

Pfizer SARS-COV2 Vaccine (BNT162b2) Cumulative analysis of Post Authorization Adverse Event reports.

Released by the US Food and Drug Administration Nov 17 2021

Adhering to a FOIA request, the FDA released the first 91 pages of data related to the Covid Vaccines. This release contains a 38 page document describing post authorization AE (adverse event) reports compiled by **Pfizer from Dec 10 thru Feb 28**. Pfizer then submitted this initial 2.5 months of real world data to the FDA.

Find the document here: https://phmpt.org/pfizers-documents/

Document name: 5.3.6 postmarketing experience.pdf

The FDA is requesting 55 years to process and release the remaining documents.

(Source: Reuters) **CLICK HERE**

Page 5 – "Magnitude of underreporting is unknown"

Page 6 - Reports of Adverse Events (AE) from EUA authorization through 28 February 2021. Voluntary and literature reported events. In just under three months, there were 42,086 case reports containing 158,893 AEs (out of a total number of doses that is undisclosed, as this was redacted from the report). Pfizer has hired additional personnel to cope with the volume of AE reports being received (the total number of employees hired is also redacted).

2. METHODOLOGY

Pfizer is responsible for the management post-authorization safety data on behalf of the MAH BioNTech according to the Pharmacovigilance Agreement in place. Data from BioNTech are included in the report when applicable.

Pfizer's safety database contains cases of AEs reported spontaneously to Pfizer, cases reported by the health authorities, cases published in the medical literature, cases from Pfizer-sponsored marketing programs, non-interventional studies, and cases of serious AEs reported from clinical studies regardless of causality assessment.

The limitations of post-marketing adverse drug event reporting should be considered when interpreting these data:

- Reports are submitted voluntarily, and the magnitude of underreporting is unknown.
 Some of the factors that may influence whether an event is reported include: length of time since marketing, market share of the drug, publicity about a drug or an AE, seriousness of the reaction, regulatory actions, awareness by health professionals and consumers of adverse drug event reporting, and litigation.
- Because many external factors influence whether or not an AE is reported, the spontaneous reporting system yields reporting proportions not incidence rates. As a result, it is generally not appropriate to make between-drug comparisons using these

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• Among adverse event reports received into the Pfizer safety database during the cumulative period, only those having a complete workflow cycle in the safety database (meaning they progressed to Distribution or Closed workflow status) are included in the monthly SMSR. This approach prevents the inclusion of cases that are not fully processed hence not accurately reflecting final information. Due to the large numbers of spontaneous adverse event reports received for the product, the MAH has prioritised the processing of serious cases, in order to meet expedited regulatory reporting timelines and ensure these reports are available for signal detection and evaluation activity. The increased volume of reports has not impacted case processing for serious reports, and compliance metrics continue to be monitored weekly with prompt action taken as needed to maintain compliance with expedited reporting obligations. Non-serious cases are entered into the safety database no later than 4 calendar days from receipt. Entrance into the database includes the coding of all adverse events; this allow for a manual review of events being received but may not include immediate case processing to completion.

Non-serious cases are processed as soon as possible and no later than 90 days from receipt. Pfizer has also taken a multiple actions to help alleviate the large increase of adverse event reports. This includes significant technology enhancements, and process and workflow solutions, as well as increasing the number of data entry and case processing colleagues. To date, Pfizer has onboarded approximately ^[b] additional full-time employees (FTEs). More are joining each month with an expected total of more than ^[b] (4) additional resources by the end of June 2021.

3. RESULTS

3.1. Safety Database

3.1.1. General Overview

It is estimated that approximately (b) (4) doses of BNT162b2 were shipped worldwide from the receipt of the first temporary authorisation for emergency supply on 01 December 2020 through 28 February 2021.

Cumulatively, through 28 February 2021, there was a total of 42,086 case reports (25,379 medically confirmed and 16,707 non-medically confirmed) containing 158,893 events. Most cases (34,762) were received from United States (13,739), United Kingdom (13,404) Italy (2,578), Germany (1913), France (1506), Portugal (866) and Spain (756); the remaining 7,324 were distributed among 56 other countries.

