Institute of	
vaccine India ChAdOx1 followed by an mRNA provided protection against symp	
[Covishield] infection by VOC Delta the following number of days after 2 nd dose	.•
• 67% (95% CI, 59 to 73) at least 14 days (BNT162b2)	
• 79% (95% CI, 62 to 88) at least 14 days (mRNA-1273)	
• 66% (95% CI, 41 to 80) – > 120 days (17 weeks)	
(2 Obs) [<u>114</u>][<u>121</u>]; last update 2022-01-05	
Delta Pfizer/ BNT162b2 (3 doses) provided protection against the following outc	omes
BioNTech compared to unvaccinated:	
(3 doses) Comirnaty • 81 to 93% from infection up to 30 days after 3 rd dose (RME)	
[BNT162b2] • 90% (95% CI, 89 to 90) up to 60 days after 3 rd dose	
(any time (5 Obs) [137][139][147][160][169]; last update 2022-03-16	
frame)	
BNT162b2 (3 doses) provided protection against symptomatic infec	tion
compared to unvaccinated:	(0.1)
• 94% (95% CI, 93.4 to 94.6) – at least 14 days after 3 rd dose (age 5	0+)
(1 Obs) [<u>126</u>]; last update 2021-12-15	
BNT162b2 (3 doses) provided protection against infection by VOC	Delta
compared to 2 doses:	Dena
• 84.0% (95% CI, 79 to 88) at 14 to 20 days after 3 rd dose	
• 64.0% (95% CI, 79 to 88) at 14 to 20 days after 3 dose • 45.7% (95% CI, 37.9 to 53.5) median of 30 days after 3 rd dose	
(2 Obs) [93][132]; last update 2021-12-15	
(2 ODS) [33][132], as in paint 2021-12-13	
BNT162b2 (3 doses) provided protection against the following outc	omes by
VOC Delta compared to 2 doses:	OIIICO DY

VOC	Vaccine	Findings
		Rate ratio 11.3 to 12.3 from infection at least 12 days after 3 rd dose
		• Rate ratio 17.9 to 19.5 from severe illness at least 12 days after 3 rd dose
		• Rate ratio 14.7 (95% CI, 10 to 21.4) from death at least 12 days after 3 rd
		dose
		• 90% (95% CI, 86 to 93) from death unclear number of days after 3 rd dose
		(3 Obs)[100][134][135]; last update 2022-01-05
Delta	Moderna	mRNA-1273 (3 doses) provided protection against infection by VOC Delta
	Spikevax	compared to unvaccinated:
(3 doses)	[mRNA-	• 83 to 95.7% up to 30 days after 3 rd dose (RME)
	1723]	• 92% (95% CI, 91 to 93) up to 60 days after 3 rd dose
(any time	_	(6 Obs) [137][139][147][148][160][169]; last update 2022-03-16
frame)	(up to 30	
,	days)	mRNA-1273 (3 doses) provided protection against infection by VOC Delta
		compared to 2 doses:
		• 46.6% (95% CI, 36.4 to 55.3) median of 16 days after 3 rd dose
		(1 Obs) [132]; last update 2021-12-15
Delta	AstraZeneca	ChAdOx1 (2 doses) followed by BNT162b2 provided protection against
	[ChAd0x1]	VOC Delta for the following outcomes:
2 doses	Vaxzevria	82% (95% CI, 68 to 90) from infection at least 7 days after 3rd dose
followed by 1	Serum	• 93.1 to 93.8% from symptomatic infection at least 14 days after 3 rd dose
dose of	Institute of	(RME)
another	India	(3 Obs) [126][136][139]; last update 2022-01-18
vaccine	[Covishield]	
	'	ChAdOx1 (2 doses) followed by mRNA-1273 provided protection against
(any time		VOC Delta for the following outcomes:
frame)		• 91% (95% CI, 63 to 98) from infection at least 7 days after 3rd dose
,		(1 Obs) [<u>139</u>]; last update 2022-01-05
Delta	Sinovac	CoronaVac (3 doses) provided protection against VOC Delta for the
	[CoronaVac]	following outcome ≥ 14 days after 3 rd dose:
(3 doses)		• 78.8% (95% CI, 76.8 to 80.6) from symptomatic infection
		(1 Obs) [154]; last update 2022-02-02
(any time		
frame)		
Delta	Sinovac	CoronaVac (2 doses) followed by BNT162b2 provided protection against
	[CoronaVac]	VOC Delta for the following outcomes at least 7 days after 3 rd dose:
2 doses		• 92.7 to 98% from infection (RME)
followed by 1		• 96.5% (95% CI, 96.2 to 96.7) from symptomatic infection
dose of		• 97.3% (95% CI, 96.1 to 98.1) from severe disease (hospitalization or death)
another		• 96.2% (95% CI, 94.6 to 97.3) from ICU admission
vaccine		• 96.8% (95% CI, 93.9 to 98.3) from death
		(3 Obs) [155][164][165]; last update 2022-03-02
(anytime		
frame)		CoronaVac (2 doses) followed by ChAdOx1 provided protection against
		VOC Delta for the following outcomes at least 7 days after 3 rd dose:
		• 86% (95% CI, 74 to 93) from infection
		• 93.2% (95% CI, 92.9 to 93.6) from symptomatic infection
		98.9% (95% CI, 98.5 to 99.2) from ICU admission
		• 98.1% (95% CI, 97.3 to 98.6) from death
		(2 Obs) [155][164]; last update 2022-03-02
	_1	(2 000) [100] 100], with apart 2022-07-02

VOC	Vaccine	Findings
Delta	Pfizer/	Fully vaccinated index cases by BNT162b showed VET to unvaccinated (hh
	BioNTech	contact):
Transmission	Comirnaty	• 31 to 63% (RME)
Household or	[BNT162b2]	
close contacts		Fully vaccinated index cases by BNT162b showed VET to fully vaccinated
of index case		household contacts:
		• 10 to 40% (RME)
		Fully vaccinated index cases by BNT162b showed VET to hh contacts
		(unclear status):
		• 65% (95% CI, 52 to 74)
		(3070 (3070 01, 32 10 7 1)
		Fully vaccinated hh contacts by BNT162b showed VES:
		• 46% (95% CI, 40 to 52) (vaccinated index case)
		• 61% (95% CI, 59 to 63) (unvaccinated index case)
		• 62 to 90% from infection (unclear status of index case) (RME)
		· · · · · · · · · · · · · · · · · · ·
		• 100% (95% CI, not reported) from severe disease
		(5 Obs) [105][107][108][129][149]; last update 2021-01-18
		BNT162b2 or mRNA-1273 (2 doses) hh contacts showed VES:
		• 46% (95% CI, 28 to 58) at least 7 days after 2 nd dose
		BNT162b2 or mRNA-1273 (3 doses) hh contacts showed VES:
		• 62% (95% CI, 38 to 78) at least 7 days after 3 rd dose
D-14-	M - 1	(1 Obs) [161]; last update 2022-03-02
Delta	Moderna	Fully vaccinated household contacts by mRNA-1273 showed VES (unclear
Transmission	Spikevax	status of index):
Household or	[mRNA-	• 62 to 77% from infection (RME)
close contacts	1723]	(2 Obs) [108][129]; last update 2021-12-01
of index case		DNIT1(2h2 on mDNIA 1272 (2 deces) his contacts aboved VES.
of fildex case		BNT162b2 or mRNA-1273 (2 doses) hh contacts showed VES:
		• 46% (95% CI, 28 to 58) at least 7 days after 2 nd dose
		BNT162b2 or mRNA-1273 (3 doses) hh contacts showed VES:
		• 62% (95% CI, 38 to 78) at least 7 days after 3 rd dose
Do14 -	A a t n = 17 =	(1 Obs) [161]; last update 2022-03-02
Delta	AstraZeneca	Fully vaccinated index cases by ChAdOx1 showed VET for household
Transmission	[ChAd0x1]	contacts (unclear status):
Household or	Vaxzevria Serum	• 36% (95% CI, 28 to 43) from infection
close contacts	Institute of	Fully vaccinated household contacts by ChAdOx1 showed VES (unclear
of index case	Institute of India	status of index):
of muck case	[Covishield]	• 55 to 72% from infection (RME)
Dolta	<u> </u>	(2 Obs)[107][108]; last update 2021-11-03
Delta	ChAdOx1	Fully vaccinated household contacts by ChAdOx1 followed by mRNA
Tromanical	followed by	showed VES (unclear status of index):
Transmission	mRNA	• 86% (95% CI, 45 to 97) from infection
Household or	vaccine	(1 Obs)[<u>108</u>]; last update 2021-11-03
close contacts		
of index case	Madens	
Gamma	Moderna	mRNA-1273 provided protection against VOC Gamma for the following
	Spikevax	outcomes 14 days after 1 st dose:
		• 85% (95% CI, 71 to 92) from infection

VOC	Vaccine	Findings
	[mRNA-	77% (95% CI, 63 to 86) from symptomatic infection
	1723]	• 89% (95% CI, 73 to 95) from hospitalization
		mRNA-1273 provided protection against VOC Gamma (or Beta) for the
		following outcomes 35-41 days after 1 st dose:
		• 43% (95% CI, 22 to 59) from symptomatic infection
		mRNA-1273 provided protection against VOC Gamma for the following
		outcome ate least 7 days after 2 nd dose:
		• 95% from infection (RME)
		88% (95% CI, 61 to 96) from symptomatic infection
	A . 7	(4 Obs – 5 refs) [23][47][101][122][123]; last update 2021-12-01
Gamma	AstraZeneca	ChAdOx1 provided protection against VOC Gamma for the following
	[ChAd0x1]	outcomes at least 14 days after 1 st dose:
	Vaxzevria	• 60% (95% CI, 48 to 69) from infection
	Serum	• 42 to 48% from symptomatic infection (RME)
	Institute of	• 83% (95% CI, 66 to 92) from hospitalization
	India	
	[Covishield]	ChAdOx1 provided protection against VOC Gamma for the following
		outcomes at least 14 days after 2 nd dose:
		• 90% (95% CI, 61 to 98) from infection
		• 65.4% (95% CI, 64.6 to 66.2) from symptomatic infection at 56 to 63 days
		after 2 nd dose
		• 58.7% (95% CI, 56.7 to 60.5) from symptomatic infection at 112 to 119
		days after 2 nd dose
		• 75.6% (95% CI, 73.4 to 77.6) from severe disease at 56 to 63 days after 2 nd
		dose
		• 50.5% (95% CI, 43.4 to 56.6) from severe disease at 112 to 119 days after
		2 nd dose
		(5 Obs)[47][116][122][123][142]; last update 2022-01-05
Gamma	Johnson &	Ad26.COV2-S provided protection against VOC Gamma for the following
Gaiiiiia	Johnson	outcomes 28 days after dose:
	[AD26.COV	• 50.9% (95% CI, 35.5 to 63.0) from symptomatic infection
	2.S]	
	2.0]	• 92.5% (95% CI, 54.9 to 99.6) from ICU admission
		• 90.5% (95% CI, 31.5 to 99.6) from death
	0.	(1 Obs) [117], last update 2021-11-17
Gamma	Sinovac	CoronaVac provided protection against VOC Gamma for the following
	[CoronaVac]	outcome ≥ 14 days after 2^{nd} dose:
		• 65.9% (95% CI, 65.2 to 66.6) from infection
		CoronaVac provided protection against VOC Gamma for the following
		outcome \geq 14 days after 2 nd dose for people over age 70:
		• 41.6% (95% CI, 26.9 to 63.3) from symptomatic infection
		(2 Obs) [30][49]; last update 2021-07-14
Gamma	ChAdOx1	ChAdOx1 followed by either BNT162b2 or mRNA-1273 at least 14 days after
	followed by	2 nd dose provided protection against VOC Gamma for the following
	mRNA	outcomes:
	vaccine	• 96% (95% CI, 70 to 99) against infection
		(1 Obs) [123]; last update 2021-11-17
Beta	Moderna	mRNA-1273 provided protection against VOC Beta for the following
	Spikevax	outcomes 14 days after 1 st dose:
	1	• 61.3% (95% CI, 56.5 to 65.5) from infection
L	I	1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2

VOC	Vaccine	Findings
	[mRNA-	• 77% (95% CI, 63 to 86) from symptomatic infection
	1723]	• 89% (95% CI, 73 to 95) from hospitalization
		• 81.6% (95% CI, 71.0 to 88.8) from severe, critical, or fatal disease
		(combined with Alpha)
		mRNA-1273 provided protection against VOC Beta for the following
		outcomes 35-41 days after 1 st dose:
		• 43% (95 CI, 22 to 59) from symptomatic infection
		mRNA-1273 provided protection against VOC Beta for the following
		outcome 7 days after 2 nd dose:
		• 96.4% (95% CI, 91.9 to 98.7) from infection
		88% (95% CI, 61 to 96) from symptomatic infection
		• 95.7% (95% CI, 73.4 to 99.9) from severe, critical, or fatal disease
		(combined with Alpha)
		(2 Obs – 3 refs) [23][47][50]; last update 2021-07-14
Beta	AstraZeneca	ChAdOx1 provided protection against VOC Beta for the following outcome
	[ChAd0x1]	14 days after 1 st dose:
	Vaxzevria	• 48% (95% CI, 28 to 63) from symptomatic infection
	Serum	• 83% (95% CI, 66 to 92) from hospitalization
	Institute of	ChAdOx1 provided protection against VOC Beta for the following outcome
	India	after 2 doses:
	[Covishield]	• 10.4% (95% CI, -76.8 to 54.8) from mild to moderate disease
D.	N.T.	(1 RCT, moderate quality; 1 Obs) [4] [47]; last update 2021-07-07
Beta	Novavax	NVX-CoV2373 provided protection against VOC Beta for the following
	[NVX- CoV2373	outcome after 7 days after 2 nd dose:
	C0V2575	• Post-hoc: 43% (95% CI, -9.8 to 70.4) from symptomatic infection
A11	Moderna	(1 RCT, moderate quality), [17]; last update 2021-07-14
Alpha	Spikevax	mRNA-1273 provided protection against VOC Alpha for the following outcomes 14-41 days after 1 st dose:
	[mRNA-	• 58.9 to 88.1% from infection (RME)
	1723]	60 to 61% from symptomatic infection (RME)
	1723]	81.6% (95% CI, 71.0 to 88.8) from severe, critical, or fatal disease
		(combined with Beta)
		mRNA-1273 provided protection against VOC Alpha for the following
		outcomes at least 7 days after 2 nd dose:
		86 to 100% from infection (RME)
		90 to 95.7% from symptomatic infection (RME)
		• 95.7% (95% CI, 73.4 to 99.9) from severe, critical, or fatal disease
		(combined with Beta)
		(10 Obs – 11 refs) [8][23][31][34][37][47][50][60][74][101][102]; last update
		2021-10-20
Alpha	AstraZeneca	ChAdOx1 provided protection against VOC Alpha for the following outcome
_	[ChAd0x1]	14 days after 1 st dose:
	Vaxzevria	• 64% (95% CI, 60 to 68) from symptomatic infection
	Serum	• 85% (95% CI, 81 to 88) from hospitalization
	Institute of	ChAdOx1 provided protection against VOC Alpha for the following outcome
	India	21 to 28 days after 1 st dose:
	[Covishield]	• 44 to 74% from infection (RME)
		ChAdOx1 provided protection against confirmed VOC Alpha for the
		following outcome at least 14 days after 2 doses:

VOC	Vaccine	Findings
		• 62 to 79% from infection (RME)
		(1 RCT, moderate quality; 5 Obs)[9][10][5][47][70][71][]; last update 2021-08-25
Alpha	Novavax	NVX-CoV2373 provided protection against VOC Alpha for the following
	[NVX-	outcome after 2 doses:
	CoV2373	• 89.7% (95% CI, 80.2 to 94.6) from symptomatic infection.
		No hospitalizations or deaths in vaccinated group
		• Post hoc: 86.3% (95% CI, 71.3 to 93.5) from confirmed Alpha
		symptomatic infection
		(1 RCT, moderate quality), [19]; last update 2021-06-16
Alpha	ChAdOx1	ChAdOx1 followed by BNT162b2 or mRNA-1273 at least 14 days after 2 nd
	followed by	dose provided protection against VOC Alpha for the following outcomes:
	mRNA	• 88% (95% CI, 83 to 92) against infection
	vaccine	(1 Obs) [70]; last search date 2021-08-25
Alpha	Pfizer/	BNT162b2 reduced transmission of VOC Alpha (VET) from a vaccinated
	BioNTech	index case (14 to 21 days after 1st dose) to household contacts compared to
Transmission	Comirnaty	households of unvaccinated index cases:
Household or	[BNT162b2]	• 30 to 49% from infection (RME)
close contacts		BNT162b2 reduced transmission of VOC Alpha (VET) from a vaccinated
of index case		HCW (10 weeks after 1st dose) to household spouse:
		• 42.9% (95% CI, 22.3 to 58.1) from infection
		Fully vaccinated index cases showed VET for household contacts (unclear
		status):
		• 70 to 82% from infection (RME) <u>Fully vaccinated household contacts</u> showed VES (unclear status of index):
		• 65 to 94% from infection (RME)
		(8 Obs) [6][14][33][40][48][104][107][108]; last update 2021-11-03
Alpha	Moderna	mRNA-1273 reduced transmission of VOC Alpha (VET) from a vaccinated
124710	Spikevax	HCW (10 weeks after 1 st dose) to household spouse:
Transmission	[mRNA-	• 42.9% (95% CI, 22.3 to 58.1) from infection
Household or	1723]	Fully vaccinated index cases by mRNA-1273 showed VET for household
close contacts	_	contacts (unclear status):
of index case		• 88% (95% CI, 50 to 97) from infection
		Fully vaccinated household contacts by mRNA-1273 showed VES (unclear
		status of index):
		86 to 91% from infection (RME)
		(3 Obs)[<u>33</u>][<u>104</u>][<u>108</u>]; last update 2021-11-03
Alpha	AstraZeneca	ChAdOx1 reduced transmission of VOC Alpha (VET) from a vaccinated
/m	[ChAd0x1]	index case (14 to 21 days after 1st dose) to household contacts compared to
Transmission	Vaxzevria	households of unvaccinated index cases:
Household or	Serum	• 30 to 47% from infection (RME)
close contacts	Institute of	Fully vaccinated index cases by ChAdOx1 showed VET to household
of index case	India	contacts (unclear status):
	[Covishield]	• 58 to 63% from infection (RME)
		Fully vaccinated household contacts by ChAdOx1 showed VES (unclear status of index case):
		status of index case):
		• 38 to 87% from infection (RME)
Alpha	Johnson &	(6 Obs) [6][14][40][104][107][108]; last update 2021-12-01 Fully vaccinated index cases by Ad26.COV2.S showed VET for household
Alpha	Johnson & Johnson	contacts (unclear status):
	Joinison	Contacts (unclear status).

VOC	Vaccine	Findings
Transmission	[AD26.COV	• 77% (95% CI, 6 to 94) from infection
Household or	2.S]	Fully vaccinated household contacts by Ad26.COV2.S showed VES (unclear
close contacts		status of index):
of index case		• 12% (95% CI, -71 to 54) from infection
		(1 Obs) [<u>104</u>]; last update 2021-11-03

VOC) Alpha to Delta	Pfizer/	BNT162b2 provided protection against infection by VOC Alpha to
inpila to Della	BioNTech	Delta at least 7 days after 2 nd dose:
		• 69.7% (95% CI, 68.6 to 70.8)
	Comirnaty	
	[BNT162b2]	BNT162b2 or mRNA-1273 provided protection against VOC Alpha
		to Delta for the following outcomes \geq 14 days after 2 nd dose:
		• 57% (95% CI, 53 to 60) from infection at 144 days after 2nd dose
		• 68% (95% CI, 64 to 71) from symptomatic infection at 42 to 69 days after 2 nd dose
		• 39% (95% CI, 29 to 48) from symptomatic infection at 98 to 148 days after 2 nd dose
		• 92% (95% CI, 85 to 96) from severe disease in people with no risk conditions
		• 72% (95% CI, 51 to 84) from severe disease with very high risk conditions
		• 95% (95% CI, 88 to 98) from death at 14 to 41 days after 2 nd dose
		• 86 to 93% from death at 70 to 148 days after 2 nd dose(RME)
		BNT162b2 showed OR 1.61 (95% CI, 1.45 to 1.79) for infection
		comparing <u>fully vaccinated Jan to Feb</u> (VOC_Alpha) vs <u>fully</u>
		vaccinated Mar to May (VOC Delta).
Alpha to Delta	Pfizer/	(5 Obs) [95] [96] [127] [144] [145]; last update 2022-12-01 BNT162b2 (3 doses) provided protection against VOC Alpha to
Alpha to Delta	BioNTech (3	Delta for the following outcomes compared to unvaccinated:
	doses)	• 88% (95% CI, 86 to 89) from infection at least 14 days after 3rd dose (age>18)
	Comirnaty	4000 (480-10)
	[BNT162b2]	BNT162b2 (3 doses) provided protection against VOC Alpha to
		Delta for the following outcomes:
		• 75% (95% CI, 71 to 78) from infection at least 14 days after 3rd
		dose compared to 2 doses (given at least 6 months previously)
		(age>18)
	3.5.5	(1 Obs) [146]; last update 2022-01-05
Alpha to Delta	Moderna	mRNA-1273 provided protection against infection by VOC Alpha
	Spikevax	to Delta at least 7 days after 2 nd dose:
	[mRNA-1723]	• 78.2% (95% CI, 76.7 to 79.6)
		mRNA-1273 or BNT162b2 provided protection against VOC Alpha
		to Delta for the following outcomes \geq 14 days after 2 nd dose:
		• 73% (95% CI, 70 to 76) from infection at 144 days after 2 nd dose
		• 92% (95% CI, 85 to 96) from severe disease in people with no
		risk conditions

