Weekly COVID-19 Vaccine Updates

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Introduction

This document summarises the vaccine efficacy and effectiveness, the vaccine specifications, the vaccine development pipeline and the timeline for World Health Organization (WHO) review of the various COVID-19 vaccines in late phase development. This document is updated weekly.

- Vaccine efficacy refers to the performance of a vaccine in a controlled clinical trial (study) situation
- · Vaccine effectiveness refers to the performance of a vaccine in a population under real-world conditions

Key messages

- COVID-19 vaccine efficacy results from different trials cannot be directly compared against each other. They must be interpreted in the context of study designs (including case definitions, clinical endpoints, access to testing), target populations, and COVID-19 epidemiologic conditions (including circulation of variants of concern)
- All COVID-19 vaccines in late phase development report high vaccine efficacy against severe COVID-19 and favourable safety profiles
- Most vaccines have high effectiveness against infection, including with the Delta variant
- The Pfizer/BioNTech vaccine has been authorised in children aged ≥5 years by the US FDA, EU EMA, Australian TGA and Health Canada; and in those ≥12 years by the UK MHRA and NZ Medsafe, The EMA, MHRA and TGA have authorised the Moderna vaccine in those ≥12 years.
- WHO SAGE recommends that 1) immunocompromised persons should be offered an additional dose of all WHO EUL COVID-19 vaccines as part of an extended primary series; and 2) following the Sinopharm inactivated vaccines, a third dose of the same vaccine or a different vaccine should be offered as part of an extended primary series.¹ ATAGI has recommended that immunocompromised persons in Australia receive a third dose as part of the primary series and booster doses of Pfizer/BioNTech be offered to all irrespective of the primary COVID-19 vaccine given.^{2,3} Boosters are recommended for all adults ≥18 years of age in the US and will be offered to all adults in the UK by January 2022.^{4,5}
- Mixed vaccine schedules (i.e. delivering different types of vaccine for the first and second dose) are under investigation as these could facilitate better protection against variants of concern and enable vaccination programs to continue if a particular vaccine is unavailable
- Seven intranasal vaccines are in development (6 live-attenuated viruses or virus-vectored vaccines; 1
 protein subunit.⁶ These may be beneficial in preventing transmission (Page 15)
- A very rare clotting disorder with low platelets (Thrombosis with Thrombocytopaenia Syndrome TTS) has been associated with the AstraZeneca and Johnson & Johnson vaccines.^{7–9} The majority of cases fully recover with adequate treatment. The risk following the first dose of AstraZeneca vaccine has been estimated by the EMA as 1 in 100,000 and by the Australian Technical Advisory Group on Immunisation (ATAGI) as 1 in 50,000.^{10,11} Risk of TTS is much lower following the *second* dose of AstraZeneca vaccine: estimate in the UK is 1 in 1.5 million second doses.¹²
- The risk of TTS following the first dose of Johnson & Johnson vaccine has been estimated as 1 in 319,000 in the USA $^{\rm 13}$
- The risk of myocarditis/pericarditis is increased following the second dose of Pfizer/BioNTech and Moderna vaccines, particularly in younger males, occurring in >1 in 20,000 males under 25 years of age.¹⁴ Highest rate in males 16-17 years of age following Pfizer/BioNTech vaccine but no clear difference in risk between Moderna and Pfizer/BioNTech.¹⁵ There is a small increase in risk of myocarditis in females <30 and males >50 years of age. Data from Ontario, Canada, and the UK suggest higher rates following Moderna than Pfizer/BioNTech vaccine. ATAGI in Australia continue to review the data.
- Appropriate communication on the benefit-risk profile of COVID-19 vaccines (Page 15) remains crucial to
 maintain confidence in immunisation programmes and to avoid vaccine hesitancy.

New updates

Key updates include (also highlighted in yellow text in the document):

- Effectiveness of Pfizer/BioNTech and Moderna boosters against hospitalisation following AstraZeneca, Pfizer/BioNTech and Moderna primary series in the UK:
 - 2-9 weeks post-booster: 94% (89-97)
 - ≥10 weeks post-booster: 89% (80-95)
- Effectiveness of Moderna against infection and hospitalisation with Delta and Omicron in USA (Pages 6, 7, 8, 9, 10, 27, 29, 31 and 32):
 - Infection O Sir
 - Single dose:
 - Delta: 60.2% (42.6-72.3)
 - Omicron: 20.3% (0.0-39.8)
 - Two doses:
 - Delta 14-90 days: 82.8% (69.6-90.3)
 - Delta 181-270 days: 61.4% (56.8-65.5)
 - Omicron 14-90 days: 30.4% (5.0-49.0)
 - Omicron 181-270 days: 0.0% (0.0-1.2)
 - Three doses:
 - Delta ≤2 months: 95.7% (94.2-96.9)
 - Delta >2 months: 90.7% (81.4-95.3)
 - Delta ≥65 years: 97.2% (94.1-98.7)
 - Omicron ≤2 months: 63.6% (57.4-68.9)
 - Omicron >2 months: 39.1% (3.8-61.5)
 Omicron ≥65 years: 57.1% (14.2-78.6)
 - Hospitalisation
 - auon
 - Single dose Delta: 68.9% (0.0-99.4)
 Two doses Delta: 98.0% (87.2-99.7)
 - Two doses Delta: 98.0% (87.2-99.7)
- Effectiveness of AstraZeneca and mRNA vaccines against infection and hospitalisation with Omicron in the UK (updated 13 Jan 2022):
 - Infection:
 - AstraZeneca (2-4 weeks): 45-50%
 - AstraZeneca: No effect beyond 20 weeks
 - Pfizer/BioNTech or Moderna 2 doses (2-4 weeks): 65-70%
 - Pfizer/BioNTech or Moderna 2 doses (20 weeks): ~10%
 - Pfizer/BioNTech or Moderna booster (2-4 weeks): 65-75%
 - Pfizer/BioNTech or Moderna booster (5-9 weeks): 55-65%
 - Pfizer/BioNTech or Moderna booster (≥10 weeks): 45-50%
 - Hospitalisation (pooled analysis of AstraZeneca, Pfizer/BioNTech and Moderna):
 - Single dose (≥4 weeks): 58% (37-72)
 - Single dose (24 weeks): 36% (37-72)
 2 doses (2-24 weeks): 64% (54-71)
 - 2 doses (≥25 weeks): 04% (34-71)
 2 doses (≥25 weeks): 44% (30-54)
 - 3 doses (2-4 weeks): 92% (89-94)
 - 3 doses (2-4 weeks): 32 % (03-54)
 3 doses (5-9 weeks): 88% (84-91)
 - 3 doses (3-3 weeks): 00% (04-31)
 3 doses (≥10 weeks): 83% (78-87)
- Preliminary reports from Israel (data not released) indicate that a fourth dose of Pfizer/BioNTech produces a
 greater antibody than after the third dose but has very little effect on infection
- The Novavax vaccine has been authorised by the European Medicines Agency and Australian TGA (Page 4)

COVID-19 Vaccine Specifications

	ASTRAZENECA	GAMALEYA	JOHNSON & JOHNSON	MODERNA	NOVAVAX	PFIZER/ BIONTECH	SINOVAC	SINOPHARM	BHARAT BIOTECH	CLOVER
VACCINE TYPE	Viral vector (chimpanzee adenovirus ChAdOx1)	Viral vector (recombinant adenovirus types 5 and 26)	Viral vector (recombinant adenovirus type 26)	mRNA	Protein subunit	mRNA	Inactivated virus	Inactivated virus	Inactivated virus	Protein
Available Through COVAX	~	-	 	-	~	~	-	-	-	-
Doses Required	4-12 weeks apart	3 weeks apart	2 doses with 2-6 month interval also recommended	4 weeks apart*	3 weeks apart	3-4 weeks apart*	2-4 weeks apart*	3-4 weeks apart*	3 weeks apart	3 weeks apart
Third dose/ boosters	As part of primary series for those with immunocomp.			As part of primary series for those with immunocomp. USA: at least 6 months after primary series in at-risk groups and ≥65 years	As part of primary series for those with immunocomp.	As part of primary series for those with immunocomp. USA: at least 6 months after primary series in at-risk groups and ≥65 years	As part of primary series for ≥60 years	As part of primary series for ≥60 years	-	-
Shipping, Storage & Presentation	Normal cold chain requirements (2-8°C); 10-dose vials	-18.5°C (liquid form); 2-8°C (dry form)	Shipped at -20°C; 2-8°C for up to 3 months; 5-dose vials	-25°C to -15°C; 10-dose vials	2-8°C; 10-dose vials	-80°C to -60°C; 2-8°C for up to 1 month; 6-dose vials	2-8°C; Single-dose vials	2-8°C; Single-dose vials/ pre- filled syringes	2-8°C; 10-dose or 20-dose vials	2-8°C
Approval by a Stringent Regulatory Authority (SRA)	WHO EUL, EMA, TGA, MHRA	Under review by WHO SAGE	WHO EUL, EMA, FDA, MHRA	WHO EUL, EMA, FDA, TGA	WHO EUL, EMA, TGA	WHO EUL, EMA, FDA, TGA, MHRA	WHO EUL	WHO EUL	WHO EUL	-
WHO EUL: EMA: FDA: TGA:	WHO Emergency U European Medicine Food and Drug Adn Therapeutic Goods	lse Listing s Agency ninistration (US) Administration (Au	stralia)							

MHRA: Medicines and Healthcare Products Regulatory Agency (UK)



COVID-19 Vaccine Efficacy

VACCINE	VACCINE EFFICACY						
VACCINE	SYMPTOMATIC INFECTION	MODERATE-SEVERE	SEVERE	HOSPITALISATION/DEATH			
AstraZeneca	UK: 66.7% (57.4-74.0) ¹⁶ USA, Chile, Peru: 76% ¹⁷ (not peer-reviewed) Single dose in UK (22-90 days post-vaccination): 76.0% (59.3 to 85.9) ¹⁶ Efficacy with different interval between doses in UK: 12+ weeks: 82.4% (2.7-91.7) <6 weeks: 54.9% (32.7-69.7) ¹⁶	-	Severe/critical and hospitalisation in USA, Chile, Peru: 100% ¹⁷ (not peer-reviewed) UK: 100% (15 cases in the placebo group) ¹⁶	Hospitalisation in UK: 100% (9 cases in placebo group) ¹⁶			
Bharat Biotech	India: 77.8% (65.2-86.4) ¹⁸	-	India: 93.4% (57.1-99.8) ¹⁸	-			
Clover	Philippines, Colombia, Brazil, South Africa and Belgium: Overall: 67.2% (54.3-76.8); Delta: 78.7% (57.3-90.4) ¹⁹	Philippines, Colombia, Brazil, South Africa and Belgium: Overall: 83.7% (55.9-95.4); Delta: 81.7% (35.9-96.6) ¹⁹		Hospitalisation in Philippines, Colombia, Brazil, South Africa and Belgium: 100% (42.7-100) ¹⁹			
Gamaleya	Russia: 91.6% (85.6–95.2) ²⁰ Single dose (Sputnik Light) in Argentina: 78.6% ²¹	Moderate-severe: 100% (20 cases in the placebo group) ²⁰	-				
Johnson & Johnson	USA: 93.2% (91.0-94.8) ²²	Moderate to severe/critical: All sites: 66.1% (55.0-74.8) USA: 72.0% (58.2-81.7) Latin America: 61.0% (46.9-71.8) South Africa: 64.0% (41.2-78.7) ^{23.24} South Africa: 67-71% ²⁵	85.4% (54.2-96.9) ²⁴ USA: 98.2% (92.8-99.6) ²²	100% (5 deaths in placebo group) ²⁴ Death in South Africa: 96% ²⁵			
Moderna	USA: 94.1% (89.3-96.8) ²⁶ USA: >90% ²⁷ Efficacy in USA: 12-17 years: Symptomatic: 92.7% (67.8-99.2) Infection: 69.8% (49.9-82.1) Asymptomatic infection: 59.5% (28.4-77.3) ²⁸	-	USA: 100% (30 cases in placebo group) ²⁶ US: >95% ²⁷	USA: 100% (1 death in placebo group) ²⁶			
Novavax	UK: 89.7% (80.2-94.6) ²⁹ US and Mexico: 90.4% (82.9-94.6) ³⁰	US and Mexico: 100% (87.0-100) ³⁰	-				
Pfizer/BioNTech	Argentina, Brazil, Germany, South Africa, Turkey and the USA: 94.6% (89.9–97.3) ³¹ Infection over 6 months: 91.3% (89.0-93.2) ³² Single dose: 100%	-	Argentina, Brazil, Germany, South Africa, Turkey and the USA: 88.9% (20.1–99.7) ³¹ Severe disease: 96.7% (80.3-99.9) ³²				
Sinopharm	UAE, Bahrain, Egypt and Jordan: 78.1% (64.9-86.3) ³³	-		Hospitalisation in UAE, Bahrain, Egypt and Jordan: 78.7% (26.0-93.9) ³³			
Sinovac	Brazil: 50.7% (35.9-62.0) Chile: 67% (65-69) Indonesia: 65% (20-85) ³³ Turkey: 83·5% (65·4-92·1) ³⁴	Requiring medical assistance in Brazil: 83.7% (58.0-93.7) Moderate-severe: 100% (56.4-100.0) ³⁵		Hospitalisation: Brazil: 100% (56-100) Chile: 85% (83-97) Turkey: 100% (20-100) ³³			
CanSino Biologics	Pakistan, Mexico, Argentina, Russia and Chile: 57.5% (39.7-70.0) ³⁶	-	·	•			



Vaccine Effectiveness Summary at-a-glance

Detailed summary available in Appendix 1.

