COVID-END COVID-19 Evidence Network to support Decision-making ... in Canada

COVID-19 Living Evidence Synthesis #6

(Version 20: 22 September 2021)

Question

What is the efficacy and effectiveness of available COVID-19 vaccines for variants of concern?

Findings

For vaccine effectiveness in variants of concern (VOC), we present a <u>visual summary of evidence in Table 1</u> and <u>detailed statements in Table 2</u>.

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, 197 studies were appraised and 89 used to complete this summary. The reasons for excluding the remaining 108 studies are reported in the second section of Appendix 2.

13 new studies have been added since the previous edition of this living evidence synthesis, all of which are signaled by a last-updated date of 22 September 2021 (highlighted in yellow). The new studies included results for VOC Alpha¹ [B.1.1.7] (7), VOC Beta [B.1.351] (1), and VOC Delta [B.1.617.2] (10).

Pfizer/Comirnaty [BNT162b2]

We have moderate certainty evidence that 2 doses of BNT162b2 prevented infection (range of mean estimates: 70 to 97%), prevented severe

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.

¹ As of August 9, inclusion of Alpha studies may be temporarily delayed to permit resource allocation to Delta.

disease (range of mean estimates: 92 to 98%), prevented death (range of mean estimates: 91 to 99%), and reduced transmission of VOC **Alpha** to close contacts (range of mean estimates: 65 to 80%).

We have moderate certainty evidence that 2 doses of BNT162b2 prevented symptomatic infection from VOC **Beta** (range of mean estimates: 84 to 88%).

We have low certainty evidence that 2 doses of BNT162b2 prevented infection from VOC **Delta** (range of mean estimates: 42 to 80%); moderate certainty evidence it prevented symptomatic infection from VOC Delta (range of mean estimates: 62 to 94%); low certainty evidence it prevented severe, critical, or fatal disease from VOC Delta (range of mean estimates: 93 to 98%) and low certainty evidence it prevented death due to VOC Delta (90% [95% CI not reported]).

We have low certainty evidence that BNT162b2 prevented symptomatic disease from VOC **Gamma** (range of mean estimates: 84 to 88% - 2 reports from the same study population).

Moderna/Spikevax [mRNA-1273]

We have moderate certainty evidence that 2 doses of mRNA-1273 prevented infection from VOC **Alpha** (range of mean estimates: 86 to 100%) and low certainty evidence it prevented infection from VOC Beta (96.4% [95% CI, 92 to 99] – 1 Obs). We have low certainty evidence that it prevented severe, critical, or fatal disease from VOC Alpha (combined with Beta) (95.7% [95% CI, 73.4 to 99.9] – 1 Obs).

We have moderate certainty evidence that 2 doses of mRNA-1273 prevented infection from VOC **Delta** (range of mean estimates: 76 to 86%) and low certainty evidence that it prevented severe, critical, or fatal disease (range of mean estimate: 93 to 100%).

We have low certainty evidence that 2 doses of mRNA-1273 prevented symptomatic infection from VOC **Delta** (90.3% [95% CI, 67.2 to 97.1] – 1 Obs).

We have low certainty evidence that 2 doses of mRNA-1273 prevented symptomatic infection from VOC **Gamma** (88% [95% CI, 61 to 96] – 1 Obs).

AstraZeneca/Vaxzevria [ChAdOx1]

We have moderate certainty evidence that 2 doses of ChAdOx1 prevented infection from VOC **Alpha** (range of mean estimates: 62 to 79%) and moderate certainty evidence that it provided limited protection from infection by VOC Beta (10.4% [95% CI, -76.8 to 54.8]- 1 RCT).

We have low certainty evidence that 2 doses of ChAdOx1 prevented infection from VOC **Delta** (range of mean estimates: 60 to 67%) and moderate certainty evidence it prevented symptomatic infection from VOC Delta (range of mean estimates: 61 to 70%). We have low certainty evidence that 2 doses of ChAdOx1 prevented ICU admission (99.2% [95% CI, 97.6 to 99.7] – 1 Obs*) and low certainty evidence it prevented death (range of mean estimates: 97 to 99.6%).

We have low certainty evidence one dose of ChAdOx1 provided limited protection against symptomatic infection against VOC **Gamma** (48% [95% CI, 28 to 63] – 1 Obs).

^{*}combined with Alpha

Other vaccines

We have moderate certainty evidence that **Johnson & Johnson [AD26.COV2.S]** prevented severe disease from VOC **Beta** (81.7% [95% CI, 46.2 to 95.4] - 1 RCT).

We have low certainty evidence that **AD26.COV2.S** prevented infection from VOC **Delta** (67% [95% CI, 60 to 73] – 1 Obs).

We have moderate certainty evidence that 2 doses of **Novavax [NVX-Co2373]** prevented symptomatic infection from VOC **Alpha** (86.3% [95% CI, 71.3 to 93.5] - 1 RCT) and moderate certainty evidence that it prevented symptomatic infection from VOC **Beta** (43% [95% CI, -9.8 to 70.4] - 1 RCT).

We low certainty evidence that 2 doses of **Sinovac** [**CoronaVac**] prevented symptomatic infection due to VOC **Delta** (59% [95% CI, 16 to 81.6] – 1 Obs) and prevented severe infection (range of mean estimates: 89 to 100%) due to VOC **Delta**.

We have low certainty evidence that 2 doses of CoronaVac prevented infection from VOC **Gamma** (65.9% [95% CI, 65.2 to 66.6] - 1 Obs).

We have low certainty evidence that 2 doses of Sinopharm [BBIBP-CorV] prevented ICU admission (95.4% [95% CI, 94.6 to 96.2] – 1 Obs*) from VOC **Delta** and low certainty evidence it prevented death (94.3% [95% CI, 93.1 to 95.4] – 1 Obs*).

We have low certainty evidence that 2 doses of Gamaleya [Sputnik V] prevented ICU admission (100% [95% CI, 99.2 to 100] – 1 Obs*) from VOC **Delta** and low certainty evidence it prevented death (99.5% [95% CI, 98.5 to 99.9] – 1 Obs*).

Combinations of vaccines

We have low certainty evidence that 1 dose of **AstraZeneca [ChAdOx1]** followed by 1 dose of **Pfizer [BNT162b2]** or **Moderna [mRNA-1273]** prevented infection by VOC **Alpha** (88% [95% CI, 83 to 92] – 1 Obs).

Table 1: Visual summary of evidence for COVID-19 vaccines for variants of concern

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 = pooling of moderate to high quality RCTs or pooling of observational studies with low risk of bias and with consistent findings

Moderate certainty evidence = single RCT of moderate to high quality or ≥ one observational study with low to moderate risk of bias and with at least partially consistent findings

Low certainty evidence = single RCT of low quality or single observational study of any quality or multiple low or moderate observational studies with inconsistent findings

Outcome	Vaccine Effectiveness (2 doses unless otherwise stated) for			
(and vaccine)	each combination of vaccine, variant, and outcome			
	Alpha	Beta	Gamma	Delta
Any Infection				
Pfizer	70 to 97%			42 to 80%
Moderna	86 to 100%	96%		76 to 86%
AstraZeneca	62 to 79%	10%**		60 to 67%
Johnson & Johnson				67%
Novavax				
CoronaVac			66%	
AZ/PF or MOD	88%			
Symptomatic Infect	cion (reported wher	n data on "any infec	tion" is limited)	
Pfizer		84 to 88%	84 to 88%	62 to 94%
Moderna			88%	90%
AstraZeneca			48%*	61 to 70%
Johnson & Johnson				
Novavax	86%	43%**		
CoronaVac				59%
Transmission				
Pfizer	65 to 80%			
Moderna				
AstraZeneca				
Johnson & Johnson				
Novavax				
CoronaVac				
Severe Disease (ma	y include death fo	r some studies)		
Pfizer	92 to 98%			93 to 98%
Moderna	96%	96%		93 to 100%
AstraZeneca				99% ICU admit
Johnson & Johnson		82%*		
Novavax				
CoronaVac				89 to 100%
Sinopharm				95% ICU admit

Sputnik V				100% ICU admit
Outcome	Vaccine Ef	fectiveness (2 dose	es unless otherwis	e stated) for
(and vaccine)	each co	mbination of vacc	ine, variant, and o	outcome
	Alpha	Beta	Gamma	Delta
Death				
Pfizer	91 to 99%			90%
Moderna				
AstraZeneca				97 to 100%
Johnson & Johnson				
Novavax				
CoronaVac				
Sinopharm				94%
Sputnik V				99%

^{*}single dose

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

AZ, AstraZeneca; MOD, Moderna; PF, Pfizer

Table 2: Key findings about vaccine effectiveness

Vaccine	Effectiveness	Findings
Pfizer/	From COVID-NMA	Compared to placebo, vaccination with BNT162b2 reduces
BioNTech		the incidence of symptomatic cases of COVID-19 and
		probably reduces severe and critical disease substantially,
Comirnaty		although there remains uncertainty about the effect on
		mortality; it may increase the incidence of severe adverse
[BNT162b2]		events. Review of RCTs (AMSTAR 10/11); last search date
		2021-09-03; GRADE evidence profile updated on 2021-09-
		17.
		[BNT162b2 to complete vaccination scheme started with
		Astra Zeneca vaccine] Synthesis pending. Review of RCTs
		(AMSTAR 8/9); last search date 2021-09-17.
		[BNT162b2 to complete vaccination scheme started with
		Astra Zeneca at 28 days vs two doses Astra Zeneca
		separated by 28 days Compared to vaccination with Astra
		Zeneca vaccine, having a second dose of BNT16b2 after a
		first dose of Astra Zeneca may not increase the risk of any
		adverse event, while the incidence of serious adverse events
		is uncertain. Review of RCTs (AMSTAR 10/11); last search
		date 2021-09-17; GRADE evidence profile updated on 2021-
		07-19
	By variant of concern	
	• Alpha	BNT162b2 provided protection against VOC Alpha for the
		following outcomes 14 days after 1 st dose:
		• 46 to 78% from infection (RME)
		BNT162b2 provided protection against VOC Alpha for the
		following outcomes 42 to 49 days after at least one dose:
		• 93% (95% CI, 89 to 96) from death
		BNT162b2 provided protection against VOC Alpha for the
		following outcomes at least 7 days after 2 nd dose:
		• 70 to 97% from infection (RME)
		• 92 to 98% from severe disease (RME)
		• 91 to 99% from death (RME)
		Unvaccinated vs BNT162b2 at least 14 days after 2 nd dose
		against VOC Alpha for severe disease:
		• HR 2.84 (95% CI, 1.80 to 4.47)
		(24 Obs)
		[1][2][3][8][9][10][15][21][22][23][28][31][34][36][37]*[41][43]
		[44]* [53][60][74][75][79][80][86][88]; last update 2021-09-22
	• Alpha, VE over	BNT162b2 provided protection against symptomatic
	time	infection by VOC Alpha when the 2 nd dose was given the
		following number of days after 1st dose:
		• 77% (95% CI, 66 to 85) at 19-29 days (age 65 to 79)
		• 86% (95% CI, 70 to 94) at 85+ days (age 65 to 79)
		(1 Obs) [79]; last update 2021-09-22