_	e Frame for More	than One VOC (insufficient data to divide them into separate
VOC)		- 720/ /050/ CT 54 + 0 A C
		• 72% (95% CI, 51 to 84) from severe disease with very high risk conditions
		• 93% (95% CI, 81 to 97) from death at 144 days after 2 nd dose
Alpha to Delta	AstraZeneca	(3 Obs) [95][127][145]; last update 2022-01-05 ChAdOx1 provided protection against infection by VOC Alpha to
Aiplia to Delta	[ChAd0x1]	Delta at least 7 days after 2 nd dose:
	Vaxzevria	• 43.4% (95% CI, 4.4 to 66.5)
	Serum Institute	13.170 (7370 C1; 1.1 to 00.3)
	of India	ChAdOx1 provided protection against VOC Alpha to Delta for the
	[Covishield]	following outcomes \geq 14 days after 2 nd dose:
		• 94% (95% CI, 90 to 96) from severe disease in people with no
		risk conditions
		• 63% (95% CI, 46 to 75) from severe disease with very high risk
		conditions
		• 33% (95% CI, 23 to 42) from symptomatic infection at 42 to 69
		days after 2 nd dose
		• 34% (95% CI, 10 to 52) from symptomatic infection at 70 to 140
		days after 2 nd dose
		• 95% (95% CI, 90 to 97) from death at least 14 days after 2 nd dose
41.1 . 10.1	T 1	(2 Obs) [95][127][144]; last update 2022-01-05
Alpha to Delta	Johnson &	Ad26.COV2.S provided protection against VOC Alpha to Delta for
	Johnson [AD26.COV2.S]	the following outcomes ≥ 14 days after 2^{nd} dose:
	[AD20.CO v2.5]	 36% (95% CI, 30 to 42) from infection at 144 days after 2nd dose 72% (95% CI, 49 to 85) from death at 144 days after 2nd dose
		(1 Obs) [145]; last update 2022-01-05
Alpha to Delta	Heterologous	Heterologous mRNA vaccines provided protection against infection
rupila to Delta	mRNA	by VOC Alpha to Delta at least 7 days after the 2 nd dose:
	vaccines	• 84.7% (83.1 to 86.1)
	ChAdOx1	ChAdOx1 followed by either BNT162b2 or mRNA-1273 provided
	followed by	protection against infection by VOC Alpha to Delta at least 7 days
	mRNA vaccine	after 2 nd dose:
		• 60.7% (95% CI, 57.5 to 63.6)
		(1 Obs) [127]; last update 2021-12-01
Alpha to Delta	Moderna	mRNA-1273 or BNT162b showed OR of 8.89 (95% CI, 5.92 to
36.	Spikevax	13.34) for unvaccinated vs fully vaccinated against infection (VOC
Maintenance	[mRNA-1723]	Alpha)
hemodialysis		mPNIA 1273 or RNIT162b abound OP of 2.27 (050/ CL 1.72 to
(not updated after		mRNA-1273 or BNT162b showed OR of 2.27 (95% CI, 1.72 to 3.00) for unvaccinated vs fully vaccinated against infection (VOC
Nov 5, 2021)		Delta)
1 10 v 2, 2021)		(1 Obs) [106]; last update 2021-11-03
Alpha or Beta	Pfizer/	BNT162b2 or mRNA-1273 provided protection against infection by
1	BioNTech	VOC Alpha or Beta at the following number of days after 2 nd dose:
Immunosuppressed,		• 46.6% (95% CI, 0.0 to 73.7) ≥14 days
renal transplant	Comirnaty	• 66.0% (95% CI, 21.3 to 85.3) ≥42 days
_	[BNT162b2]	• 73.9% (95% CI, 33 to 98.9) ≥56 days
(not updated after		BNT162b2 or mRNA-1273 provided protection against severe,
Nov 5, 2021)		critical, or fatal disease by VOC Alpha or Beta at the following
		number of days after 2 nd dose:

Studies Covering Time VOC)	e Frame for More	e than One VOC (insufficient data to divide them into separate
		 72.3% (95% CI, 0.0 to 90.9) ≥14 days 85% (95% CI, 35.7 to 96.5) ≥42 days 83.8% (95% CI, 31.3 to 96.2) ≥56 days (1 Obs) [90]; last update 2021-09-22
Alpha or Beta Immunosuppressed, renal transplant (not updated after Nov 5, 2021)	Moderna Spikevax [mRNA-1723]	mRNA-1273 or BNT162b2 provided protection against infection by VOC Alpha or Beta at the following number of days after 2 nd dose: • 46.6% (95% CI, 0.0 to 73.7) ≥14 days • 66.0% (95% CI, 21.3 to 85.3) ≥42 days • 73.9% (95% CI, 33 to 98.9) ≥56 days mRNA-1273 or BNT162b2 provided protection against severe, critical, or fatal disease by VOC Alpha or Beta at the following number of days after 2 nd dose: • 72.3% (95% CI, 0.0 to 90.9) ≥14 days
		 85% (95% CI, 35.7 to 96.5) ≥42 days 83.8% (95% CI, 31.3 to 96.2) ≥56 days (1 Obs) [90]; last update 2021-09-22
Alpha or Beta Previously infected (not updated after Nov 5, 2021)	Pfizer/ BioNTech Comirnaty [BNT162b2]	BNT162b2 (2 doses) <u>after prior infection</u> provided protection against VOC Alpha (or Beta) for the following outcomes: • 85% (95% CI, 80 to 89) against re-infection compared to BNT162b2 without prior infection (1 Obs) [72]; last update 2021-08-25
Nov 5, 2021) Alpha or Beta Previously infected (not updated after	Moderna Spikevax [mRNA-1723]	mRNA-1273 (2 doses) <u>after prior infection</u> did not offer additional protection against VOC Alpha (or Beta) for the following outcomes: • 15% (95% CI, -105 to 66) against re-infection compared to mRNA-1273 without prior infection (1 Obs) [72]; <i>last update 2021-08-25</i>
Nov 5, 2021) Beta to Delta	Pfizer/ BioNTech Comirnaty [BNT162b2]	BNT162b2 provided protection against infection by VOC Beta to VOC Delta for the following number of days after the 2 nd dose: • 65.8% (95% CI, 63.8 to 67.7) at 5 to 9 weeks • 29.7% (95% CI, 21.7 to 36.9) at 15 to 19 weeks • 0% (95% CI, 0 to 0) 20 to 24 weeks BNT162b2 provided protection against hospitalization or death by VOC Beta to VOC Delta for the following number of days after the 2 nd dose: • 94.2% (95% CI, 91.0 to 96.5) at 5 to 9 weeks • 86.4% (95% CI, 69.9 to 94.8) at 15 to 19 weeks • 95.3% (95% CI, 70.5 to 99.9) at 20 to 24 weeks (1 Obs) [98]; <i>last update</i> 2021-10-06
HCW (not updated after Nov 5, 2021)	Pfizer/ BioNTech Comirnaty [BNT162b2]	BNT162b2 provided protection against VOC Beta or Gamma for the following outcomes 14 to 42 days after 1 st dose: • 37.2% (95% CI, 16.6 to 52.7) from infection BNT162b2 provided protection against VOC Beta or Gamma for the following outcome 7 days after 2 nd dose: • 79.2% (95% CI, 64.6 to 87.8) from infection (1 Obs)[27]; last update 2021-06-01

Studies Covering Time Frame for More than One VOC (insufficient data to divide them into separate		
VOC)		
Beta or Gamma	Pfizer/	BNT162b2 reduced transmission of VOC Beta or Gamma from
	BioNTech	vaccinated HCW (VET) compared to unvaccinated community ≥14
Transmission	Comirnaty	days after 1 st dose:
Vaccinated HCW vs	[BNT162b2]	• 54.7% (95% CI, 44.8 to 62.9) from infection
unvaccinated		BNT162b2 reduced transmission of VOC Beta or Gamma from
community		vaccinated HCW (VETompared to unvaccinated community ≥7
		days after 2 nd dose:
		• 84.8% (95% CI, 75.2 to 90.7) from infection
		(1 Obs) [<u>27]</u> ; last update 2021-06-08

Special Populations	(will not be updated a	after November 5, 2021)
Delta	Pfizer/	BNT162b2 provided protection against VOC Delta for the
	BioNTech	following outcomes at least 14 days after 1 st dose:
Adolescents	Comirnaty	• 59% (95% CI, 52 to 65) from infection
	[BNT162b2]	BNT162b2 provided protection against VOC Delta for the
(moved to		following outcomes at least 7 days after 2 nd dose:
Pediatric/Adolescent		• 90 to 92% against infection (RME)
LES)		(2 Obs) [112][120]; last update 2021-11-17
Delta	Pfizer/	BNT162b2 provided protection against VOC Delta for the
	BioNTech	following outcomes \geq 14 days after 2 nd dose:
HCW	Comirnaty	• 66% (95% CI, 26 to 84)
	[BNT162b2]	(1 Obs) [81]; last update 2021-09-22
Delta	AstraZeneca	ChAdOx1 provided protection against VOC Delta for the
	[ChAd0x1]	following outcomes at least 14 days after 2nd dose:
HCW	Vaxzevria	• 54 to 85% from infection (RME)
	Serum Institute of	• 64% (95% CI, 38 to 78) from symptomatic infection
	India	(2 Obs) [59][66]; last update 2021-10-06
	[Covishield]	
Delta	Pfizer/	BNT162b2 (2 doses) provided protection against VOC Delta for
	BioNTech	the following outcomes compared to <u>natural immunity</u> <u>after prior</u>
Previously	Comirnaty	infection:
infected,	[BNT162b2]	• 66% (95% CI, 22 to 86) from infection
(65+)		(1 Obs) [<u>103</u>]; last update 2021-10-20
Delta	Moderna	mRNA-1273 (2 doses) provided protection against VOC Delta for
	Spikevax	the following outcomes compared to <u>natural immunity</u> <u>after prior</u>
Previously infected	[mRNA-1723]	infection:
(65+)		• 68% (95% CI, 30 to 86) from infection
		• 30% (-11 to 1) from death
		(1 Obs) [<u>103</u>]; last update 2021-10-20
Delta	Moderna	mRNA-1273 provided protection against VOC Delta for the
	Spikevax	following outcomes at least 14 days after 2 nd dose:
Prison		57% (95% CI, 42 to 67.5)
	[mRNA-1723]	(1 Obs) [113]; last update 2021-11-03
Gamma	Sinovac	CoronaVac provided protection against VOC Gamma for the
	[CoronaVac]	following outcomes ≥14 days after 1 st dose:
HCW		• 35.1% (95% CI, -6.6 to 60.5) from infection
		• 49.6% (95% CI, 11.3 to 71.4) from symptomatic infection
		(1 Obs)[18]; last update 2021-05-07

Special Populations	s (will not be updated a	after November 5, 2021)
Gamma	Pfizer/	BNT162b2 (or mRNA-1273) provided protection against VOC
	BioNTech	Gamma 14 days after 2 nd dose:
LTC residents	Comirnaty	• 52.5% (95% CI, 26.9 to 69.1) against infection
	[BNT162b2]	• 78.6% (95% CI, 47.9 to 91.2) against severe disease
		(1 Obs) [61]; last update 2021-08-11
Gamma	Moderna	mRNA-1273 (or BNT162b2) provided protection against VOC
	Spikevax	Gamma for the following outcomes 14 days after 2 nd dose:
LTC residents	[mRNA-1723]	• 52.5% (95% CI, 26.9 to 69.1) against infection
	'	• 78.6% (95% CI, 47.9 to 91.2) against severe disease
		(1 Obs) [61]; last update 2021-08-11
Gamma	Pfizer/	BNT162b2 provided protection against VOC Gamma for the
	BioNTech	following outcomes ≥ 21 days after 1 st dose:
Over 70 years	Comirnaty	• 61% (95% CI, 45 to 72) from infection
j	[BNT162b2]	(1 Obs)[<u>35</u>]; last update 2021-07-07
Gamma	Moderna	mRNA-1273 provided protection against VOC Gamma for the
	Spikevax	following outcome ≥21 days after 1 st dose:
Over 70 years	[mRNA-1723]	• 61% (95% CI, 45 to 72) from infection
		(1 Obs) [35]; last update 2021-06-23
Alpha	Pfizer/	BNT162b2 provided protection against VOC Alpha for the
r	BioNTech	following outcomes 14 to 21 days after 1 st dose:
HCW	Comirnaty	• 64 to 84% from infection (RME)
	[BNT162b2]	BNT162b2 provided protection against VOC Alpha for the
	'	following outcomes at least 7 days after 2 nd dose:
		• 90 to 97% from infection (RME)
		BNT162b2 provided protection against VOC Alpha for the
		following outcome 7 days after 2 nd dose:
		• 86% (95% CI, 69 to 93) from asymptomatic infection [25]
		BNT162b2 provided protection against infection by VOC Alpha
		for the following number of days after 2 nd dose:
		• 85% (95% CI, 68 to 93) at 14 to 119 days
		• 73% (95% CI, 49 to 86) ≥150 days
		(6 Obs)[11][34][45][46][56][81]; last update 2021-11-17
Alpha	AstraZeneca	ChAdOx1 provided protection against VOC Alpha for the
прпа	[ChAd0x1]	following outcomes at least 14 days after 1st dose:
HCW	Vaxzevria	• 64% (95% CI, 50 to 74) from infection
IIC W	Serum Institute of	ChAdOx1provided protection against VOC Alpha for the
	India	following outcomes at least 14 days after 2 nd dose:
	[Covishield]	• 90% (95% CI, 62 to 98) from infection
	[Covisincia]	(1 Obs) [46]; last update 2021-07-07
Alpha	Pfizer/	BNT162b2 provided protection against VOC Alpha for the
Аірпа	BioNTech	following outcomes 7 days after 2 nd dose:
LTC residents	Comirnaty	• 53% (95% CI, 29 to 69) from infection
LI O ICSIGCIIIS	[BNT162b2]	• 89% (95% CI, 81 to 93) from death
	[D11110202]	
A11	DC: - · /	(1 Obs)[32]; last update 2021-10-06
Alpha	Pfizer/	BNT162b2 provided protection against VOC Alpha for the
0	BioNTech	following outcomes 7 days after 2 nd dose:
Over 65 years,	Comirnaty	• 86% (95% CI, 78 to 91) from infection
requiring home	[BNT162b2]	• 97% (95% CI, 88 to 99) from death
support		(1 Obs)[<u>32</u>]; last update 2021-07-07

Special Populations	(will not be updated a	after November 5, 2021)		
Alpha	Pfizer/	BNT162b2 provided protection against VOC Alpha for the		
	BioNTech	following outcomes at least 21 days after 1st dose:		
Over 70 years	Comirnaty	• 41 to 67% from infection (RME)		
	[BNT162b2]	BNT162b2 provided protection against VOC Alpha for the		
		following outcomes at least 7 days after 2 nd dose:		
		• 75 to 90% from infection (RME)		
		(3 Obs)[<u>28</u>][<u>35</u>][<u>51</u>]; last update 2021-10-06		
Alpha	Moderna	mRNA-1273 provided protection against VOC Alpha for the		
	Spikevax	following outcome ≥21 days after 1 st dose:		
Over 70 years	[mRNA-1723]	• 67% (95% CI, 57 to 75) from infection		
		(1 Obs) [<u>35</u>]; last update 2021-06-23		
Alpha	AstraZeneca	ChAdOx1 provided protection against VOC Alpha for the		
	[ChAd0x1]	following outcomes at least 14 days after 2 nd dose:		
Over 80 years	Vaxzevria	• 88% (95% CI, 48 to 97) from symptomatic infection		
	Serum Institute of	te of (1 Obs) [79]; last update 2021-10-20		
	India			
	[Covishield]			
Alpha	Pfizer/	BNT162b2 provided protection against VOC Alpha for the following outcomes at least 28 days after 1st dose:		
	BioNTech	following outcomes at least 28 days after 1st dose:		
Pregnant	Comirnaty	• 78% (95% CI, 57 to 89) from infection		
	[BNT162b2]	BNT162b2 provided protection against VOC Alpha for the		
		following outcomes 7 to 56 days after 2 nd dose:		
		• 86.1% (95% CI, 82.4 to 89.1) from infection		
		• 89% (95% CI, 43 to 100) from hospitalization		
		(2 Obs) [<u>52</u>][<u>54</u>]; last update 2021-07-28		
Epsilon	Pfizer/	BNT162b2 provided protection against VOC Epsilon for the		
	BioNTech	following outcome 15 days after 1 st dose:		
	Comirnaty	• 58.9% (95% CI, -9.7 to 84.5) from infection		
	[BNT162b2]	BNT162b2 provided protection against VOC Epsilon for the		
		following outcome 15 days after 2 nd dose:		
		• 85.7% (67.2 to 93.9) from infection		
	36.1	(2 Obs) [8][31]; last update 2021-06-08		
Epsilon	Moderna	mRNA-1273 provided protection against VOC Epsilon for the		
	Spikevax	following outcome 15 days after 1 st dose:		
	[mRNA-1723]	• 58.9% (95% CI, -9.7 to 84.5) from infection		
		mRNA-1273 provided protection against VOC Epsilon for the		
		following outcome 15 days after 2 nd dose:		
		• 85.7% (67.2 to 93.9) from infection		
T :1 +		(2 Obs) [8][31]; last update 2021-06-08		

Links to references are provided in Appendix 1

Iorio A, Little J, Linkins L, Abdelkader W, Bennett D, Lavis JN. COVID-19 living evidence synthesis #6 (version 6.32): What is the efficacy and effectiveness of available COVID-19 vaccines in general and specifically for variants of concern? Health Information Research Unit (HIRU); McMaster and Ottawa Knowledge Synthesis and Application Unit, 2 March 2022.

To help Canadian decision-makers as they respond to unprecedented challenges related to the COVID-19 pandemic, COVID-END in Canada is preparing rapid evidence responses like this one. The development and continued updating of this living evidence synthesis has been funded by the Canadian Institutes of Health Research (CIHR) and the Public Health Agency of Canada. The opinions, results, and conclusions are those of the team that prepared the living evidence synthesis, and independent of the Government of Canada, CIHR and the Public Health Agency of Canada. No endorsement by the Government of Canada, CIHR or Public Health Agency of Canada is intended or should be inferred.

Appendix 1: Summary of Study Findings and Appraisals

	Section 1: included studies				
Ref	Author	Bottom line	ROBINS- I*	Design, Notes	
		*Note: ROBINS-I score risk of bias: Low risk of	_	s high quality	
1	<u>Dagan</u>	BNT162b2 showed VE 46% (95% CI, 40 to 51) against infection 14 to 20 days after 1 st dose and VE 92% (95% CI, 88 to 95) 7 days after 2 nd dose. BNT162b2 showed VE 92% (95% CI, 75 to 100) for severe disease at 7 days after 2 nd dose.	Moderate	Data-linkage study in Israel; .5 M matched participants (2 M excluded – also (possible overlap with Haas); time and setting for VOC Alpha (estimated 80%).	
2	Haas	BNT162b2 showed VE 95.3% (95% CI, 94.9 to 95.7) against infection; VE 97.5% (95% CI, 97.1 to 97.8) against severe or critical COVID-19-related hospitalization; VE 96.7% (95% CI, 96.0 to 97.3) against death 7 days after 2 nd dose.	Serious	Data-linkage study in Israel; >6.5 M matched participants (possible overlap with Dagan) Updated May 14 due to final publication; sample confirmed VOC Alpha (estimated 94%).	
3	*Delayed exclusion- only included infected	BNT162b2 showed lower relative VE (2.4:1) against Alpha. after 1 st dose; and lower VE (8:1) against Beta after 2 nd dose in a population with >90% of Alpha and <1% Beta	Moderate	Case-control study in Israel; small sample for Beta (no overlap CHS cohort); confirmed VOC Alpha and Beta.	
4	<u>Madhi</u>	ChAdOx1 nCoV-19 showed VE 10.4% (95% CI, -76.8 to 54.8) against mild to moderate disease 14 days after 2 nd dose.	Moderate quality (RCT)	RCT in South Africa; Underpowered for 20% efficacy (42 cases); VOC Beta.	
5	Emary	ChAdOx1nCoV-19 showed VE 61.7% (95% CI, 36.7 to 76.9) against infection by VOC Alpha \geq 15 days after 2 nd dose.	Moderate quality (RCT)	RCT in UK; neutralization of Alpha 9 times lower; no sequencing for 45% of cases; 52 cases (19%) had VOC Alpha.	
6	Shah	ChAdOx1nCoV-19 or BNT162b2 reduced infection in unvaccinated household contacts of vaccinated HCW by about 30% (HR, 0.70, 95% CI, 0.63 to 0.78) ≥ 14 days after 1 st dose; ChAdOx1nCoV-19 or BNT162b2 reduced infection in HCW by about 55% (HR 0.45, 95% CI, 0.42 to 0.49) and hospitalization by 84% (HR 0.16, 95% CI, 0.09 to 0.27) ≥ 14 days after 1 st dose.	Moderate	Data-linkage study in Scotland - (25% of cases had received 2 doses); time and setting for VOC Alpha.	
7	Sadoff	Single dose Ad26.COV2.S showed VE 38.1% (95% CI, 4.2 to 60.4) at 14 days and VE 51.9% (95% CI, 19.1 to 72.2) at 28 days against moderate to severe disease and VE 81.7% (95% CI, 46.2 to 95.4) at 28 days	Moderate quality (RCT) Updated 2022/03/16	RCT; over 40,000 participants; Argentina, Brazil, Chile, Colombia, Mexico, Peru, South Africa, and the United States; sequenced for VOC Alpha, Beta, Delta, Gamma.	