VACCINE	ANY INFECTION	SYMPTOMATIC INFECTION	HOSPITALISATION/ SEVERE DISEASE	DEATH	EFFECTIVENESS 4 to <6 MONTHS	EFFECTIVENESS ≥6 MONTHS
AstraZeneca	45-73% ³⁷⁻⁴³ Single dose 30-67% ^{37,39,42,44}	-39-78% ^{40,45-51*} Single dose: 38-68% ^{45,51-53} Moderate-Severe: 82%; Single dose: 79% ⁴²	84-100% ^{40,43,46-48,51,54-57} Single dose: 49-94% ^{51,52,56,58}	90-100% ^{43,46,47,59}	Infection: 45% ^{49,51} Hospitalisation or death: 42-64% ⁵¹	Infection: 42% ⁴⁹
Bharat Biotech	-	50% ⁶⁰	-	-	-	-
Johnson & Johnson	50-88% ^{40,61-64}	54% ⁴⁰	71-91% ^{25,40,57,62,65}	-	-	Infection: 13% ⁶⁶ Death: <65 years: 73%; ≥65 years: 52% ⁶⁶
Moderna	37-92% ^{40,63,67-71} Single dose: 20-72% ^{44,72}	82-95% ^{40,47,68,69,73} Single dose: 72% ^{73,74}	92-98% ^{40,47,57,65,67-70,72} Single dose: 69-96% ^{44,72}	98% ⁶⁸	-	Infection: 22-58% ^{66,69} Death: <65 years: 82%; ≥65 years: 76% ⁶⁶ Hospitalisation and death: 56% ⁶⁹
Pfizer/BioNTech	55-95% ^{37,38,40,41,52,63,67,71,75–82} Single dose: 36-57% ^{37,39,44}	-18-97% ^{40,45-} 47,49,50,52,73,78,82-84* Single dose: 49-61% ^{45,73,74}	85- 98%40.46,47,52,56,57,65,67,75,78,80,82,83,85- 87 Single dose: 85-94% ^{56,58}	90-100%46,47,59,75,78,82,85,86	Infection: 47-65% ^{49,80}	Infection: 43-64% ^{49,66} Death: <65 years: 84%; ≥65 years: 70% ⁶⁶
Sinopharm	-	90% ³³	-	-	-	-
Sinovac	53-60% ^{43,85}	59 ⁴⁶	73-91% ^{43,46,85}	74-95 ^{%43,46,85}	-	-

* Low/negative values are against the Omicron variant



Vaccine Efficacy/Effectiveness Against Delta at-a-glance

Detailed summary and vaccine efficacy/effectiveness against other variants available in Appendix 2.

	VACCINE EFFECTIVENESS UNLESS OTHERWISE STATED							
VACCINE	ANY INFECTION	SYMPTOMATIC INFECTION	HOSPITALISATION AND DEATH	EFFECTIVENESS 4 to <6 MONTHS	EFFECTIVENESS ≥6 MONTHS			
AstraZeneca	60-67% ^{37-39,42} Single dose 30-67% ^{37,39,42,44}	25-76% ⁴⁷⁻⁵¹ Moderate-Severe: 82%; Single dose: 38-79% ^{42.51}	84-94% ^{47,48,51,55,56,59} Single dose: 49-88% ^{44,51,56}	Infection: 45% ^{49,51} Hospitalisation or death: 64% ⁵¹	Infection: 42% ⁴⁹			
Bharat Biotech	Efficacy: 65% ¹⁸	50% ⁶⁰	-	-	-			
Clover	-	Efficacy: 79% ¹⁹	Efficacy (moderate-severe): Delta: 82% ¹⁹	-	-			
Johnson & Johnson	78% ⁶²	-	71-85% ^{25.62}	-	Infection: 13% ⁶⁶ Death: <65 years: 73%; ≥65 years: 52% ⁶⁶			
Moderna	76-88% ^{67,70,71} Single dose: 60-72% ^{44,72}	95% ⁴⁷	81-98% ^{47,67,70,72} Single dose: 69-96% ^{44,72}	-	Infection: 22-58% ^{66,69} Death: <65 years: 82%; ≥65 years: 76% ⁶⁶ 56% ⁶⁹			
Pfizer/BioNTech	39-93% ^{37,38,67,71,79,80} Single dose: 36-57% ^{37,39,44}	56-90% ^{47,49,50} Single dose: 33% ⁵⁰	75-100% ^{47,56,59,67,79,80} Single dose: 78-94% ^{44,56}	Infection: 53-65% ^{49,80}	Infection: 43-64% ^{47,49,66} Death: <65 years: 84%; ≥65 years: 70% ⁶⁶			

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Vaccine Efficacy/Effectiveness and Immunogenicity of Boosters Against Delta at-a-glance

Booster refers to an additional dose administered at least 5 months after completion of the primary series. Booster efficacy/effectiveness reported relative to primary series (not unvaccinated) unless otherwise specified. Detailed summary available in Appendix 3.

VACCINE		BOOSTER VACCINE EFFECTIVENESS (UNLESS OTHERWISE STATED)					
VACCINE	ACCINE IMMUNOGENICITY ANY INFECTION		SYMPTOMATIC INFECTION	HOSPITALISATION	DEATH		
Johnson & Johnson	Strong antibody response ⁸⁸	-	-	-	-		
Moderna	Strong antibody response ⁸⁸	Effectiveness: 83-96%* ^{71,72}	-	-	-		
Pfizer/BioNTech	Strong antibody response ⁸⁸	Effectiveness: 81%* ⁷¹	Efficacy: 84-95% ^{89,90} Effectiveness: 89-93% ^{*49,50} After AZ primary series: 90-94% ^{*49,50}	93% ⁹¹	81-90% ^{91,92}		
Sinopharm	Strong antibody response but stronger with a Sinopharm recombinant protein vaccine booster ⁹³	90% ³³	-	-	-		
Sinovac	Strong antibody response ⁹⁴	-	-	-	-		

*Relative to unvaccinated



Vaccine Efficacy/Effectiveness Against Omicron at-a-glance

Detailed summary available in Appendix 2.

VACCINE	VACCINE EFFECTIVENESS (AFTER 2 DOSES) UNLESS OTHERWISE STATED				
VACCINE	ANY INFECTION	SYMPTOMATIC INFECTION	HOSPITALISATION AND DEATH		
AstraZeneca	-	No protective effect beyond 15 weeks ⁴⁹ -39% ⁵⁰	-		
Johnson & Johnson	Strong antibody response ⁸⁸	-	Booster: Hospitalisation: 85% (54-95) ⁹⁵		
Moderna	30-37% ^{71,72} Single dose: 20% ⁷² Booster: 64% ⁷²	-	-		
Pfizer/BioNTech	55% ⁷¹ Booster: 55% ⁷¹	-18-88% ^{49,50,96} ≥15 weeks: 34-37% ⁴⁹ Booster: 54-76% ^{49,50} Single dose: 34% ⁴⁹	Hospitalisation: 70% ⁹⁶		



Vaccine Efficacy/Effectiveness in High-Risk Groups at-a-glance

Detailed summary available in Appendix 4.

	VACCINE EFFICACY/EFFECTIVENESS					
VACONE	DIABETES	OBESITY	AT RISK FOR SEVERE COVID-19	ELDERLY*		
AstraZeneca	-	-	Efficacy against symptomatic infection: 76% ¹⁷ Effectiveness of single dose against: Symptomatic infection: 60% ⁴⁵ Effectiveness against: Symptomatic infection: 80% ⁴⁵ Hospitalisation: 63% ⁵⁴	Efficacy against infection: 85% ¹⁷ Effectiveness of single dose against: Symptomatic infection: 53-61% ^{45,52} Hospitalisation: 80% ⁹⁷ Death: 83% ⁷⁴ Effectiveness against: Symptomatic infection: 39-81% ^{43,45,47,48,98} Hospitalisation: 37-94% ^{43,46,98,99} Death: 65-94% ^{43,59,74}		
Bharat Biotech	-	-	Efficacy against infection: 66% ¹⁸	Efficacy against symptomatic infection: 68% ¹⁸		
Gamaleya	-	-	-	Symptomatic infection: 92% ²⁰		
Johnson & Johnson	Efficacy: 23% ²³	Efficacy: 66% ²³	Efficacy: 59% ²³	Efficacy 66% ²³		
Moderna	-	-	Efficacy against symptomatic infection: 84-91% ²⁶ Effectiveness against hospitalisation: 84% (80-87) ⁵⁷	Efficacy against symptomatic infection: 86% ²⁶ Effectiveness against infection: pre-Omicron: 75-97% ^{68,70,72} ; Omicron: 57% ⁷²		
Novavax			Efficacy against infection: 91% ³⁰			
Pfizer/BioNTech	Effectiveness against infection: 82% ⁸¹ 89% ⁸⁶	Effectiveness against infection: 90% ⁸⁶	Efficacy against symptomatic infection: 95% ³¹ Effectiveness of single dose against symptomatic infection: 56% ⁴⁵ Effectiveness against: Infection: 71-90% ^{75,86} Symptomatic infection: 89% ⁴⁵ Hospitalisation: 72-81% ⁷⁵	Efficacy against symptomatic infection: 95-100% ^{31,32} Effectiveness of single dose against: Infection: 76% ⁵² Symptomatic infection: 40-56% ^{45,73} Hospitalisation: 71-81% ^{97,99} Death: 77% ⁷⁴ Effectiveness against: Infection: 70-89% ^{75,81,86,100} Smptomatic infection: 61-93% ^{45,47,73,98} Hospitalisation: 43-93% ⁹⁸⁻¹⁰⁰ Death: 87-98% ^{59,74}		
Sinopharm	-	81% ³³	-	Effectiveness against symptomatic infection 91% ³³		
Sinovac	-	Infection: 75% ³³	Infection: 49% ³³	Infection: 28% ⁴³ Hospitalisation: 32% ⁴³ Death: 34% ⁴³		

*Estimates in those ≥60 years to ≥90 years



Vaccine Efficacy/Effectiveness in Children

VACCINE			
VAGCINE	<12 years	12-18 years	COUNTRIES VACCINATING CHILDREN BTAGE GROUP
Moderna	Well tolerated and produced strong antibody response in 6-11 year olds in USA (Moderna press release) ¹⁰¹	Efficacy in USA, 12-15 years: 96% ¹⁰²	≥12 years: Authorised by EMA, MHRA, TGA ≥12 years: France, Italy, Japan, Australia, Canada, Guinea, Philippines ≥3 years: Colombia
Novavax	7-12 years trial underway in India	Study in 12-18 years has started recruitment	
Pfizer/BioNTech	 5-11 years: Antibody response and safety profile for reactogenicity similar to 16-25 year-olds¹⁰³ Efficacy against symptomatic infection in 5-11 year olds in USA: 90.9% (68.3-98.3)¹⁰⁴ <5 years: Evaluation of a 3 dose schedule underway due to low immunogenicity of 2 doses¹⁰⁵ 	Efficacy in USA, 12-15 years: 100% ¹⁰⁶ ; Up to 4 months after second dose: 100% (87.5-100) ¹⁰⁷ Effectiveness in Israel 12-18 years: Any infection: 90% (88-92); Symptomatic infection: 93% (88-97) ¹⁰⁸ Effectiveness against hospitalisation in USA: 12-18 years: 93% (83-97) ¹⁰⁹ Effectiveness: Symptomatic infection 12-15 years: Single dose 2 weeks: 75.4% (73.9-76.9); Single dose 8-9 weeks: 46.8% (14.9-66.7); 16-17 years: Single dose 2 weeks: 75.9% (74.3-77.4); Single dose 8-9 weeks: 34.3 (30.7 to 37.7); 2 doses 2-9 weeks: 94.6% (92.8-95.9); Hospitalisation: 16-17 years: Single dose: 84.5% (64.6-93.2) ¹¹⁰ USA: Effectiveness against: Hospitalisation: 94% (90-96); ICU admission: 98% (88-100) ¹¹²	 ≥5 years: Authorised by FDA, EMA, TGA ≥5 years: USA, Israel, Canada, Oman, Saudi Arabia, Bahrain, UAE, Costa Rica ≥12 years: Authorised by EMA, MHRA TGA, Medsafe ≥12 years: USA, Canada, UK Israel, France, Spain, Italy, Netherlands, Germany, South Africa, Singapore, Japan, Australia, Estonia, Denmark, Greece, Ireland, Lithuania, Sweden, Finland, Norway, Switzerland, Jordan, Morocco, Egypt, Guinea, Namibia, South Korea, Philippines, Brazil 12-15 years single dose: South Africa 12-17 years single dose: Hong Kong ≥3 years: Colombia
Sinovac	Phase I/II studies complete in 3-17 year olds in China ¹¹³ ; Phase 3 studies underway in Chile, Kenya, Malaysia, the Philippines, and South Africa	-	≥12 years: Indonesia ≥6 years: Chile, El Salvador, Ecuador Indonesia ≥3 years: China, Colombia, Hong Kong
Sinopharm	Phase I/II studies i	≥12 years: Morocco ≥3 years: China, UAE, Venezuela, Argentina, Bahrain	
AstraZeneca	Trials suspended when evidence emerged of the hig	≥3 years: Colombia	
Bharat Biotech	Phase 2/3 tri	-	