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General Overview – Outcomes - AEs

Page 7 – GENDER: The proportion of reported AEs show a *majority of AEs affecting females,* with nearly 3 times as many females reporting compared to males.

AGE & OUTCOME: Average age and case outcome data mirrors what has been found from an independent patient-led survey with over 500 respondents with persistent symptoms post-vaccine. This detailed report was also submitted to the FDA and can be found HERE.

OUTCOME: It is important to note the number of reports indicating the ongoing nature of the AE, as well as the reports indicating a fatal outcome, from *one vaccine brand, in 2.5 months.*

When breaking out the AEs in System Organ Classes (SOCs), we see that after general and administration site conditions, the most common reports are *nervous system disorders*, followed by *musculoskeletal* and *connective tissue disorders*.

Table 1 below presents the main characteristics of the overall cases.

Table 1. General Overview: Selected Characteristics of All Cases Received During the Reporting Interval

Characteristics		Relevant cases (N=42086)
Gender:	Female	29914
	Male	9182
	No Data	2990
Age range (years):	≤ 17	175*
0.01 -107 years	18-30	4953
Mean = 50.9 years	31-50	13886
n = 34952	51-64	7884
	65-74	3098
	≥ 75	5214
	Unknown	6876
Case outcome:	Recovered/Recovering	19582
	Recovered with sequelae	520
	Not recovered at the time of report	11361
	Fatal	1223
	Unknown	9400

a. in 46 cases reported age was <16-year-old and in 34 cases <12-year-old.

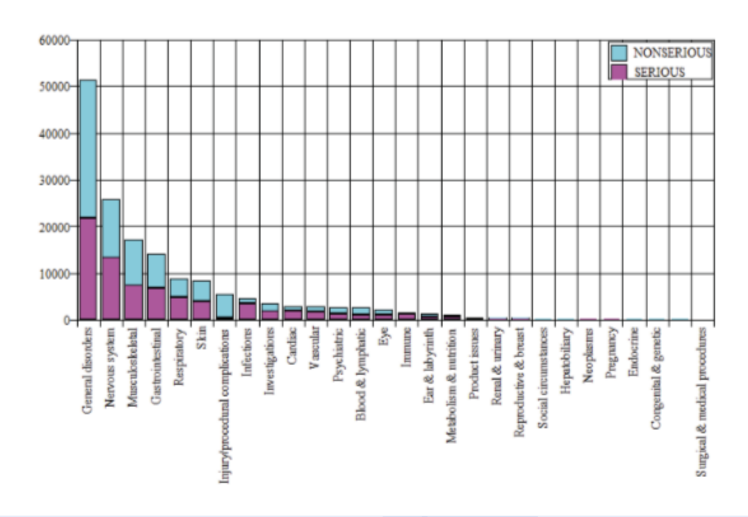
As shown in Figure 1, the System Organ Classes (SOCs) that contained the greatest number (≥2%) of events, in the overall dataset, were General disorders and administration site conditions (51,335 AEs), Nervous system disorders (25,957), Musculoskeletal and connective tissue disorders (17,283), Gastrointestinal disorders (14,096), Skin and subcutaneous tissue disorders (8,476), Respiratory, thoracic and mediastinal disorders (8,848), Infections and infestations (4,610), Injury, poisoning and procedural complications (5,590), and Investigations (3,693).



Page 5 – A large portion of the reported events were described as "Serious."

BNT162b2 5.3.6 Cumulative Analysis of Post-authorization Adverse Event Reports

Figure 1. Total Number of BNT162b2 AEs by System Organ Classes and Event Seriousness



Nervous System Disorders

Page 9 –Under "Nervous System Disorders" we see that headache is 24.1% of the total reported AEs.

3.6% parasthesias (tingling, burning), 2.4% hypoesthesia (numbness), totaling 5%. Reports indicate this is an early symptom of *neuropathy* and other nerve damage.

