Myocarditis and myopericarditis cases following COVID-19 mRNA vaccines administered to 12–17-year olds in Victoria, Australia

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ABSTRACT

**Importance** COVID-19 mRNA vaccine-associated myocarditis has previously been described; however specific features in the adolescent population are currently not well understood.

**Objective** To describe myocarditis adverse events following immunisation reported following any COVID-19 mRNA vaccines in the adolescent population in Victoria, Australia.

**Design** Statewide, population-based study.

**Setting** Surveillance of Adverse Events Following Vaccination in the Community (SAEFVIC) is the vaccine-safety service for Victoria, Australia.

**Participants** All SAEFVIC reports of myocarditis and myopericarditis in 12–17-year-old COVID-19 mRNA vaccinees submitted between 22 February 2021 and 22 February 2022, as well as accompanying diagnostic investigation results where available, were assessed using Brighton Collaboration criteria for diagnostic certainty.

**Exposures** Any mRNA COVID-19 vaccine.

**Main outcomes** Confirmed myocarditis associated with Brighton Collaboration criteria (levels 1–3).

**Results** Clinical review demonstrated definitive (Brighton level 1) or probable (level 2) diagnoses in 75 cases. Confirmed myocarditis reporting rates were 8.3 per 100 000 doses in this age group. Cases were predominantly male (n=62, 82.7%) and post dose 2 (n=61, 81.3%). Rates peaked in the 16–17-year-old group and were higher in males than females (17.7 vs 3.9 per 100 000, p<0.001). The most common presenting symptoms were chest pain, dyspnoea and palpitations. A large majority of cases who had a cardiac MRI had abnormalities (n=33, 91.7%). Females were more likely to have ongoing clinical symptoms at 1-month follow-up (p=0.02).

**Conclusion** Accurate evaluation and confirmation of episodes of COVID-19 mRNA vaccine-associated myocarditis enabled understanding of clinical phenotypes in the adolescent age group. Any potential vaccination and safety surveillance policies needs to consider age and gender differences.

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Previous case studies suggest that there is an association between COVID-19 mRNA vaccines and increased rates of myocarditis, most commonly in the following 2 weeks.

⇒ Recommendations on use of COVID-19 mRNA vaccines in different age groups vary globally and may not take into account their different risk profiles.

WHAT THIS STUDY ADDS

⇒ Incidence of myocarditis post COVID-19 mRNA vaccines are higher after the second dose and appears to differ by age and gender, with adolescent and young adult males being at a higher risk.

⇒ Females were more likely to have ongoing clinical symptoms at 1-month follow-up compared with males.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE AND/OR POLICY

⇒ Potential policy and safety surveillance adjustments may need to take into account age and gender differentials in myocarditis when reviewing COVID-19 primary (two-dose) vaccine roll-out to the adolescent population.

INTRODUCTION

Australia has utilised two mRNA vaccines as part of its COVID-19 vaccine strategy in 12–17-year olds, namely Comirnaty BNT162b2 COVID-19 (Pfizer-BioNTech) and Spikevax mRNA-1273 (Moderna). Comirnaty was initially provisionally licensed by the Australian regulator, the Therapeutic Goods Administration (TGA) for those aged 16 years and older, and administered from 25 January 2021 in adults 18+ years. Spikevax received provisional TGA approval for administration in individuals from 18 years from 9 August 2021. Following the availability of age-specific clinical trial data, further approvals were
added for the 12–15-year-old group for Comirnaty (22 July 2021) and 12–17-year old for Spikevax (4 September 2021).

Of particular interest in the young adult population is postvaccination myocarditis and pericarditis causally associated with COVID-19 mRNA vaccines. These adverse events of special interest (AESI) were first flagged in Israel, which implemented Comirnaty at a population level as soon as it was available (20 December 2020).2 In a linked electronic health record observational case–control study design from Israel’s largest healthcare insurer, Barda et al described an elevated risk of myocarditis following Comirnaty (risk ratio (RR): 3.24; 95% CI: 1.55 to 12.44), while noting a substantially higher risk following COVID-19 disease (RR: 18.28; 95% CI: 3.95 to 25.12).3

Subsequent postlicensure observational military and report-based case studies confirmed the highest risk group for post mRNA COVID-19 vaccine myocarditis in young males (<24 years old) following the second vaccine dose.4–6 Both the spontaneous Vaccine Adverse Event Reporting System and active Vaccine Safety Datalink Surveillance System in the USA have confirmed a myocarditis signal safety,7 along with similar findings in Canada, the UK and various Nordic countries.6