Maternal Vaccination

VACCINE	VACCINE EFFICACY/EFFECTIVENESS IN PREGNANT WOMEN	OTHER OUTCOMES
AstraZeneca		In four clinical trials in the UK, Brazil and South Africa, fertility was unaffected by vaccination and there was no increased risk of miscarriage and no instances of stillbirth in women vaccinated before pregnancy ¹¹⁴
Moderna	-	 Following maternal vaccination with mRNA vaccines, 100% of neonates (n=36) had protective antibodies at birth¹¹⁵ Comparison in USA of 35,691 participants who received an mRNA vaccine in pregnancy and nonpregnant women (v-safe registry and VAERS): Injection-site pain more frequent in pregnancy; headache, myalgia, chills, and fever less frequent. Adverse pregnancy and neonatal outcomes similar in pregnant women to studies conducted before the Covid-19 pandemic¹¹⁶
Novavax	-	-
Pfizer/BioNTech	Effectiveness in Israel: Any infection: 96% (89-100); Symptomatic infection: 97% (91-100); Hospitalisation: 89% (43-100) ¹¹⁷ Effectiveness in Israel: Any infection: 78% (57-89) ¹¹⁸	 Following maternal vaccination with mRNA vaccines, 100% of neonates (n=36) had protective antibodies at birth¹¹⁵ NIH-funded MOMI-VAX study will assess antibody responses in 750 pregnant women and 250 women vaccinated with any US-licensed vaccine within 2 months of birth, and their infants¹¹⁹ Comparison in USA of 35,691 participants who received an mRNA vaccine in pregnancy and nonpregnant women (v-safe registry and VAERS): Injection-site pain more frequent in pregnancy; headache, myalgia, chills, and fever less frequent. Adverse pregnancy and neonatal outcomes similar in pregnant women to studies conducted before the Covid-19 pandemic¹¹⁶
Sinopharm		· ·
Sinovac		e de la construcción de la constru



Vaccine Efficacy/Effectiveness Against Asymptomatic Infection at-a-glance

Some of these studies assessed multiple variants, including Delta but none analysed the Delta variant alone. Detailed summary of vaccine efficacy/effectiveness against transmission available in Appendix 5.

VACCINE	VACCINE EFFECTIVENESS UNLESS OTHERWISE STATED
AstraZeneca	Efficacy: 54% ¹⁶
Bharat Biotech	Efficacy: 64 ¹⁸
Johnson & Johnson	Efficacy: 60% ²³
Moderna	73% ⁶⁸
Pfizer/BioNTech	65-92% ^{69,82-84,120,121}



Mixed Dose Vaccine Safety and Immune Responses Mixed vaccine schedules (i.e. delivering different types of vaccine for the first and second dose) could be particularly useful to facilitate better protection against variants of concern and enable vaccination programs to continue if a particular vaccine is unavailable.

SCHEDULE	SAFETY	IMMUNE RESPONSES OR EFFECTIVENESS	COUNTRIES USING SCHEDULE
AZ-PF	 Spain: Similar side effects to those receiving 2 doses of the same vaccine; no safety concerns (not peer reviewed)¹²² UK: Greater systemic side effects (mild-moderate symptoms) following the booster dose than with 2 doses of the same vaccine; no safety concerns¹²³ Germany: greater reactogenicity with first dose of AstraZeneca than with the Pfizer/BioNTech booster¹²⁴ Increased reactogenicity (54.4%; 49.4-59.5) vs AstraZeneca-AstraZeneca (33.5%; 28.0-39.2)¹²⁵ Total adverse event reporting in Korea: 0.28% (vs AZ-AZ: 0.22%; and PF-PF: 0.31%) 	 Spain: ≥8 week dose interval: Stronger immune response following Pfizer/BioNTech than after 2 doses of AstraZeneca vaccine (not peer reviewed)¹²² Spain: 8-12 week dose interval: robust antibody response¹²⁸ UK: 4 week dose interval: stronger antibody and cellular response than after 2 doses of AstraZeneca vaccine¹²⁷ Germany: 9-12 week dose interval: Significantly stronger immune response following Pfizer/BioNTech booster than AstraZeneca, and slightly stronger than after 2 doses of Pfizer/BioNTech (not peer reviewed)¹²⁹ Germany: 4-fold greater immune response than 2 doses of AstraZeneca South Korea: 6-fold greater neutralising antibody response than 2 doses of AstraZeneca Germany: Higher neutralising antibody response against wild-type, Alpha, Beta, Gamma and Delta variants than AZ-AZ¹⁵⁰ 	Canada, Denmark, Finland, France, Germany, Sweden, Norway, Spain and South Korea ¹³¹
PF-AZ	UK: Greater systemic side effects (mild-moderate symptoms) following the booster dose than with 2 doses of the same vaccine; no safety concerns ¹²³ Greater reactogenicity with first of homologous and heterologous prime-boost immunisation with BNT162b2 and ChAdOx1-nCoV19: a prospective cohort study increased reactogenicity (55.2%; 46.1-64.1) vs Pfizer/BioNTech- Pfizer/BioNTech (33.3%; 23.4-44.5) ¹²⁵	UK: 4 week dose interval: weaker antibody response than after 2 doses of Pfizer/BioNTech vaccine (but stronger than after 2 doses of AstraZeneca vaccine) ¹²⁷	-
AZ-mRNA (PF or MO)	-	Effectiveness in Canada: Infection: 90% (89-91); 87% (85-89); Hospitalisation: 99% (98-100); 98% (95-99) ⁴⁸	
Primary series of PF, J&J or MO followed by PF, J&J or MO booster	Reactogenicity for all combinations similar to primary series ⁸⁸	Homologous boosters increased neutralising antibody titres 4.2 to 20-fold; Heterologous boosters increased neutralising antibody titres 6 to 76-fold ⁸⁸	-
AZ, MO and PF		Canada: Trial underway mixing and matching all three vaccines with study arms assessing 4 week and 16 week dose intervals ¹³² Denmark: Vaccine effectiveness against infection: AZ-PF or AZ-MO: 88% (83-92) ¹³³	AstraZeneca followed by either Moderna or Pfizer/BioNTech: Denmark, Finland, France, Germany, Sweden, Norway and Spain ¹³¹
Sinovac-AZ	-		Thailand



Adverse Events Following Immunisation with WHO EUL Vaccines

Adverse events following immunisation (AEFIs) are any reactions occurring after immunisation. They can be either expected or unexpected. The vaccine may not actually cause the AEFI; it may occur coincidentally as millions of people are being vaccinated so some people may get sick after vaccination but this does not necessarily mean that it is due to the vaccine. Special investigations determine whether they are due to the vaccine. Adverse events of special interest (AESIs) are of scientific and medical concern that are found through active surveillance, that have the potential to be causally associated with a vaccine and that need to be carefully monitored and confirmed by further special studies.

For all injectable vaccines, appropriate medical treatment and supervision should always be readily available in case of an anaphylactic event following administration.

	ASTRAZENECA	MODERNA	PFIZER/BIONTECH	JOHNSON & JOHNSON	SINOPHARM	SINOVAC	CLOVER	BHARAT BIOTECH
Adverse events following immunisation (AEFIs)*	Very common (more than 1 in 10 people): headache, nausea, muscle pain, joint pain, injection site tenderness/ pain/ warmth/ itch, fatigue, malaise, fever, chills Common (between 1 in 10 and 1 in 100 people): injection site swelling/ redness ¹³⁴	Injection site pain (92%)/ swelling (15%)/ redness (10%), fatigue (70%), headache (65%), muscle pain (62%), joint pain (46%), fever (16%), chills (45%), nausea/vomiting (23%), axillary swelling/tenderness (20%) ¹³⁵	Very common: headache, muscle pain, joint pain, injection site pain/ swelling, fatigue, fever, chills; Common: nausea, injection site redness ¹³⁴ Uncommon (between 1 in 100 and 1 in 1000 people): lymphadenopathy, insomnia, pain in extremity of vaccinated arm, malaise, injection site itch; Rare: (between 1 in 1000 and 1 in 10,000): acute peripheral facial paralysis ¹³⁶	Injection site pain/ redness/ swelling, headache, fatigue, muscle pain, nausea, fever ¹³⁷	Injection site pain (16%)/ itch (1%)/ swelling (2%)/ redness (1%), fever (4%), fatigue (3%), nausea (1%), headache (1%), diarrhoea (1%), muscle pain (<1%), itch (non- injection site) (1%) ¹³⁸	Fatigue (8.3%), fever (3.3%), diarrhoea (0.8%), nausea (1.7%), headache (2.5%), muscle pain (1.7%), injection site pain (10.0%)/ redness (0%)/ swelling (0%) ¹³⁹	Very common: Injection site pain, fatigue, headache Common: Injection site erythema, myalgia, arthralgia, loss of appetite, nausea, chills Uncommon: Injection site swelling, fever ¹⁹	Common: Injection site pain/redness/ itching, headache, fever, malaise, fatigue, body ache
Adverse events of special interest (AESIs)	Thrombosis with thrombocytopaenia syndrome (TTS) (see page 13 for estimated risk); EMA PRAC: Guillain- Barre syndrome (GBS) ¹⁴⁰ Australia: Guillain-Barre syndrome: 52 cases (10.4 per million doses) ¹⁴¹ Australia: Immune thrombocytopaenia (ITP)	Myopericarditis (most common in younger males) USA VAERS: myocarditis cases per million second doses: 18-24 year males 38.5, females: 5.3; 25-29 year males: 17.2, females: 5.7 ¹⁴² ITP** ¹⁴³	Myopericarditis (most common in younger males) USA VAERS: myocarditis cases per million second doses: 12-15 year males: 39.9, females: 3.9; 16-17 year males: 69.1, females: 7.9; 18-24 year males: 36.8, females: 2.5; 25-29 year males 10.8, females: 1.2 ¹⁴² >1 in 20,000 males under 25 years of age ¹⁴ Israel: 1 to 5 cases of myocarditis per 100,000 persons ^{144,145} ITP**143	TTS (see page 14 for estimated risk) USA: Guillain-Barre Syndrome: 100 preliminary reports of GBS following 12.5 million doses of vaccine administered (mostly males >50 years) ¹⁴⁶	-	-	-	-

*Details for AstraZeneca, Moderna, Pfizer/BioNTech and Johnson & Johnson from product information sheets in SRA countries, based on data from clinical trials; Sinopharm, Sinovac and Bharat Biotech details from published clinical trials **The ITP cases are mostly without the thrombotic events characteristic of TTS



Serious Adverse Events

Caution is required when comparing safety profiles as definitions and reporting systems vary in trials and in particular phase IV studies. Risk of TTS in PICs available in Appendix 6.