Vaccine	Effectiveness	Findings
	• Beta	BNT162b2 provided protection against VOC Beta (or
		Gamma) for the following outcomes 35-41 days after 1st
		dose:
		• 43% (95% CI, 22 to 59) from symptomatic infection
		BNT162b2 provided protection against VOC Beta (or
		Gamma) for the following outcome 7 days after 2 nd dose:
		• 84 to 88% from symptomatic infection (RME)
		• 95% (95% CI, 81 to 99) from hospitalization
		BNT162b2 provided protection against VOC Beta for the
		following outcomes \geq 14 days after 2 nd dose:
		• 75% (95% CI, 70.5 to 78.9) from infection
		• 100% (95% CI, 73.7 to 100) from severe, critical, or fatal disease
		(2 Obs – 3 refs)[23][36][47]; last update 2021-07-14
	• Delta	BNT162b2 provided protection against VOC Delta for the
		following outcome at least 14 to 21 days after 1st dose:
		• 30 to 65% from infection (RME)
		• 33 to 47.5% from symptomatic infection (RME)
		• 87 to 94% from hospitalization (RME)
		• 100% (95% CI not reported) from 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 80% from infection (RME)
		• 62 to 93.7% from symptomatic infection (RME)
		• 96% (95% CI, 86 to 99) from hospitalization
		• 93 to 98% from severe, critical, or fatal disease (RME)
		• 90% (95% CI not reported) from death
		(15 Obs)
		[29][38][42][47][57][63][64][65][71][74][76][84][86][88][92];
		last update 2021-09-22
	Delta, VE over time	BNT162b2 showed a higher risk of infection by VOC Delta
		in participants <u>fully vaccinated</u> (≥14 days after 2 nd dose)
		longer than or equal to 146 days ago vs fully vaccinated less
		than 146 days ago [OR 2.06 (95% CI, 1.69 to 2.51)]
		(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:
		• 93% (95% CI, 85 to 87) at 7 to 30 days
		• 53% (95% CI, 39 to 65) at ≥127 days
		BNT162b2 provided protection against infection by VOC
		Delta 5 months after 2 nd dose:
		• 50% (95% CI, 45 to 55) - age 16 to 39
		• 58% (95% CI, 54 to 62) - age 40 to 59
		• 57% (95% CI, 52 to 62) - age 60+
		BNT162b2 provided protection against symptomatic
		infection by VOC Delta for the following number of days
		after 2 nd dose:

Vaccine	Effectiveness	Findings
		• 62.7% (95% CI, 61.7 to 63.8) – at 1 week
		• 47.3% (95% CI, 45 to 49.6) – at 20+ weeks
		BNT162b2 provided protection against severe, critical, or
		fatal disease by VOC Delta 5 months after 2 nd dose:
		• 94% (95% CI, 87 to 97) - age 40 to 59
		• 86% (95% CI, 82 to 90) - age 60+
		(2 Obs) [76][<u>84][92];</u> last update <mark>2021-09-22</mark>
	Delta, prior	BNT162b2 (2 doses) provided protection against VOC
	infection	Delta for the following outcomes:
		• OR 13.06 (95% CI, 8.08 to 21.11) against infection
		compared to previously infected (unvaccinated)
		• OR 27.02 (95% CI, 12.7 to 57.5) against symptomatic
		infection compared to previously infected (unvaccinated)
		(1 Obs) [73]; last update 2021-09-02
	• Delta, 3 doses	BNT162b2 (3 doses) provided protection against infection
	Derta, 5 doses	by VOC Delta compared to 2 doses:
		• 3% (95% CI, -5 to 10) – at 0 to 6 days after 3rd dose
		• 84.0% (95% CI, 79 to 88) – at 14 to 20 days after 3rd
		dose
		(1 Obs) [<u>93</u>]; last update <mark>2021-09-22</mark>
	• Gamma	BNT162b2 provided protection against VOC Gamma (or
	Valillia	Beta) for the following outcomes 35-41 days after 1st dose:
		• 43% (95% CI, 22 to 59) from symptomatic infection
		BNT162b2 provided protection against VOC Gamma (or
		Beta) for the following outcome 7 days after 2 nd dose:
		,
		• 84 to 88% from symptomatic infection (RME)
		• 95% (95% CI, 81 to 99) from hospitalization
	"	(1 Obs – 2 refs)[23][47]; last update 2021-07-14
	• Epsilon	BNT162b2 provided protection against VOC Epsilon for
		the following outcome 15 days after 1 st dose:
		• 58.9% (95% CI, -9.7 to 84.5) from infection
		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
		(2 Obs) [8][31]; last update 2021-06-08
	By special population	Divitivo de la constanta de la
	• HCW, Alpha	BNT162b2 provided protection against VOC Alpha for the
		following outcomes 14 to 21 days after 1 st dose:
		• 64 to 84% from infection (RME)
		BNT162b2 provided protection against VOC Alpha for the
		following outcomes at least 7 days after 2 nd dose:
		• 80 to 96% 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
		[<u>25</u>]
		BNT162b2 provided protection against infection by VOC
		Alpha for the following number of days after 2 nd dose:

Vaccine	Effectiveness	Findings
		• 85% (95% CI, 68 to 93) at 14 to 119 days
		• 73% (95% CI, 49 to 86) ≥150 days
		(7 Obs)[<u>11][26][32][45][46][56][81];</u> last update <mark>2021-09-22</mark>
	• Over 65 years,	BNT162b2 provided protection against VOC Alpha for the
	requiring support at	following outcomes 7 days after 2 nd dose:
	home, Alpha	• 86% (95% CI, 78 to 91) from infection
		• 97% (95% CI, 88 to 99) from death
		(1 Obs)[<u>32</u>]; last update 2021-07-07
	• Over 70 years,	BNT162b2 provided protection against VOC Alpha for the
	Alpha	following outcomes at least 21 days after 1st dose:
		• 41 to 67% from infection (RME)
		BNT162b2 provided protection against VOC Alpha for the
		following outcomes at least 7 days after 2 nd dose:
		• 75 to 90% from infection (RME)
		• 69% (95% CI, 31 to 86) from symptomatic infection
	0.00	(3 Obs)[28][35][51]; last update 2021-09-22
	• Over 80 years,	BNT162b2 provided protection against VOC Alpha for the
	Alpha	following outcomes at least 14 days after 1 st dose:
		• 42 to 55.2% from infection (RME)
		• 71 to 81% from hospitalization (RME) BNT162b2 provided protection against VOC Alpha for the
		following outcomes >14 days after 2 nd dose:
		• 94% (95% CI, 73 to 99) from symptomatic infection
		• 93% (95% CI, 89 to 95) from hospitalization
		• 81% (95% CI, 74 to 87) from death
		BNT162b2 provided protection against death by VOC
		Alpha for the following number of days after 2 nd dose:
		86% (95% CI, 68 to 93) at 14 to 41 days
		• 74% (95% CI, 60 to 83) ≥98 days
		(5 Obs)[13][20][55][79][83]; last update 2021-09-22
	• LTC, Alpha	BNT162b2 provided protection against VOC Alpha for the
	, 1	following outcomes 35-48 days after 1st dose:
		• 65% (95% CI, 29 to 83) from infection
		BNT162b2 provided protection against VOC Alpha for the
		following outcomes 7 days after 2 nd dose:
		• 53% (95% CI, 29 to 69) from infection
		• 89% (95% CI, 81 to 93) from death
		(2 Obs)[<u>12][32];</u> last update 2021-07-07
	Pregnant, Alpha	BNT162b2 provided protection against VOC Alpha for the
		following outcomes at least 28 days after 1st dose:
		• 78% (95% CI, 57 to 89) from infection
		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][54</u>]; last update 2021-07-28

Vaccine	Effectiveness	Findings
	• Previously infected,	BNT162b2 (2 doses) after prior infection provided
	Alpha or Beta	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
	• Immunosuppressed,	BNT162b2 or mRNA-1273 provided protection against
	renal transplant,	infection by VOC Alpha or Beta at the following number of
	Alpha or Beta	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
		BNT162b2 or mRNA-1273 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, 33.7 to 96.2) ≥56 days
		(1 Obs) [90]; last update 2021-09-22
	• Over 70 years,	BNT162b2 provided protection against VOC Gamma for
	Gamma	the following outcomes ≥ 21 days after 1 st dose:
	Gamma	• 61% (95% CI, 45 to 72) from infection
		(1 Obs)[<u>35</u>]; last update 2021-07-07
	• HCW, Delta	BNT162b2 provided protection against VOC Delta for the
		following outcomes \geq 14 days after 2 nd dose:
		• 66% (95% CI, 26 to 84)
		(1 Obs) [<u>81</u>]; last update <u>2021-09-22</u>
	• HCW, Beta or	BNT162b2 provided protection against VOC Beta or
	Gamma	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
	. ITC D	(1 Obs)[27]; last update 2021-06-01
	• LTC, Beta	BNT162b2 provided protection against VOC Beta for the following outcome >28 days after 2 doses:
		• 50% (95% CI, 34 to 73) from infection
		(1 Obs)[24]; last update 2021-06-01
	LTC, Gamma	BNT162b2 (or mRNA-1273) provided protection against
	(residents)	VOC Gamma 14 days after 2 nd dose:
	(residents)	• 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
	Transmission	
	Household of	BNT162b2 reduced transmission of VOC Alpha from a
	vaccinated	vaccinated index case (14 to 21 days after 1st dose) to
	individual, Alpha	