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		against severe disease (confirmed VOC		
		Beta).		
		C: 1 1 A 107 COMO C 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		
		Single dose Ad26.COV2.S showed VE		
		36.4% (95% CI, 13.9 to 53.2) at 14 days and		
		VE 36.5% (95% CI, 14.1 to 53.3) at 28 days		
		against moderate to severe disease		
		(confirmed VOC Gamma)		
8	<u>Andrejko</u>	BNT162b2 or mRNA-1273 showed VE	Serious	Test-negative study in
		58.9% (95% CI, -9.7 to 84.5) at 15 days		California; 645 participants;
		after 1 st dose, and VE 85.7% (95% CI, 67.2		69% of population at time had
		to 93.9) 15 days after 2 nd dose against		VOC Alpha or Epsilon.
		infection.		
9	Glampson	ChAdOx1nCoV-19 showed VE 74% (95%	Serious	Retrospective cohort in UK;
	-	CI, 65 to 81) against infection 28 days after		2M participants; time and
		1 st dose.		setting for VOC Alpha.
		BNT162b2 showed VE 78% (95% CI, 73 to		
		82) against infection 28 days after 1 st dose.		
10	Pritchard	ChAdOx1nCoV-19 or BNT162b2 showed	Serious	Survey of randomly selected
		VE 66% (95% CI, 59 to 72%) 21 days after	0 2220 410	private households with
		1 st dose and 78% (95% CI, 68 to 85%) after		longitudinal follow-up in UK;
		2 nd dose against infection.		370,000 participants; sample
		2 dose against infection.		confirmed VOC Alpha.
11	Hall	BNT162b2 vaccine showed VE of 70%	Moderate	Prospective cohort with
11	(SIREN)	(95% CI, 55 to 85) 21 days after 1 st dose and	Moderate	standardized testing for HCW
	(OTTELLY)	85% (95% CI, 74 to 96) 7 days after 2 nd dose		over all of England; 23,000
		against infection in HCW.		participants; time and setting
		against infection in Frew.		for VOC Alpha
12	Shrotri	Similar effect sizes were seen for ChAdOx1	Critical	Prospective cohort in
12	<u>51110t11</u>	(aHR 0.32, 95% CI, 0.15 to 0.66) and	Citicai	England: 9160 of 10412 frail
	*D-1 J	BNT162b2 (aHR 0.35, 95% CI, 0.17 to		LTC residents; routine
	*Delayed	0.71) at 35-48 days after 1 st dose.		screening; time and setting for
	exclusion –	0.71) at 33-46 days after 1 dose.		
1.2	critical ROB	DN/F4 (01 0 1 1 1 1 1 1 7 4 40 / (050 / C)		VOC Alpha
13	<u>Hyams</u>	BNT162b2 showed VE 71.4% (95% CI,		Test negative case-control
		43.1 to 86.2) against hospitalization 14 days		study in Scotland. Single
	WD 1 1	after 1 st dose; ChAdOx1nCoV-19 showed		center; 466 participants, 80+;
	*Delayed	VE 80.4% (95% CI, 36.4 to 94.5) against		time and setting for VOC
	exclusion –	hospitalization 14 days after 1 st dose for		Alpha
	did not	80+.		
	report			
	clinical	When effectiveness analysis for BNT162b2		
	outcomes of	was restricted to the period covered by		
	interest for	ChAdOx1nCoV-19, the estimate was 79.3%		
	this LES	(95% CI, 47.0 to 92.5).		
14	<u>Harris</u>	BNT162b2 or ChAdOx1 reduced likelihood	Serious	Data-linkage and case-control
		of VET by vaccinated HCW to household		study in England; 338,887
		contacts by 40-50% 21 days after 1st dose.		participants; time and setting
				for VOC Alpha
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15	Goldberg	Prior infection (in unvaccinated) has similar VE against infection [94.8%], and severe illness [96.4%] as two doses of BNT162b2.	Serious	Data-linkage study in Israel; 6,351,903 participants; likely overlaps with Dagan and Haas; time and setting for VOC Alpha
16	*Delayed exclusion – VOI instead of VOC	VE 66.2% (95% CI, 40.5% to 80.8%) against infection among LTC residents and 75.9% (95% CI, 32.5% to 91.4%) among HCW. VE 94.4% (95% CI, 73.9% to 98.8%) against hospitalization among residents; no HCW were hospitalized. Three residents died, two of whom were unvaccinated (VE 94.4%; 95% CI, 44.6% to 99.4%).	Critical	Outbreak analysis in LTC in Kentucky; small number of events; VOI R.1
17	Shinde	NVX-CoV2372 VE showed VE 50.4% (95% CI, 16.6 to 70.5) against symptomatic infection 7 days after 2 nd dose.	Moderate quality (RCT)	RCT in South Africa; 4387 participants; 38/41 cases VOC Beta
18	Hitchings	CoronaVac showed VE of 35.1% (95% CI, -6.6 to 60.5) against infection in HCW after 1 st dose.	Serious	Case-control study in HCWs in Manaus; 53,176 participants; 75% prevalence of Gamma; 776 (28%) of 2797 PCR were used for the case-controls; rate of previous infection high in the population
19	<u>Heath</u>	NVX-CoV2373 showed VE 89.7% (95% CI, 80.2 to 94.6) against symptomatic infection after 2 nd dose. No hospitalizations or deaths in vaccinated group.	Moderate quality (RCT)	RCT; 15,187 participants in UK Post hoc: VE 86.3% (95% CI, 71.3 to 93.5) against Alpha variant; 10 cases in vaccinated participants; 66 infections confirmed Alpha; 11 infections no sequencing available
20	*Delayed exclusion – did not report clinical outcomes of interest for this LES	BNT162b2 showed VE 81% (95% CI, 76 to 85) against hospitalization 28 days after 1st dose and 93% (95% CI, 89 to 95) 14 days after the 2nd dose for people 80+. ChAdOx1 showed VE 73% (95% CI, 60 to 81) against hospitalization 28 days after 1st dose; sample size too small to report VE after 2nd dose for people 80+.		Screening study in UK; 13,907 hospitalized patients; results for age 80+; time and setting for VOC Alpha
21	*Delayed exclusion – critical ROB	BNT162b2 showed VE 44% (95% CI, 32 to 53) after 1 st dose and 69% (95% CI, 31 to 86) after 2 nd dose against symptomatic infection in 70+. Single dose ChAdOx1 showed VE 55% (95% CI, 41 to 66) against death.	Critical	Data-linkage study in England; 48,096 cases above age 70+; 12.7% BNT162b2 and 8.2% ChAdOx1; VE also reported for 80+ and LTC; time and setting for VOC Alpha
22	Chodick	BNT162b2 showed VE 90% (95% CI, 79 to 95) against infection and VE 94% (95% CI,	Serious	Data-linkage study in Israel (Maccabi Health Care

	T			
		88 to 97) against death 7-27 days after 2 nd		Organization); 1,178,597
		dose; 71% (95% CI, 37 to 87) in		participants; time and setting
		immunosuppressed.		for VOC Alpha
23	Chung	BNT162b2 or mRNA-1273 showed VE 61% (95% CI, 56 to 66) against symptomatic infection by VOC Alpha 14 days after 1 st dose and 90% (95% CI, 85 to	Moderate	Test-negative study in Ontario 324,033 participants; screening for variants started 2 months into study period; results also
		94) 7 days after 2 nd dose; 43% (95% CI, 22 to 59) against symptomatic infection by VOC Beta or Gamma 14 days after 1 st dose and 88% (95% CI, 61 to 96) 7 days after 2 nd dose.		reported for age>70 and according to vaccine (but not according to confirmed variant)
24	*Delayed exclusion – critical ROB	BNT162b2 showed VE 50% (95% CI, 34 to 73) against infection with VOC Beta >28 days after 2 doses.	Critical	Outbreak in a single LTC in France; 90 participants; all samples genome sequenced for VOC Beta; 2 deaths in vaccinated group
25	Angel	BNT162b2 showed VE 97% (95% CI, 94 to 99) against symptomatic infection and 86% (95% CI, 69 to 93) against asymptomatic infection ≥ 7 days after 2 doses in HCW.	Serious	Retrospective cohort at a single centre tertiary medical centre in Israel, 6,710 participants; testing strategy was different between vaccinated and unvaccinated; time and setting for VOC Alpha
26	*Delayed exclusion – critical ROB	BNT162b2 showed VE 61.9% (95% CI, 19.2 to 82) against infection 14 to 20 days after 1 st dose; 96% (95% CI, 82.2 to 99.1) ≥ 7 days after 2 nd dose in HCW.	Critical	Data-linkage, single centre medical centre in Italy, 2,034 participants; time and setting for VOC Alpha
27	Yassi	BNT162b2 (93%) or mRNA-1273 showed VE 37.2% (95% CI, 16.6 to 52.70) against infection by VOC Beta or Gamma 14 to 42 days after 1 st dose and 79.2% (95% CI, 64.6 to 87.8) 7 days after 2 nd dose in HCW.	Serious	Data-linkage, 25,558 Canadian HCW; evenly split between VOC Gamma and VOC Beta by end of study period
28	Bernal (1)	BNT162b2 showed VE 60% (95% CI, 40 to 73) against confirmed symptomatic infection by VOC Alpha at least 28 days after 1 st dose and 90% (95% CI, 84 to 94) at least 14 days after 2 nd dose for people 70+.	Serious	Test-negative in England, 156,930 participants; spike gene target failure as proxy for confirmed VOC Alpha
29	Bernal (3)	BNT162b2 showed VE 47.5% (95% CI, 41.6 to 52.8) at least 21 days after 1st dose and VE 93.7% (95% CI, 91.6 to 95.3) at least 14 days after 2nd dose against symptomatic infection by confirmed VOC Alpha. ChadOx1showed VE 48.7% (95% CI, 45.2 to 51.9) at least 21 days after 1st dose and VE 74.5% (95% CI, 68.4 to 79.4) at least 14 days after 2nd dose against symptomatic infection by confirmed VOC Alpha.	Serious	Test-negative in England; 19,109 sequenced cases: 14,837 VOC Alpha and 4,272 VOC Delta.

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		BNT162b2 showed VE 35.6% (95% CI, 22.7 to 46.4) at least 21 days after 1 st dose and VE 88% (95% CI, 85.3 to 90.1) at least 14 days after 2 nd dose against symptomatic infection by confirmed VOC Delta. ChAdOx1 showed VE 30% (95% CI, 24.3 to 35.3) at least 21 days after 1 st dose and VE 67% (95% CI, 61.3 to 71.8) at least 14 days after 2 nd dose against symptomatic infection by confirmed VOC Delta.		
30	Ranzani	CoronaVac reduced risk of symptomatic infection by VOC Gamma VE 41.6% (95% CI, 26.9 to 63.3) ≥ 14 days after 2 nd dose for people 70+.	Serious	Test-negative in Brazil; 44,055 participants; sequencing not performed; effectiveness declined with age; time and setting for VOC Gamma
31	Andrejko (2)	BNT162b2 and mRNA-1273 showed VE 86.8% (95% CI, 68.6 to 94.7) and VE 86.10% (95% CI, 69.1 to 93.9), respectively, against infection 15 days after 2 nd dose.	Serious	Test-negative in California; 1,023 participants; expansion of sample size and timeline since previous study by same authors; VOC Alpha, Epsilon
32	Emborg	BNT162b2 showed VE 53-86% against infection across high-risk groups, VE 75-87% against hospitalization across high-risk groups, VE 89% (95% CI, 81 to 93) against death in LTCF residents and VE 97% (95% CI, 88 to 99) against death in 65+ requiring personal care 7 days after 2 nd dose.	Serious	Data-linkage population study of high-risk groups in Denmark; 864,096 participants; sample confirmed VOC Alpha
33	Salo	BNT162b2 showed VE 42.9% (95% CI, 22.3 to 58.1) against infection in unvaccinated household members of vaccinated HCW 10 weeks after 1 st dose.	Moderate	Data-linkage for household contacts of HCW in Finland; 52,766 spouses of vaccinated HCW; time and setting for VOC Alpha
34	Shrestha	BNT162b2 or mRNA-1273 showed VE 97.1% (95% CI, 94.3 to 98.5) against infection ≥14 days after 2 nd dose (based on multivariable model).	Moderate	Retrospective cohort of employees of a health care system in Ohio; 46,866 participants (60%) vaccinated by end of study; time and setting for VOC Alpha
35	Skowronski	BNT162b2 (85%) or mRNA-1273 showed VE 67% (95% CI, 57 to 75) against infection by confirmed VOC Alpha ≥21 days after 1 st dose for 70+. BNT162b2 (85%) or mRNA-1273 showed VE 61% (95% CI, 45 to 72) against infection by confirmed VOC Gamma ≥21 days after 1 st dose for 70+.	Serious	Test-negative in Canada; 16,993 specimens; out of 1,131 genetically sequenced: 45% VOC Alpha and 28% Gamma; results reported by vaccine but not according to confirmed variant
36	Abu-Raddad	BNT162b2 showed VE 89.5% (95% CI, 85.9 to 92.3) against infection, VE 100% (95% CI, 81.7 to 100) against any severe,	Serious	Test-negative in Qatar; 17,293 cases; sequencing showed 50% VOC Beta and 45% VOC

37	Akhrass *Delayed exclusion - failure to	critical, or fatal disease by VOC Alpha ≥ 14 days after 2 nd dose. BNT162b2 showed VE 75% (95% CI, 70.5 to 78.9) against infection, VE 100% (95% CI, 73.7 to 100) against severe, critical, or fatal disease by VOC Beta ≥ 14 days after 1 st dose. BNT162b2 or mRNA-1273 showed overall VE 60.4% (95% CI, 30 to 77.6) against symptomatic infection ≥ 14 days after 1 st dose; BNT162b2 or mRNA-1273 showed	Critical	Alpha between February-March 2021 Retrospective cohort of HCW at a single centre in Kentucky, USA; 2,134 participants; time and setting for VOC Alpha
	report outcomes of interest for this LES	overall VE 95.7% (95% CI, 90 to 98.2) against symptomatic infection \geq 14 days after 2 nd dose.		r a sava g a a a a a a
38	Sheikh	BNT162b2 showed VE 30% (95% CI, 17 to 41) against confirmed VOC Delta infection and VE 33% (95% CI, 15 to 47) against symptomatic infection at least 28 days after 1st dose; VE 79% (95% CI, 75 to 82) against infection and VE 83% (95% CI, 78 to 87) against symptomatic infection at least 14 days after 2nd dose. ChAdOx1 showed VE 18% (95% CI, 9 to 25) against confirmed VOC Delta infection and VE 33% (95% CI, 23 to 41) against symptomatic infection at least 28 days after 1st dose; VE 60% (95% CI, 53 to 66) against infection and VE 61% (95% CI, 51 to 70%) against symptomatic infection at least 14 days after 2nd dose.	Serious	Test-negative in Scotland; 626,900 specimens; also compared hospitalization rates between S gene positive (VOC Delta) and S gene negative specimens within 14 days of positive test result (not summarized here)
39	Furer *Delayed exclusion – critical risk of bias	BNT162b2 reported no symptomatic infections in the vaccinated group (0/686) compared to 0.83% infections in the vaccinated general population control group.	Critical	Prospective cohort of adults with autoimmune inflammatory rheumatic diseases in Israel; 686 participants; time and setting for VOC Alpha
40	Martinez- Baz	BNT162b2 showed VE 65% (95% CI, 56 to 73) against infection and VE 94% (95% CI, 60 to 99) against hospitalization at least 14 days after 2 nd dose in close contacts of COVID+ index cases. ChAdOx1 showed VE 44% (95% CI, 31 to 54) against infection and VE 92% (95% CI, 46 to 99) against hospitalization at least 14 days after 1 st dose in close contacts of index cases. Second dose results not reported.	Serious	Prospective cohort of close contacts of COVID+ people in Spain; 20,961 participants; VOC Alpha confirmed for small sample; sample size for Moderna too small to report results separately

41	Chodick (2)	BNT162b2 showed VE 51.4% (95% CI, 16.3 to 71.8) against infection 13 to 24 days after 1 st dose.	Serious	Data-linkage study in Israel (Maccabi Health Care Services); 351,897 participants; time and setting for VOC Alpha
42	Stowe	BNT162b2 showed VE 94% (95% CI, 46 to 99) at least 21 days after 1 st dose and VE 96% (95% CI, 86 to 99) at least 14 days after 2 nd dose against hospitalization by confirmed VOC Delta. ChAdOx1 showed VE 71% (95% CI, 51 to 83) at least 21 days after 1 st dose and VE 92% (95% CI, 75 to 97) 14 days after 2 nd dose against hospitalization by confirmed VOC Delta.	Serious	Same cohort as Bernal (3) with extended time frame for symptomatic infection and adding in data-linkage to hospitalization; 14,019 participants; sample confirmed VOC Delta
43	<u>Saciuk</u>	BNT162b2 showed VE 93% (95% CI, 92.6 to 93.4) against infection, VE 93.4% (95% CI, 91.9 to 94.7) against hospitalization and VE 91.1% (95% CI, 86.5 to 94.1) against death at least 7 days after 2 nd dose	Serious	Retrospective cohort of members of a health management organization in Israel; 1,650,885 participants; time and setting for VOC Alpha
44	*Delayed exclusion – critical risk of bias	BNT162b2 showed VE 61% (95% CI, 49 to 71) at least 14 days after 1 st dose and VE 89% (95% CI, 82 to 94) at least 7 days after 2 nd dose against infection	Serious	Retrospective cohort of a subpopulation of members of a health management organization in Israel who had undergone repeated PCR testing; 6,286 participants; time and setting for VOC Alpha
45	Azamgarhi	BNT162b2 showed VE 70% (95% CI, 6 to 91) against infection at least 14 days after 1 st dose	Serious	Single centre cohort study of HCW in UK; 2,260 participants; time and setting for VOC Alpha
46	Lumley	BNT162b2 (63%) or ChAdOx1showed VE 64% (95% CI, 50 to 74) 14 days after 1 st dose and VE 90% (95% CI, 62 to 98) 14 days after 2 nd dose against infection	Serious	Prospective cohort of HCWs in Oxfordshire, UK; 13,109 participants; confirmed VOC Alpha
47	Nasreen	BNT162b2 showed VE 89% (95% CI, 86 to 91) against symptomatic infection and VE 95% (95% CI, 92 to 97) against hospitalization at least 7 days after 2 nd dose (VOC Alpha); VE 84% (95% CI, 69 to 92) against symptomatic infection and VE 95% (95% CI, 81 to 99) against hospitalization at least 7 days after 2 nd dose (VOC Beta/Gamma); VE 87% (95% CI, 64 to 95) against symptomatic infection at least 7 days after 2 nd dose (VOC Delta).	Moderate	Test-negative study in Ontario 421,073 participants (same population as for Chung but extended to May 2021 and more detailed with respect to reporting of VOC); screening for VOC Alpha, Beta/Gamma and Delta varied during study period

		BNT162b2 showed VE 78% (95% CI, 65 to 86) against hospitalization at least 7 days after 2 nd dose (VOC Delta). mRNA-1273 showed VE 92% (95% CI, 86 to 96) against symptomatic infection and VE 94% (95% CI, 89 to 97) against		
		hospitalization at least 7 days after 2 nd dose (VOC Alpha).		
		mRNA-1273 showed VE 77% (95% CI, 63 to 86) against symptomatic infection and VE 89% (95% CI, 73 to 95) against hospitalization at least 14 days after 1st dose (VOC Beta/Gamma); VE 72% (95% CI, 57 to 82) against symptomatic infection and VE 96% (95% CI, 72 to 99) against hospitalization at least 14 days after 1st dose (VOC Delta).		
		ChAdOx1 showed VE 64% (95% CI, 60 to 68) against symptomatic infection and VE 85% (95% CI, 81 to 88) against hospitalization at least 14 days after 1 st dose (VOC Alpha); VE 48% (95% CI, 28 to 63)		
		against symptomatic infection and VE 83% (95% CI, 66 to 92) against hospitalization at least 14 days after 1 st dose (VOC Beta/Gamma); VE 67% (95% CI, 44 to 80) against symptomatic infection and VE 88% (95% CI, 60 to 96) against hospitalization at least 14 days after 1 st dose (VOC Delta).		
48	Gazit	BNT162b2 showed VE 80% (95% CI, 73 to 85) at least 7 days after 2 nd dose against infection in vaccinated household members of a confirmed COVID+ case.	Serious	Retrospective cohort of household members (household = 2 adults with no children) of a health management organization in Israel; 173,569 households; time and setting for VOC Alpha
49	Jara	CoronaVac showed VE 65.9% (95% CI, 65.2 to 66.6) against infection and VE 86.3% (95% CI, 84.5 to 87.9) against death at least 14 days after 2 nd dose.	Moderate	Prospective cohort in Chile; 10.2 million participants; time and setting for VOC Gamma
50	Chemaitelly	mRNA-1273 showed VE 88.1% (95% CI, 83.7 to 91.5) and VE 100% (95% CI, 91.8 to 100) against infection by confirmed VOC Alpha at least 14 days after 1 st and 2 nd dose, respectively.	Serious	Test-negative in Qatar; >75,000 participants; sample sequenced for VOC Alpha and VOC Beta
		mRNA-1273 showed VE 61.3% (95% CI, 56.5 to 65.5) and VE 96.4% (95% CI, 91.9		

		to 98.7) against infection by confirmed VOC Beta at least 14 days after 1 st and 2 nd dose, respectively.		
		mRNA-1273 showed VE 81.6% (95% CI, 71.0 to 88.8) and VE 95.7% (95% CI, 73.4 to 99.9) against severe, critical, or fatal disease at least 14 days after 1 st and 2 nd dose, respectively (combined VOC Alpha and Beta).		
51	Baum	BNT162b2 or mRNA-1273 showed VE 41% (95% CI, 25 to 54) against infection ≥ 21 days after 1 st dose; BNT162b2 or mRNA-1273 showed VE 75% (95% CI, 65 to 82) against infection ≥ 7 days after 2 nd dose in age 70+. BNT162b2 or mRNA-1273 showed VE 41% (95% CI, 17 to 58) against infection ≥ 21 days after 1 st dose; BNT162b2 or mRNA-1273 showed VE 77% (95% CI, 65 to 85) against infection ≥ 7 days after 2 nd dose in chronically ill (age 16-69). ChAdOx1 showed VE 24% (95% CI, -1 to 43) against infection ≥ 21 days after 1 st dose in chronically ill (age 16-69).	Serious	Data-linkage study in Finland; 901,092 participants age 70+ and 774,526 participants age 16 to 69 years with chronic illness; time and setting for VOC Alpha; results for mRNA vaccines not reported separately
52	Balicer	BNT162b2 showed VE 86.1% (95% CI, 82.4 to 89.1) against infection; VE 89% (95% CI, 43 to 100) against hospitalization 7 to 56 days after 2 nd dose. Too few events to report VE for severe disease or death.	Serious	Data-linkage study of pregnant women over age 16 in Israel (same database as Dagan); 21,722 participants; time and setting for VOC Alpha.
53	Mateo- Urdiales	BNT162b2 (61%) or ChAdOx1 (31%) or mRNA-1273 (7%) or Ad26.COV ₂ -S (0.6%) showed VE 78% (95% CI, 76 to 79) against infection 42 to 49 days after at least 1 st dose; VE 93% (95% CI, 89 to 96) against death 35 to 42 days after at least 1 st dose.	Serious	Data-linkage study in Italy; 13,721,506 participants; time and setting for VOC Alpha. Results not reported by vaccine and some participants (42%) who also received 2 nd dose were included in estimates.
54	Goldshtein	BNT162b2 showed VE 78% (95% CI, 57 to 89) against infection at least 28 days after 1 st dose.	Serious	Data-linkage study of pregnant women in Israel (same database as Gazit); 15,060 participants; time and setting for VOC Alpha.
55	<u>Mason</u>	BNT162b2 showed VE 55.2% (95% CI, 40.8 to 66.8) and VE 70.1% (95% CI, 55.1 to 80.1) against infection 21 to 27 days and 35 to 41 days after 1 st dose, respectively.	Moderate	Case-control study of age 80-83 vs 76-79 community-dwelling unvaccinated residents in England; time and setting for VOC Alpha

56	Fabiani	BNT162b2 showed VE 84.1% (95% CI,	Serious	Retrospective cohort of HCW
		39.7 to 95.8) and VE 85.4% (95% CI, -35.3		in Italy; 6,423 participants;
		to 98.4) against infection 14 to 21 days and \geq 21 days after 1 st dose, respectively in		time and setting for VOC Alpha
		HCW.		Στιριια
		DN/H4 (01.0.1		
		BNT162b2 showed VE 95.1% (95% CI, 62.4 to 99.4) against infection ≥7 days after		
		2 nd dose in HCW.		
57	Chia	BNT162b2 or mRNA-1273 showed VE	Serious	Retrospective cohort of
		92.7% (95% CI, 65.7 to 98.4) against severe		confirmed VOC Delta
		disease (defined as requiring supplemental		admitted to hospital (including
		oxygen) > 14 days after 2^{nd} dose.		asymptomatic) in Singapore;
				218 participants; not reported by vaccine
58	<u>Kaur</u>	Two doses of Covishield showed VE 87%	Critical	Preliminary report of
	*Delayed	(95% CI, 33 to 97) against severe disease		prospective cohort in India;
	exclusion –	when compared with one dose (timing of		1500 participants; time and
59	critical ROB Pramod	doses not reported). Covishield showed VE 49% (95% CI, 17 to	Critical	setting for VOC Delta Test-negative study in a single
39	<u>1 1411100</u>	68) against infection 21 days after 1st dose	Citucai	hospital site in India; 360
	*Delayed	and VE 54% (95% CI, 27 to 71) against		matched pairs (203
	exclusion –	infection 14 days after 2 nd dose.		symptomatic pairs); time and
	critical ROB			setting for VOC Delta
		Covishield showed VE 58% (95% CI, 28 to		
		75) against symptomatic infection 21 days after 1 st dose and VE 64% (95% CI, 38 to		
		78) against symptomatic infection 14 days		
		after 2 nd dose.		
60	Carazo	BNT162b2 or mRNA-1273 showed VE	Serious	Test-negative study in
		60% (95% CI, 53.6 to 65.5) against infection		Quebec, Canada; 58,476
		by confirmed VOC Alpha 14 days after 1 st dose.		participants; sample confirmed VOC Alpha; reported
		dosc.		according to vaccine but not
		BNT162b2 or mRNA-1273 showed VE		concurrently for VOC Alpha
		92.6% (95% CI, 87.1 to 95.8) against		
		infection by confirmed VOC Alpha 7 days		
61	Williams	after 2 nd dose. BNT162b2 or mRNA-1273 showed VE	Serious	Outbreak in a single LTCF in
01	wimailis	52.5% (95% CI, 26.9 to 69.1) against	Senous	Ontario; 60 residents and 83
		infection and VE 78.6% (95% CI, 47.9 to		staff; sample confirmed VOC
		91.2) against severe disease 14 days after 2 nd		Gamma
		dose in residents at LTCF. Two deaths in		
		vaccinated residents but were palliative prior		
		to infection.		
		BNT162b2 or mRNA-1273 showed VE		
		66.2% (95% CI, 2.3 to 88.3) against		
		infection 14 days after 2 nd dose in staff at		
		LTCF. None of the staff developed severe		
		disease.		