VACCINE	VACCINE SAFETY
AstraZeneca	108 SAEs in 12,282 (0.9%) vaccine recipients and 127 in 11,962 (1.1%) placebo recipients; 7 deaths all considered unrelated to vaccination (2 vaccine, 5 placebo) ¹⁶ US Phase III study: No serious safety concerns involving 32,449 participants ¹⁷ (not peer-reviewed) EMA investigation: possible link between the AstraZeneca vaccine and Thromboots with Thrombootspatenia Syndrome (TTS) Blood clots affected the brain (central venous sinus thrombosis, CVST) and abdomen (splanchnic vein thrombosis) There have been reports of 169 cases of CVST and 53 cases of splanchnic vein thrombosis in ~34 million vaccinated people inctrope The EMA confirmed the overall benefits of the vaccine in preventing COVID-19 outweigh the risks of side effects ⁸ UK: Risk factors for death in patients with TTS following the AstraZeneca vaccine: baseline platelet count; and intracranial haemorrhage ¹⁴⁷ TTS reported to occur in ~1 in 50,000 vaccinated adults in Australia ¹ Several countries introduced age recommendations for the vaccine in 0000 vaccinated adults in Australia ¹ Several countries introduced age recommendations for the vaccine in 000 lowing 5 cases of this very rare disorder post vaccination ¹⁵¹ WHO GACVS reports Guillain Bare Syndrome (GBS) rates following adenovirus vector vaccines: EU/EEA: 44; AUS: 9.7; KOR: 0.4; PHL: <1 ¹⁹² Guillain-Barre Syndrome in England: IRR 15-21 days: 2.90 (2.15-3.92); Scotland: IRR 1-28 days: 2.32 (1.08-5.02); following SARS-CoV-21 ¹⁶³ Immune thrombocytopaenia (ITP) in Victoria, Australia: 8 cases per million doses (17 cases; 15 after second dose) (Expected background rate: 20-49 years: 1.9; ≥50 years: 4.1) ¹⁹⁴
Gamaleya	45 SAEs in 16,427 (0.3%) vaccine recipients and 23 in 5,435 (0.4%) placebo recipients; 4 deaths all considered unrelated to vaccination (3 vaccine, 1 placebo) ²⁰
Johnson & Johnson	83 SAEs in 21,895 (0.4%) vaccine recipients and 96 in 21,888 placebo recipients (0.4%); 19 deaths all considered unrelated to vaccination (3 vaccine, 16 placebo) ²³ EMA investigation of 8 reports of TTS. Most cases occurred in women <60 years of age but specific risk factors have not been confirmed ⁹ USA: Cases of TTS per million doses: Overall: 3.1; Female: 5.2; Male: 1.5 ¹⁴² Deaths: 5 (4 female, 1 male)Guillain-Barre Syndrome: 100 preliminary reports of GBS following 12.5 million doses of vaccine administered in USA (mostly males >50 years) ¹⁴⁶ WHO GACVS reports Guillain Barre Syndrome (GBS) rates following adenovirus vector vaccines: USA: 7.8; KOR: 0.9; EU/EEA: A2: 2.1 ¹⁵²
Moderna	153 SAEs in 15,166 (1.0%) placebo recipients and 147 in 15,185 (1.0%) vaccine recipients; 5 deaths considered unrelated to vaccine (2 vaccine, 3 placebo) ³⁶ Anaphylaxis reported in the US at a rate of 2.5 per million doese ¹⁵⁵ No obvious safety signals among pregnant women who received mRNA COVID-19 vaccines in USA ¹¹⁶ USA: Myo/pericarditis reported in 40.6 males and 4.2 females aged 12-29 years per million second doese of mRNA vaccine; and 2.4 males and 1.0 females aged 30+ ¹⁵⁶ USA VAERS: myocarditis cases per million second doses: 18-24 years of mRNA vaccine; and 2.4 males and 1.0 females aged 30+ ¹⁵⁶ USA VAERS: myocarditis cases per million second doses in those aged 18-24 years. Males 198.6; Females 5.6 ¹⁵⁷ Otrain, Canada; Myo/pericarditis cases per million second doses: Myocarditis: 28.3; Pericarditis: 17.2 ¹⁵⁸ EMA PRAC: French study: 57 extra cases of myocarditis no 4.9 per million 16-24-year-old males compared to unexposed; Nordic study: 190 extra cases of myocarditis per million 16-24-year-old males ¹⁵⁹ Myo/pericarditis in Ontario, Canada: Rates in males 18-24 years in the UK; First dose: 8 (4-9) per million; Second dose: 15 (12-16) ¹⁵¹ Myo/pericarditis excess cases in people 40 years in the UK; First dose: 8 (4-9) per million; Second dose: 15 (12-16) ¹⁵¹ Myo/pericarditis following boosters: 4 cases ¹⁶³
Novavax	SAEs at low levels and similar between vaccine and placebo groups ⁴⁶⁴
Pfizer/BioNTech	SAEs and deaths were low and comparable between vaccine and placebo groups (total 37,586 participants) ³¹ Anaphylaxis reported in the US at a rate of 4.7 per million doses ¹⁵⁵ No obvious safety signals among pregnant women who received mRNA COVID-19 vaccines in USA ¹¹⁶ Brazil: SAEs 5. 4/100,000 doses USA: Myopericarditis reported in 40.6 males and 4.2 females aged 12-29 years per million second doses of mRNA vaccine; and 2.4 males and 1.0 females aged 30+ ¹⁵⁶ USA: Myopericarditis reported in 40.6 males and 4.2 females aged 12-29 years per million second doses of mRNA vaccine; and 2.4 males and 1.0 females aged 30+ ¹⁵⁶ USA: VAERS: myocarditis cases per million doses: 12-15 year males: 39.9, females: 3.9, females: 7.9; 18-24 year males: 36.8, females: 2.5; 25-29 years per males: 0.8, females: 1.2 ⁴⁶ Ontario, Canada; Myolpericarditis cases per million second doses in those aged 18-24 years: Males 35.5, females 39.9 ¹⁵⁷ Overall rates in the UK per million second doses: Myocarditis 7.3 (81.1-194.6) cases per million people aged 16-19 ¹⁵⁶ Israel: Myo/pericarditis: 16-9 (69.3-1144 6) cases per million in those aged 18-29 ¹⁶⁵ , 137.3 (81.1-194.6) cases per million people aged 16-19 ¹⁵⁶ Haemorrhagic structs in England: IRR 15-21 days: 1.38 (1.1-21.71) (not replicated in Sootland data) ¹⁵³ Israel: Myo/pericarditis: 16-19 year old males: Second dose: 161 cases per million; Third dose: 52 cases per million 12-29-year-old males ¹⁵⁹ Hong Kong: Myo/pericarditis in 12-17 year-olds: After first dose: 34 (1-19-5) cases per million; Third dose: 56 (24-125); Males, second dose: 373 (270-513) ¹⁶⁸ Myo/pericarditis in 0ntario, Canada: Rates higher with shorter dose interval (i.e. 530 days) ¹⁶⁰ Myocarditis secress cases in people -40 years in the UK. First dose: 2 (1.3) per million; Second dose: 3 (24-1 ³⁵¹) Myo/pericarditis in Ontario, Canada: Rates higher with shorter dose interval (i.e. 530 days) ¹⁶⁰ Myocarditis 5-11 years: 11 confirmed cases following -60 million doses administered ¹⁶⁹ LSA: Myocar
Sinovac	Brazil: SAEs: 79.7/100,000 doses Safety in Chile 6-11 years: Adverse events following 0.011% of doses administered (most common: yomiting, itching, injection site pain and erythema) ⁷⁰



Risk of Myo/Pericarditis in 12-17 Year Olds Following mRNA Vaccines

Estimated number of myo/pericarditis cases that potentially might occur in Pacific Island Countries if all 12-17 year olds received the Pfizer/BioNTech or Moderna vaccine, based on official country population estimates and incidence myo/pericarditis (highest and lowest estimates) in other settings. Nearly all myo/pericarditis cases related to mRNA vaccination are mild and managed conservatively.

COUNTRY	TOTAL POPULATION	ESTIMATED POPULATION AGED 12-17 YEARS*	POTENTIAL NUMBER OF MYO/PERICARDITIS CASES IF ALL 12-17 YEAR OLDS RECEIVED THE PFIZER/BIONTECH VACCINE**	POTENTIAL NUMBER OF MYO/PERICARDITIS CASES IF ALL 12-17 YEAR OLDS RECEIVED THE MODERNA VACCINE ***
American Samoa	55,519	7,217	0.3-1.5	0.1-1.4
Cook Islands	15,300	1,989	<1	<1
Federated States of Micronesia	102,300	13,299	0.5-2.8	0.3-2.6
Fiji	867,000	112,710	4.2-23.9	2.3-22.0
French Polynesia	275,918	35,869	1.3-7.6	0.7-7.0
Guam	159,358	20,717	0.8-4.4	0.4-4.1
Kiribati	113,400	14,742	0.6-3.1	0.3-2.9
Marshall Islands	54,900	7,137	0.3-1.5	0.1-1.4
Nauru	10,900	1,417	<1	<1
New Caledonia	271,407	35,283	1.3-7.5	0.7-6.9
Niue	1,611	209	<1	<1
Northern Mariana Islands	53,883	7,005	0.3-1.5	0.1-1.4
Palau	18,000	2,340	<1	<1
Papua New Guinea	7,744,700	1,006,811	37.7-213.4	20.8-196.8
Samoa	195,979	25,477	1.0-5.4	0.5-5.0
Solomon Islands	642,000	83,460	3.117.7	1.7-16.3
Tokelau	1,160	151	<1	<1
Tonga	99,419	12,924	0.5-2.7	0.3-2.5
Tuvalu	10,507	1,366	<1	<1
Vanuatu	272,173	35,382	1.3-7.5	0.7-6.9
Wallis and Futuna	11,558	1,503	<1	<1
All Pacific Island Countries	10,976,992	1,427,009	53.4-302.5	29.5-279.0

* Based on estimate of 13% of population aged 12-17 years

** Based on estimates of myo/pericarditis occurring in 212 per million second doses of Pfizer/BioNTech in 12-17 year olds in Hong Kong and 37.4 per million second doses in 16-17 year olds in USA (VAERS data)^{142,168} *** Based on estimates of myo/pericarditis occurring in 195.5 per million doses in 18-24 year olds in Canada (data not available for <18 years) and 20.7 per million second doses in 18-24 year olds in USA (VAERS data; not available for <18 years)^{142,160}

Who Can be Vaccinated Based on WHO SAGE Recommendations?

WHO SAGE have made recommendations for use of AstraZeneca, Moderna, Pfizer/BioNTech, Johnson & Johnson, Sinopharm, Sinovac, Bharat Biotech and Novavax vaccines: https://www.who.int/groups/strategic-advisory-group-of-experts-on-immunization/covid-19-materialss

	ASTRAZENECA	MODERNA	PFIZER/BIONTECH	JOHNSON & JOHNSON	SINOPHARM	SINOVAC	BHARAT BIOTECH	NOVAVAX
Minumum Age	18 years	18 years	12 years	18 years	18 years	18 years	18 years	18 years
Maximum Age (SAGE WHO)	None	None	None None		None	None	None	None
Pregnancy	Yes if high priority group & approved by health provider	Yes if high priority group & approved by health provider	es if high priority up & approved by lealth provider		Yes if high priority group & approved by health provider	Yes if high priority group & approved by health provider	Yes if the benefits outweigh the potential risks	Yes if the benefits outweigh the potential risks
Breastfeeding	Yes	Yes	Yes	Yes	Yes	Yes	Yes	
People previously infected with SARS- CoV-2	May delay 6 months; Within 90 days if VOCs associated with reduced effectiveness are circulating (e.g. Beta)	May delay 6 months; Within 90 days if VOCs associated with reduced effectiveness are circulating (e.g. Beta)	May delay 6 months; Within 90 days if VOCs associated with reduced effectiveness are circulating (e.g. Beta)	May delay 6 months; <6 months may be advisable if VOCs with reduced neutralisation activity are circulating	May delay 6 months; <6 months may be advisable if VOCs associated with reduced effectiveness are circulating	May delay 6 months; <6 months may be advisable if VOCs associated with reduced effectiveness are circulating	May delay 6 months; <6 months may be advisable if VOCs associated with reduced effectiveness are circulating	May delay 6 months; <6 months may be advisable if VOCs associated with reduced effectiveness are circulating
Immunocompromised Including HIV	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series	An additional dose at least 1 month, and within 3 months, after completing the primary series
History of Anaphylaxis (Severe Allergy)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)	Yes (unless the allergy is to the vaccine or its components)



Vaccine Development Pipeline

WHO has recommended that vaccines adopted by countries have WHO SAGE EUL and/or Stringent Regulatory Approval. Last updated 6 January 2021.

	NUMBER OF VACCINE CANDIDATES AT EACH PHASE OF DEVELOPMENT						
VACCINE ITTE	PRE-CLINICAL	PHASE I/II	PHASE III	PHASE IV	IN USE*		
RNA	25	12	3	2	2 (Pfizer/BioNTech, Moderna)		
DNA	16	9	4	0	1 (Zydus Cadila Healthcare Limited)		
Vector (non- replicating)	26	8	2	3	4 (CanSino, Gamaleya, Johnson & Johnson, AstraZeneca)		
Vector (replicating)	18	6	1	0	0		
Inactivated	5	8	7	3	9 (Sinopharm/BIBP; Sinopharm/WIBP; Sinovac; Bharat; Chumakov; Research Instutute for Biological Safety Problems; Shenzhen Kangtai Biological Products; Shifa Pharmed, Erciyes University)		
Live-attenuated	2	1	0	0	0		
Protein subunit	72	20	16	1	 10 (Novavax, Vector institute; Anhui Zhifei Longcom Biopharmaceutical Chinese Academy of sciences; Center for Genetic Engineering and Biotechnology; Instituto Finlay de Vacunas, Cuba [peptides 1 and 2]; Medigen Vaccine Biologics, Taiwan; Vaxine Pty Medytox CinnaGen Co, Biological E Ltd, Razi Vaccine and Serum Research Institute) 		
Virus-like particle	20	4	2	0	0		
Other/unknown	32	6	0	0	0		

*Not all vaccines in use have SRA (as recognised by WHO) approval (see Vaccine specifications table and WHO SAGE Emergency Use Listing and prequalification timeline for approval status of vaccines).

