

# 483 More Deaths After COVID Vaccines Reported to VAERS, as Pfizer and Moderna Push for More Boosters

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*VAERS data released Friday by the Centers for Disease Control and Prevention included a total of 1,183,495 reports of adverse events from all age groups following COVID vaccines, including 25,641 deaths and 208,209 serious injuries between Dec. 14, 2020, and March 11, 2022.*

by [Megan Redshaw](#), [Children's Health Defense](#)

March 18, 2022

The Centers for Disease Control and Prevention (CDC) today released new data showing a total of [1,183,495 reports of adverse events](#) following COVID-19 vaccines were submitted between Dec. 14, 2020, and March 11, 2022, to the Vaccine Adverse Event Reporting System (VAERS). VAERS is the primary government-funded system for reporting adverse vaccine reactions in the U.S.

The data included a total of [25,641 reports of deaths](#) – an increase of 483 over the previous week – and [208,209 reports of serious injuries](#), including deaths, during the same time period – up 4,321 compared with the previous week.

Excluding “[foreign reports](#)” to VAERS, [788,624 adverse events](#), including [11,728 deaths](#) and [76,231 serious injuries](#), were

reported in the U.S. between Dec. 14, 2020, and March 11, 2022.

[Foreign reports](#) are reports foreign subsidiaries send to U.S. vaccine manufacturers. Under U.S. Food and Drug Administration (FDA) regulations, if a manufacturer is notified of a foreign case report that describes an event that is both serious and does not appear on the product's labeling, the manufacturer is required to submit the report to VAERS.

Of the 11,728 U.S. [deaths reported](#) as of March 11, 17% occurred within 24 hours of vaccination, 22% occurred within 48 hours of vaccination and 60% occurred in people who experienced an [onset of symptoms](#) within 48 hours of being vaccinated.

In the U.S., 556 million COVID vaccine doses had been administered as of March 11, [including](#) 328 million doses of Pfizer, 209 million doses of Moderna and 19 million doses of Johnson & Johnson (J&J).



## Search Results

From the 3/11/2022 release of VAERS data:

**Found 1,183,495 cases where Vaccine is COVID19**

[Government Disclaimer on use of this data](#)

Table

↓	↑ ↓	
Event Outcome	Count	Percent
Death	25,641	2.17%
Permanent Disability	47,676	4.03%
Office Visit	181,686	15.35%
Emergency Room	103	0.01%
Emergency Doctor/Room	123,732	10.45%
Hospitalized	140,759	11.89%
Hospitalized, Prolonged	353	0.03%
Recovered	328,324	27.74%
Birth Defect	992	0.08%
Life Threatening	29,135	2.46%
Not Serious	526,937	44.52%
<b>TOTAL</b>	<b>↑ 1,405,338</b>	<b>↑ 118.74%</b>

† Because some cases have multiple vaccinations and symptoms, a single case can account for multiple entries in this table. This is the reason why the Total Count is greater than 1183495 (the number of cases found), and the Total Percentage is greater than 100.

Every Friday, [VAERS](#) publishes vaccine injury reports received as of a specified date. Reports submitted to VAERS require further investigation before a causal relationship can be confirmed.

Historically, VAERS has been shown to report only [1% of actual vaccine adverse events](#).

**U.S. VAERS data from Dec. 14, 2020, to March 11, 2022, for 5- to 11-year-olds show:**

- [9,161 adverse events](#), including [217 rated as serious](#) and [5 reported deaths](#).

The most recent death involves a 7-year-old boy (VAERS I.D. [2152560](#)) from Washington who died 13 days after receiving his first dose of Pfizer's COVID vaccine when he went into shock and suffered cardiac arrest. He was unable to be resuscitated and died in the emergency department.

- [17 reports](#) of myocarditis and pericarditis (heart inflammation).

The CDC uses a [narrowed case definition](#) of “myocarditis,” which [excludes cases](#) of cardiac arrest, [ischemic strokes](#) and deaths due to heart problems that occur before one has the chance to go to the emergency department.