Small Fiber Neuropathy - CLICK HERE

CNS Inflammation Harvard Study - CLICK HERE

Neuro AutoImmune - CLICK HERE

POTS - CLICK HERE - CLICK HERE

5.3.6 Cumulative Analysis of Post-authorization Adverse Event Reports

Table 2. Events Reported in ≥2% Cases

		Cumulatively Through 28 February 2021
MedDRA SOC	MedDRA PT	AEs (AERP%) N = 42086
	Pain	3691 (8.8%)
	Malaise	2897 (6.9%)
	Asthenia	2285 (5.4%)
	Drug ineffective	2201 (5.2%)
	Vaccination site erythema	930 (2.2%)
	Vaccination site swelling	913 (2.2%)
	Influenza like illness	835 (2%)
Infections and infestations		
	COVID-19	1927 (4.6%)
Injury, poisoning and proce	dural complications	, , , , , , , , , , , , , , , , , , , ,
	Off label use	880 (2.1%)
	Product use issue	828 (2.0%)
Musculoskeletal and connec	tive tissue disorders	
	Myalgia	4915 (11.7%)
	Pain in extremity	3959 (9.4%)
	Arthralgia	3525 (8.4%)
Nervous system disorders	,	1 222 (31112)
	Headache	10131 (24.1%)
	Dizziness	3720 (8.8%)
	Paraesthesia	1500 (3.6%)
	Hypoaesthesia	999 (2.4%)
Respiratory, thoracic and m		
	Dyspnoca	2057 (4.9%)
	Cough	1146 (2.7%)
	Oropharyngeal pain	948 (2.3%)
Skin and subcutaneous tissu		3.40 (2.0.10)
	Pruritus	1447 (3.4%)
	Rash	1404 (3.3%)
	Erythema	1044 (2.5%)
	Hyperhidrosis	900 (2.1%)
	Urticaria	862 (2.1%)
Total number of events		93473

3.1.2. Summary of Safety Concerns in the US Pharmacovigilance Plan

Table 3. Safety concerns

Important identified risks	Anaphylaxis	
Important potential risks	Vaccine-Associated Enhanced Disease (VAED), Including Vaccine-associated Enhanced Respiratory Disease (VAERD)	
Missing information	Use in Pregnancy and lactation Use in Paediatric Individuals <12 Years of Age Vaccine Effectiveness	

Page 21 – Neurological AESIs

Serious neurological AEs of Special Interest (AESIs) were 1.2% (501) of the reports and include: Convulsions, Demyelination, PTs, Ataxia, Cataplexy, Encephalopathy, Fibromyalgia, Trigeminal Neuralgia, Intracranial pressure increased, Meningitis, aseptic Meningitis, Guillain Barre Syndrome, etc.

These are conditions that have a test, a name and a diagnosis code. 3.19% were reported as fatal, 17.7% not resolved and 32.1% unknown outcome.