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62	*Delayed exclusion – critical ROB	ChAdOx1 showed VE 33.4% (95% CI, 26.4 to 39.7) against symptomatic infection and VE 50.9% (95% CI, 33.6 to 63.8) against ICU admission and VE 61.8% (95% CI, 48.9 to 71.4) against death at least 28 days after 1st dose for 60+. ChAdOx1 showed VE 77.9% (95% CI, 69.2 to 84.2) against symptomatic infection and VE 89.9% (95% CI, 70.9 to 96.5) against ICU admission and VE 93.6% (95% CI, 81.9 to 97.7) against death at least 14 days after 2nd dose	Critical	Test-negative study in Sao Paulo, Brazil; 61,164 participants over age 60; time and setting for VOC Gamma
63	Tang	after 2 nd dose. BNT162b2 showed VE 65.5% (95% CI, 40.9 to 79.9) against infection ≥ 14 days after 1 st dose; BNT162b2 showed VE 59.6% (95% CI, 50.7 to 66.9) against infection ≥ 14 days after 2 nd dose. BNT162b2 showed VE 100% (95% CI, not reported) against severe, critical or fatal disease ≥ 14 days after 1 st dose; BNT162b2 showed VE 97.3% (95% CI, 84.4 to 99.5) against severe, critical or fatal disease ≥ 14 days after 2 nd dose. mRNA-1273 showed VE 79.7% (95% CI, 60.8 to 89.5) against infection ≥ 14 days after 1 st dose; mRNA-1273 showed VE 86.1% (95% CI, 78.0 to 91.3) against infection ≥ 14 days after 2 nd dose. mRNA-1273 showed VE 100% (95% CI, not reported) against severe, critical or fatal disease ≥ 14 days after 1 st dose; mRNA-1273 showed VE 100% (95% CI, not reported) against severe, critical or fatal	Serious	Test-negative study in Qatar; 1,140,337 participants; weekly random sequencing of positive samples for VOC Delta
64	<u>Puranik</u>	disease ≥ 14 days after 2 nd dose. BNT162b2 showed VE 42% (95% CI, 13 to 62) against infection 14 days after 2 nd dose. mRNA-1273 showed VE 76% (95% CI, 58 to 87) against infection 14 days after 2 nd dose.	Serious	Data-linkage study involving Mayo Clinic Health in USA; 25,859 matched triples from Minnesota only; time and setting for Delta at end of study time frame so only last month of data (July 2021) reported here
65	*Delayed exclusion – critical ROB	BNT162b2 or ChAdOx1 showed VE 64% (95% CI, 11 to 85) against infection unreported number of days after 2 nd dose (Round 12: 2021-05-20 to 2021-06-07).	Critical	Surveillance study in England; 121,872 participants; time and setting for VOC Delta; only included data from aged 18 to 64 years due to lowest risk for misclassification bias due to

		BNT162b2 or ChAdOx1 showed VE 49% (95% CI, 22 to 67) against infection unreported number of days after 2 nd dose (Round 13: 2021-06-24 to 2021-07-12).		self-reported vaccination status
66	<u>Issac</u>	ChAdOx1 showed VE 85% (95% CI, 71 to 92) against infection 14 days after 2 nd dose.	Serious	Prospective cohort of HCW at a single hospital in India; 342 participants; time and setting for VOC Delta.
67	Marco *Delayed exclusion – critical ROB	ChAdOx1 showed VE 23% (95% CI, not reported) against infection at least 21 days after 1 st dose.	Critical	Outbreak study of prison inmates in Barcelona; 217 participants (184 inmates); sequenced for VOC Alpha
68	Kale *Delayed exclusion – critical ROB	ChAdOx1 showed VE 60% (95% CI, 45 to 70) against infection at least 14 days after 2 nd dose.	Critical	Prospective cohort of HCW at a single hospital in India; 1858 participants; sample sequenced for VOC Delta
69	<u>Israel</u>	BNT162b2 showed OR 2.06 (95% CI, 1.69 to 2.51) for infection comparing fully vaccinated ≥146 days vs fully vaccinated less than 146 days.	Moderate	Retrospective cohort of fully vaccinated members of a health management organization in Israel who underwent testing; 33,993 participants; time and setting for VOC Delta
70	Gram	ChAdOx1 showed VE 44% (95% CI, 29 to 56) against infection 21 to 27 days after 1 st dose. No deaths in vaccinated participants. First dose ChAdOx1 followed by second dose BNT162b2 or mRNA-1273 showed VE 88% (95% CI, 83 to 92) against infection ≥ 14 days after 2 nd dose.	Serious	Data-linkage study in Denmark; 5,542,079 participants; sequenced for VOC Alpha (includes heterologous vaccines)
71	Pouwels	BNT162b2 showed VE 59% (95% CI, 52 to 65%) against infection ≥21 days after 1 st dose and VE 78% (95% CI, 68 to 84) against infection ≥ 14 days after 2 nd dose (VOC Alpha age 18+). BNT162b2 showed VE 57% (95% CI, 50 to 63) against infection ≥21 days after 1 st dose and VE 80% (95% CI, 77 to 83) against infection ≥ 14 days after 2 nd dose (VOC Delta age 18+). ChAdOx1 showed VE 63% (95% CI, 55 to 69) against infection ≥21 days after 1 st dose and VE 79% (95% CI, 56 to 90) against infection ≥ 14 days after 2 nd dose (VOC Alpha age 18+). ChAdOx1 showed VE 46% (95% CI, 35 to 55) against infection ≥21 days after 1 st dose and VE 67% (95% CI, 62 to 71) against	Serious	Survey of randomly selected private households with longitudinal follow-up in UK; 743,526 participants; also reported for 18-64 years; sample sequenced for VOC Alpha and VOC Delta

		infection \geq 14 days after 2 nd dose (VOC		
		Delta age 18+).		
		DNIA 4050 1 1115 550/ (050/ CL / 4		
		mRNA-1273 showed VE 75% (95% CI: 64		
		to 83) against infection ≥21 days after 1 st		
70	A1 D 11 1	dose (VOC Delta age 18 to 64).	С.	D 1 1
72	Abu-Raddad	BNT162b2 <u>after prior infection</u> showed VE	Serious	Retrospective matched
	<u>(2)</u>	85% (95% CI, 80 to 89) against re-infection		cohorts (2) of fully vaccinated
		compared to BNT162b2 without prior infection.		in Qatar; 151,076 participants; sample sequenced for VOC
		intection.		Alpha and VOC Beta
		mRNA-1273 after prior infection showed		Tupna and VOC Beta
		VE 15% (95% CI, -105 to 66) against re-		
		infection compared to mRNA-1273 without		
		prior infection.		
73	Gazit (2)	BNT162b2 showed OR 13.06 (95%	Moderate	Retrospective matched
'3	<u> </u>	CI, 8.08 to 21.11) against infection and OR	Moderate	cohorts of fully vaccinated in
		27.02 (95% CI, 12.7 to 57.5) against		Israel; 778,658 participants;
		symptomatic disease compared to prior		time and setting for VOC
		infection.		Delta
74	Rosenberg	BNT162b2 (51%), mRNA-1273 (40%) or	Serious	Surveillance report in New
' '		Ad26.COV2.S (9%) showed VE 91.7%	0 0 2 2 3 6 7 6	York, USA; >13 million
		against infection ≥14 days after 2 nd dose		participants; time and setting
		(Week of May 3, 2021: VOC Alpha).		for VOC Delta (from 2% to
				80% during study period)
		BNT162b2 (51%), mRNA-1273 (40%) or		
		Ad26.COV2.S (9%) showed VE 79.8%		
		against infection ≥14 days after 2 nd dose		
		(Week of July 19, 2021: VOC Delta).		
75	<u>Al-Qahtani</u>	BNT162b2 \geq 14 days after 2 nd dose, showed	Critical	Retrospective cohort of fully
		VE 99.9% (95% CI, 99.2 to 100) against		vaccinated (>14 days after 2 nd
		ICU admission, and VE 99.5% (95% CI,		dose) in Bahrain; 1,242,279
		98.4 to 99.8) against death (VOC Alpha and		participants; time and setting
	*Delayed	Delta).		for VOC Alpha (dominant
	exclusion			before May 2021) and Delta
	due to	$ChAdOx1 \ge 14$ days after 2^{nd} dose, showed		(dominant after May 2021).
	critical ROB	VE 99.2% (95% CI, 97.6 to 99.7) against		
		ICU admission, and VE 99.6% (95% CI,		
		97.2 to 100) against death (VOC Alpha and		
		Delta).		
		DDIDD C V >14.1 C and 1		
		BBIBP-CorV ≥14 days after 2 nd dose,		
		showed VE 95.4% (95% CI, 94.6 to 96.2)		
		against ICU admission, and VE 94.3% (95%)		
		CI, 93.1 to 95.4) against death (VOC Alpha		
		and Delta).		
		Sputnik V \geq 14 days after 2 nd dose, showed		
		VE 100% (95% CI, 99.2 to 100) against		
		ICU admission, and VE 99.5% (95% CI,		
		98.5 to 99.9) against death (VOC Alpha and		
		Delta).		
		Day.		

76	Goldberg (2)	BNT162b2 showed VE 50% (95% CI, 45 to 55) for those vaccinated in January 2021, and VE 73% (95% CI, 67 to 78) for those vaccinated in May 2021 against infection after the 2 nd dose (VOC Delta age 16 to 39). BNT162b2 showed VE 58% (95% CI, 54 to 62) for those vaccinated in January 2021, and VE 80% (95% CI, 71 to 86) for those vaccinated in May 2021 against infection after the 2 nd dose (VOC Delta age 40 to 59). BNT162b2 showed VE 57% (95% CI, 52 to 62) for those vaccinated in January 2021,	Serious	Data-linkage study of fully vaccinated in Israel; 4,785,245 participants; sequenced for VOC Delta (dominant after May 2021) (results over varying time periods since vaccination reported)
		and VE 75% (95% CI, 58 to 85) for those vaccinated in May 2021 against infection after the 2 nd dose (VOC Delta age 60+). BNT162b2 showed VE 94% (95% CI, 87 to 97) for those vaccinated in January 2021, and VE 98% (95% CI, 94 to 99) for those vaccinated in March 2021 against severe, critical, or fatal disease after the 2 nd dose (VOC Delta age 40 to 59). BNT162b2 showed VE 86% (95% CI, 82 to		
		90) for those vaccinated in January 2021, and VE 91% (95% CI, 85 to 95) for those vaccinated in March 2021 against severe, critical, or fatal disease after the 2 nd dose (VOC Delta age 60+).		
77	*Delayed exclusion – critical risk of bias	BNT162b2, mRNA-1273, or Ad26.COV2.S showed VE 78% (95% CI, 71 to 84) in Mesa County and VE 89% (95% CI, 88 to 91) in other Colorado counties against symptomatic infection an unreported number of days after 2 nd dose (VOC Delta).	Critical	Surveillance report in Mesa County-Colorado, USA; 37,439 cases participants; sample sequenced for VOC Delta (43% to 88% during study period)
78	*Delayed exclusion – critical risk of bias	ChAdOx1 showed unadjusted VE 75.2% (95% CI, 73.8 to 76.8) against infection ≥14 days after 1st dose, and unadjusted VE 54.6% (95% CI, 52.6 to 56.6) ≥14 days after 2nd dose against infection in HCW (VOC Alpha to Delta).	Critical	Retrospective cohort of Armed Forces HCW and frontline workers in India; 1,595,630 participants; time and setting for VOC Delta at end of study only.
79	Amirthaling am	BNT162b2 showed VE 77% (95% CI, 56 to 88) against symptomatic infection when 2 nd dose given 19-29 days after 1 st dose, and VE 94% (95% CI, 73 to 99) against symptomatic infection when 2 nd dose given 85+ days after 1 st dose (VOC Alpha age 80+).	Moderate	Test-negative study in England; 750 participants; time and setting for VOC Alpha (dominant before May 2021) and Delta (dominant after May 2021).

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		BNT162b2 showed VE 77% (95% CI, 66 to		(results over varying time
		85) against symptomatic infection when 2 nd		periods since vaccination
		dose given 19-29 days after 1st dose, and VE		reported)
		86% (95% CI, 70 to 94) against		
		symptomatic infection when 2 nd dose given		
		85+ days after 1 st dose (VOC Alpha age 65		
		to 79).		
		ChAdOx1 showed VE 96% (95% CI, 72 to		
		100) against symptomatic infection when 2 nd		
		dose given 19-29 days after 1 st dose, and VE		
		88% (95% CI, 48 to 97) against		
		symptomatic infection when 2 nd dose given 85+ days after 1 st dose after 2 nd dose (VOC		
		Alpha age 80+).		
		Tupita age 60 1).		
		ChAdOx1 showed VE 66% (95% CI, 47 to		
		77) against symptomatic infection when 2 nd		
		dose given 19-29 days after 1 st dose, and VE		
		73% (95% CI, 56 to 83) against		
		symptomatic infection when 2 nd dose given		
		85+ days after 1 st dose after 2 nd dose (VOC		
		Alpha age 65 to 79).		
80	Butt (2)	Unvaccinated participants had HR 2.84	Critical	Case-control study in Qatar;
		(95% CI, 1.80 to 4.47) of severe disease		456 matched cases; time and
	*Delayed	compared to BNT162b2 ≥14 days after 2 nd		setting for VOC Alpha
	exclusion –	dose.		
	critical ROB			
81	<u>Fowlkes</u>	BNT162b2 (65%), mRNA-1273 (33%), or	Moderate	Prospective cohort of HCW
		Ad26.COV2.S (2%) showed VE 91% (95%		and other essential frontline
		CI, 81 to 96) against infection \geq 14 days		workers in 6 states in the
		after 2 nd dose (during time of VOC Alpha).		USA; 7,112 participants;
		DNT162b2 (650/) mDNIA 1272 (220/) or		updated report to cover VOC
		BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 66% (95%		Delta period
		$ C = 26$ to 84) against infection ≥ 14 days		
1		CI, 26 to 84) against infection \geq 14 days after 2 nd dose (during time of VOC Delta)		
		CI, 26 to 84) against infection \geq 14 days after 2 nd dose (during time of VOC Delta).		
		after 2 nd dose (during time of VOC Delta).		
		after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or		
		after 2 nd dose (during time of VOC Delta).		
		after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95%		
		after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days		
		after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI,		
		after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta).		
82	Bhattachary	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE	Critical	Cross-sectional cohort of
82	Bhattachary a	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against	Critical	HCW and their families at a
82	<u>a</u>	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against symptomatic infection ≥ 14 days after 2 nd	Critical	HCW and their families at a single site in India; 638
82	<u>a</u> *Delayed	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against	Critical	HCW and their families at a single site in India; 638 participants (55 inpatients);
82	*Delayed exclusion	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against symptomatic infection ≥ 14 days after 2 nd	Critical	HCW and their families at a single site in India; 638
82	<u>a</u> *Delayed	after 2 nd dose (during time of VOC Delta). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 85% (95% CI, 68 to 93) against infection 14-119 days after full vaccination) and VE 73% (95% CI, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta). Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against symptomatic infection ≥ 14 days after 2 nd	Critical	HCW and their families at a single site in India; 638 participants (55 inpatients);

	T			T
		Covaxin (94%) and Covishield showed VE		
		93% (95% CI, 64 to 99) against ICU		
		admission or death \geq 14 days after 2 nd dose.	25.1	
83	Nunes	BNT162b2 (45%) or mRNA-1273 (8%) showed VE 96% (95% CI, 92 to 98) against COVID-related death ≥14 days after 2 nd dose (age 65 to 79).	Moderate	Data-linkage study of community-dwelling adults≥65 in Portugal; 2,050,950 participants; time and setting for VOC Alpha to
		BNT162b2 (80%) or mRNA-1273 (2%) showed VE 81% (95% CI, 74 to 87) against COVID-related death ≥14 days after 2 nd dose (age ≥80).		VOC Delta
		BNT162b2 (80%) or mRNA-1273 (2%) showed VE 86% (95% CI, 68 to 93) against COVID-related death 14 to 41 days after 2 nd		
		dose and VE 74% (95% CI, 60 to 83)		
		against COVID-related death \geq 98 days after 2 nd dose for HR 1.80 (0.77 to 4.25) (age \geq 80).		
84	Tartof	BNT162b2 showed VE 75% (95% CI, 71 to 78) against infection 7 days after 2 nd dose (confirmed VOC Delta).	Moderate	Retrospective cohort of members of a health management organization in California; 3,436,957
		BNT162b2 showed VE 91% (95% CI, 88 to 92) against infection 7 days after 2 nd dose (confirmed non-VOC Delta).		participants; VOC Alpha to VOC Delta (only 28% confirmed Delta)
		BNT162b2 showed VE 93% (95% CI, 85 to 87) against infection 7 to 30 days after 2 nd dose and VE 53% (95% CI, 39 to 65) against infection ≥ 127+ days after 2 nd dose (confirmed VOC Delta).		(results over varying time periods since vaccination reported)
		BNT162b2 showed VE 97% (95% CI, 95 to 99) against infection 7 to 30 days after 2 nd dose and VE 67% (95% CI, 45 to 80) against infection ≥ 127+ days after 2 nd dose (confirmed non-VOC Delta).		
85	*Delayed exclusion – critical ROB	CoronaVac (combined with other inactivated vaccines) showed VE 59% (95% CI, 16 to 81.6) against symptomatic infection and VE 100% against severe infection ≥14 days after 2 nd dose.	Critical	Test-negative study in Guangzhou, China; 366 participants; sample sequenced for VOC Delta
86	Scobie *Delayed exclusion – critical ROB	BNT162b2 or mRNA-1273 (92%), or Ad26.COV2.S showed VE 90% (95% CI not reported) against infection and VE 93% (95% CI not reported) against death ≥ 14 days after 2 nd dose (April to June: VOC Alpha).	Critical	Surveillance study in 13 states in the USA; 615,454; time and setting for VOC Alpha to VOC Delta

	1			
		BNT162b2, mRNA-1273, or Ad26.COV2.S		
		showed VE 76% (95% CI not reported)		
		against infection and VE 90% (95% CI not		
		reported) against death ≥ 14 days after 2^{nd}		
		dose (June to July: VOC Delta>50%).		
87	Satwik	ChAdOx1 showed VE 18% (95% CI, -10 to	Critical	Retrospective cohort study of
		38) against symptomatic infection; VE 37%		HCW at a single hospital in
		(-24 to 68) against moderate to severe		New Delhi, India; 4276
		disease and VE 69% (95% CI, -160 to 97)		participants; sample sequenced
		against death ≥21 days after 1 st dose.		for VOC Delta
	*Delayed	,		
	exclusion	ChAdOx1 showed VE 28% (95% CI, 10 to		
	due to	41) against symptomatic infection; VE 67%		
	critical ROB	(44 to 81) against moderate to severe disease		
		and VE 97% (95% CI, 43 to 99.8) against		
		death ≥ 14 days after 2 nd dose.		
88	Seppala	BNT162b2 (74%) or ChAdOx1 (22%) or	Serious	Population cohort in Norway;
	Seppara	mRNA-1273 (10%) showed VE 84.4%	5511543	4,204,859 participants;
		$(95\% \text{ CI, } 81.8 \text{ to } 86.5) \text{ against infection } \ge 7$		sequenced for VOC Alpha
		days after 2 nd dose (VOC Alpha).		and VOC Delta
		days after 2 dose (VOC mpna).		and voc Beta
		BNT162b2 (74%) or ChAdOx1 (22%) or		
		mRNA-1273 (10%) showed VE 64.6%		
		$(95\% \text{ CI}, 60.6 \text{ to } 68.2)$ against infection ≥ 7		
		days after 2^{nd} dose (VOC Delta).		
89	Polinski	Ad26.COV2.S showed VE* 67% (95% 60	Serious	Data-linkage of members of a
07	<u>1 OIII13KI</u>	to 73) against infection unknown number of	Scrious	medical insurance group in
		days after dose (June to July: VOC Delta in		USA; 1,914,670 participants;
		high prevalence states). *unadjusted for substantial		time and setting for VOC
		under-reporting of vaccination status		Alpha to Delta (only data for
				VOC Delta reported here)
90	Chemaitelly	BNT162b2 or mRNA-1273 showed VE	Serious	Retrospective cohort of
70		46.6% (95% CI, 0.0 to 73.7) against	Schous	immunosuppressed kidney
	<u>(2)</u>	infection \geq 14 days after 2 nd dose, VE 66.0%		transplant recipients in Qatar;
		$(95\% \text{ CI}, 21.3 \text{ to } 85.3) \ge 42 \text{ days after } 2^{\text{nd}}$		782 participants; time and
		dose, and VE 73.9% (95% CI, 33 to 98.9)		setting for VOC Alpha and
		\geq 56 days after 2 nd dose (VOC Alpha and		VOC Beta.
		Beta).		VOC Beta.
		Deta).		
		BNT162b2 or mRNA-1273 showed VE		
		72.3% (95% CI, 0.0 to 90.9) against severe,		
		critical, or fatal disease ≥ 14 days after 2 nd		
		dose, VE 85% (95% CI, 35.7 to 96.5) ≥42		
		dose, VE 83% (95% CI, 35.7 to 90.3) 242 days after 2 nd dose, and VE 83.8% (95% CI,		
		days after 2 dose, and VE 63.6 /6 (93/6 CI, 31.3 to 96.2) ≥56 days after 2^{nd} dose (VOC		
		Alpha and Beta).		
91	Hu	Inactivated vaccines (CoronaVac) showed	Serious	Outbreak report of
71	114	VE 89% (95% CI, 55 to 98) against severe,	Scrious	hospitalized cases in China;
		critical, or fatal disease ≥ 14 days after 2^{nd}		476 participants; PCR
		dose (VOC Delta).		population for VOC Delta.
92	Andrews	BNT162b2 showed VE 62.7% (61.7 to 63.8)	Moderate	Test-negative study in
1/2	AHUIEWS	against symptomatic infection 1 week after	Moderate	England; 1,475,391
		against symptomatic infection I week after		England, 1,473,391