The case phenotype includes chest pain as the most common symptom, but also includes fever, shortness of breath and other non-specific symptoms such as headache, myalgia and vomiting. Common investigation findings include a raised troponin, abnormal ECG with ST or T wave changes and late gadolinium enhancement or myocardial oedema seen on cardiac MRI.8–11

| Table 1 | Brighton collaboration criteria for myocarditis |
|----------------|-----------------------------------------------|
| **Level 1: ‘definitive’ case** | **Level 2: ‘probable case’** | **Level 3: ‘possible case’** |
| Histopathologic examination of myocardial tissue (autopsy or endomyocardial biopsy) showed myocardial inflammation | Clinical symptoms and exclusion as per Level three case AND Elevated myocardial biomarkers ≥1 new finding of Troponin T level above upper limit of normal OR Troponin I level above upper limit of normal OR CK myocardial band | Presence of ≥1 new or worsening of the following clinical symptoms: ▶ Chest pain/pressure ▶ Dyspnoea/shortness of breath/pain breathing ▶ Diaphoresis ▶ Palpitations ▶ Sudden death OR Presence of ≥2 new or worsening of the following clinical symptoms: ▶ Fatigue ▶ Abdominal Pain ▶ Syncope ▶ Oedema ▶ Cough AND | ≥1 new supported finding of inflammation Elevated CRP/ESR or D-Dimer AND Presence of ≥1 new abnormal ECG such as: ▶ ST-segment or T-wave abnormalities (elevation or inversion) ▶ PACs and PVCs AND ▶ No other identifiable cause of the symptoms and findings |
| OR ≥1 new finding of Troponin T or I level above upper limit of normal AND ≥1 New cMRI findings consistent with Oedema on T2-weighted study, typically patchy in nature OR Late gadolinium enhancement on T1-weighted study with an increased enhancement ratio between myocardial and skeletal muscle typically involving at least one non-ischaemic regional distribution with recovery (myocyte injury) | ▶ Focal or diffuse left or right ventricular function abnormalities (eg, decreased ejection fraction) ▶ Segmental wall motion abnormalities ▶ Global systolic or diastolic function depression/abnormality ▶ Ventricular dilation ▶ Wall thickness change ▶ Intracavitary thrombi | ▶ Paroxysmal or sustained atrial or ventricular arrhythmias (premature atrial or ventricular beats, and/or supraventricular or ventricular tachycardia, interventricular conduction delay, abnormal Q waves, low voltages) ▶ AV nodal conduction delays or intraventricular conduction defects (atrioventricular block (grade I–III), new bundle branch block) ▶ Continuous ambulatory electrocardiographic monitoring that detects frequent atrial or ventricular ectopy | ▶ Echocardiogram (ECHO) abnormalities ≥1 new finding of ▶ Echocardiogram (ECHO) abnormalities ≤1 global new finding of OR ECG abnormalities ≥1 new finding of ▶ Oedema on T2-weighted study with an incr | ▶ Echocardiogram (ECHO) abnormalities ≥1 new finding of OR ECG abnormalities ≥1 new finding of ▶ Oedema on T2-weighted study with an incr | ▶ Echocardiogram (ECHO) abnormalities ≥1 new finding of OR ECG abnormalities ≥1 new finding of ▶ Oedema on T2-weighted study with an incr |
| Echocardiogram (ECHO) abnormalities biomarkers ≥1 new finding of as per level 2 case | ▶ Troponin I level above upper limit of normal ▶ Troponin T level above upper limit of normal | ▶ Troponin I level above upper limit of normal ▶ Troponin T level above upper limit of normal | ▶ Troponin I level above upper limit of normal ▶ Troponin T level above upper limit of normal |
| | ▶ Histopathologic examination of myocardial tissue (autopsy or endomyocardial biopsy) showed myocardial inflammation | ▶ Histopathologic examination of myocardial tissue (autopsy or endomyocardial biopsy) showed myocardial inflammation | ▶ Histopathologic examination of myocardial tissue (autopsy or endomyocardial biopsy) showed myocardial inflammation |
| AV, atrioventricular; cMRI, cardiac MRI; PAC, premature atrial complex; PVC, premature ventricular complex. | | | |
Due to this AESI signal, the risk, clinical manifestations and follow-up of myocarditis and pericarditis following mRNA COVID-19 in younger populations has been of particular interest. Some regions (eg, Hong Kong) are only administering a single dose of mRNA vaccine in adolescents aged 12–17 years and notably, there is limited use of Spikevax in this age group internationally. Spikevax is not currently licensed by the US Food and Drug Administration (FDA) for 12–17-year olds, while Canada and several European countries have issued preferential recommendations for Comirnaty over Spikevax for young adults. This study describes clinical presentation and evaluation of myocarditis AESI following mRNA COVID-19 vaccination in 12–17-year-old adolescents in Victoria, Australia.