Vaccine	Effectiveness	Findings
		household contacts compared to households of
		unvaccinated index cases:
		• 30 to 49% from infection (RME)
		BNT162b2 reduced transmission of VOC Alpha from a
		vaccinated HCW (10 weeks after 1st dose) to household
		spouse:
		• 42.9% (95% CI, 22.3 to 58.1) from infection
		(3 Obs) [6][14][33]; last update 2021-07-07
	 Vaccinated close 	BNT162b2 reduced transmission to close contacts
	contacts of	COVID+ index cases at least 7 to 14 days after 2 nd dose:
	COVID+, Alpha	• 65 to 80% from infection (RME)
		• 94% (95% CI, 60 to 99) from hospitalization
		(2 Obs)[<u>40</u>][<u>48</u>]; last update 2021-07-14
	 Vaccinated HCW vs 	BNT162b2 reduced transmission of VOC Beta or Gamma
	unvaccinated	from vaccinated HCW compared to unvaccinated
	community, Beta	community ≥14 days after 1 st dose:
	and Gamma	• 54.7% (95% CI, 44.8 to 62.9) from infection
		BNT162b2 reduced transmission of VOC Beta or Gamma
		from vaccinated HCW compared to unvaccinated
		community ≥ 7 days after 2^{nd} dose:
		• 84.8% (95% CI, 75.2 to 90.7) from infection
		(1 Obs) [27]; last update 2021-06-08
Moderna	From COVID-NMA	Compared to placebo, vaccination with mRNA-1723
0.11		probably reduces the incidence of symptomatic cases of
Spikevax		COVID-19 substantially and it may reduce severe disease.
[DNIA 1702]		while the incidence of serious adverse events is probably not
[mRNA-1723]		increased. Review of RCTs (AMSTAR 10/11); last search
		date 2021-09-17; GRADE evidence profile updated on 2021-01-25
	By variant of concern	01-23
	Alpha	mRNA-1273 provided protection against VOC Alpha for
	тирна	the following outcomes 14-41 days after 1 st dose:
		• 58.9 to 88.1% from infection (RME)
		• 60 to 61% from symptomatic infection (RME)
		• 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 to 15 days after 2 nd dose:
		• 86 to 100% from infection (RME)
		• 90 to 95.7% from symptomatic infection (RME)
		• 93% (95% CI not reported) from death
		• 95.7% (95% CI, 73.4 to 99.9) from severe, critical, or
		fatal disease (combined with Beta)
		(9 Obs – 10 refs) [8][23][31][34][37][47][50][60][74][86]; last
		update 2021-09-22
	• Beta	mRNA-1273 provided protection against VOC Beta for the
		following outcomes 14 days after 1 st dose:
		• 61.3% (95% CI, 56.5 to 65.5) from infection

Vaccine	Effectiveness	Findings
		• 77% (95% CI, 63 to 86) from symptomatic infection
		• 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
	Delta	mRNA-1273 provided protection against VOC Delta for
	• Delta	the following outcomes at least 14 days after 1 st dose:
		• 75 to 80% from infection (RME)
		` '
		• 72% (95% CI, 57 to 82) from symptomatic infection
		• 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:
		• 76 to 86% from infection (RME)
		• 93 to 100% from severe, critical or fatal disease (RME)
		(6 Obs) [47][57][63][64][71][74]; last update 2021-09-22
	• Delta, VE over time	mRNA-1273 provided protection against symptomatic
		infection by VOC Delta the following number of days after
		2 nd dose:
		• 95.2% (95% CI, 94.4 to 95.9) – at 1 week
		• 90.3% (95% CI, 67.2 to 97.1) – at 10 to 14 weeks
		(1 Obs) [92]; last update 2021-09-22
	• Gamma	mRNA-1273 provided protection against VOC Gamma for
		the following outcomes 14 days after 1 st dose:
		• 77% (95% CI, 63 to 86) from symptomatic infection
		• 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 (or
		Beta) for the following outcome 7 days after 2 nd dose:
		• 88% (95% CI, 61 to 96) from symptomatic infection
	Г 1	(1 Obs – 2 refs) [23] [47]; last update 2021-07-07
	Epsilon	mRNA-1273 provided protection against VOC Epsilon for
		the following outcome 15 days after 1 st dose:
		• 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

Vaccine	Effectiveness	Findings
		(2 Obs) [8][31]; last update 2021-06-08
	Special population	
	• Over 70 years,	mRNA-1273 provided protection against VOC Alpha for
	Alpha	the following outcome ≥21 days after 1 st dose:
		• 67% (95% CI, 57 to 75) from infection
		(1 Obs) [<u>35</u>]; last update 2021-06-23
	 Previously infected, 	mRNA-1273 (2 doses) after prior infection did not offer
	Alpha or Beta	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]; last update 2021-08-25
	 Immunosuppressed, 	mRNA-1273 or BNT162b2 provided protection against
	<mark>renal transplant,</mark>	infection by VOC Alpha or Beta at the following number of
	Alpha or Beta	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) \geq 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) [<u>90</u>]; last update <u>2021-09-22</u>
	• Over 70 years,	mRNA-1273 provided protection against VOC Gamma for
	Gamma	the following outcome ≥21 days after 1 st dose:
		• 61% (95% CI, 45 to 72) from infection
	TTC C	(1 Obs) [35]; last update 2021-06-23
	• LTC, Gamma	mRNA-1273 (or BNT162b2) provided protection against
	(residents)	VOC Gamma for the following outcomes 14 days after 2 nd dose:
		• 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
	Transmission	(1 000) [<u>01]</u> ; wit upunt 2021-00-11
	Household of	mRNA-1273 reduced transmission of VOC Alpha from a
	vaccinated	vaccinated HCW (10 weeks after 1 st dose) to household
	individual, Alpha	spouse:
	1110111100001, 11111111	• 42.9% (95% CI, 22.3 to 58.1) from infection
		(1 Obs)[<u>33</u>]; last update 2021-07-07
AstraZeneca	From COVID-NMA	Compared to vaccinating with MedACWY (meningitis
[ChAd0x1]		vaccine), vaccination with ChAd0x1 probably reduces the
1		cases of symptomatic COVID-19 infection. The effects on
Vaxzevria		severe or critical disease and mortality are uncertain.
		(*)Review of RCTs (AMSTAR 10/11); last search date 2021-
Serum Institute		09-17; GRADE evidence profile updated on 2021-01-25.
of India		(*) Rare cases of serious blood clots associated with a low

Vaccine	Effectiveness	Findings
[Covishield]		platelet count known as vaccine-induced thrombotic
		thrombocytopenia (VITT or VIPIT) have been reported.
		The frequency of VITT varies by age and country.
		AstraZeneca to complete vaccination scheme started with
		BNT16b2 at 28 days vs two doses of BNT16b2 separated
		by 28 days] Compared to vaccination with BNT16b2
		vaccine, having a second dose of AstraZeneca after a first
		dose of BNT 16b2 may increase the risk of any adverse
		event, while the incidence of serious adverse events is
		uncertain. Review of RCTs (AMSTAR 10/11); last search date
		2021-09-17; GRADE evidence profile updated on 2021-07-
	D . · · · C	19
	By variant of concern	Ch AdOv1 provided protection assigns VOC Alpha for the
	• Alpha	ChAdOx1 provided protection against VOC Alpha for the following outcome 14 days after 1 st dose:
		• 64% (95% CI, 60 to 68) from symptomatic infection
		• 85% (95% CI, 81 to 88) from hospitalization
		ChAdOx1nCoV-19 provided protection against VOC Alpha
		for the following outcome 21 to 28 days after 1st dose:
		• 44 to 74% from infection (RME)
		ChAdOx1provided protection against confirmed VOC
		Alpha for the following outcome at least 14 days after 2
		doses:
		• 62 to 79% from infection (RME)
		(1 RCT, moderate quality; 5 Obs)[9][10][5][47][70][71][]; last
		update 2021-08-25
	• Alpha, VE over	ChAdOx1 provided protection against symptomatic
	<mark>time</mark>	infection by VOC Alpha when the 2 nd dose was given the following number of days after 1 st dose:
		• 66% (95% CI, 47 to 77) at 19-29 days (age 65 to 79)
		• 73% (95% CI, 56 to 83) at 85+ days (age 65 to 79)
		(1 Obs) [79]; last update 2021-09-22
	• Beta	ChAdOx1 provided protection against VOC Beta for the
		following outcome 14 days after 1 st dose:
		• 48% (95% CI, 28 to 63) from symptomatic infection
		• 83% (95% CI, 66 to 92) from hospitalization
		ChAdOx1 provided protection against VOC Beta for the
		following outcome after 2 doses:
		• 10.4% (95% CI, -76.8 to 54.8) from mild to moderate
		disease (1 PCT producets quality 1 Obs) [4][47]; left ut data 2021 07
		(1 RCT, moderate quality; 1 Obs) [4][47]; last update 2021-07-07
	• Delta	ChAdOx1 provided protection against VOC Delta for the
		following outcome at least 21 days after 1st dose:
		• 18 to 49% from infection (RME)
		• 33 to 58% from symptomatic infection (RME)
		• 71% (95% CI, 51 to 83) from hospitalization