	T			
		2 nd dose and VE 47.3% (45.0 to 49.6) 20+		participants; VOC Alpha to
		weeks after 2 nd dose (VOC Delta).		VOC Delta (only data for VOC Delta reported here)
		ChAdOx1showed VE 92.4% (92.1 to 92.7)		, o d Bena reported nere)
		against symptomatic infection 1 week after		
		2 nd dose and VE 69.7% (68.7 to 70.5) 20+		
		weeks after 2 nd dose (VOC Delta).		
		mRNA-1273 showed VE 95.2% (94.4 to		
		95.9) against symptomatic infection 1 week		
		after 2 nd dose and VE 90.3% (67.2 to 97.1)		
		10 to 14 weeks after 2 nd dose (VOC Delta).		
93	<u>Patalon</u>	BNT162b2 (3 doses) showed relative VE	Moderate	Test-negative study of fully
		3% (95% CI, -5 to 10) against infection 0 to		vaccinated in Israel comparing
		6 days after 3 rd dose; relative VE 84.0%		(2 doses versus 3 doses);
		(95% CI, 79 to 88) 14 to 20 days after 3 rd		182,076 participants; time and
		dose compared to 2 doses.		setting for VOC Delta
94	Kissling	BNT162b2 showed VE 87% (95% CI, 74 to	Serious	Test-negative study of adults
		93) against symptomatic infection 14 days		>65 years in primary care
		after 2 nd dose.		setting in I-MOVE group
				(England, France, Ireland, the
				Netherlands, Portugal,
				Scotland, Spain and Sweden);
				4,964 participants; sample
0.5	3.5.77.1	D) //// (01 0 D) 1 4 0 7 0 1 1 1 1 1 1 1 1	0 :	sequenced for VOC Alpha.
95	<u>McKeigue</u>	BNT162b2 or mRNA-1273 showed VE	Serious	Case-control study of people
		92% (95% CI, 85 to 96) against severe		with clinical risk conditions in
		disease in people with no risk conditions		Scotland; 50,935 participants;
		and VE 72% (95% CI, 51 to 84) against		time and setting for VOC
		severe disease in people eligible for shielding at least 14 days after 2 nd dose.		Alpha to VOC Delta
		,		
		ChAdOx1 showed VE 94% (95% CI, 90 to		
		96) against severe disease in people with no		
		risk conditions and VE 63% (95% CI, 46 to		
		75) against severe disease in people eligible		
		for shielding \geq 14 days after 2 nd dose.		
96	<u>Kertes</u>	BNT162b2 showed OR 1.61 (95% CI, 1.45	Serious	Data-linkage study of people
		to 1.79) for infection comparing <u>fully</u>		fully vaccinated 6 months
		vaccinated Jan to Feb vs fully vaccinated		previously in Israel; 1,423,098
		Mar to May.		participants; time and setting for VOC Alpha to VOC Delta
97	Barlow	BNT162b2 or mRNA-1273 showed VE	Serious	Test-negative study in Oregon;
- '		74% (95% CI, 65 to 82) against infection ≥	221040	1000 participants; time and
		14 days after 2 nd dose.		setting for VOC Delta
		Ad26.COV2.S showed VE 51% (95% CI, -2		
		to 76) against infection \geq 14 days after 2 nd		
		dose.		
98	Chemaitelly	BNT162b2 showed VE 65.8% (95% CI,	Serious	Test-negative study in Qatar;
	<u>(3)</u>	63.8 to 67.7) against infection 5 to 9 weeks	Cerrous	1,472,761 participants; time
	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	after 2 nd dose; VE 29.7% (95% CI, 21.7 to		-,, . or participants, unic
	I	arcer 2 dose, 11 27.170 (7570 CI, 21.7 to		

		36.9) against infection 15 to 19 weeks after 2 nd dose and VE 0% (95% CI, 0 to 0) against infection 20 to 24 weeks after 2 nd dose. BNT162b2 showed VE 94.2% (95% CI, 91.0 to 96.5) against hospitalization or death 5 to 9 weeks after 2 nd dose; VE 86.4% (95% CI, 69.9 to 94.8) against hospitalization or death 15 to 19 weeks after 2 nd dose and VE 95.3% (95% CI, 70.5 to 99.9) against hospitalization or death 20 to 24 weeks after 2 nd dose.		and setting for VOC Beta to VOC Delta (results over varying time periods since vaccination reported)
99	Thompson (3)	BNT162b2 or mRNA-1273 showed VE 90% (95% CI, 86 to 93) against ICU admission ≥14 days after 2 nd dose. BNT162b2 showed VE 92% (95% CI, 88 to 94) against hospitalization at 28 to 41 days after 2 nd dose and VE 86% (95% CI, 74 to 93) ≥112 days after 2 nd dose.	Serious	Test-negative study of adults ≥50 years in the USA; 76,463 participants; time and setting for VOC Alpha (results over varying time periods since vaccination reported)
100	Bar-On	BNT162b2 (3 doses) showed adjusted rate ratio of 11.3 (95% CI, 10.4 to 12.3) against any infection and adjusted rate ratio of 19.5 (95% CI, 12.9 to 29.5) against severe illness ≥12 days after 3 rd dose compared to 2 doses.	Serious	Data-linkage study of fully vaccinated (age>60) (2 doses versus 3 doses) in Israel; 1,137,804 participants; time and setting for VOC Delta
101	Bruxvoort (2)	mRNA-1273 showed VE 98.4% (95% CI, 96.9 to 99.1) against infection ≥14 days after 2 nd dose (VOC Alpha). mRNA-1273 showed VE 95.5% (95% CI, 90.9 to 97.8) against infection ≥14 days after 2 nd dose (VOC Gamma). mRNA-1273 showed VE 86.7% (95% CI, 84.3 to 88.7) against infection ≥14 days after 2 nd dose (VOC Delta). mRNA-1273 showed VE 94.1% (95% CI, 90.5 to 96.3) against infection 14 to 60 days after 2 nd dose (VOC Delta). mRNA-1273 showed VE 80.0% (95% CI, 70.2 to 86.6) against infection 151 to 180 days after 2 nd dose (VOC Delta).	Serious	Test-negative study in Kaiser Permanente group in California; 48,918 participants; sequenced for VOC Alpha, VOC Delta, VOC Gamma and VOI Mu (results not included in this LES) (results over varying time periods since vaccination reported)
102	Tande (2)	BNT162b2 or mRNA-1273 showed VE 91% (95% CI, 72 to 98) against infection ≥14 days after 2 nd dose (January to March – VOC Alpha).	Serious	Point prevalence screening study in Mayo Clinic, USA; 46,008 participants; time and setting for VOC Alpha to VOC Delta

		BNT162b2 or mRNA-1273 showed VE		
		63% (95% CI, 44 to 76) against infection		
		≥14 days after 2 nd dose (June to August –		
		VOC Delta).		
103	Young-Xu	Two doses of BNT162b2 reduced risk of	Moderate	Retrospective cohort study of
103		infection by HR 66% (95% CI, 22 to 86)	Moderate	previously infected adults
	<u>(2)</u>	,		1
		compared to previously infected adults age		followed by Veterans Affairs
		65+ (June to August VOC Delta).		in USA; 47,102 participants;
				time and setting for VOC
		Two doses of mRNA-1273 reduced risk of		Delta
		infection by HR 68% (95% CI, 30 to 86)		
		and death by HR 30% (95% CI, -11 to 1)		
		compared to previously infected adults age		
		65+ (June to August VOC Delta).		
104	de Gier (1)	Fully vaccinated index to unvaccinated (hh	Serious	Retrospective cohort of
101	<u>40 0101 (1)</u>	contact) showed VET 73% (95% CI: 65 to	5511545	household and close contacts
		79).		in the Netherlands; 113,582
		(7).		1
		DNTT1/21 /) 1 1 1 1 1 7 7 7 7 7 7 7 7 7 7 7 7 7 7		cases and 253,168 contacts;
		BNT162b (case) showed VET 70% (95%		time and setting for VOC
		CI, 61 to 77) when fully vaccinated.		Alpha
		mRNA-1273 (case) showed VET 88% (95%		(hh = household)
		CI, 50 to 97) when fully vaccinated.		
		ChAdOx1 (case) showed VET 58% (95%		
		CI, -12 to 84) when fully vaccinated.		
		,		
		Ad26.COV2.S (case) showed VET 58%		
		(95% CI, -12 to 84) when fully vaccinated.		
		(0 / 1 32, 12 18 8 // 11 18 18 18 18 18 18 18 18 18 18 18 18		
		BNT162b showed VE 65% (95% CI, 60 to		
		70) when hh contact was fully vaccinated.		
		70) when the contact was rully vaccinated.		
		mRNA-1273 showed VE 91% (95% CI, 79		
		to 97) when hh contact was fully vaccinated.		
		ChAdOx1 showed VE 87% (95% CI, 77 to		
		93) when hh contact was fully vaccinated.		
		Ad26.COV2.S showed VE 12% (95% CI, -		
		71 to 54) when hh contact was fully		
		vaccinated.		
105	de Gier (2)	Fully vaccinated index to unvaccinated (hh	Serious	Retrospective cohort of
		contact) showed VET 63% (95% CI: 46 to		household and close contacts
		75).		in the Netherlands; 4,921
		1 - 7 -		cases and 7,771 contacts; time
		BNT162b (>50%) or mRNA-1273 or		and setting for VOC Delta
				and setting for VOC Delta
		ChAdOx1 or Ad26.COV2.S (case) showed		
		VET 40% (95% CI, 20 to 54) when both		
401	3.5 1	case and contacts are fully vaccinated.		D : 1 2
106	<u>Manley</u>	mRNA-1273 (50%) or BNT162b (48%) or	Serious	Retrospective cohort of
		Ad26.COV2.S (2%) showed OR of 8.89		maintenance dialysis patients

		(95% CI, 5.92 to 13.34) for unvaccinated vs fully vaccinated against infection (VOC Alpha)		in USA; 15,251 participants; time and setting for VOC Alpha to VOC Delta
		mRNA-1273 (50%) or BNT162b (48%) or Ad26.COV2.S (2%) showed OR of 2.27 (95% CI, 1.72 to 3.00) for unvaccinated vs fully vaccinated against infection (VOC Delta)		
107	Eyre	BNT162b2 (cases) showed VET 82% (95% CI, 71 to 88) against transmission after 2 nd dose. (VOC Alpha) ChAdOx1 (cases) showed VET 63% (95% CI, 37 to 78) against transmission after 2 nd dose. (VOC Alpha) BNT162b2 (contacts) showed VE 94% (95% CI, 90 to 96) against infection after 2 nd dose. (VOC Alpha) ChAdOx1 (contacts) showed VE 71% (95% CI, 51 to 83) against infection after 2 nd dose. (VOC Alpha) BNT162b2 (cases) showed VET 65% (95% CI, 52 to 74) against transmission after 2 nd dose. (VOC Delta) ChAdOx1 (cases) showed VET 36% (95% CI, 28 to 43) against transmission after 2 nd dose. (VOC Delta) BNT162b2 (contacts) showed VE 90% (95% CI, 87 to 92) against infection after 2 nd dose. (VOC Delta) ChAdOx1 (contacts) showed VE 72% (95% CI, 68 to 75) against infection after 2 nd dose. (VOC Delta).	Serious	Retrospective cohort of contacts in England; 99,597cases and 151,821 contacts; S-gene proxy for VOC Alpha and VOC Delta
108	Martinez- Baz (2)	BNT162b2 (contacts) showed VE 71% (95% CI, 61 to 78) against infection after 2 nd dose (VOC Alpha) mRNA-1273 (contacts) showed VE 86% (95% CI, 56 to 95) against infection after 2 nd dose (VOC Alpha) ChAdOx1 (contacts) showed VE 38% (95%	Serious	Prospective cohort of close contacts in Spain; 12,263 cases and 30,240 contacts; sequenced for VOC Alpha to VOC Delta (includes heterologous vaccines)
		CI, -42 to 73) against infection after 2 nd dose (VOC Alpha)		

		BNT162b2 (contacts) showed VE 67% (95% CI, 59 to 74) against infection after 2 nd dose (VOC Delta)		
		mRNA-1273 (contacts) showed VE 77% (95% CI, 64 to 85) against infection after 2 nd dose (VOC Delta)		
		ChAdOx1 (contacts) showed VE 55% (95% CI, 39 to 67) against infection after 2 nd dose (VOC Delta)		
		ChAdOx1 followed by BNT162b2 (contacts) showed VE 86% (95% CI, 45 to 97) against infection (VOC Delta)		
109	<u>Cohn</u>	BNT162b2 showed VE 49% (95% CI, 47 to 52) against infection at least 15 days after last dose (August: VOC Delta)	Serious	Data-linkage study of veterans in USA; 619,755 participants; time and setting for VOC Alpha to VOC Delta (only
		mRNA-1273 showed VE 64% (95% CI, 62 to 66) against infection at least 15 days after last dose (August: VOC Delta)		Delta reported here)
		Ad26.COV2.S showed VE 3% (95% CI, - 0.1 to 12) against infection at least 15 days after last dose (August: VOC Delta)		
110	Rosenberg (2)	BNT162b2 showed VE 69% (95% CI, 67.4 to 70.6) against infection at least 15 days after last dose (August: VOC Delta; age 18-49)	Serious	Prospective study in New York; 8,834,604 participants; time and setting for VOC Alpha to VOC Delta (only Delta reported here). Also
		mRNA-1273 showed VE 78.4% (95% CI, 75.9 to 79.6) against infection at least 15 days after last dose (August: VOC Delta; age 18-49)		compared VE over time since vaccination (results not reported here)
		Ad26.COV2.S showed VE 70.2% (95% CI, 67.4 to 73.0) against infection at least 15 days after last dose (August: VOC Delta; age 18-49)		
		BNT162b2 showed VE 77.8% (95% CI, 67.4 to 70.6) against infection at least 15 days after last dose (August: VOC Delta; age		
		65+)		
		, ,		

		1 10 (OOT 10 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		
		Ad26.COV2.S showed VE 70.8% (95% CI, 65.7 to 76.0) against infection at least 15 days after last dose (August: VOC Delta; age 65+)		
111	Robles- Fontan	BNT162b2 showed VE 56% (95% CI, 53 to 59) against infection at least 15 days after 2 nd dose (October: VOC Delta) mRNA-1273 showed VE 71% (95% CI, 68 to 74) against infection at least 15 days after 2 nd dose (October: VOC Delta) Ad26.COV2.S showed VE 27% (95% CI, 17 to 37) against infection at least 15 days after last dose (October: VOC Delta)	Serious	Data-linkage study in Puerto Rico; 1,913,454 person-years; time and setting for VOC Alpha to VOC Delta (only results for Delta reported here)
112	Glatman- Freedman (2)	BNT162b2 showed VE 91.5% (95% CI, 88.2 to 93.9) against infection at least 8 days after 2 nd dose in adolescents age 12 to 15 years. There were no deaths in either group.	Serious	Population cohort in Israel of adolescents age 12 to 15 years; 2,034,591 vaccinated persondays and 13,623,714 unvaccinated person-days; time and setting for VOC Delta
113	Chin	mRNA-1273 showed VE 56.6% (95% CI, 42 to 67.5) against infection at least 14 days after 2 nd dose.	Serious	Outbreak report from a prison in California; 827 participants; sample sequenced for VOC Delta
114	Nordstrom Panyani (2)	BNT162b2 showed VE 47% (95% CI, -39 to 55) against symptomatic infection 121 to 180 days after second dose. mRNA-1273 showed VE 71% (95% CI, 56 to 81) against symptomatic infection 121 to 180 days after second dose. ChAdOx1 showed VE 41% (95% CI, 29 to 51) against symptomatic infection to 120 days after second dose. ChAdOx1 followed by mRNA vaccine showed VE 66% (95% CI, 41 to 80) against symptomatic infection >120 days after second dose. BNT162b2 or mRNA-1273 or ChAdOx1 showed VE 42% (95% CI, -35 to 75) against severe disease (hospitalization or death) >180 days after second dose	Serious	Case-control study in Sweden; 1,684,958 participants; time and setting for VOC Alpha to VOC Delta (only Delta results reported here) (includes heterologous vaccines) (results over varying time periods since vaccination reported)
116	Ranzani (2)	ChAdOx1 showed VE 42.4% (95% CI, 24.6 to 56.0) against symptomatic infection 21 days after 1 st dose.	Low	Test-negative study in Brazil; 9,197 tests; time and setting for VOC Gamma to Delta

117	D '(2)	A 107 COMO C 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	C :	и
117	Ranzani(3)	Ad26.COV2.S showed VE 50.9% (95% CI, 35.5 to 63.0) against symptomatic infection, VE 92.5% (95% CI, 54.9 to 99.6) against ICU admission, and VE 90.5% (95% CI, 31.5 to 99.6) against death 28 days after dose.	Serious	Test-negative study in Brazil; 11,817 tests; time and setting for VOC Gamma to Delta
118	Chadeau- Hyam	BNT162b2 showed VE 71.3% (95% CI, 56.6 to 81.0) against infection unreported number of days after 2 nd dose (Round 13 and Round 14) mRNA-1273 showed VE 75.1% (95% CI, 22.7 to 92.0) against infection unreported number of days after 2 nd dose (Round 13 and Round 14) ChAdOx1showed VE 44.8% (95% CI, 22.5 to 60.7) against infection unreported	Serious	Surveillance study in England; 87,966 participants who consented to data-linkage for vaccine status; sequenced for VOC Delta
		number of days after 2 nd dose (Round 13 and Round 14)		
119	Sheikh (2)	BNT162b2 showed VE 90% (95% CI, 86 to 94) against death at least 14 days after 2 nd dose (confirmed VOC Delta)	Serious	Retrospective cohort in Scotland; 114,706 participants; proxy for VOC Delta
		ChAdOx1 showed VE 91% (95% CI, 83 to 94) against death at least 14 days after 2 nd dose (confirmed VOC Delta)		
120	Reis	BNT162b2 showed VE 59% (95% CI, 52 to 65) against infection 14 to 20 days after 1 st dose (age 12 to 18) BNT162b2 showed VE 90% (95% CI, 88 to 92) against infection 7 to 21 days after 2 nd	Moderate	Case-control study in Israel; 94,354 vaccinated matched to 94,354 unvaccinated adolescents age 12 to 18; time and setting for VOC Delta
		dose (age 12 to 18)		
121	Nordstrom (2)	BNT162b2 showed VE 78% (95% CI, 78 to 79) against symptomatic infection at least 14 days after 2 nd dose. mRNA-1273 showed VE 87% (95% CI, 84 to 88) against symptomatic infection at least 14 days after 2 nd dose. ChAdOx1 showed VE 50% (95% CI, 41 to 58) against symptomatic infection at least 14 days after 2 nd dose.	Serious	Retrospective cohort study in Sweden; 721,787 participants; time and setting for VOC Delta (includes heterologous vaccines)
		ChAdOx1 followed by BNT162b2 showed VE 67% (95% CI, 59 to 73) against symptomatic infection at least 14 days after 2 nd dose.		

		ChAdOx1 followed by mRNA-1273 showed VE 79% (95% CI, 62 to 88) against symptomatic infection at least 14 days after 2 nd dose.		
122	Skowronski (2)	BNT162b2 showed VE 79% (95% CI, 73 to 84) against infection at least 21 days after 1 st dose (VOC Gamma) mRNA-1273 showed VE 85% (95% CI, 71 to 92) against infection at least 21 days after 1 st dose (VOC Gamma) ChAdOx1 showed VE 60% (95% CI, 48 to 69) against infection at least 21 days after 1 st dose (VOC Gamma)	Serious	Test-negative study in Canada; 68,074 participants; sample sequenced for VOC Alpha, Gamma and Delta (only VOC Gamma reported here)
123	Skowronski (3)	Delta BNT162b2 showed VE 89% (95% CI, 88 to 89) against infection at least 14 days after 2 nd dose (Quebec- VOC Delta) mRNA-1273 showed VE 91% (95% CI, 90 to 92) against infection at least 14 days after 2 nd dose (Quebec- VOC Delta) ChAdOx1 showed VE 73% (95% CI, 69 to 78) against infection at least 14 days after 2 nd dose (Quebec- VOC Delta) ChAdOx1 followed by mRNA vaccine showed VE 88% (95% CI, 85 to 89) against infection at least 14 days after 2 nd dose (Quebec- VOC Delta) Gamma BNT162b2 showed VE 93% (95% CI, 89 to 95) against infection at least 14 days after 2 nd dose (BC- VOC Gamma) mRNA-1273 showed VE 95% (95% CI, 85 to 99) against infection at least 14 days after 2 nd dose (BC- VOC Gamma) ChAdOx1 showed VE 90% (95% CI, 61 to 98) against infection at least 14 days after 2 nd dose (BC- VOC Gamma) ChAdOx1 followed by mRNA vaccine showed VE 96% (95% CI, 70 to 99) against infection at least 14 days after 2 nd dose (BC- VOC Gamma) ChAdOx1 followed by mRNA vaccine showed VE 96% (95% CI, 70 to 99) against infection at least 14 days after 2 nd dose (BC- VOC Gamma) Time since vaccination (Delta)	Serious	Test-negative study in Canada; 380,532 British Columbia and 854,915 Quebec participants; sequenced for VOC Alpha, Gamma and Delta (selected data only reported here due to space constraints) (includes heterologous vaccines) (results over varying time periods since vaccination reported)

BNT162b2 showed VE 85% (95% CI, 84 to 86) against infection at 4 months after 2nd dose (Quebec – VOC Delta)

mRNA-1273 showed VE 88% (95% CI, 86 to 90) against infection at 4 months after 2nd

ChAdOx1 showed VE 72% (95% CI, 66 to 77) against infection at 4 months after 2nd dose (Quebec – VOC Delta)

dose (Quebec – VOC Delta)

ChAdOx1 followed by mRNA vaccine showed VE 86% (95% CI, 81 to 89) against infection at 4 months after 2nd dose (Quebec – VOC Delta)

Time since vaccination and interval between doses (VOC Alpha to Delta)

BNT162b2 showed VE 92% (95% CI, 91 to 93) at 14 to 27 days after 2nd dose (interval 7+ weeks) and VE 90% (95% CI, 88 to 91) at 4 months after 2nd dose (interval 7+ weeks) (Quebec)

mRNA-1273 showed VE 92% (95% CI, 90 to 94) at 14 to 27 days after 2nd dose (interval 7+ weeks) and VE 91% (95% CI, 87 to 94) at 112+ days after 2nd dose (interval 7+ weeks) (Quebec)

ChAdOx1 showed VE 85% (95% CI, 60 to 94) at 14 to 27 days after 2nd dose (interval 7+ weeks) and VE 72% (95% CI, 66 to 77) at 84 days after 2nd dose (interval 7+ weeks) (Quebec)