Source: London School of Hygiene and Tropical Medicine COVID-19 vaccine tracker.



WHO SAGE Emergency Use Listing and Prequalification Timeline

MANUFACTURER	NAME OF VACCINE	PLATFORM	STATUS OF ASSESSMENT	ANTICIPATED DECISION DATE
Pfizer/BioNTech	BNT162b2/COMIRNATY Tozinameran (INN)	mRNA	Final decision made	Authorised 31/12/20
AstraZeneca	AZD1222	Adenoviral vector Final decision made		SK Bio: Authorised 15/02/21 EU nodes: Authorised 16/04/21 CSL, Australia: Authorised 09/07/21 Daiichi Sankyo, Japan: Authorised 09/07/21
Serum Institute of India	Covishield (ChAdOx1_nCoV19)	Adenoviral vector	Final decision made	Authorised 15/02/21
Sinopharm/Beijing Institute of Biological Products (BIBP)	SARS-CoV-2 Vaccine (Vero Cell), Inactivated (InCoV)	Inactivated In progress		Authorised: 07/05/2021
Sinovac	SARS-CoV-2 Vaccine (Vero Cell), Inactivated	Inactivated	Inactivated In progress	
Moderna	mRNA-1273	mRNA	In progress (to use abridged procedure relying on EMA)	Authorised 30/04/2021
Johnson & Johnson	Ad26.COV2.S	Adenoviral vector	Final decision made	Authorised 12/03/21
The Gamaleya National Center	I Center Sputnik V Adenoviral vector		On hold, awaiting completion of rolling submission	Will be determined when all data are submitted
Bharat Biotech	Covaxin; BBV152	Inactivated	Rolling data assessment started 6 July 2021	Authorised 03/11/2021
CanSino Biologics	Ad5-nCoV	Adenoviral vector	Rolling data assessment started 9 August 2021	ТВС
Novavax	NVX-CoV2373	Protein subunit	Rolling data assessment started 19 August 2021	Authorised 20/12/2021
CureVac	Zorecimeran	mRNA	Expression of interest accepted; Application withdrawn by manufacturer	-
Clover Biopharmaceuticals	SCB-2019 (CpG 1018/Alum)	Protein subunit	Rolling data starting 20 September	ТВС

Source: WHO Guidance Document: Status of COVID-19 Vaccines within WHO EUL/PQ evaluation process. Available at: https://www.who.int/teams/regulation-prequalification/eul/covid-19



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Other resources on COVID-19 vaccines:

WHO COVID-19 vaccines website: <u>https://www.who.int/emergencies/diseases/novel-coronavirus-2019/covid-19-vaccines</u> EMA COVID-19 vaccines website: <u>https://www.ema.europa.eu/en/human-regulatory/overview/public-health-threats/coronavirus-disease-covid-19/treatments-vaccines/covid-19-vaccines</u>

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Appendix 1: COVID-19 Vaccine Effectiveness

VACCINE	SEVERE / HOSPITALISATION / DEATH	INFECTION AND OTHER OUTCOMES
AstraZeneca	Single dose in Scotland: 94% (73-99) ⁵⁸ Risk of death in vaccine failures compared to unvaccinated cases in England reduced by: 55% (41-66) ¹⁷¹ (not peer reviewed) Single dose against hospitalisation in Spain: 92% (46-99) ⁵² Pooled analysis of AstraZeneca, Pizer/BioNTech and Moderna vaccines in Italy: Hospitalisation: 89% (85-91); Death: 93% (89-96) ¹⁷² Chile: Hospitalisation: 100%; ICU admission: 100%; Death: 100% ⁴⁶ Scotland: Hospitalisation: 94% (90-99) ⁵⁴ Netherlands: Hospitalisation: 94% (92-95) ⁵⁷ Spain: Hospitalisation: 95% (79-99) ⁴⁰ Scotland: Death: 91% (86-94) ⁵⁹ British Colombia and Quebec, Canada: Hospitalisation: 94% (90-96); 94% (89-97) ⁴⁸ Scotland and Brazil, <i>respectively</i> (delta variant dominant in Scotlang gamma variant common in Brazil): Hospitalisation or death: Single dose: 49.3% (43.3-54.6); 57.9% (56.9-58.9); 2 doses: 2-3 weeks: 83.7% (79.7-87.0); 86.4% (85.4-87.3); 2 doses 18-19 weeks: 63.7% (59.6-67.4); 42.2% (32.4-50.6) ⁵¹	Pooled analysis of Pfizer/BioNTech and AstraZeneca vaccines in elderly care home residents in UK: Reduction in risk of infection 4 weeks after-single dose: 56% Reduction in risk of infection 5 weeks after single dose: 62% ¹⁷³ Pooled analysis of Pfizer/BioNTech and AstraZeneca vaccines: reduced odds of infection post-second dose: 70% (62-77) ¹⁷⁴ Single dose in Spain: Any infection: 44% (31-54); Symptomatic infection: 50% (37-61) ⁵² Pooled analysis of AstraZeneca, Pfizer/BioNTech and Moderna vaccines in Italy: Infection: 78% (76-79) ¹⁷² Single dose against symptomatic infection in multiple European countries: 68% (39-83) ⁵³ Symptomatic infection in 16-64 years in UK: single dose: 50.2% (40.8-58.2); 2 doses: 78.0% (69.7-84.0) ⁴⁵ Symptomatic infection in Chile: 68.7% (39.8-83.7) ⁴⁶ Spain: Any infection: 54% (48-60); Symptomatic infection: 71% (69-74); 73% (69-77) ⁴⁸ England REACT-1 study: Any infection: 44.8% (22.5-60.7) ⁴¹ India: Any infection: 2 doses: 76.2% (63.7-84.4); 15-19 weeks: 46.5% (46.1-94·0) ⁴² UK: Symptomatic infection: 2-9 weeks: 76.2% (63.7-84.4); 15-19 weeks: 48.5% (44.7-52.0) 20-24 weeks: 45.4% (43.0-47.6); ≥25 weeks: 41.8% (39.4-44.1) ⁴⁹ Brazil: Infection: 72.9% (71.9-73.8); Hospitalisation: 88.0% (86.8-92.); Death: 90.2% (88.5-91.5) ⁴³ England: Omicron: Symptomatic infection: 39% (-5030); Delta: 25.0% (24.3-25.7) ⁵⁰ Scotland and Brazil, <i>respectively</i> (delta variant dominant in Scotland; gamma variant common in Brazil): Symptomatic infection: Single dose: 37.6% (34.6-40.5); 37.6% (37.3-37.9); 2 doses 2-3 weeks: 67.9% (65.9-69.8); 69.8% (69.3-70.4); 2 doses 18-19 weeks: 44.6% (41.5-47.6); 57.7% (55.4-60.0) ⁵¹
Johnson & Johnson	USA: Hospitalisation: 81% (79-84) ⁶² USA: 71% (56-81) ⁶⁵ Netherlands: Hospitalisation: 91% (88-94) ⁵⁷ Spain: Hospitalisation: 74% (43-88) ⁴⁰ Death in veterans in USA: <65 years: 73.0% (52.0-84.8); ≥65 years: 52.2% (37.2-63.6) ⁶⁶ South Africa: Booster against hospitalisation during Omicron wave: 85% (54-95) ⁶⁵	USA: Any infection: 76.7% (30.3-95.3) ⁶¹ USA: Infection: 79% (77-80) ⁶² Efficacy following booster 2 months after first dose: Moderate-Severe infection in USA: 94% (58-100); worldwide: 75% (55- 87) ¹⁷⁵ Spain: Any infection: 50% (42-57); Symptomatic infection: 54% (45-62) ⁴⁰ Symptomatic infection in veterans in USA: 88% (87-89) ⁶³ Any infection in USA: 73.6% (65.9-79.9) ⁹⁴ Infection in veterans in USA: 73.6% (65.9-79.9) ⁹⁴
Moderna	Pooled analysis of AstraZeneca, Pfizer/BioNTech and Moderna in Italy: Hospitalisation: 89% (85- 91); Death: 93% (89-96) ¹⁷² Pooled analysis of Pfizer/BioNTech and Moderna against hospitalisation: 2-12 weeks after second dose: 86% (82%-90%) 13-24 weeks after second dose: 86% (82%-90%) USA: Hospitalisation: 95.8% (90.7-98.1); Death: 97.9% (66.9-99.9) ⁶⁸ Pooled Pfizer/BioNTech and Moderna against hospitalisation in Scotland: 92% (85-99) ⁵⁴ USA: 93% (91-95) ⁵⁵ Spain: Hospitalisation: 98% (82-100) ⁴⁰ Qatar: Decline in effectiveness accelerated beyond the fourth month after the second dose; First month after second dose: 96.0% (93.9-97.4); ≥7 months: 55.6% (-44.3-86.3) ⁶⁹ USA: Hospitalisation: 97.6% (92.8-99.2) ⁷⁰ Moderna and Pfizer/BioNTech in British Colombia and Quebec, Canada: Hospitalisation:98% (97-98); 97% (96-97) ⁴⁶ Death in veterans in USA: <65 years: 81.5% (70.7-88.4); ≥65 years:75.5% (71.8-78.7) ⁶⁶ USA: Hospitalisation with Delta: Single dose: 63.9% (0.0-99.4); 2 doses Delta: 98.0% (87.2- 99.7) ⁷²	Pooled analysis of Moderna and Pfizer/BioNTech vaccines in USA: Infections in nonvaccinated: 234 of 8969; 2.61% (2.29-2.96) Fully vaccinated: 4/8121; 0.05% (0.01-0.13) ¹⁷⁷ Pooled analysis of Moderna and Pfizer/BioNTech vaccines against infection in USA: Fully vaccinated: 90% (68–97) Two weeks after first dose: 80% (59–90) ¹⁷⁸ Pooled analysis of AstraZeneca, Pfizer/BioNTech and Moderna vaccines in Italy: Infection: 78% (76-79) ¹⁷² Single dose against symptomatic disease in the UK: Age 15-39 years: 72% (46-86) ⁷⁴ Minnesota, USA: January to July 2021 (Delta variant <0.7% in May): Infection: 86% (81-91); Hospitalisation: 92% (81-97) July (Delta variant <70%): Infection: 76% (58-87); Hospitalisation: 81% (33-96) ⁶⁷ Infection in USA (98% vaccines used Pfizer/BioNTech and Moderna): Pre-Delta variant predominant: 91% (81–96); Delta variant predominant: 66% (26–84) ¹⁷⁹ Infection in Canada: 1 dose: 72% (63-80); 2 doses: 94% (86-97) ⁷³ USA: Any infection: 82% (78-86); Symptomatic infection: 88.3% (86.1-90.2) ⁶⁸ Spain: Any infection: 82% (76-478.6); 27 months: 22.3% (-1.7-40.7) ⁶⁹ USA: Any infection: 82.7% (76.4-78.6); 77 months: 22.3% (-1.7-40.7) ⁶⁹ USA: Any infection: 82.7% (88.8-89.6); September: 58.0% (56.9-59.1) ⁶⁸ Denmark: Effectiveness against infection: Omicron: First month: 36.7% (48.3-88.7) ⁷⁰ Moderna and Pfizer/BioNTech in British Colombia and Quebec. Canada: Symptomatic infection: 90% (89-90); 88% (88-89) ⁴⁸ Infection in veterans in USA: March: 89.2% (70.4-74.0); Booster first month: 4.2% (-30.8-29.8); Delta: First month: 88.2% (58.1-91.8); Third month: 72.2% (70.4-74.0); Booster first month: 82.8% (58.8-92.9) ⁷¹ USA: Any infection: Single dose: Delta: 60.2% (42.6-72.3); Omicron: 20.3% (0.0-39.8); 2 doses: Delta 14-90 days: 82.8% (69.6-90.3); 181-270 days: 61.4% (56.8-65.5);