Vaccine	Effectiveness	Findings
		• 69% (95% CI, -160 to 97) from death
		ChAdOx1 provided protection against VOC Delta for the
		following outcome 14 to 21 days after 2 nd dose:
		• 60 to 67% from infection (RME)
		• 28 to 67% from symptomatic infection (RME)
		• 99.2% (95% CI, 97.6 to 99.7) from ICU admission*
		• 92% (95% CI, 75 to 97) from hospitalization
		• 97 to 99.6% from death (RME)
		ChAdOx1 provided protection against VOC Delta for the
		following outcome after 2 doses compared to one dose
		(uncertain timing):
		• 87% (95% CI, 33 to 97) from severe disease
		(9 Obs) [29][38][42][47][58][65][71][75][87]; last update 2021-
		09-22
	D.L. III.	*combined with VOC Alpha
	• Delta, VE over time	ChAdOx1 provided protection against symptomatic
		infection by VOC Delta the following number of days after
		2 nd dose:
		• 92.4% (95% CI, 92.1 to 92.7) – at 1 week
		• 69.7% (95% CI, 68.7 to 70.5) – at 20 weeks
	0	(1 Obs) [92]; last update 2021-09-22
	• Gamma	ChAdOx1nCoV-19 provided protection against VOC
		Gamma for the following outcome 14 days after 1 st dose:
		• 48% (95% CI, 28 to 63) from symptomatic infection
		• 83% (95% CI, 66 to 92) from hospitalization
	- T 1	(1 Obs)[47]; last update 2021-07-07
	• Epsilon	no data
	Special populations	Ch AdOv I provided protection accions VOC Alpha for the
	• HCW, Alpha	ChAdOx1provided protection against VOC Alpha for the following outcomes at least 14 days after 1 st dose:
		,
		• 64% (95% CI, 50 to 74) from infection ChAdOx1provided protection against VOC Alpha for the
		following outcomes at least 14 days after 2 nd dose:
		• 90% (95% CI, 62 to 98) from infection
		(1 Obs) [46]; last update 2021-07-07
	• Over 70 years,	ChAdOx1 provided protection against VOC Alpha for the
	Alpha	following outcomes 28 days after 1 st dose:
	11171111	• 55% (95% CI, 41 to 66) from death
		(1 Obs) [21]; last update 2021-07-07
	• Over 80 years,	ChAdOx1 provided protection against VOC Alpha for the
	Alpha	following outcomes at least 14 days after 1 st dose:
	F **	• 73 to 80% from hospitalization (RME)
		• 42% (95% CI, 29 to 53) from infection
		ChAdOx1provided protection against VOC Alpha for the
		following outcomes at least 14 days after 2 nd dose:
		88% (95% CI, 48 to 97) from symptomatic infection
		(3 Obs) [13][20][79]; last update 2021-09-22

Vaccine	Effectiveness	Findings
	• LTC, Alpha	ChAdOx1 provided protection against VOC Alpha for the following outcomes 35-48 days after 1 st dose:
		• 68% (95% CI, 34 to 85) from infection
		(1 Obs)[12]; last update 2021-07-07
	Prison, Alpha	ChAdOx1 did not provide protection against VOC Alpha
		for the following outcome 21-23 days after 1st dose:
		• 23% (95% CI, not reported) against infection
		(1 Obs) [<u>67</u>]; last update 2021-08-18
	HCW, Delta	ChAdOx1 provided protection against VOC Delta for the
		following outcomes at least 14 days after 2nd dose:
		• 54 to 85% from infection (RME)
		• 64% (95% CI, 38 to 78) from symptomatic infection (3 Obs) [59][66][68][]; <i>last update</i> 2021-08-25
	Over 60 years,	ChAdOx1 provided protection against VOC Alpha for the
	Gamma	following outcomes at least 28 days after 1 st dose:
	U 1122222111	• 33.4% (95% CI, 26.4 to 39.7) from symptomatic
		infection (lower than minimal acceptable protective
		effect per WHO)
		• 50.9% (95% CI, 33.6 to 63.8) from ICU admission
		• 61.8% (95% CI, 48.9 to 71.4) from death
		ChAdOx1 provided protection against VOC Alpha for the
		following outcomes at least 14 days after 2 nd dose:
		• 77.9% (95% CI, 69.2 to 84.2) from symptomatic infection
		• 89.9% (95% CI, 70.9 to 96.5) from ICU admission
		• 93.6% (95% CI, 81.9 to 97.7) from death
		(1 Obs) [62]; last update 2021-08-11
	Transmission	(1 0 00) (1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
	Household of	ChAdOx1nCoV-19 reduced transmission of VOC Alpha
	vaccinated	from a vaccinated index case (14 to 21 days after 1st dose) to
	individual, Alpha	household contacts compared to households of
		unvaccinated index cases:
		• 30 to 47% from infection (RME)
	T	(2 Obs) [6] [14]; last update 2021-06-08
	Vaccinated close contacts of	ChAdOx1nCoV-19 reduced transmission to close contacts COVID+ index cases at least 14 days after 2 nd dose:
	contacts of COVID+, Alpha	• 44% (95% CI, 31 to 54) from infection
	, Tupita	• 92% (95% CI, 46 to 99) from hospitalization
		(1 Obs)[40]; last update 2021-06-23
Johnson &	From COVID-NMA	[Johnson & Johnson's Janssen vaccine] Vaccination with
Johnson		AD26.COV2.S probably reduces the incidence of
[AD26.COV2.S]		symptomatic cases of COVID-19 by around 67%, and it
		probably reduces severe disease and mortality, while the
		incidence of serious adverse events may not increase.
		Review of RCTs (AMSTAR 10/11); <i>last search update</i> 2021-09-17. GRADE evidence profile updated on 2021-05-28
		07-17. GRADE evidence profile updated on 2021-05-28
	1	L

Vaccine	Effectiveness	Findings
		Interim summary, provided by VOC-study group: Ad26.COV2.S VE in ~40,000 randomized subjects was 66.9%; adjusted (95% CI, 59.0 to 73.4) at 14 days and 66.1% (95% CI, 55.0 to 74.8) at 28 days. For severe cases VE was 76.7% (95% CI, 54.6 to 89.1) at ≥14 days and 85.4% (95% CI, 54.2 to 96.9) at ≥28 days). (1 RCT, moderate quality of evidence) [Z] Rare cases of serious blood clots associated with a low platelet count known as vaccine-induced thrombotic thrombocytopenia (VITT, VIPIT) have been reported. The frequency of VITT varies by age and country. (data not systematically reviewed); <i>last update 2021-05-17</i>
	By variant of concern	systematically reviewed), itsi uputit 2021-05-17
	Alpha	no data
	Beta	VE against VOC 20H/501Y.V2 variant (Beta) was 52.0% and 64.0% at 14 days and 28 days for moderate, and 73.1% and 81.7% for severe cases. (1 RCT) [7]; last update 2021-04-22
	• Delta	Ad26.COV2.S provided protection against VOC Delta for the following outcomes an unknown number of days after dose: • 67% (95% 60 to 73) against infection (1 Obs)[89]; last update 2021-09-22
	• Gamma	no data
	Epsilon	no data
Sinovac [CoronaVac]	• Overall	[Coronavac vaccine] Compared to placebo, vaccination with CoronaVac may reduce the incidence of symptomatic cases of COVID-19 by 50%, close to the lowest level deemed effective by the WHO and it may substantially reduce the incidence of severe disease due to COVID-19; the evidence for any difference in serious adverse events is uncertain, although the vaccination probably increases the incidence of any adverse event. Review of RCTs (AMSTAR 10/11); last search date 2021-09-17; GRADE evidence profile updated 2021-06-25
	By variant of concern	
	• Delta	CoronaVac provided protection against VOC Delta for the following outcome ≥ 14 days after 2 nd dose: • 59% (95% CI, 16 to 81.6) from symptomatic infection • 89 to 100% from severe infection (RME) (2 Obs) [85] [91]; last update 2021-09-22
	• Gamma	CoronaVac provided protection against VOC Gamma for the following 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 ≥ 14 days after 2 nd dose for people over age 70:

Vaccine	Effectiveness	Findings
		• 41.6% (95% CI, 26.9 to 63.3) from symptomatic
		infection
		(2 Obs) [30] [49]; last update 2021-07-14
	• Epsilon	no data
	By special population	
	HCW, Gamma	Corona Vac provided protection against VOC Gamma for
		the following outcomes ≥14 days after 1 st dose:
		• 35.1% (95% CI, -6.6 to 60.5) from infection
		• 49.6% (95% CI, 11.3 to 71.4) from symptomatic
		infection (4 Ob) [18], but an data 2021, 05, 07
C:	E COMP	(1 Obs)[18]; last update 2021-05-07
Sinopharm (Wuhan)	From COVID- NMA	[Sinopharm - strain HBO2] Vaccination with Sinopharm HBO2 probably reduces the incidence of symptomatic cases
[WIV04]	INIVLA	of COVID-19, and it may reduce severe disease, while the
[WIVOT]		incidence of adverse events is probably not increased.
Sinopharm		Review of RCTs (AMSTAR 10/11); last search date 2021-09-
(Beijing)		17. GRADE evidence profile updated on 2021-06-11
[HBO2]		
[BBIBP-CorV]		[Sinopharm - strain WIV04] Vaccination with Sinopharm
		WIV04 probably reduces the incidence of symptomatic
		cases of COVID-19, and it may reduce severe disease, while
		the incidence of adverse events is probably not increased.
		Review of RCTs (AMSTAR 10/11); last search date 2021-09-
		17. GRADE evidence profile updated on 2021-06-11
	• Delta	BBIBP-CorV provided protection against VOC Delta for
		the following outcomes ≥14 days after 2 nd dose:
		• 95.4% (95% CI, 94.6 to 96.2) against ICU admission*
		• 94.3% (95% CI, 93.1 to 95.4) against death* (1 Obs) [75]; <i>last update</i> 2021-09-02
		*combined with VOC Alpha
Novavax	From COVID-	[Novavax vaccine] The effects of vaccination against
[NVX-	NMA	COVID-19 with the Novavax vaccine are currently
CoV2373]		uncertain; it probably slightly increase the risk of any
		adverse events Review of RCTs (AMSTAR 10/11); last
		search date 2021-09-17; GRADE evidence profile updated
	D	on 2021-07-01
	By variant of concern	NVV CoV2272 provided protection assignt VOC Alpha for
	• Alpha	NVX-CoV2373 provided protection against VOC Alpha for the following outcome after 2 doses:
		 89.7% (95% CI, 80.2 to 94.6) from 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
	• Beta	NVX-CoV2373 provided protection against VOC Beta for
	2000	the following outcome after 7 days after 2 nd dose:
		• Post-hoc: 43% (95% CI, -9.8 to 70.4) from symptomatic
		infection

