

Significant events in COVID-19 vaccination practice in Australia

Year	Month	Intervention
2020	October	Vaxzevria (AstraZeneca adenovirus viral vector) COVID-19 vaccine granted provisional determination by the Therapeutic Goods Administration (TGA), making it eligible for provisional registration
	October	Comirnaty (mRNA) COVID-19 vaccine granted provisional determination by the TGA, making it eligible for provisional registration
	November	COVID-19 vaccine Janssen (adenovirus viral vector) granted provisional determination by the TGA, making it eligible for provisional registration
2021	January	Nuvaxovid (protein-based) COVID-19 vaccine granted provisional determination by the TGA, making it eligible for provisional registration Comirnaty provisionally registered for use in individuals aged ≥ 16 years
	February	<p>Vaxzevria provisionally registered for use in individuals aged ≥ 18 years</p> <p>Nationally funded COVID-19 vaccination program began. The rollout was carried out in phases, with population groups prioritised according to ATAGI advice</p> <p><i>Phase 1a</i></p> <ul style="list-style-type: none"> Quarantine and border workers Frontline healthcare workers Aged care and disability care staff Aged care and disability care residents <p><i>Phase 1b</i></p> <ul style="list-style-type: none"> Healthcare workers currently employed and not included in Phase 1a Household contacts of quarantine and border workers Critical and high-risk workers who are currently employed, including defence, police, fire, emergency services and meat processing Essential outbound travellers with a travel exemption Elderly people aged ≥ 80 years Elderly people aged ≥ 70 years Aboriginal and Torres Strait Islander people aged ≥ 50 years Adults with an underlying medical condition or significant disability <p><i>Phase 2a</i></p> <ul style="list-style-type: none"> People aged ≥ 50 years Aboriginal and Torres Strait Islander people aged 16–49 years Other critical and high-risk workers <p><i>Phase 2b</i></p> <ul style="list-style-type: none"> People aged 16–49 years <p><i>Phase 3</i></p> <ul style="list-style-type: none"> People aged less than 16 years

Year	Month	Intervention
2021 (cont.)	March	Phase 1b began
	April	Recommendations for use of Vaxzevria changed due to an association with thrombosis with thrombocytopenia syndrome (TTS): <ul style="list-style-type: none"> • Comirnaty is preferred over Vaxzevria in individuals aged <50 years • Those who have received their first dose of Vaxzevria with no TTS could still receive their second dose of Vaxzevria
	May	Phase 2a began in people aged ≥50 years
	June	Recommendations for use of Vaxzevria changed due to further emerging data on the association with TTS: <ul style="list-style-type: none"> • Comirnaty preferred over Vaxzevria in individuals aged <60 years
	June	COVID-19 vaccine Janssen provisionally registered for use in individuals aged ≥18 years
	June	Spikevax (mRNA) COVID-19 vaccine granted provisional determination by the TGA, making it eligible for provisional registration
	July	Comirnaty indication age extended to include use in individuals aged ≥12 years
	August	Spikevax provisionally registered for use in individuals aged ≥18 years
	September	Spikevax indication age extended to include use in individuals aged ≥12 years
	October	<p>A 3rd primary dose of COVID-19 vaccine recommended in severely immunocompromised populations 2–6 months after the 2nd dose of vaccine. An mRNA vaccine is preferred to Vaxzevria for this 3rd dose</p> <p>Comirnaty provisionally registered for a booster dose 6 months after the 2nd dose in immunocompetent individuals aged ≥18 years</p> <p>A booster dose was recommended for immunocompetent individuals aged ≥18 years who had their primary COVID-19 vaccine course ≥6 months ago. The highest-priority groups recommended to receive booster doses are those with risk factors for severe COVID-19 and/or those at increased occupational risk of COVID-19. Comirnaty is preferred irrespective of the primary COVID-19 vaccine used</p>
	December	<p>Comirnaty indication age extended to include use in individuals aged ≥5 years</p> <p>Spikevax provisionally registered for a booster dose 6 months post-dose 2 in immunocompetent individuals aged ≥18 years</p> <p><i>12 December:</i> Due to ongoing transmission of Omicron and Delta variants, the recommended minimum interval between the primary course and the booster dose was shortened from 6 months to 5 months</p> <p><i>24 December:</i> The recommended minimum interval between the primary course and the booster dose was shortened from 5 months to 4 months and, when capacity permits (late January 2022), 3 months</p>