		-
		3 doses: Delta ≤2 months: 95.7% (94.2-96.9); >2 months: 90.7% (81.4-95.3);
		Omicron ≤2 months: 63.6% (57.4-68.9); >2 months: 39.1% (3.8-61.5) ⁷²
Pfizer/BioNTech	Severe in Israel: 92% (75-100) ⁸³ Severe/critical in Israel: 97.5% (97.1-97.8) ⁸² Single dose against hospitalisation in Scotland: 85% (76-91) ⁵⁸ Risk of death in vaccine failures compared to unvaccinated cases in England reduced by: Single dose: 44% (32-53) Fully vaccinated: 69% (31-86) ¹⁷¹ (not peer reviewed) Israel: Hospitalisation: 97.2% (96.8-97.5); Death: 96.7% (96.0-97.3) ⁸² Hospitalisation in Spain: 94% (60-99) ⁹² Priority groups in Denmark: Hospitalisation: 93% (89-96); Death: 94% (90-96) ⁷² Pooled analysis of AstraZeneca, Pfizer/BioNTech and Moderna vaccines in Italy: Hospitalisation: 89% (85-91); Death: 93% (80-96) ⁷⁷² USA care facility: Hospitalisation: 94.4 (73.9-98.8); Death 94.4 (44.6-99.4) ⁷⁸ Uruguay: Hospitalisation: 97.4% (96.0-98.8); Death: 94.2 (95.4-96.8) ⁸⁶ Israel: Hospitalisation: 97.4% (97.6-98.8); Death: 91.1% (86-5-94.1) ⁸⁶ Chile: Hospitalisation: 97.2% (96.6-97.6); IC ul admission: 98.3% (97.6-98.8); Death: 91.0% ⁴⁶ Pooled analysis of Pfizer/BioNTech and Moderna against hospitalisation: 2.12 weeks after second dose: 86% (82%-90%) 13-24 weeks after second dose: 86% (82%-90%) 15-24 weeks after second dose: 86% (82%-90%) 15-24 weeks after second dose: 86% (82%-90%) 16-24 weeks after second dose: 86% (82%-90%) 17-24 weeks a	3 doses: Delfa \$2 months: 95.7% (94.2-96.9): >2 months: 90.7% (81.4-95.3); Omicron \$2 months: 83.6% (57.4-68.9); >2 months: 39.1% (8.6-61.5)? Pooled analysis of Moderna and Prizer/BioNTech vaccines in USA: Infections in nonvaccinated: 234 of 9699; 2.61% (2.29-2.96) Fully vaccinated: 4/8121: 0.05% (0.01-0.103) ¹⁷¹ Pooled analysis of Moderna and Prizer/BioNTech vaccines in USA: Fully vaccinated: 90% (68-97); Two weeks after first dose: 80% (59-90) ¹⁷⁶ Symptomatic infection in Israel: 90% (79-95) ¹⁶¹ Israel: Any infection: 95.3% (94.9-95.7); Pooled analysis of Prizer/BioNTech and AstraZeneca vaccines in elderly care home residents in UK: 4 weeks after first dose: 65%, 5 weeks after first dose: 62%, ¹⁷³ Documented infection in Israel: end AstraZeneca vaccines in elderly care home residents in UK: 4 weeks after first dose: 65%, 5 weeks after first dose: 62%, ¹⁷³ Documented infection is ratel: incidence decreased from 9.4 infections per 1,000 HCWs in the week following first dose to <1.0 infection astrael: encidence avaccines: reduced odds of infection post-second dose. ¹⁷⁰ Voled analysis of Prizer/BioNTech and AstraZeneca vaccines: reduced odds of infection post-second dose. ¹⁷⁰ Voled analysis of Prizer/BioNTech and AstraZeneca vaccines: reduced odds of infection post-second dose. ¹⁷⁰ Voled analysis of AstraZeneca. Prizer/BioNTech and Moderna vaccines in taly: Infection: 82% (73-88) ⁵² Infection in priority groups in Demark: 82% (79-84) ¹⁷³ USA: Single dose against infection: 10 2 care facilities: 63% (33-79) ⁷⁷ A care facility residents: >14 days after first dose: 17% (42-8); >7 days after second dose: 64% (14-84) ¹⁸ USA: Single dose against symptomatic disease in the UK: Age 15-39 years: 61% (56-65) ¹⁴ Symptomatic infection in nultiple European countries: single dose: 61% (39-76); 2 doses: 67% (76-79) ¹⁷² Uruguay: Infection: 78.1% (77.0-79.1) ¹⁶ Symptomatic infection in USA (96 years in UK: single dose: 46.8% (27-94.3); 2 doses: 67% (74-93) ⁵³ Symptomatic on Houter 10.7% (16-49 ara): 10.5% (69-63
		Denta: Single dose: 30.1% (22.735.6), 2 dose: 30.3% (23.536.6) Denmark: Effectiveness against infection: Omicron: First month: 55.2% (23.5-73.7); Third month: 9.8% (-10.0-26.1); Booster first month: 54.6% (30.4-70.4); Delta: First month: 86.7% (84.6-88.6); Third month: 72.8% (71.7-73.8); Booster first month: 81.2% (79.2-82.9) ⁷¹
	Uruguay: Hospitalisation: 90.9% (88.6-92.7):	
Sinovac	Death: 94.7% (93.4-95.7) ⁸⁵ Chile: Hospitalisation: 86.0% (85.6-86.5); ICU admission: 89.7% (89.1-90.2); Death: 86.4% (85.6-8 <u>7.2)⁴⁶</u>	Uruguay: Infection: 59.9% (59.1-60.7) ⁸⁵ Symptomatic infection in Chile: 58.5% (58.0-59.0) ⁴⁶ Brazil: Infection: 52.7% (52.1-53.4); Hospitalisation: 72.8% (71.8-73.7); Death: 73.7% (72.3-75.0) ⁴³
Sinopharm	-	Symptomatic infection in Bahrain: 90% (88-91) ³³
Bharat Biotech		India: Symptomatic infection: 50% (33-62) ⁶⁰

Appendix 2: Vaccine Efficacy/Effectiveness Against Variants

Refer to Appendix 1 for vaccine effectiveness results for the Pfizer/BioNTech vaccine in Scotland, England and Israel, where all locations had predominant B.1.1.7 circulation. There are four Variants of Concern listed by WHO.¹⁸¹ The WHO recommends labelling SARS-CoV-2 variants with letters of the Greek alphabet, as in the tables below.¹⁸²

VACONIE	VACCINE EFFICACY/EFFECTIVENESS (EFFECTIVENESS AGAINST INFECTION UNLESS SPECIFIED)						
VACCINE	B.1.1.7 (ALPHA) VARIANT	B.1.351 (BETA) VARIANT	P.1 (GAMMA) VARIANT				
AstraZeneca	UK: 70-4% (43-6-84-5) (vs. 81-5% (67-9-89-4) against wild variant) ^{#3} England: ≥21 days after one dose: 48.7% (45.2-51.9); ≥14 days after two doses: 74.5% (68.4-79.4) ³⁷ Scotiand: 73% (66-78) ^{#3} Canada: Single dose: 64% (60-68) ⁴⁴ UK: Single dose: 63% (55-68); 2 doses: 75% (66-60) ⁵⁹ Severe disease in Canada: Single dose: 85% (81-88) ⁴⁴	South Africa: 10.4% (~76.8 to 54.8) ¹⁸⁴ Study against severe disease underway ²³	Brazil: Symptomatic infection: Single dose: 7.6% (37.3-37.9; 2 doses 2-3 weeks: 60.6% (69.3-70.4); 2 doses 18-19 weeks: 57.7% (55.4-60.0); Hosphalisation or death: Single dose: 57.9% (55.9-56.9); 2 doses 2-3 weeks: 86.4% (65.4-67.3); 2 doses 18-19 weeks: 42.2% (32.4-50.6) ⁵¹				
Johnson & Johnson	• • • • • • • • • • • • • • • • • • •	Moderate to severe/critical: 64.0% (41.2-78.7) Severe/critical: 81.7% (46.2-95.4) ²³ Efficacy against hospitalisation in South Africa: 67% ²⁵	Moderate to severe/critical: 68.1% (48.8-80.7); Severe/critical: 87.6% (7.8-99.7) ²³				
Moderna	Canada: Single dose: 83% (80-86); 2 doses: 92% (86-96) ⁴⁴ Severe disease in Canada: Single dose: 79% (74-83); 2 doses: 94% (89-97) ⁴⁴	-	-				
Novavax	UK: 86.3% (71.3-93.5) (vs. 96.4% (73.8-99.5) against wild variant) ²⁹	South Africa: 51.0% (-0.6 to 76.2) ¹⁸⁵					
Pfizer/BioNTech	Case-control study in Israel: After one dose, vaccinees were disproportionally infected with B.1.1.7 (OR: 26:10) ¹⁸⁸ Qatar: 89.5% (85.9-92.3) ¹⁹⁷ England: ≥21 days after social dose: 47.5% (41.6 to 52.8) ≥14 days after social dose: 37% (91.6-95.3) ²⁷ Sotiland: 92% (90-93) ⁸⁸ Canada: Single dose: 66% (64-68); 2 doses: 78% (66-81) ⁹⁴ UK: Single dose: 59% (52-65); 2 doses: 78% (66-84) ⁹⁸ Severe disease in Qatar: 100% (81.7-100) ⁸⁹⁷ Severe disease in Canada: Single dose: 85% (78-82); 2 doses: 95% (92-97) ⁴⁴	Israel case-control study: Vaccinees infected at least 1 week after the second dose were disproportionally infected with B.1.351 (odds ratio: 8:1) ¹⁹⁶ Qatar: 75.0% (70.5-78.9) ¹⁸⁷ South Africa: 100% (53.5-100) ²² Severe disease in Qatar: 100% (73.7-100) ¹⁸⁷					
Sinovac	Chile: 67% (65-69)* ³³		Brazii: 1 or 2 doses: 37.9% (~46.4-73.6) ¹⁸⁸ Chile: 67% (65-69) ⁴⁵³ Brazii: ≿70 years: 41.6% (26.9-53.3); 70-74 years: 61.8% (34.8-77.7); 75-79 years: 48.9% (23.3-66.0); ≥80 years: 20% (0 6.47.9) ¹⁸⁹				

* While it is known P.1. and B.1.1.7 were circulating at the time of the study, the extent is unknown based on available surveillance