**METHODS**

Victoria is a south-eastern Australian state with a population of approximately 6.6 million. Adverse events following immunisation (AEFI) are spontaneously reported by patients, caregivers or healthcare providers to Surveillance of Adverse Events Following Vaccination in the Community (SAEFVIC), the state-wide vaccine safety service. SAEFVIC comprises central reporting enhanced passive and active surveillance systems integrated with clinical services and has been operating since 2007.

Identified reports of myocarditis and myopericarditis in 12–17-year-old vaccinees submitted to SAEFVIC between 22 February 2021 and 22 February 2022 were assessed. The majority of these were practitioner reported. Myocarditis and myopericarditis reports (henceforth summarised as myocarditis) were systematically followed up and diagnostic test results (where available) obtained to confirm the diagnoses, including potential alternative causes such as viral, autoimmune or medication-related myocarditis. Data gathered included a list of symptoms such as chest pain, shortness of breath, dizziness and fatigue (table 1), as well as investigations undertaken by treating clinicians, including ECG, cardiac biomarkers, echocardiogram and cardiac MRI scans. All troponin levels obtained were high sensitivity troponin assays, with troponin levels reported as fold increase from the upper limit of normal to facilitate comparison between different assays.

Once data were available, each case was categorised by at least two independent experts utilising the Brighton Collaboration definition with graded levels of certainty (table 1). All reports were forwarded to the national regulator, the TGA, who report weekly on spontaneous AEFI reports at a national level.

Cases where vaccination was the most likely cause of their diagnosis were followed up after 1 month to answer a series of questions about ongoing symptoms and clinical management. Cases were declared lost to follow-up after three unsuccessful attempts to contact.

**Patient and Public Involvement**

**Statistical analysis**

Data were analysed using Microsoft PowerBI (V.2.91.701.0) with 90% Poisson CIs calculated for rates. Vaccine doses administered and population estimates were obtained from the Australian Immunisation Registry. Age groups were defined by the vaccine roll-out in Victoria, whereby 16–17-year-olds were eligible for vaccination prior to 12–15-year olds. Mood’s median test was used to compare median fold increase in troponin levels.

**RESULTS**

As of 22 February 2022, 454974 12–17-year-old Victorians had received 871 689 mRNA doses (782964 Comirnaty and 88725 Spikevax). This equated to approximately 97.9% first dose and 93.7% second dose coverage of this age cohort.

At this timepoint, there were 75 reports of confirmed myocarditis (44 myocarditis, 31 myopericarditis) as per Brighton Collaboration level 1 (definitive) or 2 (probable) criteria. This equates to a rate of 8.3 per 100 000 doses (90% CI: 6.8 to 10.1) in this age group (table 1).

Cases were predominantly in males (n=62, 82.7%) and post dose 2 (n=61, 81.3%). Presentation was temporally related to Comirnaty in 63 (84.1%) and to Spikevax in 12 (16.0%) cases, respectively. Higher rates were observed