Vaccine	Effectiveness	Findings
		(1 RCT, moderate quality), [17]; last update 2021-07-14
FBRI [EpiVacCorona]	From COVID- NMA	[EpiVacCorona] The effects of using vaccination with EpiVacCorona are uncertain. Review of RCTs (AMSTAR 10/11); last search date 2021-09-17; GRADE evidence profile updated on 2021-06-11
Bharat Biotech [Covaxin]	From COVID- NMA	[COVAXIN] Vaccination with BBV152 probably reduces the incidence of symptomatic cases of COVID-19, and it may reduce severe disease, while the incidence of serious adverse events is probably not increased. Review of RCTs (AMSTAR 10/11); last search date 2021-09-17. GRADE evidence profile updated on 2021-07-29.
	By special population	
Gamaleya [Sputnik V]	HCW, Delta	Covaxin provided protection against VOC Delta for the following outcomes ≥14 days after 2 nd dose: • 83% (95% CI, 73 to 89) from symptomatic infection • 93% (95% CI, 64 to 99) from ICU admission or death (1 Obs); [82]; last update 2021-09-22
[Gam-COVID- Vac]		
	• Delta	Gam-COVID-Vac provided protection against VOC Delta for the following outcomes ≥14 days after 2 nd dose: • 100% (95% CI, 99.2 to 100) against ICU admission* • 99.5% (95% CI, 98.5 to 99.9) against death* (1 Obs) [75]; last update 2021-09-02 *combined with VOC Alpha
Combinations	of Vaccines	
AstraZeneca followed by Pfizer or Moderna	• Alpha	First dose ChAdOx1 followed by second dose BNT162b2 or mRNA-1273 (≥ 14 days) provided protection against VOC Alpha for the following outcomes: • 88% (95% CI, 83 to 92) against infection (1 Obs) [70]; last search date 2021-08-25

^{*}delayed exclusion (see Section 2: excluded studies for reason)

Links to references are provided in Appendix 1

Pan American Health Organization/World Health Organization. Pharmacovigilance for COVID-19 Vaccines. https://covid-19pharmacovigilance.paho.org

Iorio A, Little J, Linkins L, Abdelkader W, Bennett D, Lavis JN. COVID-19 living evidence synthesis #6 (version 6.20): What is the efficacy and effectiveness of available COVID-19 vaccines in general and specifically for variants of concern? Hamilton: Health Information Research Unit, 22 September 2021.

The COVID-19 Evidence Network to support Decision-making (COVID-END) is supported by an investment from the Government of Canada through the Canadian Institutes of Health Research (CIHR). 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 opinions, results, and conclusions are those of the evidence-synthesis team that prepared the rapid response, and are independent of the Government of Canada and CIHR. No endorsement by the Government of Canada or CIHR is intended or should be inferred.

Appendix 1: Reference list

	Section 1: included studies					
Ref	Author	Bottom line	ROBINS-I*	Design, Notes		
		*Note: ROBINS-I score risk of bias: Low r	isk of bias indica	tes 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.	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	<u>Haas</u>	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.	Moderate	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	Kustin	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 52.0% (95% CI, 30.3 to 67.4) at 14 days and VE 64.0% (95% CI, 41.2 to 78.7) at 28 days against moderate to severe disease and VE 81.7% (95% CI, 46.2 to 95.4) at 28 days against severe disease (VOC Beta in South Africa).	Moderate quality (RCT)	RCT; over 40,000 participants; Argentina, Brazil, Chile, Colombia, Mexico, Peru, South Africa, and the United States; 86 of 91 cases sequenced for VOC Beta.		

8	Andrejko	BNT162b2 or mRNA-1273 showed VE 58.9% (95% CI, -9.7 to 84.5) at 15 days after 1 st dose, and VE 85.7% (95% CI, 67.2 to 93.9) 15 days after 2 nd dose against infection.	Moderate	Test-negativease-positive random sampling matched control study in California; 645 participants; 69% of population at time had VOC Alpha or Epsilon.
9	Glampson	ChAdOx1nCoV-19 showed VE 74% (95% CI, 65 to 81) against infection 28 days after 1 st dose. BNT162b2 showed VE 78% (95% CI, 73 to 82) against infection 28 days after 1 st dose.	Moderate	Retrospective cohort in UK; 2M participants; 389,587 vaccinated (58% Pfizer, 42 AZ); time and setting for VOC Alpha.
10	Pritchard	ChAdOx1nCoV-19 or BNT162b2 showed VE 66% (95% CI, 59 to 72%) 21 days after 1 st dose and 78% (95% CI, 68 to 85%) after 2 nd dose against infection.	Moderate	Prospective cohort in UK; 370,000 participants; sample confirmed VOC Alpha.
11	Hall (SIREN)	BNT162b2 vaccine showed VE of 70% (95% CI, 55 to 85) 21 days after 1 st dose and 85% (95% CI, 74 to 96) 7 days after 2 nd dose against infection in HCW.	Moderate	Prospective cohort with standardized testing for HCW over all of England; 23,000 participants; time and setting for VOC Alpha
12	Shrotri	Similar effect sizes were seen for ChAdOx1 (aHR 0.32, 95% CI, 0.15 to 0.66) and BNT162b2 (aHR 0.35, 95% CI, 0.17 to 0.71) at 35-48 days after 1 st dose.	Moderate	Prospective cohort in England: 9160 of 10412 frail LTC residents, 66% Pfizer, 33% AZ; routine screening; time and setting for VOC Alpha
13	Hyams	1st BNT162b2 showed VE 71.4% (95% CI, 43.1 to 86.2) against hospitalization 14 days after 1st dose; ChAdOx1nCoV-19 showed VE 80.4% (95% CI, 36.4 to 94.5) against hospitalization 14 days after 1st dose for 80+. When effectiveness analysis for BNT162b2 was restricted to the period covered by ChAdOx1nCoV-19, the estimate was 79.3% (95% CI, 47.0 to	Moderate	Test negative case-control study in Scotland. Single center; 466 participants, 80+; time and setting for VOC Alpha
14	<u>Harris</u>	92.5). BNT162b2 or ChAdOx1 reduced likelihood of transmission by 40-50% for household contacts of HCW 21 days after 1 st dose.	Moderate	Data-linkage and case-control study in England; 338,887 participants; time and setting for VOC Alpha
15	Goldberg	Prior infection (in unvaccinated) has similar VE against infection [94.8%], and severe illness [96.4%] as two doses of BNT162b2.	Moderate	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	<u>Hitchings</u>	CoronaVac showed VE of 35.1% (95% CI, -6.6 to 60.5) against infection in HCW after 1 st dose.	Moderate	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 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	Ismail	BNT162b2 showed VE 81% (95% CI, 76 to 85) against hospitalization 28 days after 1 st dose and 93% (95% CI,89 to 95) 14 days after the 2 nd dose for people 80+. ChAdOx1 showed VE 73% (95% CI, 60 to 81) against hospitalization 28 days after 1 st dose; sample size too small to report VE after 2 nd dose for people 80+.	Moderate	Screening study in UK; 13,907 hospitalized patients; results for age 80+; time and setting for VOC Alpha
21	Bernal (2)	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.	<u>Critical</u>	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, 88 to 97) against death 7-27 days after 2 nd dose; 71% (95% CI, 37 to 87) in immunosuppressed.	Moderate	Data-linkage study in Israel (Maccabi Health Care Organization); 1,178,597 participants; compared time

				frames to estimate effectiveness against 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 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.	Moderate	Test-negative study in Ontario 324,033 participants; limitations in symptom collection; screening for variants started 2 months into study period; results also reported for age>70 and according to vaccine (but not according to confirmed variant)
24	Bailly	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.	Moderate	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	<u>Bianchi</u>	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.	Moderate	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.	Moderate	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+.	Moderate	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 1 st dose and VE 93.7% (95% CI, 91.6 to 95.3) at least 14 days after 2 nd dose against symptomatic infection by confirmed VOC Alpha. ChadOx1showed VE 48.7% (95% CI, 45.2 to 51.9) at least 21 days after 1 st dose and VE 74.5% (95% CI, 68.4 to 79.4) at least 14 days after 2 nd dose	Moderate	Test-negative in England; 19,109 sequenced cases: 14,837 Alpha and 4,272 Delta.

		against symptomatic infection by confirmed VOC Alpha. 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+.	Moderate	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.	Moderate	Test-negative in California; 1,023 participants; expansion of sample size and timeline since previous study by same authors; self-reported vaccine receipt; 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.	Moderate	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	<u>Shrestha</u>	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+.	Moderate	Test-negative in Canada; 16,993 specimens; out of 1,131 genetically sequenced: 45% VOC Alpha and 28% Gamma;