VACCINE	VACCINE EFFICACY/EFFECTIVENESS (EFFECTIVENESS AGAINST IN	FECTION UNLESS SPECIFIED)
VACCINE	B.1.617.2 (DELTA) VARIANT	OMICRON VARIANT
AstraZeneca	England: ≥21 days after one dose: 30.0% (24.3-53.0); ≥14 days after second dose: 67.0% (61.3-71.8) ³⁷ Scotland: 60% (53-66) ³⁸ Canada: Single dose: 67% (44-80) ⁴⁴ UK: Single dose: 46% (53-65); 2 doses: 67% (62-71) ³⁹ Symptomatic infection in England: 67% (64.3-67.0) ¹⁷ Hospitalisation in England: 1 dose; 17% (51-83); 2 doses: 92% (75-97) ⁴⁵ , 93.9% (91.3-95.7) ⁴⁷ Death in England: 94.1% (91.8-95.8) ¹⁷ Severe disease in Canada: Single dose: 88% (60-96) ⁴⁴ Hospitalisation and death in Scotland: 88% (85-90) ⁵⁹ Scotland: Death: 91% (86-94) ⁵⁹ India: Any infection: 2 doses: 61.1% (51-57-27); Single dose: 42.% (31-6-57-7); Moderate-severe: 2 dose: 81.5% (9-99-90): Single dose: 79.2% (46.1-59-0) ⁴² UK: Symptomatic infection: 29 weeks: 76.2% (63.7-84.4); 15-19 weeks: 40.5% (44.7-52.0) 2 doses: 67.9% (65.9-69.8); 2 doses: 37.6% (24.6-40.5); 2 doses: 67.9% (65.9-69.8); 2 doses: 37.6% (24.6-40.5); 2 doses: 67.9% (65.9-69.8); 2 doses: 46.4% (41.5-47.6); Hospitalisation or death: Single dose: 40.3% (43.3-54.6); 2 doses: 57.9% (65.9-69.8); 2 doses: 46.4% (46.% (41.5-47.6); Hospitalisation or death: Single dose: 40.3% (43.3-54.6); 2 doses: 3.7% (79.7-87.0); 2 doses: 48.19 weeks: 46.9% (59.6-67.4) ⁵¹	UK: No protective effect beyond 15 weeks ⁴⁹ England: Symptomatic infection: -39% (-50–30) ⁵⁰ England: Hazard ratio for hospitalisation with Omicoro relative to primary Delta infection in unvaccinated: Single dose: 0.42; 2 doses: 0.37, 2 doses plus Pfizer booster: 0.21 ¹⁹⁰
Johnson & Johnson	Efficacy against hospitalisation in South Africa: 71% ²⁵ USA: Infection: 78% (73-82): Hospitalisation: 85% (73-91) ⁶²	South Africa: Booster against hospitalisation: 85% (54-95) ⁹⁵
Moderna	Canada: Single dose; 72% (57-82) ⁴⁴ Minnesota, USA: 70% (56-87) ⁶⁷ England: 95.2% (94.4-95.9) ⁴⁷ Severe disease in Canada: Single dose; 96% (72-99) ⁴⁴ Severe disease in Canada: Single dose; 96% (72-99) ⁴⁴ Severe disease in Minnesota: 81% (33-96) ⁶⁷ Hospitalisation in England: 97.5% (82.3-99.7) ⁴⁷ Pooled Pfizer/BiOHZech and Moderna against hospitalisation and death in Scotland: 91% (88-93) ⁵⁵ USA: Infection: 86.7% (94.3-86.7), Hospitalisation: 97.6% (92.8-99.2) ⁷⁰ Denmark: First month: 88.2% (83.1-91.8); Third month: 72.2% (70.4-74.0); Booseff first month: 82.8% (58.8-92.9) ⁷¹ USA: Infection: Single dose, 60 2% (42.6-72.3); 2 doses: 14% (60.6-80.3); 181-270 days: 61.4% (56.8-5.5); 3 doses: s2 months: 95.7% (94.2-96.9); s2 months: 90.7% (67.2-99.7) ⁷²	Denmark: First month: 36.7% (-69.9-76.4); Third month: 4.2% (-30.8-29.8) ⁷¹ USA: Infection: Single dose: 20.3% (0.0-39.8); 2 doses: 14-90 days: 30.4% (5 0-49.0); 181-270 days: 0.0% (0.0-1.2); 3 doses: ≤2 months: 63.6% (57.4-68.9); >2 months: 39.1% (3.8-61.5) ⁷²
Pfizer/BioNTech	England: ≥21 days after one dose: 36.5% (22.7-46.4); ≥14 days after second dose: 88.0% (85.3-90.1) ³⁷ Scotland: 79% (75-62) ³⁸ Canada: Single dose: 56% (45-64), 2 doses: 87% (64-95) ⁴⁴ Effectiveness in Israel: Infection: 64%; Symptomatic illness: 64% ¹⁹¹ Israel 6m after roll out: 39.0% (9.0-59.0) ⁷⁹ Minnesota, USA 42% (15-452) ⁶⁷ UK: Single dose: 57% (50-63); 2 doses: 80% (77-83) ³⁹ England: 89.8% (89.6-90.0) ⁴⁷ Hospitalisation in England: 1 dose: 94% (46-99); 2 doses: 95% (66-99) ⁵⁶ , 96.00,0 ¹⁷ Hospitalisation in Israel: 99.2% (65-99.2) ⁴⁷ Severe disease in Canada: Single dose: 78% (65-86) ⁴⁴ Hospitalisation in Israel: 93% ¹⁶¹ Severe disease in Canada: Single dose: 78% (65-86) ⁴⁴ Hospitalisation in Israel: 91.4% (82.5-95.7) ⁷⁹ Severe disease in Israel: 91.4% (82.5-95.7) ⁷⁹ Severe disease in Canada: 75% (24-94) ⁵⁷ Pooled Pfizer/BioNTech and Moderna against hospitalisation and death in Scotland: 91% (88-93) ⁵⁶ Scotland: Death: 90% (80-49) ⁵⁹ UK: Symptomatic infercion: 29.2% (65-79.5); 15-19 weeks: 72.2% (71-73.4); 20-24 weeks: 64.8% (62.6-66.9); ≥25 weeks: 63.5% (614-65.5) ⁴⁹ Denmark: First month: 86.7% (84.6-88.6); Third month: 72.8% (71-73.4); Booster first month: 81.2% (79.2-82.9) ⁷¹	 UK: Symptomatic infection: Single dose: 34.2% (-3.5-58.1); 2 doses: 2-9 weeks: 88.0% (65.9-95.8); 15-19 weeks: 34.1% (9.7-52.0); Booster after PF primary series: 75.5% (56.1-86.3); Booster after AZ primary series: 71.4% (41.8-86.0)⁴⁹ South Africa: Symptomatic infection: 33%; Hospitalisation: 70% (risk of hospital admission 29% lower for Omicron compared to first wave in mid-2020) adjusted for vaccination status)⁴⁶ England: Hazard ratio for hospitalisation with Omicron relative to primary Delta infection in unvaccinated: Single dose: 0.66: 2 doses: 0.26: 3 doses: 0.37¹⁶⁹ Denmark: First month: 55.2% (23.5-73.7); Third month: 9.8% (-10.0-26.1); Booster first month: 54.6% (30.4-70.4)⁷¹
Sinovac	China (combined Sinovac and Sinopharm): Single dose: 13.8% (-60.2-54.8);; 2 doses: 59.0% (16.0-81.6) Severe disease: 100% ¹⁹²	• • • • • • • • • • • • • • • • • • •
Sinopharm	China (combined Sinovac and Sinopharm); Single dose: 13.8% (+60.2-54.8); 2 doses: 59.0% (+6.0-81.6) Severe disease: 100%; ^{rea}	
Bharat Biotech	Efficacy against infection in India: 65.2% (33.1-83.0) ¹⁸ India: Symptomatic infection: 50% (33-62) ⁸⁰	
Clover	Efficacy in Philippines, Colombia, Brazil, South Africa and Belgium: Symptomatic infection: 78.7% (57.3-90.4); Mod-Severe: 81.7% (35.9- 96 8) ¹⁹	



Appendix 3: Vaccine Efficacy/Effectiveness and Immunogenicity of Boosters Against Delta

Booster refers to an additional dose administered at least 5 months after completion of the primary series. Booster efficacy/effectiveness reported relative to primary series (not unvaccinated) unless otherwise specified.

VACCINE		VACCINE EFFECTIVENESS UNLESS OTHERWISE STATED					
VACCINE	IMMONOGENICITY	ANY INFECTION	SYMPTOMATIC INFECTION	HOSPITALISATION	DEATH		
Johnson & Johnson	Strong antibody response ⁸⁸			-			
Moderna	Strong antibody response ⁸⁸	USA (relative to unvaccinated): Delta: ≤2 months: 95.7% (94.2- 96.9); >2 months: 90.7% (81.4- 95.3) ⁷²	-	-			
Pfizer/BioNTech	Strong antibody response ⁸⁸	Israel; Infection ~10 times lower in boosted group (range across five age groups, 9.0 to 17.2) ⁹² Denmark (relative to unvaccinated): First month: 54.6% (30.4-70.4) ⁷¹	Efficacy: ≥6 months (median 10.8 months) in USA, South Africa, Brazil: 95.3 (89.5- 98.3) ⁸⁹ UK: Booster relative to primary series at least 4.6 months earlier: PF primary: 84.4% (82.8-85.8); AZ primary: 87.4% (84.9-89.4); Relative to unvaccinated: PF primary: 93.1% (91.7-94.3) ⁸⁰ UK: relative to unvaccinated: PF primary: 93.6% (92.0-93.1); AZ primary: 93.8% (93.2-94.3) ⁴⁹ England: PF primary: 88.6% (88.1-89.1) AZ primary: 89.7% (88.9-90.4) ⁵⁰	Israel ≥5 months: 93% (88- 97) ⁹¹ Israel: Severe disease: ≥60 years: 17.9 times lower in boosted group (15.1-21.2); 40-59 years: 21.7 times lower (10.6-44.2) ⁹²	Israel ≥5 months: 81% (59-97) ⁹¹ Israel ≥5 months: 90% (86-93) ¹⁹³ Israel ≥60 years: mortality 14.7 times lower in boosted group: (10.0-21.4) ⁹²		
Sinovac	Strong antibody response ⁹⁴	-	-	-			



Appendix 4: Vaccine Efficacy/Effectiveness in High-Risk Groups

MACONE		VACCINE EFFICACY UNLESS OTHERWISE STATED					
VACCINE	DIABETES	OBESITY	AT RISK FOR SEVERE COVID-19	ELDERLY			
AstraZeneca	-		76% against symptomatic infection in a sample where 60% had comorbidities, including diabetes, severe obesity or cardiac disease ¹⁷ (not peer- reviewed) Effectiveness against symptomatic infection in the UK in those with comorbidities and ≥65 years: Single dose: 60.0% (46.5-70.1); 2 doses: 79.7% (61.6-89.3) ⁴⁵ Hospitalisation in Scotland: 63% (46-75) ⁵⁴	In ≥65 years: 85% ¹⁷ (not peer-reviewed) Effectiveness against hospitalisation at 28-34 days after a single dose (pooled analysis of AstraZeneca and Pfizer vacines): 18-64 years: 85% (68-93); 65-79 years: 79% (77-95); ≥80 years: 81% (65-90) ⁵⁸ Effectiveness of single dose against hospitalisation in England: ≥80 years: 73% (60-81) ⁵⁰ Effectiveness in England: Symptomatic infection ≥70 years: 73% (27-90); Hospitalisation ≥80 years: 37% (3-59) ⁵⁸ Hospitalisation following single dose in the UK: ≥80 years: 50% (34-62) ⁵² Single dose in Spain: ≥60 years: 53% (19-72) vs. 18-59 years: 50% (34-62) ⁵² Effectiveness against death in the UK; ≥65 years: single dose: 83% (78-86); Two doses: 94% (80-98) ⁷⁴ Effectiveness against death in the UK; ≥65 years: single dose: 60.9% (49.0-70.0); 2 doses: 76.4% (58.8-86.5) ⁴⁵ Effectiveness against death in Scotland ≥60 years: 90% (84-94) ⁵⁶ British Colombia and Quebec, Canada: Symptomatic in8c+09 years: 73% (42-88); 81% (74-86) ⁴⁸ Brazil: Infection: 80-89 years: 73.1% (71-274.9); 290 years: 90.9% (87.7-91.7); 290 years: 65.4% (46.1-77.8) ⁴³			
Gamaleya	-	-	-	Symptomatic infection >60 years: 91.8% (67.1–98.3) ²⁰			
Johnson & Johnson	Moderate to severe/critical: 23.0% (-90.1-69.8) ²³	Moderate to severe/critical: 65.9% (47.8-78.3) ²³	Moderate to severe/critical: With any comorbidity: 58.6% (40.6-71.6) ²³ No comorbidity: 68.8% (59.0-76.6) ²³	Moderate-severe/critical disease ≥28 post vaccination: 18-59 years: 66.1% (53.3-75.8) 60+ years: 66.2% (36.7-83.0) ²³			
Moderna	-		Symptomatic infection, comorbidities, including diabetes and obesity: In low risk: 95.1% (89.6-97.7) In high risk: 90.9% (74.7-96.7) ²⁶ Pooled Pfizer/BioNTech and Moderna against hospitalisation in Scotland: 72% (51-84) ⁵⁶ Netherlands: Hospitalisation in a population at high risk for severe COVID-19: 84% (80-87) ⁵⁷	Symptomatic infection: 18-64 years: 95.6% (90.6-97.9) ≥65 years: 86.4% (61.4-95.2) ³⁶ Pooled Moderna and Pfizer vaccines against hospitalisation ≥65 years: 94% (49-99) ¹⁹⁴ Infection in Canada: 1 dose ≥70 years: 54% (31-69); 2 doses ≥70 years: 95% (83-98) ⁷³ Pooled Moderna and Pfizer vaccines in Portugal: Hospitalisation 65-79 years: 94% (88-97); ≥80 years: 82% (72-89); Death 65-79 years: 96% (92-98); Death ≥80 years: 81% (74-87) ¹⁹⁵ USA: Hospitalisation: ≥65 years: 75.2% (59.6-84.8) vs 18-64 years: 87.9% (85.5-89.9) ⁷⁰ USA: Effectiveness against infection: Delia ≥65 years: 97.2% (94.1-98.7); Omicron ≥65 years: 57.1% (14.2-78.6) ⁷²			
Pfizer/BioNTech	Effectiveness in Israel: Diabetes or cardoiovascular disease: 82% (62-92) ⁸¹ Effectiveness against infection in Israel: (88-9% (87-3-90-2) ⁹⁸	Effectiveness against infection in Israel: (89·7% (88·6-90·7) ⁹⁶	Symptomatic infection: With any comorbidity or obesity: 95.3% With no comorbidity 94.7% ³¹ Denmark: Infection: 71% (58-80); Hospitalisation: 81% (49-93) ⁷⁵ Effectiveness against infection in Israel: Hypertension: (89.7% (88.6-91.7) ⁸⁶ Effectiveness against symptomatic infection UK in those with comorbidities and 265 years: Single dose: 56.4% (46.2-64.6) 2 doses: 88.5% (81.5-20.9) ⁴⁶ Pooled Pfizer/BioNTech and Moderna against hospitalisation in Scotland: 72% (51-84) ⁵⁴	Effectiveness against infection ≥75 years: 96.2% (76 9-99.9)»2 Mymptomatic infection: >55 years: 93.7% (80.6-98.8); >65 years: 94.7% (66.7-99.9); >75 years: 100% (-13.1-100) ³¹ Effectiveness against hospitalisation 28-34 days after a single dose (pooled analysis of AstraZeneca and Pfizer vacines): 18-64 years: 85% (68.93); 65-79 years: 79% (17-95); >80 years: 81% (65-90) ³⁸ England 80-83 years: Documented infection: 70.1% (56.1-80.1) Hospital attendance: 78.9% (60.0-89.9); Hospital admission: 75.6% (52.8-87.6) ¹⁰⁰ Reduction in incidence of infection in vaccinated people aged >60 years and urvaccinated people aged 20-39 years, respectively: Documented infection: 45% versus 28%; Hospitalisation: 66% versus 22% ¹⁰⁹ Pooled Moderna and Pfizer vaccines against hospitalisation ≈65 years: 94% (49-99) ¹⁹⁴ Effectiveness in England: Symptomatic infection ≥70 years: 61% (51-69); Hospitalisation ≥80 years: 43% (33-52); Death ≥80 years (vaccine failure vs non-vaccinated); 51% (37-62) ¹⁰⁰ Fully vaccinated: 93% (89-95) ¹⁰⁰ (61-69); Hospitalisation ≥80 years: 43% (33-52); Death ≥80 years (vaccine failure vs non-vaccinated); 51% (37-62) ¹⁰⁰ Fully vaccinated: 93% (89-95) ¹⁰⁰ (not peer reviewed) Effectiveness in Israel: 65-74 years: 82% (63-92); ≥75 years: 82% (61-91) ⁸¹ Hospitalisation following single dose in the UK: ≥80 years: 71.4% (43.1-86.2) ¹⁰⁷ Single dose in Spain: ≥60 years: 76% (55-87) ys. 18-59 years: 85% (74-91) ⁵² Effectiveness against infection in Israel: >77% (50-89) ¹⁵⁶ Effectiveness against infection in Israel: >77% (50-89) ¹⁵⁶ Effectiveness against symptomatic infection in Israel: >70 years: 89 ¹ % (72-81); Two doses: 98% (94-99) ⁷⁴ Effectiveness against symptomatic infection in Israel: >70 years: 89 ¹ % (72-81); Two doses: 98.7% (80.1-91.1) ⁴⁵ Infection in Canada: 1 dose >70 years: 40% (29-50); 20 sees >70 years: 99.1% (82-9) ⁷⁴ Effectiveness against symptomatic infection in the UK, ≥65 years: single dose: 77% (72-81); Two doses: 98.7% (80.1-91.1) ⁴⁵ Infection in Scala z			
Novavax			Any infection with comorbidity, age ≥65 years or frequent COVID-19 exposure in USA and Mexico: 91.0% (83.6-95.0) ³⁰				
Sinovac	-	74.9% (53.7-86.4) ³³	Any comorbidity: 48.9% (26.6-64.5) ³³	Brazil: Infection: 80-89 years: 55.6% (53.3-57.8); ≥90 years: 27.6% (20.5-34.0); Hospitalisation: 80-89 years: 63.5% (60.4-66.3); ≥90 years: 31.8% (21.9-40.5); Death: 80-89 years: 67.2% (63.6-70.5); ≥90 years: 33.6% (21.9-43.5) ⁴³			
Sinopharm	-	80.7% (56.7-91.4) ³³		Effectiveness against symptomatic infection in Bahrain: ≥60 years: 91% (87-94) ³³			
Bharat Biotech	-	-	Any infection with comorbidity: 66.2% (33.8-84.0) ¹⁸	Symptomatic infection in India: ≥60 years: 67.8% (8.0-90.0) vs 18-59 years: 79.4% (66.0-88.2)¹8			