| Table 2 | Count and rate of cases by sex, age group, dose number and Brighton Collaboration level | Male | Female | Total |
|---|---|---|---|---|
| | Count | Rate per 100 000 doses (90% CI) | Count | Rate per 100 000 doses (90% CI) | Count | Rate per 100 000 doses (90% CI) |
| Age group (years) | | | | | | |
| Total | 62 | 13.6 (10.9 to 16.8) | 13 | 2.9 (1.7 to 4.7) | 75 | 8.3 (6.8 to 10.1) |
| Age | 12–15 | 34 | 11.4 (8.4 to 15.2) | 7 | 2.4 (1.1 to 4.6) | 41 | 7.0 (5.3 to 9.1) |
| | 16–17 | 28 | 17.7 (12.6 to 24.2) | 6 | 3.9 (1.7 to 7.6) | 34 | 10.8 (8.0 to 14.4) |
| Dose | 1 | 10 | 4.4 (2.4 to 7.5) | 4 | 1.8 (0.6 to 4.2) | 14 | 3.2 (1.9 to 4.9) |
| | 2 | 52 | 24.2 (19.0 to 30.5) | 9 | 4.3 (2.3 to 7.5) | 61 | 14.4 (11.5 to 17.8) |
| Brighton collaboration level | Definitive | 27 | 5.9 (4.2 to 8.2) | 3 | 0.7 (0.2 to 1.8) | 30 | 3.3 (2.4 to 4.5) |
| | Probable | 35 | 7.7 (5.7 to 10.2) | 10 | 2.3 (1.2 to 3.8) | 45 | 5.0 (3.8 to 6.4) |
in the 16–17-year-old age group, and in males compared with females (p=0.001; table 2).

Of the 75 cases in the study period, 5 of them were considered by the treating clinician to have an alternative cause for myocarditis more likely than COVID-19 vaccination. One of these cases had evidence of historical COVID-19 infection, with a positive COVID-19 respiratory PCR 3 weeks prior to receiving first dose Comirnaty vaccination, with subsequent onset of myocarditis 2 days later.

For the 70 cases related to COVID-19 mRNA vaccination, onset of symptoms ranged from 0 to 49 days after vaccination, with a median of 2 days (IQR: 1–3 days), in males from 0 to 34 days (median 2 days) and in females from 1 to 49 days (median 2 days). Fifty-one (77.1%) cases required hospital admission with a median length of stay of two nights. No cases required intensive care unit (ICU) admission and no deaths were recorded. All admissions were discharged to home.

All cases had chest pain as a presenting symptom. Other common symptoms included dyspnoea (21 cases, 30%), palpitations (14, 20%) and diaphoresis. Thirty-three (47.1%) cases had concomitant non-specific symptoms such as dizziness, vomiting and fatigue (table 3).

ECG abnormalities were observed in 49 (70.0%) cases, with the most common finding being ST-elevation. An echocardiogram was performed in 70 cases and was abnormal in 8 (11.4%). An initial diagnostic cardiac MRI was performed in 36 cases with abnormalities documented in the majority of these (33 cases, 91.7%) (table 3).

There was a trend for males towards higher and more variable increases in troponin, with a median fold rise of 144 times above normal levels compared with females at 33 times above normal (p=0.222; figure 1).

Follow-up at 1 month was completed for 64 of the 70 cases where COVID-19 vaccination was the most likely cause of their diagnosis (table 4). The remaining six cases were lost to follow-up and excluded from analysis.

Symptoms remained in 50.0% of those who participated in follow-up, with a higher percentage of females having ongoing symptoms (44.6 vs 87.5%, p=0.02) including chest pain (48.0 vs 71.4%, p=0.01; table 4).

**DISCUSSION**

We describe 75 cases of myocarditis reported following COVID-19 mRNA vaccination in the adolescent age group. Our rate of cases was similar, although slightly higher than other international studies in Israel, the UK and the USA.17

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**Table 3** Symptoms, laboratory, ECG and imaging data

| Symptoms (n=70), n (%) | Count |
|-----------------------|-------|
| Chest pain            | 70 (100%) |
| Palpitations          | 14 (20.0%) |
| Dyspnoea              | 21 (30.0%) |
| Diaphoresis           | 6 (8.6%) |
| Non-specific symptoms (dizziness, vomiting, fatigue) | 33 (47.1%) |

| Laboratory test | Value |
|-----------------|-------|
| Troponin (n=70), median fold increase | 138.3 (IQR 56.9–315.0) |

| Testing/imaging | Count |
|-----------------|-------|
| ECG (n=70), n (%) |       |
| Abnormal        | 49 (70.0%) |
| Normal          | 21 (30.0%) |

| Abnormal ECG findings or arrhythmias (n=49) |       |
|--------------------------------------------|-------|
| ST-wave or T-wave changes/elevation        | 28 (57.1%) |
| ST segment depression in AVR                | 9 (18.4%) |
| PR depression without reciprocal ST depression | 7 (14.3%) |
| AV node conduction delay                    | 3 (6.1%) |
| T wave inversion                            | 3 (6.1%) |
| Other                                       | 9 (18.4%) |