124	Lin	PNT162b2 showed VE 04 00/ /04 5 to 05 2	Carian-	Data linkaga atrada in NI- ud-
124	Lin	BNT162b2 showed VE 94.9% (94.5 to 95.2) against symptomatic infection and VE 95.9% (95% CI, 92.9 to 97.6) against death at 60 days months after 2 nd dose.	Serious	Data-linkage study in North Carolina; 10,600,823 participants; time and setting for VOC Alpha to Delta
		BNT162b showed VE 70.1% (95% CI, 68.9 to 71.2) against symptomatic infection and VE 88.4% (95% CI, 83 to 92.1) against death at 210 days after 2 nd dose)		(results over varying time periods since vaccination reported)
		mRNA-1273 showed VE 96% (95.6 to 96.4) against symptomatic infection at 60 days; VE 96% (95% CI, 91.9 to 98) against death at 90 days after 2 nd dose.		
		mRNA-1273 showed VE 81.9% (95% CI, 81 to 82.7) against symptomatic infection and VE 93.7% (95% CI, 90.2 to 95.9) against death at 210 days after 2 nd dose)		
		Ad26.COV2.S showed VE 79% (77.1 to 80.7) against symptomatic infection at 30 days and VE 64.3% (95% CI, 62.3 to 66.1) at 150 days months after dose.		
		Ad26.COV2.S showed VE 89.4% (95% CI, 52.3 to 97.6) against death at 120 days after dose)		
125	<u>Barda</u>	BNT162b2 (3 doses) showed VE 92% (82 to 97) against severe disease and VE 81% (95% CI, 59 to 97) against death at least 7 days after 3 rd dose compared to 2 doses (given 5 months previously).	Serious	Data-linkage study of fully vaccinated (2 doses vs 3 doses) participants in Israel; 728,321 participants in each group; time and setting for VOC Delta
126	Andrews (2)	BNT162b2 (3 doses) showed VE 94% (95% CI, 93.4 to 94.6) against symptomatic infection at least 14 days after 3 rd dose in age>50 (compared to unvaccinated) ChAdOx1 (2 doses followed by BNT162b2) showed VE 93.1% (95% CI, 91.7 to 94.3) against symptomatic infection at least 14	Moderate	Test-negative study of fully vaccinated participants (>140 days since 2 nd dose) over age 50 in England; 271,747 participants; sequencing for VOC Delta
127	Stamfalt (2)	days after 3 rd dose in age>50 (compared to unvaccinated)	Moderate	Population cohort study in
12/	Starrfelt (2)	BNT162b2 showed VE 69.7% (95% CI, 68.6 to 70.8) against infection at least 7 days after 2 nd dose (VOC Alpha to Delta) mRNA-1273 showed VE 78.2% (95% CI,	Moderate	Population cohort study in Norway; 4,293,544 participants; time and setting for VOC Alpha to VOC Delta
		76.7 to 79.6) against infection at least 7 days after 2 nd dose (VOC Alpha to Delta)		(includes heterologous vaccines)

		ChAdOx1 showed VE 43.4% (95% CI, 4.4 to 66.5) against infection at least 7 days after 2 nd dose (VOC Alpha to Delta) Heterologous mRNA showed VE 84.7% (95% CI, 83.1 to 86.1) against infection at least 7 days after 2 nd dose (VOC Alpha to Delta) ChAdOx1 followed by mRNA showed VE 60.7% (95% CI, 57.5 to 63.6) against infection at least 7 days after 2 nd dose (VOC Alpha to Delta)		
128	Preio- Alhambra	ChAdOx1 followed by BNT162b2 showed HR 0.61 (95% CI, 0.52 to 0.71) against infection vs ChAdOx1 (homologous) – unreported number of days after 2 nd dose	Serious	Retrospective cohort study in Spain; 28,650 participants aged 19 to 59 years; time and setting for VOC Delta (compared heterologous vaccines with homologous vaccines)
129	<u>Ng</u>	BNT162b2 or mRNA-1273 showed VE 61.6% (95% CI, 37.5 to 80.4) against transmission to fully vaccinated hh contacts and VE 100% (95% CI, not reported) against severe disease in fully vaccinated hh contacts	Serious	Retrospective cohort study of household contacts in Singapore; 753 contacts; index sequenced for VOC Delta
130	<u>Desai</u>	BBV152 showed VE 50% (95% CI, 33 to 62) against symptomatic infection at least 14 days after 2 nd dose	Serious	Test-negative study of HCW in India; 1,068 matched pairs; time and setting for VOC Delta
131	Thiruvengad am(pub)	ChAdOx1showed VE 46.2% (95% CI, 31.6 to 57.7) against infection at least 21 days after 1 st dose. ChAdOx1showed VE 63.1% (95% CI, 51.5 to 72.1) against infection at least 14 days after 2 nd dose.	Serious	Test-negative study in India; 5,143 participants; sequencing for VOC Delta
132	Sharma	BNT162b2 showed VE 45.7% (95% CI, 37.9 to 52.5) against infection median of 30 days after 3 rd dose compared to 2 doses (given at least 180 days previously) mRNA-1273 showed VE 46.6% (95% CI, 36.4 to 55.3) against infection median of 16 days after 3 rd dose compared to 2 doses (given at least 180 days previously)	Serious	Case-control study of fully vaccinated (2 doses versus 3 doses) in veterans in USA; 129,130 pairs; time and setting for VOC Delta

133	Cohn (2) Arbel	BNT162b2 showed VE 43% (95% CI, 42 to 45) against infection after unclear number of days after 2 nd dose (September 2021) mRNA-1273 showed VE 58% (95% CI, 57 to 59) after unclear number of days against infection after 2 nd dose (September 2021) Ad26.COV2.S showed VE 13% (95% CI, 9 to 17) against infection after unclear number of days after dose (September 2021) BNT162b2 (3 doses) showed VE 90% (95%	Serious	Retrospective cohort study of Veterans in the US; 780,225 Veterans; time and setting for VOC Delta (same population as Cohn but extended study time frame) Data-linkage study of fully
154	<u> Moci</u>	CI, 86 to 93) against death at 7 to 54 days after 3 rd dose compared to 2 doses (given at least 5 months previously)		vaccinated (>50 years) (2 doses versus 3 doses) in Israel; 843,208 participants; time and setting for VOC Delta
135	Bar-On (2)	BNT162b2 (3 doses) showed adjusted rate ratio of 12.3 (95% CI, 11.8 to 12.8) against infection and adjusted rate ratio of 17.9 (95% CI, 15.1 to 21.2) against severe disease and adjusted rate ratio of 14.7 (95% CI, 10 to 21.4) against death at least 12 days after 3 rd dose compared to 2 doses (given at least 5 months previously) (age>60). BNT162b2 (3 doses) showed adjusted rate ratio of 9.0 (95% CI, 8.4 to 9.7) against infection at least 12 days after 3 rd dose compared to 2 doses (given at least 5 months previously) (age 30-39).	Serious	Data-linkage study of fully vaccinated (>16 years) (2 doses versus 3 doses) in Israel; 4,696,865 participants; time and setting for VOC Delta (same population as Bar-On but extended end of study and additional ages and outcomes)
136	Andrews (3)	BNT162b2 (2 doses) showed VE 88% (65.9 to 95.8) against symptomatic infection at 2-9 weeks after 2 nd dose (VOC Omicron) BNT162b2 (2 doses) showed VE 34.3% (-5 to 58.7) against symptomatic infection at 25+ weeks after 2 nd dose (VOC Omicron) BNT162b2 (3 doses) showed VE 75.5% (56.1 to 86.3) against symptomatic infection at least 2+ weeks after 3 rd dose (VOC Omicron) ChAdOx1 (2 doses) showed VE 5.9% (-29.7 to 31.7) against symptomatic infection at 25+ weeks after 2 nd dose (VOC Omicron) ChAdOx1 (2 doses followed by 1 dose of BNT162b2) showed VE 71.4% (41.8 to 86) against symptomatic infection at least 2 weeks after 3 rd (VOC Omicron)	Moderate	Test-negative study of fully vaccinated participants in England; 187,887 (581 Omicron) participants; sequencing for VOC Delta and Omicron

138	McLean	BNT162b2 showed VE 53.8% (95% CI, 52.9 to 54.6) against infection up to 164 days after 2 nd dose (VOC Delta) BNT162b2 (3 doses) showed VE 81.2% (95% CI, 79.2 to 82.9) against infection up to 30 days after 3 rd dose (VOC Delta) mRNA-1273 showed VE 88.2% (95% CI, 83.1 to 91.8) against infection up to 44 days after 2 nd dose (VOC Delta) mRNA-1273 showed VE 65.0% (95% CI, 63.6 to 66.3) against infection up to 164 days after 2 nd dose (VOC Delta) mRNA-1273 (3 doses) showed VE 82.8% (95% CI, 58.8 to 92.9) against infection up to 30 days after 3 rd dose (VOC Delta) BNT162b2 showed VE 59% (95% CI, 24 to 78) against infection at least 14 days after 2 nd dose (VOC Delta - June to Dec 2021) mRNA-1273 showed VE 52% (95% CI, 20 to 71) against infection at least 14 days after 2 nd dose (VOC Delta - June to Dec 2021) BNT162b2 (3 doses) showed VE 92% (95% CI, 91 to 92) against infection at least 7 days after 3 rd dose. mRNA-1273 (3 doses) showed VE 94% (95% CI, 91 to 95) against infection at least 7 days after 3 rd dose. ChAdOx1 (2 doses) followed by BNT162b2 showed VE 82% (95% CI, 68 to 90) against infection at least 7 days after 3 rd dose. ChAdOx1 (2 doses) followed by BNT162b2 showed VE 82% (95% CI, 68 to 90) against infection at least 7 days after 3 rd dose	Serious	Prospective cohort in Wisconsin, USA; 1,518 participants; time and setting for VOC Delta Population cohort in Czech Republic; 693,579 fully vaccinated participants; time and setting for VOC Delta (includes heterologous vaccines)
		mRNA1273 showed VE 91% (95% CI, 63 to 98) against infection at least 7 days after 3 rd dose		
140	<u>Florea</u>	mRNA-1273 showed VE 86.5% (95% CI, 84.8 to 88.0) against infection at least 14 days after 2 nd dose	Serious	Prospective matched cohort study in California, USA; 1,854,008 participants; sequencing for VOC Delta
141	Kissling (2)	BNT162b2 showed VE 76% (95% CI, 72 to 81) against symptomatic infection at 30 -59 days after 2 nd dose; VE 72% (95% CI, 61 to 80) at 60-89 days after 2 nd dose and VE 65%	Serious	Test-negative study in 10 out of 14 I-MOVE countries; 14,282 participants; sample sequenced for VOC Delta

		(95% CI, 56 to 71) >90 days after 2 nd dose (age 30-59)		(results over varying time periods since vaccination reported)
		mRNA-1273 showed VE 91% (95% CI, 85 to 95) against symptomatic infection at 30 - 59 days after 2 nd dose; VE 90% (95% CI, 76 to 96) at 60-89 days after 2 nd dose (age 30-59)		reported)
		ChAdOx1 showed VE 67% (95% CI, 57 to 75) against symptomatic infection at 30 -59 days after 2 nd dose; VE 65% (95% CI, 48 to 76) at 60-89 days after 2 nd dose (age 30-59)		
		Ad26.COV2.S showed VE 50% (95% CI, 36 to 62) against symptomatic infection at 30 -59 days after dose; VE 52% (95% CI, 33 to 66) at 60-89 days after dose (age 30-59)		
142	Katikireddi	ChAdOx1 showed VE 63.3% (95% CI, 61.3 to 65.3) against symptomatic infection at 8 to 9 weeks after 2 nd dose; VE 48.7% (95% CI, 45.9 to 51.4) against symptomatic infection at 16 to 17 weeks after 2 nd dose (VOC Delta) ChAdOx1 showed VE 79.0% (95% CI, 75.9 to 81.7) against severe disease (hospitalization or death) at 8 to 9 weeks after 2 nd dose; VE 70.5% (95% CI, 67.0 to 73.7) against severe disease 16 to 17 weeks after 2 nd dose (VOC Delta) ChAdOx1 showed VE 65.4% (95% CI, 64.6 to 66.2) against symptomatic infection at 8 to 9 weeks after 2 nd dose; VE 58.7% (95% CI, 56.7 to 60.5) against symptomatic	Serious	Retrospective cohort in Scotland and Brazil; 1,972,454 fully vaccinated participants in Scotland (Delta); 42,558,839 fully vaccinated participants in Brazil (Gamma); time and setting for VOC Delta and VOC Gamma (results over varying time periods since vaccination reported)
		infection at 16 to 17 weeks after 2 nd dose (VOC Gamma) ChAdOx1 showed VE 75.6% (95% CI, 73.4 to 77.6) against severe disease (hospitalization or death) at 8 to 9 weeks after 2 nd dose; VE 50.5% (95% CI, 43.4 to 56.6) against severe disease 16 to 17 weeks after 2 nd dose (VOC Gamma)		

143	Abu-Raddad	mRNA-1273 showed VE 90.6% (95% CI,	Serious	Test-negative study in Qatar;
	<u>(4)</u>	88.7 to 92.1) against infection at 60 days after 2 nd dose; VE 80.7% (95% CI, 77 to		1,781,505 participants; time and setting for VOC Beta to
		83.8) against infection at 120 days after 2 nd dose		VOC Delta (same setting and methodology as Chemaitelly 3)
		mRNA-1273 showed VE 97.8% (95% CI, 83.7 to 99.7) against severe disease		(results over varying time periods since vaccination
		(hospitalization or death) at 60 days after 2 nd		reported)
		dose; VE 91.5% (95% CI, 60.8 to 98.1) against infection at 120 days after 2 nd dose		
144	<u>Machado</u>	BNT162b2 (majority) or mRNA-1273 showed VE 68% (95% CI, 64 to 71) against	Moderate	Retrospective cohort study of community-dwelling
		symptomatic infection at 42-69 days after 2 nd dose; VE 39% (95% CI, 29 to 48)		adults≥65 in Portugal; 2,117,002 participants; time
		against symptomatic infection at 98-148 days after 2 nd dose		and setting for VOC Alpha to VOC Delta (same population as Nunes)
		ChAdOx1 showed VE 33% (95% CI, 23 to		(same population as indies)
		42) against symptomatic infection at 42-69 days after 2 nd dose; VE 34% (95% CI, 10 to		(results over varying time
		52) against symptomatic infection at 70-140 days after 2 nd dose		periods since vaccination reported)
		BNT162b2 (majority) or mRNA-1273 showed VE 95% (95% CI, 88 to 98) against		
		death at 14-41 days after 2 nd dose; VE 93%		
		(95% CI, 87 to 96) against death at 70-148 days after 2 nd dose		
		ChAdOx1 showed VE 95% (95% CI, 90 to		
		97) against death at least 14 days after 2 nd dose		
145	<u>Irizarry</u>	BNT162b2 showed VE 57% (95% CI, 53 to 60) against infection at 144 days after 2 nd	Serious	Retrospective cohort study in Puerto Rico; 2,276,966
		dose; VE 86% (95% CI, 75 to 92) against death at 144 days after 2 nd dose		participants; time and setting for VOC Alpha to VOC Delta
		mRNA-1273 showed VE 73% (95% CI, 70		(same population as Robles-
		to 76) against infection at 144 days after 2 nd		Fontan?)
		dose; VE 93% (95% CI, 81 to 97) against death at 144 days after 2 nd dose		(results over varying time periods since vaccination
		Ad26.COV2.S showed VE 36% (95% CI,		reported)
		30 to 42) against infection at 144 days after 2 nd dose; VE 72% (95% CI, 49 to 85)		
1.4.5	T	against death at 144 days after 2 nd dose	M 1 .	D (1) 1 1
146	Tartof (2)	BNT162b2 (3 doses) showed VE 88% (95% CI, 86 to 89) against infection at least 14	Moderate	Retrospective cohort study in California, USA; 3,133,075
		days after 3 rd dose compared to		participants; time and setting
		unvaccinated (age>18)		for VOC Alpha to VOC Delta

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147	Buchan	BNT162b2 (3 doses) showed VE 75% (95% CI, 71 to 78) against infection at least 14 days after 3 rd dose compared to 2 doses (given at least 6 months previously) (age>18) BNT1652b2 or mRNA-1273 (2 doses) showed VE 6% (95% CI, -25 to 30) against infection at 7 to 59 days after 2 nd dose; VE -13% (95% CI, -38 to 8) against infection at 60 to 119 days after 2 nd dose; VE -38% (95% CI, -61 to -18) against infection at 120 to 179 days after 2 nd dose; VE -16% (95% CI, -62 to 17) against infection at >240 days after 2 nd dose (VOC Omicron) BNT162b2 (3 doses) showed VE 34% (95% CI, 16 to 49) against infection at 7 days after 3 rd dose (VOC Omicron) mRNA-1273 (3 doses) showed VE 59% (95% CI, 16 to 80) against infection at 7 days after 3 rd dose (VOC Omicron) BNT1652b2 or mRNA-1273 (2 doses) showed VE 84% (95% CI, 81 to 86) against infection at 7 to 59 days after 2 nd dose; VE 81% (95% CI, 79 to 82) against infection at 60 to 119 days after 2 nd dose; VE 80% (95% CI, 79 to 81) against infection at 120 to 179 days after 2 nd dose; VE 71% (95% CI, 66 to 75) against infection at >240 days after 2 nd dose (VOC Delta) BNT162b2 (3 doses) showed VE 93% (95% CI, 91 to 94) against infection at 7 days after 3 rd dose (VOC Delta) BNT162b2 (3 doses) showed VE 93% (95% CI, 91 to 94) against infection at 7 days after 3 rd dose (VOC Delta)	Moderate	Test-negative study in Ontario, Canada; 484,188 fully vaccinated participants; sample sequenced for VOC Delta and VOC Omicron (results over varying time periods since vaccination reported)
		mRNA-1273 (3 doses) showed VE 93% (95% CI, 90 to 96) against infection at 7 days after 3 rd dose (VOC Delta)		

148	Tseng	mRNA-1273 (2 doses) showed VE 30.4%	Serious	Test-negative study in
148	Tseng	(95% CI, 5.0 to 49.0) against infection at 14 to 90 days after 2 nd dose; VE 15.2% (0 to 30.7) against infection at 91 to 180 days after 2 nd dose; VE 0% (95% CI, 0 to 1.2) against infection at 181 to 270 days after 2 nd dose (VOC Omicron) mRNA-1273 (3 doses) showed VE 63.6% 95% CI, 57.4 to 68.9) against infection at median of 35 days after 3 nd dose (VOC Omicron) mRNA-1273 (2 doses) showed VE 82.8% (95% CI, 69.6 to 90.3) against infection at 14 to 90 days after 2 nd dose; VE 63.6% (51.8 to 72.5) against infection at 91 to 180 days since 2 nd dose; VE 61.4% (95% CI, 56.8 to 65.5) against infection at 181 to 270 days after 2 nd dose; VE 52.9% (95% CI, 43.7 to 60.5) against infection at >270 days after 2 nd dose (VOC Delta) mRNA-1273 (3 doses) showed VE 95.7% 95% CI, 94.2 to 96.8) against infection at median of 35 days after 3 rd dose (VOC	Serious	California, USA; 60,420 participants; sample sequenced for VOC Delta and VOC Omicron (results over varying time periods since vaccination reported)
149	Lyngse	Delta) BNT162b2* (cases) showed VET 10% (95% CI, 0 to 18) against transmission to vaccinated household contacts at least 7 days after 2 nd dose BNT162b2* (cases) showed VET 31% (95% CI, 26 to 36) against transmission to unvaccinated household contacts at least 7 days after 2 nd dose	Serious	Household transmission study in Denmark; 24,693 index cases; sequencing for VOC Delta
		BNT162b2* (contacts) showed VES 46% (95% CI, 40 to 52) against susceptibility to infection from vaccinated case at least 7 days after 2 nd dose BNT162b2* (contacts) showed VES 61%		
		(95% CI, 59 to 63) against susceptibility to infection from unvaccinated household contacts at least 7 days after 2 nd dose		
		*vast majority		

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150	Hitchings (3)	CoronaVac (2 doses) showed OR 1.59 (95% CI, 0.60 to 4.24) for infection comparing	Serious	Test-negative study in Brazil; 37,929 matched fully
	()	fully vaccinated ≥182 days vs fully		vaccinated participants; time
		vaccinated 14 to 41 days (age 40-64)		and setting for VOC Gamma
		vaccinated 11 to 11 days (age 10 01)		and VOC Delta
		CoronaVac (2 doses) showed OR 3.32 (95%		and voc Beta
		CI, 1.85 to 5.94) for infection comparing		
		fully vaccinated ≥182 days vs fully		
		vaccinated 14 to 41 days (age 80+)		
151	Abu-Raddad	BNT162b2 (3 doses) showed VE 50.1%	Serious	Retrospective cohort studies
	<u>(5)</u>	(95% CI, 47.3 to 52.8) against symptomatic		in Qatar; 2,232,224 fully
	\	infection; VE 100% (71.4 to 100) against		vaccinated participants;
		hospitalization and death compared to 2		sample sequenced for VOC
		doses		Omicron
		doses		Officion
		mRNA-1273 (3 doses) showed VE 50.8%		
		(95% CI, 43.4 to 57.3) against symptomatic		
		infection compared to 2 doses		
152	Zheutlin	BNT162b2 showed VE 84% (95% CI, 82 to	Serious	Matched case-control in USA;
134	Zircutiii	85) against infection ≥5 months after 2 nd	ocnous	17,017,435 fully vaccinated
		dose		participants; time and setting
		dose		for VOC Alpha to VOC Delta
		mRNA-1273 showed VE 88% (95% CI, 87		(only Delta data shown here)
		to 89) against infection ≥ 5 months after 2^{nd}		(only Delta data shown here)
		dose		(results over varying time
		dosc		periods since vaccination
		Ad26.COV2.S showed VE 74% (95% CI,		reported)
		70 to 76) against infection \geq 5 months after		reported)
		dose		
153	Cerqueira-	BNT162b2 showed VE 64.8% (95% CI,	Serious	Test-negative study in Brazil;
	Silva	54.9 to 72.4) against symptomatic infection		231,212 previously infected
	<u> </u>	≥14 days after 2 nd dose		participants; time and setting
				for VOC Gamma to VOC
		ChAdOx1 showed VE 56% (95% CI, 51.4		Delta
		to 60.2) ≥ 14 days after 2^{nd} dose		
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		CoronaVac showed VE 39.4% (95% CI,		
		36.1 to 42.6) against symptomatic infection		
		≥14 days after 2 nd dose		
		Ad26.COV2.S showed VE 44% (95% CI,		
		31.5 to 54.2) against symptomatic infection		
		≥14 days after dose		
154	Jara (2)	CoronaVac (3 doses) showed VE 78.8%	Moderate	Prospective cohort in Chile;
	- 	(95% CI, 76.8 to 80.6) against symptomatic		11,174,257 fully vaccinated
		infection; VE 92.2% (95% CI, 88.7 to 94.6)		participants; time and setting
		against ICU admission; VE 86.7% (95% CI,		for VOC Delta
		80.5 to 91.0) against death \geq 14 days after 3 rd		
		dose		(includes heterologous
				vaccines)
		BNT162b2 booster after CoronaVac (2		
		doses) showed VE 96.5% (95% CI, 96.2 to		
		,		