Appendix 5: Vaccine Efficacy/Effectiveness Against Transmission There are limitations related to the analysis and comparison of transmission data between studies and vaccines. Criteria for testing vary between studies and may include, for example, random testing, testing at defined

intervals, or retrospective serology.

VACCINE	EFFICACY/EFFECTIVENESS AGAINST ASYMPTOMATIC INFECTION	OTHER OUTCOMES
AstraZeneca	EFFICACY (UK only): 22-2% (-9-9-45-0); Symptomatic and asymptomatic combined (UK, SOUTH AFRICA & BRAZIL): 54.1% (44.7-61.9) ¹⁶ ENGLAND: Hazard ratio for single dose in vaccinated vs. unvaccinated care facility residents: 0.32 (0.15-0.66) ¹⁹⁷ ; Odds ratio for household contacts of vaccinated vs non-vaccinated health workers testing positive: 0.52 (0.43-0.62) ¹⁹⁸ Effectiveness against household transmission of Delta: 42% (14-69); Vaccination of household contacts against infection from a symptomatic household index case: 14% (-5-46) ¹⁹⁹ UK: Regular testing of randomly selected households: 79% (65-88) ²⁰⁰ ; Single dose against symptomatic and asymptomatic infection: 60% (49–68) ²⁰¹ NETHERLANDS: Effectiveness against transmission (secondary attack rate among household contacts): 58% (-12-84) ⁸⁷	SCOTLAND: POOLED ANALYSIS OF PFIZER/BIONTECH AND ASTRAZENECA: Hazard ratio for household contacts of vaccinated vs non-vaccinated health workers testing positive: 0.70 (0.63-0.78) ²⁰² PF and AZ: Secondary attack rates in household contacts in the UK: Vaccinated contacts: 25% (18-33); Unvaccinated contacts: 38% (24-53); Unvaccinated index cases: 25% (15-31); Vaccinated index cases: 25% (15-35) ²⁰³
Bharat Biotech	EFFICACY IN INDIA: Asymptomatic: 63.6% (29.0-82.4); Symptomatic and asymptomatic combined: 68.8% (46.7-82.5) ¹⁸	•
Johnson & Johnson	EFFICACY (multiple countries): Asymptomatic infection: 59.7% (32.8-76.6) ²³ UK: Single dose against symptomatic and asymptomatic infection: 60% (49–68) ²⁰¹ Netherlands: Effectiveness against transmission (secondary attack rate among household contacts): 77% (6-94) ⁹⁷	USA (Kentucky): OR for reinfection in unvaccinated vs vaccinated with Johnson & Johnson, Moderna or Pfizer/BioNTech): 2.34 (1.58–3.47) ²⁰⁴
Moderna	USA: Asymptomatic infection: 72.7% (53.4-84.0) ⁸⁸ USA: POOLED ANALYSIS OF PFIZER/BIONTECH AND MODERNA: 88.7% (68.4-97.1 ³⁰⁵ ; 90% (68%-97) ³⁰⁶ ; single dose: 80% (59-90) ³⁰⁶ ; Relative risk of infection in asymptomatic pre-surgical patients >10 days after first dose compared to unvaccinated residents: 0.21 (0.12-0.37) ³⁰⁷ ; Incident cases in unvaccinated nursing home residents decreased from 4.3% within 14 days of the first vaccination clinic to 0.3% after 42 days ³⁰⁸ MODELLING: Reduced potential for transmission: at least 61% ²⁰⁹ UK: Single dose against symptomatic and asymptomatic infection: 60% (49–68) ³⁰¹ Netherlands: Effectiveness against transmission (secondary attack rate among household contacts): 88% (50-97) ⁸⁷ USA: 63.0% (56.6-68.5) ²² Qatar: First month after second dose: 73.1% (70.3-75.5); declining to no evidence of any effect by 4 months post-vaccination ⁶⁹	USA (Kentucky): OR for reinfection in unvaccinated vs vaccinated with Johnson & Johnson, Moderna or Pfizer/BioNTech): 2.34 (1.58–3.47) ²⁹⁴
Pfizer/BioNTech	ENGLAND: 86% (76-97) 7 days after 2 doses; 72% (58-86) 21 days after 1 dose ¹²⁰ Effectiveness against household transmission of Delta: 31% (~3-61); Vaccination of household contacts against infection from a symptomatic household index case: 24% (~2-64) ¹⁹⁹ ISRAEL: 92% (88-95) ⁸⁵ ; 91.5% (90.7-92.2) ⁸² ; 95% (45-79%) ¹²¹ ; single dose: 75% (72-84) ²¹⁰ ; Effectiveness against transmission: 88.5% (80.7-92.2) ⁸² ; 91.5% (90.7-92.2) ⁸² ; 95% (45-79%) ¹²¹ ; single dose: 75% (72-84) ²¹⁰ ; Effectiveness against transmission: 88.5% (68.4-97.1) ²⁰⁵ ; 90% (68%-97) ²⁰⁶ ; single dose: 75% (59-90) ²⁰⁶ ; Relative risk of infection in asymptomatic pre-surgical patients >10 days after first dose compared to unvaccinated residents: 0.21 (0.12-0.37) ²⁰⁷ ; Incident cases in unvaccinated nursing home residents decreased from 4.3% within 14 days of the first vaccination clinic to 0.3% after 42 days ²⁰⁸ UK: single dose: 4-fold decrease in risk amongst HCWs ≥12 days post-vaccination ²¹³ ; Regular testing of randomly selected households: 80% (73-85) ²⁰⁰ ; Single dose against transmission to unvaccinated household contacts: 2 weeks after first dose: 8.7% (-28.9-35.4); 10 weeks after first dose: 42.9% (22.3-58.1) ²¹⁴ Netherlands: Effectiveness against transmission to unvaccinated household contacts of vaccinated cases: 42.9% (22.3-58.1) ²¹⁴	 ISRAEL: Lower viral load in vaccine failure cases 12-37 days after the first dose of vaccine compared to within the first 11 days, indicating potentially lower infectiousness²¹⁵; Data from 223 communities: strong correlation between community vaccination rate and a later decline in infection among children under 16 years of age who were unvaccinated²¹⁶; Substantially decreased viral load for infections occurring 12-37 days after the first dose of vaccine, indicating likely lower infectiousness²¹⁵ Detectable transmission in long-term care facilities in Spain reduced by 90% (76-93)²¹⁷ ENGLAND: Odds ratio for household contacts of vaccinated health workers vs non-vaccinated health workers testing positive: 0.54 (0.47-0.62)¹⁹⁸ SCOTLAND: POOLED ANALYSIS OF PFIZER/BIONTECH AND ASTRAZENECA: Hazard ratio for single dose in vaccinated vs. unvaccinated care facility residents: 0.35 (0.17-0.71)¹⁹⁷ USA (Kentucky): OR for reinfection in unvaccinated vs vaccinated with Johnson & Johnson, Moderma or Pfizer/BioNTech; 2.34 (1.58–3.47)²⁰⁴ PF and AZ: Secondary attack rates in household contacts in the UK: Vaccinated contacts: 25% (18-33); Unvaccinated index cases: 23% (15-31); Vaccinated index cases: 25% (15-35)²⁰³



Appendix 6: Risk of Rare Unusual Blood Clotting with Low Platelets (Thrombosis with Thrombocytopaenia Syndrome – TTS)

Estimated number of TTS that potentially might occur in Pacific Island Countries if all adults received the AstraZeneca or Johnson & Johnson vaccines, based on most recent official estimate of the adult population in each country and the incidence of these events in Europe and Australia.

COUNTRY	TOTAL POPULATION	ESTIMATED POPULATION AGED 18 YEARS AND OVER*	POTENTIAL NUMBER OF TTS CASES IF ALL ADULTS IN EACH COUNTRY RECEIVED ASTRAZENECA VACCINE**	POTENTIAL NUMBER OF TTS CASES IF ALL ADULTS IN EACH COUNTRY RECEIVED JOHNSON & JOHNSON VACCINE***
American Samoa	55,519	33,311	<1	<1
Cook Islands	15,300	9,180	<1	<1
Federated States of Micronesia	102,300	61,380	0.6-1.2	<1
Fiji	867,000	520,200	5.2-10.4	1.6
French Polynesia	275,918	165,551	1.7-3.3	<1
Guam	159,358	95,615	1.0-1.9	<1
Kiribati	113,400	68,040	0.7-1.4	<1
Marshall Islands	54,900	32,940	<1	<1
Nauru	10,900	6,540	<1	<1
New Caledonia	271,407	162,844	1.6-3.3	<1
Niue	1,611	967	<1	<1
Northern Mariana Islands	53,883	32,330	<1	<1
Palau	18,000	10,800	<1	<1
Papua New Guinea	7,744,700	4,646,820	46.5-92.9	14.6
Samoa	195,979	117,587	1.2-2.4	<1
Solomon Islands	642,000	385,200	3.9-7.7	1.2
Tokelau	1,160	696	<1	<1
Tonga	99,419	59,651	0.6-1.2	<1
Tuvalu	10,507	6,304	<1	<1
Vanuatu	272,173	163,304	1.6-3.3	<1
Wallis and Futuna	11,558	6,935	<1	<1
All Pacific Island Countries	10,976,992	6,586,195	65.9-131.7	20.8

* Based on estimate of 60% of population aged ≥18 years²¹⁸

** Based on estimates of TTS occurring in ~1 in 100,000 vaccinated adults by the European Medicines Agency and ~1 in 50,000 in Australia^{10,11}

*** Based on estimates of TTS occurring in ~1 in 319,000 vaccinated adults in USA (may be an underestimate as only cerebral venous sinus thrombosis are reported)¹³



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