Year	Month	Intervention
2022	January	<p>Nuvaxovid provisionally registered for use in individuals aged ≥ 18 years for the primary course</p> <p>Comirnaty provisionally registered for a booster dose 6 months post-dose 2 in individuals aged 16–17 years</p> <p>Funded vaccination roll-out began in children aged 5–11 years</p> <p>Severely immunocompromised children aged 5–11 years recommended to receive a 3rd primary dose of COVID-19 vaccine 2 to 6 months after their 2nd dose, in line with other severely immunocompromised age cohorts</p> <p>People aged ≥ 18 years who received a 3-dose primary course due to severe immunocompromise recommended to receive a booster (4th) dose ≥ 4 months after their 3rd dose</p>
	February	<p>A booster dose was recommended for adolescents aged 16–17 years who had their primary COVID-19 vaccine course ≥ 3 months ago</p> <p>Spikevax indication age extended to include use in individuals aged ≥ 6 years</p> <p>Vaxzevria provisionally registered for a booster dose 6 months post dose 2, in individuals aged ≥ 18 years</p>
	March	<p>A winter booster dose was recommended for the following groups ≥ 4 months after the person has received their first booster dose, or ≥ 4 months after a confirmed SARS-CoV-2 infection, if infection occurred since the person's first COVID-19 booster dose:</p> <ul style="list-style-type: none"> adults aged ≥ 65 years residents of aged care or disability care facilities individuals aged ≥ 16 years with severe immunocompromise (as defined in the ATAGI statement on the use of a 3rd primary dose of COVID-19 vaccine in individuals who are severely immunocompromised) Aboriginal and Torres Strait Islander people aged ≥ 50 years
	April	Comirnaty vaccine provisionally registered for use as a booster dose for individuals aged 12–15 years
	May	The winter booster recommendation was expanded to include people aged 16–64 years with medical risk conditions or disability
	June	<p>Nuvaxovid (Novavax) vaccine provisionally registered for use as a booster in individuals aged ≥ 18 years</p> <p>Recommendations for boosters in high-risk adolescents aged 12–15 years</p>
	July	<p>Nuvaxovid (Novavax) vaccine indication age extended to include use in individuals aged ≥ 12 years</p> <p>Spikevax vaccine indication age extended to include use in individuals aged ≥ 6 months</p> <p>A winter booster dose was recommended for individuals aged >50 years, and individuals aged 30–49 years could receive a winter booster after discussion with their regular medical provider to review their individual health needs and the benefits and risks of a second booster dose</p> <p>A 3-month interval between a recent SARS-CoV-2 infection or the first booster dose and a winter booster dose was recommended</p>
	August	Nuvaxovid (Novavax) vaccine provisionally registered for use in individuals aged 12–17 years for the primary course

Year	Month	Intervention
2022 (cont.)	August	Spikevax bivalent (original/omicron BA.1) vaccine provisionally registered for use as a booster dose in individuals aged ≥ 18 years
	September	Comirnaty vaccine provisionally registered for use as a booster dose in individuals aged 5–11 years Comirnaty vaccine indication age extended to include use in individuals aged ≥ 6 months
	October	Spikevax vaccine provisionally registered for use as a booster dose in individuals aged ≥ 12 years Spikevax vaccine was recommended as a paediatric booster dose in children aged 5–11 years with high-risk medical conditions or disability Comirnaty bivalent (original/omicron BA.1) vaccine provisionally registered for use as a booster dose in individuals aged ≥ 18 years
2023	February	A booster dose was recommended for the following groups if their last vaccine dose or confirmed infection was ≥ 6 months: <ul style="list-style-type: none"> • all adults aged ≥ 65 years • adults aged 18–64 years who have medical comorbidities or disability • the following groups, after a risk–benefit assessment: <ul style="list-style-type: none"> ○ all adults aged 18–64 years without risk factors for severe COVID-19 ○ children aged 5–17 years with high-risk medical conditions or disability Comirnaty bivalent (original/omicron BA.4/5) vaccine provisionally registered for use as a booster dose in adolescents and individuals aged ≥ 12 years Moderna bivalent (original/omicron BA.4/5) vaccine provisionally registered for use as a booster dose in adolescents and individuals aged ≥ 12 years
	March	Vaxzevria (AstraZeneca COVID-19 vaccine) no longer available
	April	Spikevax vaccine (containing ancestral strain only) granted full registration for individuals aged ≥ 6 years
	May	A BA.4/5-containing bivalent COVID-19 vaccine preferred over original (ancestral) vaccines for use as the primary course and booster doses in people aged 12–17 years Either BA.1 or BA.4/5- containing COVID-19 vaccine recommended for use as the primary course and booster doses in people aged ≥ 18 years
	June	The original Comirnaty and Spikevax vaccines (containing ancestral strain only) no longer available
	July	Comirnaty vaccine (containing ancestral strain only) granted full registration for individuals aged ≥ 6 months
	August	Spikevax bivalent (original/omicron BA.4/5) granted full registration for individuals aged ≥ 12 years

Year	Month	Intervention
2023 (cont.)	September	<p>ATAGI recommends that all adults aged ≥ 75 years receive an additional 2023 COVID-19 vaccine dose if 6 months have passed since their last dose.</p> <p>ATAGI advises the following groups should consider an additional 2023 COVID-19 vaccine dose if 6 months have passed since their last dose, after discussion with their healthcare provider:</p> <ul style="list-style-type: none"> • all adults aged 65 to 74 years; and/or • adults aged 18–64 years with severe immunocompromise <p>Within the above groups, an additional 2023 COVID-19 vaccine is likely to be of most benefit for people who:</p> <ul style="list-style-type: none"> • have no known history of SARS-CoV-2 infection (and therefore are unlikely to have protection from hybrid immunity); • have medical comorbidities that increase their risk of severe COVID-19, or disability with significant or complex health needs; or • reside in a residential aged care facility
	October	<p>Spikevax (omicron XBB 1.5) granted full registration for individuals aged ≥ 12 years</p> <p>Comirnaty (omicron XBB 1.5) granted full registration for individuals aged ≥ 5 years</p> <p>Nuvaxovid vaccine (containing ancestral strain only) granted full registration for individuals aged ≥ 12 years</p>
	November	<p>ATAGI advised the following recommendations on the use of monovalent omicron XBB 1.5 COVID-19 vaccines:</p> <ul style="list-style-type: none"> • All currently available COVID-19 vaccines are anticipated to provide benefit to eligible people, however the monovalent Omicron XBB.1.5 vaccines are preferred over other vaccines for use in individuals aged ≥ 5 years who are currently recommended primary or additional doses of COVID-19 vaccine according to the Australian Immunisation Handbook • For those who have had the recommended 2023 dose/s of COVID-19 vaccine, ATAGI is not recommending further doses or re-vaccination with an XBB.1.5-containing vaccine at this time • ATAGI notes the recent increase in COVID-19 cases across Australia since November 2023 and encourages all people who have not yet had their recommended 2023 dose/s to receive them as soon as possible