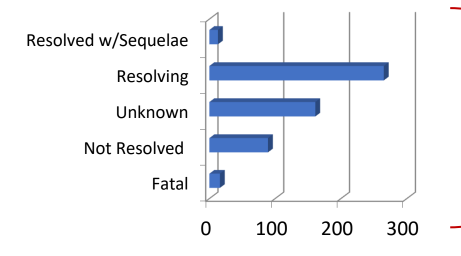


Table 7. AESIs Evaluation for BNT162b2

AESIs ^a	Post-Marketing Cases Evaluation ^b	
Category	Total Number of Cases (N=42086)	
	Relevant event outcome: resolved/resolving (1801), not resolved (959), resolved with sequelae (49), and unknown (853). Conclusion: This cumulative case review does not raise new safety issues. Surveillance will continue.	
Neurological AESIs (including demyelination) Search criteria: Convulsions (SMQ) (Broad and Narrow) OR Demyelination (SMQ) (Broad and Narrow) OR PTs Ataxia; Cataplexy; Encephalopathy; Fibromyalgia; Intracranial pressure increased; Meningitis; Meningitis aseptic; Narcolepsy	 Number of cases: 501 (1.2% of the total PM dataset), of which 365 medically confirmed and 136 non-medically confirmed. Country of incidence (≥9 cases): UK (157), US (68), Germany (49), Mexico (35), Italy (31), France (25), Spain (18), Poland (17), Netherlands and Israel (15 each), Sweden (9). The remaining 71 cases were from 22 different countries. Subjects' gender (n=478): female (328), male (150). Subjects' age group (n=478): Adult (329), Elderly (149); Number of relevant events: 542, of which 515 serious, 27 non-serious. Most frequently reported relevant PTs (>2 occurrences) included: Seizure (204), Epilepsy (83), Generalised tonic-clonic seizure (33), Guillain-Barre syndrome (24), Fibromyalgia and Trigeminal neuralgia (17 each), Febrile convulsion, (15), Status epilepticus (12), Aura and Myelitis transverse (11 each), Multiple sclerosis relapse and Optic neuritis (10 each), Petit mal epilepsy and Tonic convulsion (9 each), Ataxia (8), Encephalopathy and Tonic clonic movements (7 each), Foaming at mouth (5), Multiple sclerosis, Narcolepsy and Partial seizures (4 each), Bad sensation, Demyelination, Meningitis, Postictal state, Seizure like phenomena and Tongue biting (3 each); Relevant event onset latency (n = 423): Range from <24 hours to 48 days, median 1 day; Relevant events outcome: fatal (16), resolved/resolving (265), resolved with sequelae (13), not resolved (89) and unknown (161). 	
	Conclusion: This cumulative case review does not raise new safety issues. Surveillance will continue	

AutoImmune and Musculoskeletal AESIs

BNT162b2

5.3.6 Cumulative Analysis of Post-authorization Adverse Event Reports

Table 7. AESIs Evaluation for BNT162b2

AESIsa	Post-Marketing Cases Evaluation ^b
Category	Total Number of Cases (N=42086)
	2021). Study C4591021, pending protocol endorsement by EMA, is also intended to inform this risk.
Immune-Mediated/Autoimmune AESIs Search criteria: Immune- mediated/autoimmune disorders (SMQ) (Broad and Narrow) OR Autoimmune disorders HLGT (Primary Path) OR PTs Cytokine release syndrome; Cytokine storm; Hypersensitivity	 Number of cases: 1050 (2.5 % of the total PM dataset), of which 760 medically confirmed and 290 non-medically confirmed; Country of incidence (>10 cases): UK (267), US (257), Italy (70), France and Germany (69 each), Mexico (36), Sweden (35), Spain (32), Greece (31), Israel (21), Denmark (18), Portugal (17), Austria and Czech Republic (16 each), Canada (12), Finland (10). The remaining 74 cases were from 24 different countries. Subjects' gender (n=682): female (526), male (156). Subjects' age group (n=944): Adult (746), Elderly (196), Adolescent (2).
	 Number of relevant events: 1077, of which 780 serious, 297 non-serious. Most frequently reported relevant PTs (>10 occurrences): Hypersensitivity (596), Neuropathy peripheral (49), Pericarditis (32), Myocarditis (25), Dermatitis (24), Diabetes mellitus and Encephalitis (16 each), Psoriasis (14), Dermatitis Bullous (13), Autoimmune disorder and Raynaud's phenomenon (11 each);
	30 days, median <24 hours. • Relevant event outcome ¹ : resolved/resolving (517), not resolved (215), fatal (12), resolved with sequelae (22) and unknown (312).
	Conclusion: This cumulative case review does not raise new safety issues. Surveillance will continue
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Page 20 – Immune-Mediated/Autoimmune AESIs represented 2.5% of the reports. Of the 1077 reports, 780 were considered serious.

Hypersensitivity, <u>peripheral neuropathy</u>, pericarditis, myocarditis, encephalitis are immune mediated or autoimmune conditions.

Musculoskeletal AESIs Arthralgia, arthritis, rheumatoid arthritis, polyarthritis, polyneuropathy, post viral fatigue syndrome, chronic fatigue syndrome, etc.