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36	Abu-Raddad	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+. BNT162b2 showed VE 89.5% (95%	Moderate	limitations in symptom collection and assessment for covariates; results reported by vaccine but not according to confirmed variant Test-negative in Qatar; 17,293
		CI, 85.9 to 92.3) against infection, VE 100% (95% CI, 81.7 to 100) against any severe, critical, or fatal disease by VOC Alpha ≥ 14 days after 2 nd dose.		cases; sequencing showed 50% VOC Beta and 45% VOC Alpha between February-March 2021
		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.		
37	Akhrass	BNT162b2 or mRNA-1273 showed	Critical	Retrospective cohort of HCW at
	*Delayed exclusion -	overall VE 60.4% (95% CI, 30 to 77.6) against symptomatic infection ≥ 14 days		a single centre in Kentucky, USA; 2,134 participants; time
	failure to	after 1 st dose; BNT162b2 or mRNA-		and setting for VOC Alpha
	report	1273 showed overall VE 95.7% (95%		
	outcomes of	CI, 90 to 98.2) against symptomatic		
	interest for	infection ≥ 14 days after 2^{nd} dose.		
	this LES			
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 1 st 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 2 nd dose.	Moderate	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)
		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 1 st 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 2 nd dose.		
39	*Delayed exclusion – serious risk	BNT162b2 reported no symptomatic infections in the vaccinated group (0/686) compared to 0.83% infections in the vaccinated general population control group.	Serious	Prospective cohort of adults with autoimmune inflammatory rheumatic diseases in Israel; 686 participants; time and setting for VOC Alpha
	of bias			1

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.	Moderate	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.	Moderate	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.	Moderate	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	Saciuk	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	Moderate	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 1st dose and VE 89% (95% CI, 82 to 94) at least 7 days after 2nd 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	Moderate	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	Moderate	Prospective cohort of HCWs in Oxfordshire, UK; 13,109 participants; confirmed VOC Alpha

47	<u>Nasreen</u>	BNT162b2 showed VE 89% (95% CI,	Moderate	Test-negative study in Ontario
		86 to 91) against symptomatic infection and VE 95% (95% CI, 92 to 97) against		421,073 participants (same population as for Chung but
		hospitalization at least 7 days after 2 nd		extended to May 2021 and more
		dose (VOC Alpha); VE 84% (95% CI,		detailed with respect to
		69 to 92) against symptomatic infection		reporting of VOC); limitations
		and VE 95% (95% CI, 81 to 99) against		in symptom collection;
		hospitalization at least 7 days after 2 nd		screening for VOC Alpha,
		dose (VOC Beta/Gamma); VE 87% (95% CI, 64 to 95) against symptomatic		Beta/Gamma and Delta varied during study period
		infection at least 7 days after 2 nd dose		during study period
		(VOC Delta).		
		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).		
		dose (v o o riipiia).		
		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 1 st		
		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 1 st 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		
40	0 1	days after 1st dose (VOC Delta).	· ·	D
48	<u>Gazit</u>	BNT162b2 showed VE 80% (95% CI, 73 to 85) at least 7 days after 2 nd dose	Serious	Retrospective cohort of household members (household
		against infection in vaccinated		= 2 adults with no children) of a
				health management organization
	ı			.0

		household members of a confirmed COVID+ case.		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 1st and 2nd dose, respectively. 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 1st and 2nd 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 1st and 2nd dose, respectively (combined VOC Alpha and Beta).	Moderate	Test-negative in Qatar; >75,000 participants; sample genome sequenced for VOC Alpha and VOC 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).	Moderate	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.	Moderate	Data-linkage study of pregnant women over age 16 in Israel (same database as Dagan); 21,722 participants; time and setting for Alpha.

	1	T		<u> </u>
		Too few events to report VE for severe		
F 2	3.5	disease or death.	3.6.1	D : 1' 1 1 . 1 . 1
53	Mateo-	BNT162b2 (61%) or ChAdOx1 (31%)	Moderate	Data-linkage study in Italy;
	<u>Urdiales</u>	or mRNA-1273 (7%) or Ad26.COV ₂ -S		13,721,506 participants; time
		(0.6%) showed VE 78% (95% CI, 76 to		and setting for VOC Alpha.
		79) against infection 42 to 49 days after		Results not reported by vaccine
		at least 1 st dose; VE 93% (95% CI, 89 to		and some participants (42%)
		96) against death 35 to 42 days after at		who also received 2 nd dose were
		least 1 st dose.		included in estimates.
54	<u>Goldshtein</u>	BNT162b2 showed VE 78% (95% CI,	Moderate	Data-linkage study of pregnant
		57 to 89) against infection at least 28		women in Israel (same database
		days after 1st dose.		as Gazit); 15,060 participants;
				time and setting for VOC
				Alpha.
55	<u>Mason</u>	BNT162b2 showed VE 55.2% (95%	Moderate	Case-control study of age 80-83
		CI, 40.8 to 66.8) and VE 70.1% (95%		vs 76-79 community-dwelling
		CI, 55.1 to 80.1) against infection 21 to		unvaccinated residents in
		27 days and 35 to 41 days after 1 st dose,		England; time and setting for
		respectively.		VOC Alpha
				-
56	<u>Fabiani</u>	BNT162b2 showed VE 84.1% (95%	Moderate	Retrospective cohort of HCW
		CI, 39.7 to 95.8) and VE 85.4% (95%		in Italy; 6,423 participants; time
		CI, -35.3 to 98.4) against infection 14 to		and setting for VOC Alpha
		21 days and ≥21 days after 1 st dose,		
		respectively in HCW.		
		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		confirmed VOC Delta admitted
		severe disease (defined as requiring		to hospital (including
		supplemental oxygen) > 14 days after		asymptomatic) in Singapore; 218
		2 nd dose.		participants; not reported by
		2 4006.		vaccine and non-m-RNA
				vaccine outcomes excluded
58	Kaur	Two doses of Covishield showed VE	Serious	Preliminary report of
	13001	87% (95% CI, 33 to 97) against severe	5611043	prospective cohort in India;
		disease when compared with one dose		1500 participants; time and
		(timing of doses not reported).		setting for VOC Delta
59	Pramod	Covishield showed VE 49% (95% CI,	Serious	Test-negative study in a single
	<u> </u>	17 to 68) against infection 21 days after	octions	hospital site in India; 360
		1st dose and VE 54% (95% CI, 27 to 71)		matched pairs (203 symptomatic
		against infection 14 days after 2 nd dose.		pairs); time and setting for VOC
		agamst infection 14 days after 2 dose.		Delta
		Covishield showed VE 590/ (050/ CI		Dena
		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 60% (95% CI, 53.6 to 65.5) against infection by confirmed VOC Alpha 14 days after 1 st dose. BNT162b2 or mRNA-1273 showed VE 92.6% (95% CI, 87.1 to 95.8) against infection by confirmed VOC Alpha 7 days after 2 nd dose.	Serious	Test-negative study in Quebec, Canada; 58,476 participants; sample confirmed VOC Alpha; reported according to vaccine but not for Alpha at same time
61	Williams	BNT162b2 or mRNA-1273 showed VE 52.5% (95% CI, 26.9 to 69.1) against infection and VE 78.6% (95% CI, 47.9 to 91.2) against severe disease 14 days after 2 nd 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.	Serious	Outbreak in a single LTCF in Ontario; 60 residents and 83 staff; sample confirmed VOC Gamma
62	Hitchings(2)	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.	Serious	Test-negative study in Sao Paulo, Brazil; 61,164 participants over age 60; time and setting for VOC Gamma
63	Tang	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.	Serious	Test-negative study in Qatar; 1,140,337 participants; weekly random sequencing of positive samples for VOC Delta

		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 disease ≥ 14 days after 2 nd dose.		
64	Puranik	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.	Moderate	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	Elliot	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). 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).	Serious	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 self-reported vaccination status
66	Issac	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	ChAdOx1 showed VE 23% (95% CI, not reported) against infection at least 21 days after 1 st dose.	Moderate	Outbreak study of prison inmates in Barcelona; 217 participants (184 inmates); sequenced for VOC Alpha
68	Kale	ChAdOx1 showed VE 60% (95% CI, 45 to 70) against infection at least 14 days after 2 nd dose.	Serious	Prospective cohort of HCW at a single hospital in India; 1858 participants; sample sequenced for VOC Delta
69	Israel	BNT162b2 showed OR 2.06 (95% CI, 1.69 to 2.51) for infection comparing fully vaccinated longer than or equal to 146 days vs fully vaccinated less than 146 days.	Moderate	Retrospective cohort of fully vaccinated (>14 days after 2 nd dose) 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-	Moderate	Data-linkage study in Denmark; 5,542,079 participants; time and setting for VOC Alpha
		1273 showed VE 88% (95% CI, 83 to 92) against infection ≥ 14 days after 2 nd		
	-	dose.		
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+).	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
		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 infection ≥ 14 days after 2 nd dose (VOC Delta age 18+).		
		mRNA-1273 showed VE 75% (95% CI: 64 to 83) against infection ≥21 days after 1 st dose (VOC Delta age 18 to 64).		
72	Abu-Raddad (2)	BNT162b2 after prior infection showed VE 85% (95% CI, 80 to 89) against reinfection compared to BNT162b2 without prior infection.	Moderate	Retrospective matched cohorts (2) of fully vaccinated (>14 days after 2 nd dose) in Qatar; 151,076 participants; sample sequenced for VOC Alpha and VOC Beta
		mRNA-1273 <u>after prior infection</u> showed VE 15% (95% CI, -105 to 66) against re-infection compared to mRNA-1273 <u>without prior infection</u> .		
73	Gazit (2)	BNT162b2 showed OR 13.06 (95% CI, 8.08 to 21.11) against infection and OR 27.02 (95% CI, 12.7 to 57.5) against	Moderate	Retrospective matched cohorts of fully vaccinated (>14 days after 2 nd dose) in

		symptomatic disease compared to <u>prior</u> <u>infection.</u>		Israel; 778,658 participants; time and setting for VOC Delta
74	Rosenberg	BNT162b2 (51%), mRNA-1273 (40%) or Ad26.COV2.S (9%) showed VE 91.7% against infection ≥14 days after 2 nd dose (Week of May 3, 2021: VOC Alpha). 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	Serious	Surveillance report in New York, USA; >13 million participants; time and setting for VOC Delta (from 2% to 80% during study period)
75	Al-Qahtani	Delta). BNT162b2 ≥14 days after 2 nd dose, showed VE 99.9% (95% CI, 99.2 to 100) against ICU admission, and VE 99.5% (95% CI, 98.4 to 99.8) against death (VOC Alpha and Delta). ChAdOx1 ≥14 days after 2 nd dose, showed 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). 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 ≥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).	Serious	Retrospective cohort of fully vaccinated (>14 days after 2 nd dose) in Bahrain; 1,242,279 participants; time and setting for VOC Alpha (dominant before May 2021) and Delta (dominant after May 2021).
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).	Moderate Moderate	Data-linkage study of fully vaccinated in Israel; 4,785,245 fully vaccinated participants; time and setting for VOC Delta (dominant after May 2021).