- [34 reports](#) of blood clotting disorders.

**U.S. VAERS data from Dec. 14, 2020, to March 11, 2022, for 12- to 17-year-olds show:**

- [30,295 adverse events](#), including [1,744 rated as serious](#) and [42 reported deaths](#).

The most recent deaths involve a 17-year-old boy (VAERS I.D. [2171083](#)) from Illinois with [Duchenne muscular dystrophy](#) who died from cardiac arrest after receiving his second dose of Pfizer’s COVID vaccine, and 14-year-old boy from Guam (VAERS I.D. [2157944](#)) who died one week after his first dose of Pfizer when he suddenly committed suicide.

The boy’s VAERS report states:

“Sudden suicide one week after the vaccine. Patient was a perfectly happy child. After the vaccine, he became much more tired and achy and lost interest in doing his sports. One week later, without any warning, he hung himself.”

- [68 reports](#) of anaphylaxis among 12- to 17-year-olds where the reaction was life-threatening, required treatment or resulted in death – with 96% of cases attributed to [Pfizer’s vaccine](#).
- [646 reports](#) of myocarditis and pericarditis, with [634 cases](#) attributed to Pfizer’s vaccine.
- [162 reports](#) of blood clotting disorders, with all cases attributed to Pfizer.

**U.S. VAERS data from Dec. 14, 2020, to March 11, 2022, for all**

## age groups combined, show:

- 20% of deaths were related to cardiac disorders.
- 54% of those who died were male, 41% were female and the remaining death reports did not include the gender of the deceased.
- The [average age](#) of death was 72.7.
- As of March 11, [5,250 pregnant women](#) reported adverse events related to COVID vaccines, including 1,668 reports of [miscarriage or premature birth](#).
- Of the [3,613 cases of Bell's Palsy](#) reported, 51% were attributed to [Pfizer](#) vaccinations, 40% to [Moderna](#) and 8% to [J&J](#).
- 863 reports of [Guillain-Barré syndrome](#), with 41% of cases [attributed to Pfizer](#), 30% to [Moderna](#) and 28% to [J&J](#).
- [2,363 reports](#) of anaphylaxis where the reaction was life-threatening, required treatment or resulted in death.
- [1,683 reports](#) of myocardial infarction.
- [13,512 reports](#) of blood-clotting disorders in the U.S. Of those, [6,034 reports](#) were attributed to Pfizer, [4,818 reports](#) to Moderna and [2,617 reports](#) to J&J.
- [4,045 cases](#) of myocarditis and pericarditis with [2,483 cases](#) attributed to Pfizer, [1,377 cases](#) to Moderna and [175 cases](#) to J&J's COVID vaccine.

## Moderna asks FDA to authorize 4th dose for adults 18 and up

Moderna on Thursday asked the FDA to [amend Emergency Use Authorization](#) (EUA) of its COVID vaccine to include a fourth dose for adults 18 and older.

According to [The Associated Press](#), the request is broader than Pfizer's. Pfizer earlier this week asked the agency to authorize a fourth dose of its COVID vaccine for adults 65 and older.

In a [press release](#), Moderna said the request to include adults over 18 was made “to provide flexibility for the U.S. Centers for Disease Control and Prevention and healthcare providers to determine the appropriate use of an additional booster dose of mRNA-1273, including for those at higher risk of COVID-19 due to age or comorbidities.”

Moderna [said its decision](#) to seek FDA approval was based on studies from the U.S. and Israel about the [Omicron](#) variant, but didn't provide further information. Booster doses of Moderna are half the dose of the first and second doses.

### **Pfizer and BioNTech ask FDA to authorize fourth vaccine dose for older adults**

Pfizer and BioNTech on Tuesday [said](#) they submitted a request to the FDA for EUA of an additional booster dose of their COVID vaccine for adults 65 and older.

The companies' request was not based on robust, peer-reviewed U.S. data, but on [two recent studies from Israel](#) – both published on preprint servers without peer review.