| Echocardiogram (n=68) |       |
|----------------------|-------|
| Normal function      | 62 (91.2%) |
| Abnormal function    | 8 (8.8%) |
| Systolic dysfunction | 5 (62.5%) |
| Wall motion abnormalities | 2 (25.0%) |
| LV strain            | 1 (12.5%) |

| Cardiac MRI (n=36) |       |
|--------------------|-------|
| Abnormal findings, n (%) |       |
| Late gadolinium enhancement | 32 (97.0%) |
| Myocardial oedema    | 20 (60.6%) |
| Other abnormality on T2 imaging | 9 (27.3%) |
| Pericardial effusion or inflammation | 5 (15.2%) |
| Fibrosis             | 2 (2.4%) |

Note: LV, left ventricular.
This rate likely reflects concerted public health messaging and awareness of the AESI, coupled with the combination of active and passive surveillance methodology to ascertain these cases. While this may have led to increased reporting rates, the robust diagnostic criteria requiring laboratory and imaging tests to confirm myocarditis, together with tight COVID-19-related environmental restrictions that significantly reduced the chance of COVID-19 infection-related myocarditis, indicate that this is likely to represent an accurate rate.

Clinical findings were also similar to other international cohorts of adolescents. Dionne et al described similar symptom prevalence in a cohort of myocarditis patients <19 years. There was also similar cardiac MRI abnormalities noted in similar cohorts. In contrast to our findings, the cohort from Truong et al had a high number of patients (18.7%) requiring ICU admission, although only two required inotropic support. It is likely that the difference in ICU admission criteria, rather than true clinical severity accounts for this discrepancy.

Unique to this study is the clear clinical phenotypic differences identified between sexes. Males tended to present with symptoms and a significantly elevated peak troponin level. Just under half had no residual symptoms 4 weeks after initial onset. In contrast, females tended to have had a much lower median peak troponin level, with similar rates post dose 1 and 2. Symptoms were still experienced by a majority of females 4 weeks after onset. These differences may support a potential impact of testosterone as a risk factor for stronger cardiac inflammation in myocarditis, but with a potentially beneficial impact on duration of symptoms.

Our data indicate a higher rate of cases between the age of 16 and 17 years and a tapering in rate for the 12–17-year-old age group. Further analysis is needed and ongoing to compare rates of myocarditis between Comirnaty and Spikevax when sufficient doses are administered, although only two required inotropic support. It is likely that the difference in ICU admission criteria, rather than true clinical severity accounts for this discrepancy.

Table 4: One-month follow-up outcomes, by sex (n=64)

| Symptom          | Male (n=56) | Female (n=8) | Total (n=64) | χ² statistic (p value) |
|------------------|-------------|--------------|--------------|------------------------|
| Ongoing symptoms| 32, 50.0    | 9, 28.1      | 41, 64.1     | 5.14 (p=0.02)          |
| Chest pain       | 17, 53.1    | 9, 28.1      | 26, 41.6     | 6.05 (p=0.01)          |
| Fatigue          | 15, 46.9    | 9, 28.1      | 24, 37.5     | 3.60 (p=0.58)          |
| Palpitations     | 9, 28.1     | 9, 28.1      | 18, 28.1     | 4.16 (p=0.41)          |
| Dyspnoea         | 9, 28.1     | 9, 28.1      | 18, 28.1     | 4.16 (p=0.41)          |

LIMITATIONS
This study is based on clinical data compiled by SAEFVIC as part of vaccine safety surveillance. A passive vaccine surveillance system may under-report potential cases of myocarditis. Furthermore, there was lack of clinical data on some cases (eg, investigations not performed), making full description of clinical phenotype, evaluation and diagnosis more challenging. While patient-reported symptoms were included as part of the evaluation, Brighton diagnostic criteria were used as a benchmark to reduce recall bias. The accuracy of diagnosis was also maintained by ensuring that available patient clinical information were reviewed by at least two independent medical specialists to verify reported myocarditis cases and reduce misclassification.

CONCLUSION
Rates of myocarditis in the adolescent population differ by dose and sex in the 12–17-year-old age group. With the vaccine roll-out in younger children and adolescents in mind, these clinical phenotypic differences should be considered for future COVID-19 vaccine recommendations including ‘booster’ doses.

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