Musculoskeletal AESIs

Search criteria: PTs Arthralgia; Arthritis; Arthritis bacterial^h; Chronic fatigue syndrome; Polyarthritis; Polyneuropathy; Post viral fatigue syndrome; Rheumatoid arthritis

- Number of cases: 3600 (8.5% of the total PM dataset), of which 2045 medically confirmed and 1555 non-medically confirmed;
- Country of incidence: UK (1406), US (1004), Italy (285), Mexico (236), Germany (72), Portugal (70), France (48), Greece and Poland (46), Latvia (33), Czech Republic (32), Israel and Spain (26), Sweden (25), Romania (24), Denmark (23), Finland and Ireland (19 each), Austria and Belgium (18 each), Canada (16), Netherlands (14), Bulgaria (12), Croatia and Serbia (9 each), Cyprus and Hungary (8 each), Norway (7), Estonia and Puerto Rico (6 each), Iceland and Lithuania (4 each); the remaining 21 cases originated from 11 different countries;
- Subjects' gender (n=3471): female (2760), male (711);
- Subjects' age group (n=3372): Adult (2850), Elderly (515), Child
 (4) Adolescent (2) Infant (1):
- Number of relevant events: 3640, of which 1614 serious, 2026 non-serious;
- Reported relevant PTs: Arthralgia (3525), Arthritis (70), Rheumatoid arthritis (26), Polyarthritis (5), Polyneuropathy, Post viral fatigue syndrome, Chronic fatigue syndrome (4 each), Arthritis bacterial (1);

32 days, median 1 day;

Cardiovascular AESIs

Page 16 —As of 28 Feb '21, 3.3% of the AE reports were *cardiovascular.* This is in a mostly older population as medical professionals and the elderly were vaccinated first.

Of the 1441 reports 946 (65.6%) were considered serious. 136(9.4%) were fatal 140 (9.7%) not resolved and 380 (26.3%) unknown.

Younger people are at increased risk of cardiovascular problems related to the vaccines, so these data are likely under representative of current reports and those expected in future post marketing experience documents. Myocarditis Case Report CLICK HERE

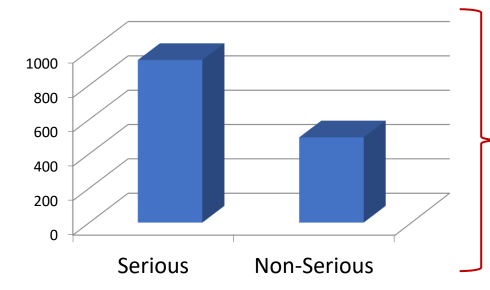


Table 7. AESIs Evaluation for BNT162b2

AESIs ^a	Post-Marketing Cases Evaluation ^b	
Category	Total Number of Cases (N=42086)	
Anaphylactic Reactions Search criteria: Anaphylactic reaction SMQ (Narrow and Broad, with the algorithm applied), selecting relevant cases according to BC criteria	Please refer to the Risk 'Anaphylaxis' included above in Table 4.	
Cardiovascular AESIs Search criteria: PTs Acute myocardial infarction; Arrhythmia; Cardiac failure; Cardiac failure acute; Cardiogenic shock; Coronary artery disease; Myocardial infarction; Postural orthostatic tachycardia syndrome; Stress cardiomyopathy; Tachycardia	 Number of cases: 1403 (3.3% of the total PM dataset), of which 241 are medically confirmed and 1162 are non-medically confirmed; Country of incidence: UK (268), US (233), Mexico (196), Italy (141), France (128), Germany (102), Spain (46), Greece (45), Portugal (37), Sweden (20), Ireland (17), Poland (16), Israel (13), Austria, Romania and Finland (12 each), Netherlands (11), Belgium and Norway (10 each), Czech Republic (9), Hungary and Canada (8 each), Croatia and Denmark (7 each), Iceland (5); the remaining 30 cases were distributed among 13 other countries; Subjects' gender: female (1076), male (291) and unknown (36); Subjects' age group (n = 1346): Adult^c (1078), Elderly^d (266) Child^c and Adolescent^f (1 each); Number of relevant events: 1441, of which 946 serious, 495 non-serious; in the cases reporting relevant serious events; Reported relevant PTs: Tachycardia (1098), Arrhythmia (102), Myocardial infarction (89), Cardiac failure (80), Acute myocardial infarction (41), Cardiac failure acute (11), Cardiogenic shock and Postural orthostatic tachycardia syndrome (7 each) and Coronary artery disease (6); Relevant event onset latency (n = 1209): Range from <24 hours to 21 days, median <24 hours; 	

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