		BNT162b2 showed VE 57% (95% CI, 52 to 62) for those vaccinated in		
		January 2021, 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	<u>Herlihy</u>	BNT162b2, mRNA-1273, or Ad26.COV2.S showed VE 78% (95%	Critical	Surveillance report in Mesa County-Colorado, USA; 37,439
	*Delayed exclusion – critical risk of bias	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).		cases participants; sample sequenced for VOC Delta (43% to 88% during study period)
78	Ghosh	ChAdOx1 showed unadjusted VE 75.2% (95% CI, 73.8 to 76.8) against	Critical	Retrospective cohort of Armed Forces HCW and frontline
	*Delayed exclusion – critical risk of bias	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		workers in India; 1,595,630 participants; time and setting for VOC Delta at end of study only.
79	Amirthaling	Delta). BNT162b2 showed VE 77% (95% CI,	Moderate	Test-negative study in England;
	<u>am</u>	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		time and setting for VOC Alpha (dominant before May 2021) and Delta (dominant after May 2021).
		(VOC Alpha age 80+).		
		BNT162b2 showed VE 77% (95% CI, 66 to 85) against symptomatic infection when 2 nd dose given 19-29 days after 1 st dose, and VE 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 infection when 2 nd dose given 19-29 days after 1 st dose, and VE 88% (95% CI, 48 to 97) against infection when 2 nd dose given 85+ days after 1 st dose after 2 nd dose (VOC Alpha age 80+). ChAdOx1 showed VE 66% (95% CI, 47 to 77) against infection when 2 nd dose given 19-29 days after 1 st dose, and		
		VE 73% (95% CI, 56 to 83) against 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 (95% CI, 1.80 to 4.47) of severe disease compared to BNT162b2 ≥14 days after 2 nd dose.	Moderate	Case-control study in Qatar; 456 matched cases; time and setting for VOC Alpha
81	Fowlkes	BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 91% (95% CI, 81 to 96) against infection ≥ 14 days after 2 nd dose (during time of VOC Alpha). BNT162b2 (65%), mRNA-1273 (33%), or Ad26.COV2.S (2%) showed VE 66%	Moderate	Prospective cohort of HCW and other essential frontline workers in 6 states in the USA; 7,112 participants; updated report to cover VOC Delta period
		(95% CI, 26 to 84) against infection ≥ 14 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, 49 to 86) against infection ≥150 days after full vaccination (during time of VOC Alpha to Delta).		
82	Bhattachary a	Covaxin (94%) and Covishield showed VE 83% (95% CI, 73 to 89) against symptomatic infection ≥ 14 days after 2 nd dose. Covaxin (94%) and Covishield showed	Serious	Prospective cross-sectional cohort of HCW and their families at a single site in India; 638 participants (55 inpatients); time and setting of VOC Delta
		VE 93% (95% CI, 64 to 99) against		

		ICU admission or death \geq 14 days after 2^{nd} dose.		
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). 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). 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 ≥ 98 days after 2 nd dose for HR	Moderate	Data-linkage study of community-dwelling adults≥65 in Portugal; 2,050,950 participants; time and setting for VOC Alpha to Delta
84	Tartof	1.80 (0.77 to 4.25) (age ≥80). BNT162b2 showed VE 75% (95% CI, 71 to 78) against infection 7 days after 2 nd dose (confirmed VOC Delta). BNT162b2 showed VE 91% (95% CI, 88 to 92) against infection 7 days after 2 nd dose (confirmed non-VOC 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). BNT162b2 showed VE 97% (95% CI, 95 to 99) against infection 7 to 30 days after 2 nd dose and VE 67% (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).	Moderate	Retrospective cohort of members of a health management organization in California; 3,436,957 participants; VOC Alpha to VOC Delta (only 28% confirmed Delta)
85	Li (3)	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.	Serious	Test-negative study in Guangzhou, China; 366 participants; sample sequenced for VOC Delta
86	Scobie	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	Serious	Surveillance study in 13 states in the USA; 615,454; time and setting for VOC Alpha to VOC Delta

	1	T		
		death \geq 14 days after 2 nd dose (April to June: VOC Alpha).		
		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 38) against symptomatic infection; VE 37% (-24 to 68) against moderate to severe disease and VE 69% (95% CI, - 160 to 97) against death ≥21 days after 1 st dose.	Serious	Retrospective cohort study of HCW at a single hospital in New Delhi, India; 4276 participants; sample sequenced for VOC Delta
		ChAdOx1 showed VE 28% (95% CI, 10 to 41) against symptomatic infection; VE 67% (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	<u>Seppala</u>	BNT162b2 (74%) or ChAdOx1 (22%) or mRNA-1273 (10%) showed VE 84.4% (95% CI, 81.8 to 86.5) against infection ≥7 days after 2 nd dose (VOC Alpha). BNT162b2 (74%) or ChAdOx1 (22%)	Moderate	Population cohort in Norway; 4,204,859 participants; sequenced for VOC Alpha and VOC Delta
		or mRNA-1273 (10%) showed VE 64.6% (95% CI, 60.6 to 68.2) against infection ≥7 days after 2 nd dose (VOC Delta).		
89	Polinski	Ad26.COV2.S showed VE* 67% (95% 60 to 73) against infection unknown number of days after dose (June to July: VOC Delta in high prevalence states). *unadjusted for substantial under-reporting of vaccination status	Serious	Data-linkage of members of a medical insurance group in USA; 1,914,670 participants; time and setting for VOC Alpha to Delta (only data for VOC Delta reported here)
90	Chemaitelly (2)	BNT162b2 or mRNA-1273 showed VE 46.6% (95% CI, 0.0 to 73.7) against infection \geq 14 days after 2^{nd} dose, VE 66.0% (95% CI, 21.3 to 85.3) \geq 42 days after 2^{nd} dose, and VE 73.9% (95% CI, 33 to 98.9) \geq 56 days after 2^{nd} dose (VOC Alpha and Beta).	Serious	Retrospective cohort of immunosuppressed kidney transplant recipients in Qatar; 782 participants; time and setting for VOC Alpha and VOC Beta.
		BNT162b2 or mRNA-1273 showed VE 72.3% (95% CI, 0.0 to 90.9) against severe, critical, or fatal disease ≥14 days		

91	Hu	after 2 nd dose, VE 85% (95% CI, 35.7 to 96.5) ≥42 days after 2 nd dose, and VE 83.8% (95% CI, 31.3 to 96.2) ≥56 days after 2 nd dose (VOC Alpha and Beta). Inactivated vaccines showed VE 89% (95% CI, 55 to 98) against severe,	Serious	Outbreak report of hospitalized cases in China; 476 participants;
		critical, or fatal disease ≥14 days after 2 nd dose (VOC Delta).		PCR population for VOC Delta.
92	Andrews	BNT162b2 showed VE 62.7% (61.7 to 63.8) against symptomatic infection 1 week after 2 nd dose and VE 47.3% (45.0 to 49.6) 20+ weeks after 2 nd dose (VOC Delta). ChAdOx1showed VE 92.4% (92.1 to 92.7) 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	Moderate	Test-negative study in England; 1,475,391 participants; VOC Alpha to VOC Delta (only data for VOC Delta reported here)
02	D . 1	(VOC Delta).	Nr. 1	
93	<u>Patalon</u>	BNT162b2 showed marginal VE 3% (95% CI, -5 to 10) against infection 0 to 6 days after 3 rd dose and marginal VE 84.0% (95% CI, 79 to 88) 14 to 20 days after 3 rd dose compared to 2 doses.	Moderate	Test-negative study in Israel comparing 2 doses of vaccine versus 3 doses of vaccine; 182,076 participants; time and setting for VOC Delta

Section 2: excluded studies			
Author	Reason for exclusion		
<u>Akhrass</u>	Delayed exclusion – clinical outcomes of interest not reported		
<u>Albahrani</u>	Prevalence of variants unknown and suspected to be <50%		
Alencar	Critical risk of bias		
<u>Alhamlan</u>	Vaccine effectiveness not reported		
Ali	Prevalence of variants unknown and suspected to be <50%		
Allen	Critical risk of bias		
Almufty	Prevalence of variants unknown and suspected to be <50%		
<u>Apisarnthanarak</u>	Vaccine effectiveness not reported		
<u>Arashiro</u>	Vaccine effectiveness not reported		