The first [study](#) was done in conjunction with Israel's Ministry of Health and involved a review of 1.1 million health records. The study concluded rates of COVID in those who received a fourth dose of Pfizer's COVID vaccine were lower compared to those who received only three doses.

According to the [preprint](#) published on medRxiv, since Jan. 2 Israel has been administering a fourth dose of the Pfizer vaccine only to people over 60 and at-risk populations.

In the [second study](#) of Israeli healthcare workers, results showed a fourth dose of either Pfizer's or Moderna's vaccine boosted antibody levels, but neither was effective at preventing infections.

### **CDC deletes thousands of reported COVID-19 deaths in children**

The CDC removed tens of thousands of deaths linked to COVID, including nearly a quarter of deaths it had attributed to those younger than 18, [The Epoch Times reported](#). The change was made on March 15 on its COVID [data tracker website](#).

“Data on deaths were adjusted after resolving a coding logic error. This resulted in decreased death counts across all demographic categories,” the CDC said on the website. The agency also acknowledged COVID death data is not complete.

Before the change, [the CDC listed](#) 1,755 deaths in children from COVID, along with 851,000 others, according to Kelley Krohnert, a Georgia resident who tracks the CDC’s updates.

The CDC removed 416 deaths among children and more than 71,000 other reported deaths – arriving at a total of about 780,000.

The CDC’s statistics are frequently cited by physicians and experts when pushing for children to receive COVID vaccines. Dr. Rochelle Walensky, the CDC’s director, referred to the tracker’s death total on November 2021 while pushing for an expert panel to advise her agency to recommend vaccination for all children 5 to 11 years old.

### **Vaccine researcher develops tinnitus 90 minutes after COVID shot, calls for more research**

A vaccinologist at the Mayo Clinic in Minnesota said he [developed tinnitus](#) after receiving his second dose of an mRNA COVID vaccine.

Dr. Gregory Poland’s symptoms began 90 minutes after receiving the vaccine. He described the condition as “fairly severe” and “extraordinarily bothersome, interfering with sleep and the ability to concentrate.”

*Dr. Gregory Poland, a vaccinologist at the Mayo Clinic in Minnesota, developed tinnitus after his second dose of an mRNA COVID-19 vaccine. He is raising questions about this*

side effect from the vaccine and suggesting more research is needed. <https://t.co/9IXh09P0cv>

– Robert F. Kennedy Jr (@RobertKennedyJr) [March 16, 2022](#)

[According to the National Institutes of Health](#), tinnitus is a sign that something is wrong with the auditory system. It is commonly described as a ringing in the ears, but it also can sound like roaring, clicking, hissing, or buzzing that accompanies soft, loud or high pitches.

According to the most recent VAERS data released on March 11, [19,851 people](#) have reported developing tinnitus after a COVID vaccine, with [12,027 cases](#) attributed to Pfizer's COVID vaccine.

### **CEO of German health insurer fired after releasing data on underreported COVID vaccine injuries**

The CEO of one of Germany's largest health insurance companies [was abruptly fired](#) last month after he released data suggesting German health authorities are significantly underreporting COVID-19 vaccine injuries.

The data, released by Andreas Schofbeck of BKK/ProVita, have since been scrubbed from the company's website.

Schofbeck, who noticed an unexpected jump in vaccine-related health insurance claims, in February [notified](#) the [Paul Ehrlich Institute](#) (PEI) – the German equivalent of the CDC – that BKK billing data indicated the PEI was underreporting adverse events to COVID vaccines.

In his [letter](#) to the PEI, Schofbeck wrote:

I'm "If these figures are extrapolated to the whole year and to the population in Germany, probably 2.5-3 million people in Germany have received medical treatment for vaccination side effects after Corona vaccination."



Dr. Dirk Heinrich, chairman of NAV-Virchow Bund, an association of private medical practitioners in Germany, said PEI and BKK would be [working closely to examine](#) the billing code data. Heinrich also stated that the conclusions from Schofbeck's letter are "[complete nonsense.](#)"

[Children's Health Defense](#) asks anyone who has experienced an adverse reaction, to any vaccine, to file a report following [these three steps](#).

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