		96.7) against symptomatic infection; VE 96.2% (95% CI, 94.6 to 97.3) against ICU admission; VE 96.8% (95% CI, 93.9 to 98.3) against death ≥14 days after 3 rd dose ChAdOx1 booster after CoronaVac (2		
		doses) showed VE 93.2% (95% CI, 92.9 to		
		93.6) against symptomatic infection; VE		
		98.9% (95% CI, 98.5 to 99.2) against ICU admission; VE 98.1% (95% CI, 97.3 to 98.6) against death ≥14 days after 3 rd dose		
155	Tan	BNT162b2 (3 doses) showed VE 73% (95% CI, 71 to 74) against infection; VE 95% (95% CI, 92 to 97) against severe disease ≥12 days after 3 rd dose compared to 2 doses mRNA-1273 (3 doses) showed VE 86% (95% CI, 81 to 90) against infection ≥12 days after 3 rd dose compared to 2 doses of BNT162b2	Serious	Retrospective cohort study in Singapore; 73,209 fully vaccinated participants (age>60); time and setting for VOC Delta (includes heterologous vaccines)
		BNT162b2 (2 doses) followed by mRNA- 1273 showed VE 82% (95% CI, 77 to 86) against infection; VE 92% (95% CI, 44 to 99) against severe disease ≥12 days after 3 rd dose compared to 2 doses of BNT162b2 mRNA-1273 (2 doses) followed by		
		BNT162b2 showed VE 90% (95% CI, 73 to 96) against infection ≥12 days after 3 rd dose compared to 2 doses of BNT162b2		
156	Suah	BNT162b2 (2 dose vaccinated July to August) showed VE 90.8% (95% CI, 89.4 to 92.0) against infection; VE 83.8% (95% CI, 78.5 to 87.8) against ICU admission; VE 90.3% (95% CI, 88.1 to 92.2) against death in September (at least 14 days after 2 nd dose)	Serious	Retrospective cohort study in Malaysia; 9,927,350 fully vaccinated participants; time and setting for VOC Delta (results over varying time periods since vaccination
		BNT162b2 (2 dose vaccinated April to June) showed VE 79.1% (95% CI, 75.8 to 81.9) against infection; VE 57.2% (95% CI, 43.4 to 67.6) against ICU admission; VE 89.3% (95% CI, 85.9 to 91.9) against death in September (at least 14 days after 2 nd dose)		reported)
		CoronaVac (2 dose vaccinated July to August) showed VE 74.4% (95% CI, 70.4 to 77.8) against infection; VE 46.1% (95% CI, 37.2 to 53.7) against ICU admission; VE 76.5% (95% CI, 72.9 to 79.6) against death in September (at least 14 days after 2 nd dose)		

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		CoronaVac (2 dose vaccinated April to		
		June) showed VE 30% (95% CI, 18.4 to		
		39.9) against infection; VE 30.2% (95% CI,		
		7.6 to 47.3) against ICU admission; VE		
		75.7% (95% CI, 67.0 to 82.1) against death		
		in September (at least 14 days after 2 nd dose)		
157	<u>Amodio</u>	mRNA-1273 showed VE 69.2% (95% CI,	Serious	Retrospective cohort study in
		67.6 to 70.8) against infection; VE 85.2%		Italy; 3,966,976 participants;
		(95% CI, 82.7 to 87.7) against severe disease		time and setting for VOC
		at 6 months after 2 nd dose		Alpha to VOC Delta (only
				Delta data shown here)
		mRNA-1273 showed VE 69.2% (95% CI,		
		67.6 to 70.8) against infection; VE 90.3%		(results over varying time
		(95% CI, 86.2 to 94.4) against severe disease		periods since vaccination
		at 8 months after 2 nd dose		reported)
158	Roberts	BNT162b2 showed VE 72.7% (95% CI,	Serious	Test-negative study in USA;
		65.4 to 78.5) against infection; VE 71.7%		170,487 participants; time and
		(95% CI, 45.1 to 85.6) against severe disease		setting for VOC Alpha to
		(21 days to <3 months after 2 nd dose)		VOC Delta (only Delta data
		(participants tested July–September 2021)		shown here)
		, d 1		,
		BNT162b2 showed VE 73.8% (95% CI,		
		63.6 to 81.2) against infection; VE 68.3%		
		(95% CI, 23.6 to 87.2) against severe disease		
		(21 days to <3 months after 2 nd dose)		
		(participants tested October–December		
		2021)		
		mRNA-1273 showed VE 79.0% (95% CI,		
		70.8 to 84.9) against infection; VE 74.5%		
		(95% CI, 42.7 to 88.9) against severe disease		
		(21 days to <3 months after 2 nd dose)		
		(participants tested July–September 2021)		
		mRNA-1273 showed VE 83.1% (95% CI,		
		68.9 to 90.9) against infection; VE 93.4%		
		(95% CI, 5.3 to 99.6) against severe disease		
		(21 days to <3 months after 2 nd dose)		
		(participants tested October–December		
		2021)		
159	Bar-On (3)	BNT162b2 (3 doses) showed a rate ratio	Serious	Data-linkage study of 4 doses
		(RR) of 1.9 (95% CI, 1.8 to 1.9) for		(>60 years) (3 doses versus 4
		infection; RR 4.0 (95% CI, 2.3 to 7.0) for		doses) in Israel; 1,138,681
		severe disease compared to 4 doses		participants; time and setting
		•		for VOC Omicron
160	<u>Willett</u>	BNT162b2 (3 doses) showed VE 43.2%	Serious	Test-negative study in
		(95% CI, 38.1 to 47.8) against infection		Scotland; 1,200,000
		(VOC Omicron)		participants; sample sequenced
				for VOC Omicron and VOC
		mRNA-1273 (3 doses) showed VE 46.3%		Delta
		(95% CI, 41.3 to 51.0) against infection		
		(VOC Omicron)		
		(VOC Omicron)		

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		BNT162b2 (2 doses) showed VE 26% (95% CI, x to x) against infection (VOC Omicron)		
		mRNA-1273 (2 doses) showed VE 23.7% (95% CI, x to x) against infection (VOC Omicron)		
		BNT162b2 (3 doses) showed VE 85.9% (95% CI, 84.2 to 87.4) against infection (VOC Delta)		
		mRNA-1273 (3 doses) showed VE 86.5% (95% CI, 84.8 to 88.0) against infection (VOC Delta)		
		BNT162b2 (2 doses) showed VE 83.5% (95% CI, x to x) against infection (VOC Delta)		
		mRNA-1273 (2 doses) showed VE 87.8% (95% CI, x to x) against infection (VOC Delta)		
161	<u>Jalali</u>	BNT162b2 or mRNA-1273 (3 doses) showed VES 47% (95% CI, 17 to 64) against transmission at least 7 days after 3 rd dose (VOC Omicron)	Serious	Retrospective cohort study in Norway; 979 primary cases and 1,888 household contacts; sample sequenced for VOC Omicron and VOC Delta
		BNT162b2 or mRNA-1273 (2 doses) showed VES 16% (95% CI, 0 to 37) against transmission at least 7 days after 2 nd dose (VOC Omicron)		Officion and VOC Delta
		BNT162b2 or mRNA-1273 (3 doses) showed VES 62% (95% CI, 38 to 78) against transmission at least 7 days after 3 rd dose (VOC Delta)		
		BNT162b2 or mRNA-1273 (2 doses) showed VES 46% (95% CI, 28 to 58) against transmission at least 7 days after 2 nd dose (VOC Delta)		
162	Chemaitelly (4)	BNT162b2 (3 doses) showed VE 56.6% (95% CI, 50.8 to 61.7) against symptomatic infection at 28 to 35 days; VE 43.7% (95% CI, 32.9 to 52.7) against symptomatic	Serious	Test negative study in Qatar; 2,193,013 participants; proxy for VOC Omicron
		infection 70 to 77 days after 3 rd dose		(results over varying time periods since vaccination
		BNT162b2 (3 doses) showed VE 90.6% (95% CI, 77.8 to 96) against severe, critical, or fatal disease at 7 to 42 days; VE 90.8%		reported)

163	Fabiani (2)	(95% CI, 81.5 to 95.5) against severe, critical, or fatal disease at 49 days+ after 3 rd dose mRNA-1273 (3 doses) showed VE 54.6% (95% CI, 41.1 to 65.0) against symptomatic infection at 28 to 35 days; VE 38.6% (95% CI, 19.4 to 53.1) against symptomatic infection at least 42 days after 3 rd dose mRNA-1273 (3 doses) showed VE 80.8% (95% CI, -51.9 to 97.6) against severe, critical, or fatal disease at 7 to 42 days after 3 rd dose BNT162b2 (2 doses) showed VE 61.9% (95% CI, 49.9 to 71.1) against symptomatic infection at 30 days; VE 45.9% (95% CI, 33.8 to 55.8) against symptomatic infection at 60 days; VE 36.3% (95% CI, 25.1 to 45.8) against symptomatic infection at 90 days after 2 nd dose mRNA-1273 (2 doses) showed VE 44.8% (95% CI, 16.0 to 63.8) against symptomatic infection at 28 to 35 days after 2 nd dose BNT162b2 showed VE 82% (95% CI, 80.5 to 83.5) against infection at 21 to 30 days after 2 nd dose; VE 67.3% (95% CI, 65.2 to 69.3) against infection at 44 to 98 days after 2 nd dose compared to non-immune period after 1 st dose BNT162b2 showed VE 96.3% (95% CI, 95 to 97.3) against severe disease at 21 to 30 days after 2 nd dose; VE 91.1% (95% CI, 95 to 97.3) against severe disease at 21 to 30 days after 2 nd dose; VE 91.1% (95% CI, 90 to 92) against severe disease at 44 to 98 days	Serious	Retrospective cohort study in Italy; 33,250,344 partially vaccinated participants; time and setting for VOC Delta (results over varying time periods since vaccination reported)
164	Stitionaltho	after 2 nd dose compared to non-immune period after 1 st dose	Serious	Test positive study in
164	Sritipsukho	CoronaVac (2 doses) + BNT162b2 showed VE 98% (95% CI, 87 to 100) against infection at least 7 days after 3 rd dose CoronaVac (2 doses) + ChAdOx1 showed VE 86% (95% CI, 74 to 93) against infection at least 7 days after 3 rd dose ChAdOx1 (2 doses) showed VE 83% (95% CI, 70 to 90) against infection at least 7 days after 2 nd dose	Serious	Test-negative study in Thailand; 3,353 participants; time and setting for VOC Delta (includes heterologous vaccines)

		CoronaVac (1 dose) + ChAdOx1 showed VE 74% (95% CI, 43 to 88) against infection at least 7 days after 2 nd dose		
		CoronaVac (2 doses) showed VE 60% (95% CI, 49 to 69) against infection at least 7 days		
165	Cerqueira- Silva(2)	after 2 nd dose CoronaVac (2 doses) + BNT162b2 showed VE 92.7% (95% CI, 91 to 94) against infection at 14 to 30 days after 3 rd dose CoronaVac (2 doses) + BNT162b2 showed VE 97.3% (95% CI, 96.1 to 98.1) against severe disease (hospitalization or death) at 14 to 30 days after 3 rd dose	Serious	Test-negative study in Brazil; 7,314,318 participants; time and setting for VOC Gamma and Delta (only booster data shown here because it is most likely to represent Delta) (results over varying time periods since vaccination reported) (includes heterologous vaccines)
166	Grima	BNT162b2 or mRNA-1273 or ChAdOx1 (3 doses) showed OR 0.60 (95% CI, 0.33 to 1.10) against transfer to ICU; OR 0.70 (95% CI, 0.27 to 1.80) against death unreported number of days after 3 rd dose (VOC Omicron) BNT162b2 or mRNA-1273 or ChAdOx1 (3 doses) showed OR 0.38 (95% CI, 0.16 to 0.92) against transfer to ICU; OR 0.80 (95% CI, 0.35 to 1.81) against death unreported number of days after 3 rd dose (VOC Delta)	Serious	Time-matched cohort in Canada; 20,064 participants hospitalized due to COVID; sequenced for variants (only VOC Omicron and VOC Delta reported here) (results not reported according to vaccine brand)
167	Monge	BNT162b2 (2 doses) followed by an mRNA vaccine showed VE 49.7% (95% CI, 48.3 to 51.1) against infection at least 7 days after 3 rd dose mRNA-1273 (2 doses) followed by an mRNA vaccine showed VE 55.3% (95% CI, 52.3 to 58.2) against infection at least 7 days after 3 rd dose ChAdOx1 (2 doses) followed by an mRNA vaccine showed VE 58.6% (95% CI, 55.5 to 61.6) against infection at least 7 days after 3 rd dose Ad26.COV2.S followed by an mRNA vaccine showed VE 48.0% (95% CI, 42.5 to 53.7) against infection at least 7 days after 3 rd dose	Serious	Retrospective cohort study in Spain; 6,222,318 fully vaccinated participants >40 years; time and setting for VOC Omicron (results over varying time periods since vaccination reported) (includes heterologous vaccines)

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168	Patalon (2)	BNT162b2 (3 doses) showed VE 35.7% (95% CI, 29.8 to 41.2) against infection up	<u>Moderate</u>	Test-negative study in Israel; 109,633 fully vaccinated
		to 90 days after 3 rd dose (Nov 2021		participants; time and setting
		compared to Aug 2021)		for VOC Omicron
169	<u>Smid</u>	BNT162b2 (3 doses) showed VE 58% (95%	Serious	Retrospective cohort study in
		CI, 57 to 58) against infection up to 60 days		Czech Republic; 4,874,253
		after 3 rd dose (VOC Omicron)		participants (for the outcomes reported here); sample
		BNT162b2 (2 doses) showed VE 49% (95%		sequenced for VOC Omicron
		CI, 48 to 50) against infection up to 60 days		and VOC Delta
		after 2 nd dose (VOC Omicron)		
		mRNA-1273 (3 doses) showed VE 61%		
		(95% CI, 60 to 62) against infection up to		
		60 days after 3 rd dose (VOC Omicron)		
		mRNA-1273 (2 doses) showed VE 48%		
		(95% CI, 44 to 52) against infection up to		
		60 days after 2 nd dose (VOC Omicron)		
		C1 A 10 A (0.1) 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		
		ChAdOx1 (2 doses) showed VE 51% (95% CI, 23 to 69) against infection up to 120		
		days after 2 nd dose (VOC Omicron)		
		days after 2 dose (voc officion)		
		Ad26.COV2.S (1 dose) showed VE 47%		
		(95% CI, 45 to 49) against infection up to		
		60 days after 2 nd dose (VOC Omicron)		
		BNT162b2 (3 doses) showed VE 90% (95%		
		CI, 89 to 90) against infection up to 60 days		
		after 3 rd dose (VOC Delta)		
		,		
		BNT162b2 (2 doses) showed VE 82% (95%		
		CI, 80 to 83) against infection up to 60 days		
		after 2 nd dose (VOC Delta)		
		mRNA-1273 (3 doses) showed VE 92%		
		(95% CI, 91 to 93) against infection up to		
		60 days after 3 rd dose (VOC Delta)		
		DNIA 4070 (0.1) 1 1475 740/		
		mRNA-1273 (2 doses) showed VE 71%		
		(95% CI, 64 to 76) against infection up to 60 days after 2 nd dose (VOC Delta)		
		oo days after 2 dose (VOC Detta)		
		ChAdOx1 (2 doses) showed VE 65% (95%		
		CI, 57 to 71) against infection up to 120		
		days after 2 nd dose (VOC Delta)		
		Ad26 COV2 \$ (1 dose) showed VE 600/		
		Ad26.COV2.S (1 dose) showed VE 60% (95% CI, 57 to 62) against infection up to		
		60 days after 2 nd dose (VOC Delta)		
		oo days areer 2 dose (v Oo Dera)		

170	Norddahl	BNT162b2 (3 doses) showed VE 47% (95% CI, 36 to 56) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) BNT162b2 (2 doses) followed by mRNA-1273 showed VE 50% (95% CI, 34 to 62) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) mRNA-1273 (3 doses) showed VE 9% (95% CI, -21 to 32) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) mRNA-1273 (2 doses) followed BNT162b2 showed VE 27% (95% CI, 9 to 61) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) ChAdOx1 (2 doses) followed by BNT162b2 showed VE 30% (95% CI, 14 to 43) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) ChAdOx1 (2 doses) followed by mRNA-1273 showed VE 7% (95% CI, -16 to 25) against infection unknown number of days after 3 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) Ad26.COV2 followed by BNT162b2 showed VE 5% (95% CI, -7 to 15) against infection unknown number of days after 2 rd dose relative to 2 doses of BNT162b2 (VOC Omicron) Ad26.COV2 followed by mRNA-1273 showed VE 70% (95% CI, 50 to 80) against infection unknown number of days after 2 rd dose relative to 2 doses of BNT162b2 (VOC Omicron)	Serious	Retrospective population cohort study in Iceland; 278,026 at least partly vaccinated participants; sequenced for VOC Omicron and VOC Delta (only Omicron data shown here) (includes heterologous vaccines)
		showed VE -70% (95% CI, -50 to -80)		
		against infection unknown number of days after 2 nd dose relative to 2 doses of BNT162b2 (VOC Omicron)		
171	Rane	BNT162b2 (2 doses) showed VE 76% (95% CI, 74 to 78) against symptomatic infection unknown number of days after 2 nd dose mRNA-1273 (2 doses) showed VE 83%	Serious	Test-negative study in New York; 1,058,493 participants; time and setting for VOC Alpha to VOC Delta (results for VOC Delta shown here)
		(95% CI, 81 to 84) against symptomatic		101 VOC Della silowii licie)

				<u> </u>
		infection unknown number of days after 2 nd		
		dose		
		Ad26.COV2.S showed VE 29% (95% CI,		
		26 to 32) against symptomatic infection		
		unknown number of days after dose		
470	****	, , , , , , , , , , , , , , , , , , ,	0 :	
172	<u>Wu</u>	BBIBP-CorV showed VES 39.4% (-20.4 to	Serious	Outbreak cohort in China;
		69.5) against symptomatic infection from 14		1,462 close-contacts of index
		to 90 days after 2 nd dose		case; sequenced for VOC
		,		Delta
		C		Delta
		CoronaVac showed VES 45.5% (-6 to 72)		
		against symptomatic infection from 14 to 90		(results over varying time
		days after 2 nd dose		periods since vaccination
		, and the second		reported)
173	Gazit (3)	BNT162b2 (single dose) after previously	Serious	Series of retrospective
		infected showed VE 82% (95% CI, 80 to		multiple nested emulated
		85) against re-infection compared to		target trials in Israel; 107,413
		, 0		
		previously infected and unvaccinated		previously infected
				participants; time and setting
				from VOC Alpha to VOC
				Delta (unable to separate
				results reported <1% Alpha so
				predominantly Delta)

Section 2: excluded studies			
Author	Reason for exclusion		
Abu-Raddad (3)	Vaccine effectiveness not reported		
<u>Akhrass</u>	Delayed exclusion – Clinical outcomes of interest for this LES not reported		
<u>Albahrani</u>	Prevalence of variants unknown and suspected to be <50%		
Alencar	Critical risk of bias		
Alhamlan	Vaccine effectiveness not reported		
<u>Alharbi</u>	Prevalence of variants unknown and suspected to be <50%		
Ali	Prevalence of variants unknown and suspected to be <50%		
Alkhafaji	Prevalence of variants unknown and suspected to be <50%		
Allen	Serious risk of bias		
Allen(2)	Results not reported according to vaccine type/brand		
Almufty	Prevalence of variants unknown and suspected to be <50%		
Al-Qahtani	Delayed exclusion – critical risk of bias		
Andeweg	Vaccine effectiveness not reported		
Andeweg (2)	Results not reported according to vaccine type/brand		
Apisarnthanarak	Vaccine effectiveness not reported		
<u>Arashiro</u>	Vaccine effectiveness not reported		
<u>Araujo</u>	Clinical outcomes of interest for this LES not reported		
Auvigne	Clinical outcomes of interest for this LES not reported		
Ayass	Clinical outcomes of interest for this LES not reported		
<u>Baden</u>	Critical risk of bias		
Bailly	Delayed exclusion – critical risk of bias		
<u>Bajema</u>	Clinical outcomes of interest for this LES not reported		
Bajema (2)	Clinical outcomes of interest for this LES not reported		
Bal	Vaccine effectiveness not reported		
Barchuk	Clinical outcomes of interest for this LES not reported		
Belayachi	Results not reported by variant		
Bergwerk	Vaccine effectiveness not reported		
Bernal (2)	Delayed exclusion – critical risk of bias		
Bhatnagar	Critical risk of bias		
Bhattacharya	Delayed exclusion – critical risk of bias		
Bianchi	Delayed exclusion – critical risk of bias		
Bjork	Prevalence of variants unknown and suspected to be <50%		
Blaiszik	Clinical outcomes of interest for this LES not reported		
Blaiszik	Clinical outcomes of interest for this LES not reported		
Borobia	Clinical outcomes of interest for this LES not reported		
Bosch	Clinical outcomes of interest for this LES not reported		
Britton	Prevalence of variants unknown and suspected to be <50%		
Britton (2)	Critical risk of bias		
Brown	Vaccine effectiveness not reported		
Brunelli	Prevalence of variants unknown and suspected to be <50%		