Bergwerk Picula	Clinical outcomes of interest not reported Vaccine effectiveness not reported
Diagla	
<u>Bjork</u>	Prevalence of variants unknown and suspected to be <50%
Borobia	Clinical outcomes of interest not reported
Britton	Prevalence of variants unknown and suspected to be <50%
Brown	Vaccine effectiveness not reported
Butt	Prevalence of variants unknown and suspected to be <50%
Butt	Serious risk of bias
Cabezas	Prevalence of variants unknown and suspected to be <50%
<u>Caillard</u>	Clinical outcomes of interest not reported
Cavanaugh	Delayed exclusion – VOI not VOC
Charmet	Serious risk of bias
Chau	Vaccine effectiveness not reported
Clemens	Prevalence of variants unknown and suspected to be <50%
Corchado-Garcia	Prevalence of variants unknown and suspected to be <50%
Dash	Critical risk of bias
de Gier Brechje	Prevalence of variants unknown and suspected to be <50%
Domi	Prevalence of variants unknown and suspected to be <50%
Ella	Prevalence of variants unknown and suspected to be <50%
<u>Farinholt</u>	Vaccine effectiveness not reported
<u>Fisher</u>	Prevalence of variants unknown and suspected to be <50%
Frenck	Prevalence of variants unknown and suspected to be <50%
<u>Furer</u>	Delayed exclusion – serious risk of bias
Geisen	Clinical outcomes of interest not reported
Gils	Clinical outcomes of interest not reported
Gorgels	Prevalence of variants unknown and suspected to be <50%
Gray	Prevalence of variants unknown and suspected to be <50%
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%
<u>Gupta</u>	Vaccine effectiveness not reported
Haas (2)	Modelling study used to estimate cases averted
<u>Hacisuleyman</u>	Critical risk of bias
<u>Hetemaki</u>	Vaccine effectiveness not reported
<u>Hollinghurst</u>	Serious risk of bias
<u>Iliaki</u>	Prevalence of variants unknown and suspected to be <50%
Jacobson	Critical risk of bias
<u>John</u>	Prevalence of variants unknown and suspected to be <50%
<u>Jones</u>	Serious risk of bias
<u>Kaabi</u>	Prevalence of variants unknown and suspected to be <50%
Keegan	Critical risk of bias
Khan	Prevalence of variants unknown and suspected to be <50%

<u>Khawaja</u>	Critical risk of bias	
<u>Kojima</u>	Prevalence of variants unknown and suspected to be <50%	
<u>Lefèvre</u>	Critical risk of bias	
<u>Li</u>	Phase 1 trial	
<u>Li (2)</u>	Clinical outcomes of interest not reported	
Ling	Prevalence of variants unknown and suspected to be <50%	
Linsenmeyer	Vaccine effectiveness not reported	
Loconsole	Vaccine effectiveness not reported	
Luo	Vaccine effectiveness not reported	
Mattar	Prevalence of variants unknown and suspected to be <50%	
Mazgatos	Critical risk of bias	
<u>Menni</u>	Serious risk of bias	
<u>Mizrahi</u>	Modelling study	
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 not reported	
Musser	Vaccine effectiveness not reported	
Mutnal	Vaccine effectiveness not reported	
Nanduri	Critical risk of bias	
Olmedo	Clinical outcomes of interest not reported	
Palacios	Prevalence of variants unknown and suspected to be <50%	
Paris	Prevalence of variants unknown and suspected to be <50%	
<u>Pawlowski</u>	Serious risk of bias	
Perry	Clinical outcomes of interest not reported	
Pilishville	Prevalence of variants unknown and suspected to be <50%	
Raches Ella	Phase 1 trial	
Rana	Critical risk of bias	
Regev-Yochay	Prevalence of variants unknown and suspected to be <50%	
Riemersma	Clinical outcomes of interest not reported	
Riley	Serious risk of bias	
Rivelli	Clinical outcomes of interest 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	
<u>Shimabukuro</u>	Clinical outcomes of interest not reported	
<u>Starrfelt</u>	Serious risk of bias	
Swift	Prevalence of variants unknown and suspected to be <50%	
<u>Tande</u>	Prevalence of variants unknown and suspected to be <50%	
<u>Tanriover</u>	Prevalence of variants unknown and suspected to be <50%	
<u>Tenforde</u>	Clinical outcomes of interest not reported	

Thiruvengadam Serious 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% Vahidy Prevalence of variants unknown and suspected to be <50% Vasileiou Clinical outcomes of interest not reported Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50% Wickert Critical risk of bias Williams (2) Serious risk of bias Young-Xu Prevalence of variants unknown and suspected to be <50%	Tenforde (2)	clinical outcomes of interest not reported
Thiruvengadam Serious 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% Vahidy Prevalence of variants unknown and suspected to be <50% Vasileiou Clinical outcomes of interest not reported Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50% Wickert Critical risk of bias Williams (2) Serious risk of bias Young-Xu Prevalence of variants unknown and suspected to be <50%	Thangaraj	Critical risk of bias
Thompson (2) Prevalence of variants unknown and suspected to be <50% Vahidy Prevalence of variants unknown and suspected to be <50% Vasileiou Clinical outcomes of interest not reported Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50% Wickert Critical risk of bias Williams (2) Serious risk of bias Young-Xu Prevalence of variants unknown and suspected to be <50%	Thiruvengadam	Serious risk of bias
Vahidy Prevalence of variants unknown and suspected to be <50%	Thompson (1)	Prevalence of variants unknown and suspected to be <50%
Vasileiou Clinical outcomes of interest not reported Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50%	Thompson (2)	Prevalence of variants unknown and suspected to be <50%
Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50%	<u>Vahidy</u>	
Veneti Clinical outcomes of interest not reported Victor Critical risk of bias Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50%	Vasileiou	Clinical outcomes of interest not reported
Volkov Modelling study Voysey Prevalence of variants unknown and suspected to be <50%	Veneti	
Voysey Prevalence of variants unknown and suspected to be <50%	<u>Victor</u>	Critical risk of bias
Wickert Critical risk of bias Williams (2) Serious risk of bias Young-Xu Prevalence of variants unknown and suspected to be <50%	Volkov	
Williams (2) Serious risk of bias Young-Xu Prevalence of variants unknown and suspected to be <50%	Voysey	
Young-Xu Prevalence of variants unknown and suspected to be <50%	Wickert	
	Williams (2)	Serious risk of bias
Zacay Delayed exclusion – critical risk of bias Delayed exclusion – critical risk of bias	Young-Xu	
	Zacay	Delayed exclusion – critical risk of bias

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

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)

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 22 Sept 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**. Low Risk of Bias indicates High Quality, and Critical Risk of Bias indicates Very Low (insufficient) Quality. 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 "serious" or "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 serious or critical risk of bias in at least one domain.

VE Study Characteristics that may introduce bias	Description
Study design	In cohort studies, people who get vaccinated may differ in health-seeking behaviour from people who do not get
ROBINS-I: Bias in selection of participants into study	vaccinated; using a test-negative study design minimizes this type of bias
	Examples and typical judgement:
	complete inception cohort with before and after comparison (low)
	• test-negative study (moderate)
	case-control or cross-sectional study (serious)
Method for confirming vaccination	Questionnaires are prone to recollection bias; Population
DODDIC I D' ' 1 'C ' C	databases developed for purpose of tracking COVID
ROBINS-I: Bias in classification of interventions	vaccines minimize this type of bias
	Examples and typical judgement:
	database linkage study (low)
	Questionnaire with confirmation by an additional method (e.g. registry) of at least a subset of study population (moderate)
	• Questionnaire without confirmation by an additional method (serious)

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

	Estimating vaccination status based on surveillance data alone (critical)
Databases used for retrieval of COVID test results, participant prognostic factors, and clinical	Databases developed for collecting data on COVID are less prone to bias due to missing information and m
outcomes	Examples and typical judgement:
ROBINS-I: Bias in classification of	database for non-COVID purpose but with individual level data (moderate)
interventions	• database for non-COVID purpose without individual level data (serious)
	• no or unclear description of database type (critical)
Assignment of infection start date	Using date of symptom onset (if within 10 days of testing) as infection start date reduces risk of
ROBINS-I: Bias in classification of interventions	misclassification bias (e.g., vaccinated participant who is reported as COVID+ may have been infected prior to receiving the vaccine or during non-immune period) and sensitivity of assays decreases over time
	Examples and typical judgement:
	• using sample date without interview or documented confirmation of symptoms ≤ 10 days (relevant for symptomatic disease only) (serious)
Verification of symptoms	Prospective, standardized collection of symptoms from patients reduces risk of missing information bias; testing
ROBINS-I: Bias in classification of	within 10 days after symptom onset reduces risk of false-
interventions	negative COVID test
	Examples and typical judgement:
	• using sample date without interview or documented confirmation of symptoms ≤ 10 days (relevant for symptomatic disease only) (serious)

Accounting for non-immune period (first 14 days after first vaccine dose)	Reported absence of vaccine effect during non-immune period reduces risk of residual confounding bias
ROBINS-I: Bias due to confounding	 Example/common case: presence of an effect during non-immune period or result not reported (moderate) unclear that non-immune period was considered (serious)
Inclusion of participants with prior	Exclusion (or separate analysis) of participants with prior
COVID infection	COVID infection reduces concern about differences in infectivity as well as risk-taking and health-seeking
ROBINS-I: Bias due to confounding	behaviour
	Examples and typical judgement:
	previously infected not excluded or analyzed
Accounting for calendar time	separately (serious) Accounting for calendar time reduces bias due to
Accounting for calcindar time	differences in vaccine accessibility and risk of exposure
ROBINS-I: Bias due to confounding	over time
(time-varying confounding)	
	Examples and typical judgement:
	• 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 prognostic factors	Adjustment for prognostic factors for COVID infection, severity of disease, and vaccination, such as age, gender,
ROBINS-I: Bias due to confounding	race, ethnicity, socioeconomic factors, occupation (HCW, LTC), and chronic medical conditions
	Examples and typical judgement:
	• no or insufficient adjustment for socioeconomic
	factors, race, ethnicity, occupation (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
ROBINS-I: Bias in measurement of	of bias introduced by detecting asymptomatic infection
outcomes	in one group but not in another (e.g. when only one group undergoes surveillance screening)
outcomes	group andergoes survemance serecting,
	Examples and typical judgement:
	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 more than one study was found, we will provide a summary statement with a <u>range of the</u> estimates across the studies.

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%.