Danserroomt	Drawalance of varients unknown and suspected to be < 50%
Bruxvoort	Prevalence of variants unknown and suspected to be <50%
Butt	Prevalence of variants unknown and suspected to be <50%
Butt	Critical risk of bias
<u>Butt (2)</u>	Delayed exclusion – critical risk of bias
<u>Cabezas</u>	Prevalence of variants unknown and suspected to be <50%
<u>Caillard</u>	Clinical outcomes of interest for this LES not reported
<u>Cardona</u>	Vaccine effectiveness not reported
<u>Cavanaugh</u>	Delayed exclusion – VOI not VOC
Chadeau-Hyams(2)	Results not reported according to vaccine type/brand
<u>Chaguza</u>	Vaccine effectiveness not reported
Charles Pon Ruban	Vaccine effectiveness not reported
<u>Charmet</u>	Serious risk of bias
<u>Chau</u>	Vaccine effectiveness not reported
Christensen	Vaccine effectiveness not reported
Chung (2)	Results not reported according to vaccine type/brand
Clemens	Prevalence of variants unknown and suspected to be <50%
Cohen	Vaccine effectiveness not reported
Cohen(2)	Vaccine effectiveness not reported
Collie	Clinical outcomes of interest for this LES not reported
Corchado-Garcia	Prevalence of variants unknown and suspected to be <50%
Corrao	Results not reported according to vaccine type/brand
<u>Dash</u>	Critical risk of bias
<u>Davies</u>	Results not reported according to vaccine type/brand
de Gier Brechje	Prevalence of variants unknown and suspected to be <50%
<u>Dickerman</u>	Results reported comparison of two vaccines (no unvaccinated or early vaccinated
	groups)
<u>Dolzhikova</u>	Critical risk of bias
<u>Domi</u>	Prevalence of variants unknown and suspected to be <50%
<u>Drawz</u>	Critical risk of bias
El Sahly	Prevalence of variants unknown and suspected to be <50%
Ella	Prevalence of variants unknown and suspected to be <50%
Elliot	Delayed exclusion – critical risk of bias
<u>El-Sahly</u>	Prevalence of variants unknown and suspected to be <50%
<u>Epaulard</u>	Clinical outcomes of interest for this LES not reported
Falsey	Prevalence of variants unknown and suspected to be <50%
Fang	Modelling study
<u>Farah</u>	Clinical outcomes of interest for this LES not reported
<u>Farinholt</u>	Vaccine effectiveness not reported
<u>Ferdinands</u>	Clinical outcomes of interest for this LES not reported
<u>Fisher</u>	Prevalence of variants unknown and suspected to be <50%
Fisman (2)	Results not reported according to vaccine type/brand
Flacco	Results not reported according to vaccine type/brand
Frenck	Prevalence of variants unknown and suspected to be <50%
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<u>Furer</u>	Delayed exclusion – critical risk of bias
Gardner	Modelling study
<u>Geisen</u>	Clinical outcomes of interest for this LES not reported
<u>Gharpure</u>	Vaccine effectiveness not reported
Ghosh	Delayed exclusion – critical risk of bias
<u>Gils</u>	Clinical outcomes of interest for this LES not reported
Goga	Vaccine effectiveness not reported
Gorgels	Prevalence of variants unknown and suspected to be <50%
Grannis	Clinical outcomes of interest for this LES not reported
Gray	Prevalence of variants unknown and suspected to be <50%
<u>Gray (2)</u>	Clinical outcomes of interest for this LES not reported
Griffin	Vaccine effectiveness not reported
<u>Guijarro</u>	Prevalence of variants unknown and suspected to be <50%
Gupta	Prevalence of variants unknown and suspected to be <50%
Gupta	Vaccine effectiveness not reported
Haas (2)	Modelling study
Hacisuleyman	Critical risk of bias
<u>Harris</u>	Modelling study
<u>Herlihy</u>	Delayed exclusion – critical risk of bias
Hetemaki	Vaccine effectiveness not reported
Hitchings (3)	Vaccine effectiveness not reported
Hitchings(2)	Delayed exclusion – critical risk of bias
Hollinghurst	Serious risk of bias
<u>Hyams</u>	Delayed exclusion - Clinical outcomes of interest for this LES not reported
Iliaki	Prevalence of variants unknown and suspected to be <50%
Iliaki	Prevalence of variants unknown and suspected to be <50%
<u>Ismail</u>	Delayed exclusion - Clinical outcomes of interest for this LES not reported
<u>Jacobson</u>	Critical risk of bias
<u>John</u>	Prevalence of variants unknown and suspected to be <50%
<u>Johnson</u>	Results not reported according to vaccine type/brand
Jones	Critical risk of bias
<u>Jucker</u>	Results not reported according to vaccine type/brand
<u>Kaabi</u>	Prevalence of variants unknown and suspected to be <50%
<u>Kahn</u>	Results not reported according to vaccine type/brand
<u>Kale</u>	Delayed exclusion – critical risk of bias
Kaur	Delayed exclusion – critical risk of bias
<u>Keegan</u>	Critical risk of bias
<u>Kemp</u>	Modelling study
Khan	Prevalence of variants unknown and suspected to be <50%
<u>Khawaja</u>	Critical risk of bias
<u>Kislaya</u>	Vaccine effectiveness not reported
Kislaya (2)	Results reported comparison of two variants
<u>Kojima</u>	Prevalence of variants unknown and suspected to be <50%

Kshirsagar	Vaccine effectiveness not reported
Kustin	Delayed exclusion - only included infected population
<u>Lamprini</u>	Clinical outcomes of interest for this LES not reported
Lan	Results not reported according to vaccine type/brand
Lauring	Clinical outcomes of interest for this LES not reported
Lee	Clinical outcomes of interest for this LES not reported
Lefèvre	Critical risk of bias
<u>León</u>	Results not reported according to vaccine type/brand
Levin-Rector	Only included previously infected
Lewis	Clinical outcomes of interest for this LES not reported
Lewnard	Clinical outcomes of interest for this LES not reported
<u>Li</u>	Phase 1 trial
<u>Li (2)</u>	Clinical outcomes of interest for this LES not reported
Li (3)	Delayed exclusion – critical risk of bias
Ling	Prevalence of variants unknown and suspected to be <50%
Linsenmeyer	Vaccine effectiveness not reported
Lippi	Results not reported according to vaccine type/brand
Lippi (2)	Critical risk of bias
Liu	Vaccine effectiveness not reported
Loconsole	Vaccine effectiveness not reported
Luo	Vaccine effectiveness not reported
Lyngse (2)	Results not reported according to vaccine type/brand
Lytras	For Waning LES
Ma	Critical risk of bias
<u>Maeda</u>	Critical risk of bias
Mallow	Results not reported according to time frame: cannot separate Alpha from Delta
Marco	Delayed exclusion – critical risk of bias
<u>Marquis</u>	Vaccine effectiveness not reported
Mattar	Prevalence of variants unknown and suspected to be <50%
<u>Mattiuzzi</u>	Results not reported according to vaccine type/brand
<u>Maurya</u>	Prevalence of variants unknown and suspected to be <50%
<u>Mazgatos</u>	Critical risk of bias
<u>McEvoy</u>	Prevalence of variants unknown and suspected to be <50%
McKeigue(2)	Results not reported according to vaccine type/brand
<u>Menni</u>	Serious risk of bias
<u>Mielke</u>	Clinical outcomes of interest for this LES not reported
<u>Mirahmadizadeh</u>	Prevalence of variants unknown and suspected to be <50%
<u>Mizrahi</u>	Modelling study
Molani	Clinical outcomes of interest for this LES not reported
Monge	Prevalence of variants unknown and suspected to be <50%
Mor	Prevalence of variants unknown and suspected to be <50%
Moustsen-Helms	Prevalence of variants unknown and suspected to be <50%
Munitz	Clinical outcomes of interest for this LES not reported

Munro	Clinical outcomes of interest for this LES not reported
Murison	Results not reported according to vaccine type/brand
Musser	Vaccine effectiveness not reported
Mutnal	Vaccine effectiveness not reported
Nanduri	Critical risk of bias
Nguyen	Results not reported according to vaccine type/brand
<u>Niessen</u>	Clinical outcomes of interest for this LES not reported
Nyberg	Clinical outcomes of interest for this LES not reported
Oduwole	Clinical outcomes of interest for this LES not reported
<u>Olmedo</u>	Clinical outcomes of interest for this LES not reported
Olson	Clinical outcomes of interest for this LES not reported
Open-SAFELY	Vaccine effectiveness not reported
Ostropolets	Not reported separately according to variant
Palacios	Prevalence of variants unknown and suspected to be <50%
<u>Paredes</u>	Clinical outcomes of interest for this LES not reported
Paris	Prevalence of variants unknown and suspected to be <50%
<u>Pattni</u>	Modelling study
<u>Pawlowski</u>	Critical risk of bias
Peralta-Santos	Clinical outcomes of interest for this LES not reported
<u>Perrella</u>	Vaccine effectiveness not reported
Perry	Clinical outcomes of interest for this LES not reported
Peter	Vaccine effectiveness not reported
Peter	Vaccine effectiveness not reported
<u>Pilishvili</u>	Prevalence of variants unknown and suspected to be <50%
Piltch-Loeb	Prevalence of variants unknown and suspected to be <50%
<u>Polinski</u>	Delayed exclusion – critical risk of bias
<u>Poukka</u>	Critical risk of bias
<u>Pulliam</u>	Modelling study
Raches Ella	Phase 1 trial
<u>Rana</u>	Critical risk of bias
Regev-Yochay	Prevalence of variants unknown and suspected to be <50%
Reynolds	Results not reported according to vaccine type/brand
<u>Riemersma</u>	Clinical outcomes of interest for this LES not reported
Riley	Critical risk of bias
Rivelli	Clinical outcomes of interest for this LES not reported
Robinson	Clinical outcomes of interest for this LES not reported
Rosero-Bixby	Clinical outcomes of interest for this LES not reported
Rovida	Critical risk of bias
Rudolph	Prevalence of variants unknown and suspected to be <50%
Salmeron Rios	Prevalence of variants unknown and suspected to be <50%
Sansone	Critical risk of bias
Satwik	Delayed exclusion – critical risk of bias
Scobie	Delayed exclusion – critical risk of bias

Self	Clinical outcomes of interest for this LES not reported
Sharma	Prevalence of variants unknown and suspected to be <50%
Sheikh (3)	Results not reported according to vaccine type/brand
Shimabukuro	Clinical outcomes of interest for this LES not reported
Shrotri	Delayed exclusion – critical risk of bias
Simon	Prevalence of variants unknown and suspected to be <50%
Simsek-Yavuz	Clinical outcomes of interest for this LES not reported
Smoliga Smoliga	Critical risk of bias
<u>Starrfelt</u>	Serious risk of bias
Suri	Vaccine effectiveness not reported
Swift	Prevalence of variants unknown and suspected to be <50%
Tande	Prevalence of variants unknown and suspected to be <50%
<u>Tande</u> <u>Tanriover</u>	Prevalence of variants unknown and suspected to be <50%
	Modelling study
Taquet Tartof (3)	Clinical outcomes of interest for this LES not reported
Tenforde	-
	Clinical outcomes of interest for this LES not reported
Tenforde (2)	Clinical outcomes of interest for this LES not reported
Tenforde (3)	Clinical outcomes of interest for this LES not reported Critical risk of bias
<u>Thangaraj</u>	
<u>Thiruvengadam</u>	Critical risk of bias
Thompson (1)	Prevalence of variants unknown and suspected to be <50%
Thompson (2)	Prevalence of variants unknown and suspected to be <50%
thompson (4)	Clinical outcomes of interest for this LES not reported
Tobolowsky	Clinical outcomes of interest for this LES not reported
Ulloa	Vaccine effectiveness not reported
<u>Uschner</u>	Critical risk of bias
<u>Vahidy</u>	Prevalence of variants unknown and suspected to be <50%
Vasileiou	Clinical outcomes of interest for this LES not reported
<u>Veneti</u>	Clinical outcomes of interest for this LES not reported
<u>Victor</u>	Critical risk of bias
Volkov	Modelling study
<u>Voysey</u>	Prevalence of variants unknown and suspected to be <50%
<u>Waldhorn</u>	Serious risk of bias
Wang	Clinical outcomes of interest for this LES not reported
<u>Ward</u>	Results not reported according to vaccine type/brand
<u>Waxman</u>	Clinical outcomes of interest for this LES not reported
Wickert	Critical risk of bias
Wijtvliet	Clinical outcomes of interest for this LES not reported
Williams (2)	Critical risk of bias
Wolff	Vaccine effectiveness not reported
Woolley	Results not reported according to vaccine type/brand
Xiang	Clinical outcomes of interest for this LES not reported
Young-Xu	Prevalence of variants unknown and suspected to be <50%

Young-Xu (4)	Critical risk of bias
Zacay	Delayed exclusion – critical risk of bias
Zhong	Clinical outcomes of interest for this LES not reported

Appendix 2: Glossary

AZ: AstraZeneca

Alpha: variant of concern B.1.1.7

Beta: variant of concern B.1.351

Delta: variant of concern B.1.617.2

Gamma: variant of concern P.1

Epsilon: variant of concern B.1.427/B.1.429

HCW: Healthcare workers

LTC: Long-term care

LTCF: Long-term care facility

MOD: Moderna

Obs: observational study

Omicron: variant of concern B.1.1.529

OR: odds ratio

PF: Pfizer

RME: range of mean estimates across 2 or more studies

VE (Vaccine effectiveness): measure of how well a vaccine protects people from getting the outcome of interest in real-world practice (For example: VE of 92% against infection means that 92% of people will be protected from becoming infected with COVID and 8% of people will still be at risk of becoming infected with COVID)

VES: vaccine effectiveness against susceptibility (vaccinated contact)

VET: vaccine effectiveness against transmission (vaccinated index case)

VOC: variant of concern

VOI: variant of interest

Appendix 3: Data-extraction template

Vaccine product	
Source	First author of study
Link	DOI or Pubmed ID
Date published	in format YYYY/MM/DD or preprint
Country	
Funding	public or industry
Study details	
Study type	RCT/cohort/data-linkage/test-negative/case-control/other
Surveillance	routine screening Y or N
Population(s)	general public/LTC/Households/HCW/Other
Control group	not vaccinated, <7day vaccinated internal control, none, other
Total (N)	number of all study participants
Female	number or %
LTC	number or %
HCW	number or %
Households	number or %
>80	number or %
>70	number or %
>60	number or %
Outcomes	outcomes separated by VOC type
Outcomes	confirmed infection/asymptomatic/mild symptomatic/severe
	symptoms/hospitalized/ICU/death
1st Dose VE	VE with 95% CI
Days post 1st dose	days post 1st dose when VE provided
2nd Dose VE	VE with 95% CI
Days post 2nd	days post 2nd dose when VE provided
dose	days post 2nd dose when viz provided
Rates per X	vaccinated vs control
person-days/years	
HR	vaccinated vs control
RR	vaccinated vs control
Adjusted	Regression, stratification, matching and associated variables
Transmission	infection rates in unvaccinated contacts of vaccinated individuals
Critical appraisal	See Appendix 5

Appendix 4: Process for assigning Variant of Concern to studies

A Variant of Concern is considered to be the dominant (≥50%) strain in a study if any of the following conditions apply:

- i) the authors make a statement about prevalence of VOC during the study time frame
- ii) time and setting of the study is consistent with a VOC being dominant according to the following open tracking sources:

Nextstrain. Real-time tracking of pathogen evolution. https://nextstrain.org/ Outbreak Info. https://outbreak.info/location-reports

Appendix 5: Research question and critical appraisal process (revised 06 Oct 2021)

Review question:

Participants	People at risk of COVID-19 (usually without but sometimes with previous
	COVID-19 infection)
Intervention	COVID-19 Vaccine
Comparator	Unvaccinated people (*)
Outcomes	PCR-diagnosis of COVID-19 infection (**); symptomatic disease;
	hospital/ICU admission; death; transmission

^(*) before-after studies, where the infection rate in the first 2 weeks after the vaccination are used as control are (**)

Critical Appraisal Process

We appraise the quality of the individual studies using an adapted version of ROBINS-I. This tool classifies the Risk of Bias of a study as **Low, Moderate, Serious, Critical, or No Information**. <u>Low Risk of Bias indicates High Quality, and Critical Risk of Bias indicates Very Low (insufficient) Quality.</u> ROBINS-I appraises 7 bias domains and judges each study against an ideal reference randomized controlled trial. To improve the utility of ROBINS-I for assessing studies reporting vaccine effectiveness, we have focused on study characteristics that introduce bias as reported in the vaccine literature. (WHO. Evaluation of COVID-19 vaccine effectiveness. Interim Guidance. 17 March 2021). Studies rated as "critical" risk of bias will not be included in the Summary statements on Page 1-2 (exception: if limited data available for an outcome for a VOC). An overall judgement of "serious" or "critical" is given when the study is judged to be at critical risk of bias in at least one domain. Three of more serious risk of bias domains is given an overall risk of bias of critical.

VE Study	Description
Characteristics that	
may introduce bias	
Study design	In cohort studies, people who get vaccinated may differ in health-seeking
	behaviour from people who do not get vaccinated; using a test-negative
ROBINS-I: Bias in	study design minimizes this type of bias
selection of participants	
into study	Examples and typical judgement:
	• test-negative design with a clearly defined symptomatic study population (low)
	• test-negative design (mixed or unclear study population) or case- control or cohort design or data-linkage with no concerns (moderate)
	cross-sectional design or case-control (concerns about whether
	controls had same access to vaccines/risk of exposure to COVID or
	unclear) or cohort design (concerns that exposed and non-exposed
	were not drawn from the same population) (serious)
Method for confirming	Questionnaires are prone to recollection bias; Population databases
vaccination	developed for purpose of tracking COVID vaccines minimize this type of
	bias
ROBINS-I: Bias in	
classification of	Examples and typical judgement:
interventions	database linkage study (low)
	• Questionnaire with confirmation by an additional method (e.g. registry) of at least a subset of study population (moderate)

^(**) commonly performed and may be appraised confirmation of specific variant, or reasonable evidence the variant was the dominant circulating strain

	• Questionnaire without confirmation by an additional method (serious)
	• Estimating vaccination status based on surveillance data alone (critical)
Databases used for	Databases developed for collecting data on COVID are less prone to bias
retrieval of COVID test	due to missing information and misclassification
results, participant	
prognostic factors, and	Examples and typical judgement:
clinical outcomes	 database for non-COVID purpose but with individual level data
	(moderate)
ROBINS-I: Bias in	 database for non-COVID purpose without individual level data
classification of	(serious)
interventions	 no or unclear description of database type (critical)
Assignment of	Using date of symptom onset (if within 10 days of testing) as infection
infection start date	start date reduces risk of misclassification bias (e.g., vaccinated participant
	who is reported as COVID+ may have been infected prior to receiving
ROBINS-I: Bias in	the vaccine or during non-immune period) and sensitivity of assays
classification of	decreases over time
interventions	
	Examples and typical judgement:
	• using a PCR positive test that was part of an ongoing standardized
	monitoring system (e.g., within a health network) (low)
	• using sample date without interview or documented confirmation of
	symptoms ≤ 10 days (relevant for symptomatic disease only) (serious)
Verification of	Prospective, standardized collection of symptoms from patients reduces
symptoms	risk of missing information bias; testing within 10 days after symptom
DODING L D'	onset reduces risk of false-negative COVID test
ROBINS-I: Bias in classification of	Examples and trained in decompositi
interventions	Examples and typical judgement:
Interventions	• using sample date without patient report/ documented confirmation of
	symptoms ≤ 10 days (relevant for symptomatic disease only) (serious)
Accounting for non-	• if symptomatic COVID is not an outcome (no information) Reported absence of vaccine effect during non-immune period reduces
immune period (first 14	risk of residual confounding bias
days after first vaccine	lisk of residual confounding bias
dose)	Example/common case:
,	presence of an effect during non-immune period or result not
ROBINS-I: Bias due to	reported (moderate)
confounding	unclear that non-immune period was considered (serious)
Inclusion of	Exclusion (or separate analysis) of participants with prior COVID
participants with prior	infection reduces concern about differences in infectivity as well as risk-
COVID infection	taking and health-seeking behaviour
ROBINS-I: Bias due to	Examples and typical judgement:
confounding	• inclusion of prior infection status as a covariate in the models
	(moderate)
	previously infected not excluded or analyzed separately (serious)
Accounting for calendar	Accounting for calendar time reduces bias due to differences in vaccine
time	accessibility and risk of exposure over time
	English and trained in decreases
	Examples and typical judgement:

ROBINS-I: Bias due to confounding (time-varying	 use of time-varying statistics without explicit mention of adjustment for calendar time (moderate)
confounding)	• not taken into account but short-time frame (e.g. ≤2 months) (serious)
	• not taken into account and time frame >2 months (critical)
Adjustment for	Adjustment for prognostic factors for COVID infection, severity of
prognostic factors	disease, and vaccination, such as age, gender, race, ethnicity,
	socioeconomic factors, occupation (HCW, LTC), and chronic medical
ROBINS-I: Bias due to	conditions
confounding	
Comountaing	Examples and typical judgement:
	• no or insufficient adjustment for occupation (or number of tests as a
	surrogate for exposure risk) -exception age>65 or LTCF resident
	(moderate)
	no or insufficient adjustment for socioeconomic factors (or
	neighborhood or income as a surrogate), race, ethnicity (serious)
	• no or insufficient adjustment for age (any study population) or chronic medical conditions (LTC)(critical)
Testing frequency	Similar frequency of testing between groups reduces risk of bias
	introduced by detecting asymptomatic infection in one group but not in
ROBINS-I: Bias in	another (e.g. when only one group undergoes surveillance screening)
measurement of outcomes	
	Examples and typical judgement:
	• no systematic screening but consistent methods for detection in one
	group vs. the other, e.g., within health networks (moderate)
	screening performed for a subset of both study groups (serious)
	 screening performed routinely in one study group but not in the other (critical)

Appendix 6: Detailed description of the narrative summary statement

We include studies with the following clinical outcomes: prevention of infection, severe disease (as defined by the study investigators), death, and prevention of transmission. These outcomes were selected because they are less susceptible to bias. If data are not available for these specific outcomes, but are available for symptomatic infection and/or hospitalization, data for these additional outcomes are provided temporarily. Studies reporting only antibody responses are excluded.

We aim at providing a lay language, standardized summary statement for each combination of vaccine and VOC for which we found evidence.

Where <u>more than one study</u> was found, we will provide a summary statement with a <u>range of the estimates across the studies</u>.

Where a <u>single study</u> provided data, we will provide the <u>estimate plus 95% confidence interval</u> for that study. As additional studies are added, the estimate plus confidence interval will be replaced by a range as described above.

In the summaries, "prevented" or "protects" will be applied to mean estimates or range of mean estimates that are greater than or equal to 50%.

Section 3: Special Groups (after 5 November 2021)	
Author	Special Group
<u>Bedston</u>	Elderly >75 years
<u>Bekker</u>	Healthcare workers
Botton	Elderly >75 years
<u>Bukatko</u>	Homeless shelter residents
Butt (2)	Veterans (on Heamodialysis)
<u>Dujmovic</u>	Nursing Home residents
<u>Embi</u>	Immunocompromised
Filon	Healthcare workers
Gaio	Healthcare workers
Goldhaber-Fiebert	Prison residents and staff
Goldin	LTCF
<u>Hall (2)</u>	Healthcare workers
Helmsdal	Healthcare workers
<u>Iskander</u>	Coast guard personnel
<u>Kawasuji</u>	Healthcare workers
Krutikov	LTCF
Lustig	Healthcare workers
<u>Malhotra</u>	Healthcare workers
<u>Manteghinejad</u>	Cancer patients only
<u>McConeghy</u>	LTCF
<u>Mohr</u>	Healthcare workers
<u>Muhsen</u>	Healthcare workers
Nunes (2)	Healthcare workers
Oliver	Maintenance dialysis patients
Paixao	Pregnant women
<u>Petráš</u>	Healthcare workers
Quach	Healthcare workers
Regev-Yochay	Healthcare workers
<u>Salvatore</u>	Prison staff and prisoners
Shen	immunosuppressed patients
Shrestha (3)	Healthcare workers
<u>Smith</u>	Renal patients only
Spensley	End-stage Kidney disease patients
<u>Spitzer</u>	Healthcare workers
<u>Subbarao</u>	LTCF
Sultan	Healthcare workers
<u>Tai</u>	special population (NBA)
Yassi (2)	Healthcare workers
Young-Xu (3)	Male Veterans