

Exhibit 471

Report 71: Musculoskeletal Adverse Events of Special Interest Afflicted 8.5% of Patients in Pfizer's Post-Marketing Data Set, Including Four Children and One Infant. Women Affected at a Ratio of Almost 4:1 Over Men.

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Report 71: Musculoskeletal Adverse Events of Special Interest Afflicted 8.5% of Patients in Pfizer's Post-Marketing Data Set, Including Four Children and One Infant. Women Affected at a Ratio of Almost 4:1 Over Men.

May 15, 2023 • by Barbara Gehrett, MD; Joseph Gehrett, MD; Chris Flowers, MD; and Loree Britt

The War Room/DailyClout Pfizer Documents Analysis Project Post-Marketing Group (Team 1) – Barbara Gehrett, MD; Joseph Gehrett, MD; Chris Flowers, MD; and Loree Britt – wrote a review of musculoskeletal adverse events of special interest (AESIs) found in Pfizer document

5.3.6 Cumulative Analysis of Post-Authorization Adverse Event Reports of PF-07302048 (BNT162B2) Received Through 28-FEB-2021 (https://www.phmpt.org/wp-content/uploads/2022/04/reissue_5.3.6-postmarketing-experience.pdf)

(a.k.a., "5.3.6"). This group of AESIs includes diagnoses of arthralgia (joint pain), arthritis (joint inflammation), arthritis/bacterial, chronic fatigue syndrome, polyarthritis (inflammation of multiple joints), post-viral fatigue syndrome, and

rheumatoid arthritis (an autoimmune and inflammatory disease (<https://www.cdc.gov/arthritis/basics/rheumatoid-arthritis.html>)).

It is important to note that the AESIs in the 5.3.6 document were reported to Pfizer for **only a 90-day period** starting on December 1, 2020, the date of the United Kingdom's public rollout of Pfizer's COVID-19 experimental mRNA "vaccine" product.

Highlights of this report include:

- 3,600 cases of musculoskeletal AESIs were reported, which equates to **8.5% of the post-marketing data set** of 42,086 cases/patients.
- The 3,600 patients reported 3,640 adverse events. **1,614 (44%) were classified as serious.**
- The time from administration to adverse event ranged from less than 24 hours to 32 days. **50% of the events started within the first 24 hours after injection.**
- Of the cases where gender was reported, 2,760 individuals were female, and 711 were male – an almost **4:1 ratio of female to male.**
- Though mostly adults were affected with these AESIs, **two adolescents, four children, and one infant** also reported musculoskeletal AESIs during a time frame when **Pfizer's BNT162b2 mRNA COVID "vaccine" was not approved for use in individuals under 16 years of age.**
- The most common adverse event was **arthralgia/joint pain (3,525 or 97%),** followed by **70 arthritis AESIs (2%), 26 rheumatoid arthritis AESIs (<1%) and 5 AESIs (<1%) polyarthritits.**
- Outcome for 3,662 of the adverse events were: 1,801 (49%) resolved or resolving, **959 (26%) not resolved, 49 (1%) resolved with sequelae,** and **853 (23%) were unknown.**

Pfizer concluded, "This cumulative case review does not raise new safety issues. Surveillance will continue."

Please read the full report below.



War Room/DailyClout Pfizer Document Analysis

Post-Marketing Team Micro-Report 14:
Musculoskeletal Adverse Events of Special Interest Review of 5.3.6

SOURCE:
https://www.phmpt.org/wp-content/uploads/2022/04/reissue_5.3.6-postmarketing-experience.pdf

5.3.6 AE REPORTING PERIOD:
"Since the first temporary authorization for emergency supply under Regulation 174 in the UK (01 December 2020) and through 28 February 2021."

5.3.6 AE CASES/EVENTS:
TOTAL AE CASES: 42,086
TOTAL AE EVENTS: 158,893

ABBREVIATIONS:
5.3.6 : Pfizer source document

SOC : System Organ Class

AE : Adverse Event

AESI : Adverse Event of Special Interest

EUA : Emergency Use Authorization by FDA

PM : Post-Marketing

BNT162b2 : Pfizer's mRNA COVID-19 vaccine

SEQUELAE: an abnormal condition resulting from a previous disease, injury, or other trauma

AGE GROUPS defined in 5.3.6 (p. 25 footnote):
Adult 18 - 64
Elderly ≥ 65
Child 2 - 11
Adolescent 12 - < 18
Infant 1 - 23 months

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<p>Musculoskeletal AESIs Search criteria: PTs, Arthralgia; Arthritis; Arthritis bacterial; Chronic fatigue syndrome; Polyarthritides; Polymyositis; Post viral fatigue syndrome; Rheumatoid arthritis</p>	<ul style="list-style-type: none"> Number of cases: 3600 (8.5% of the total PM dataset), of which 2005 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 (35), 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), Polyarthritides (5), Polymyositis, Post viral fatigue syndrome, Chronic fatigue syndrome (4 each), Arthritis bacterial (1). Relevant event onset latency (n = 2968): Range from <24 hours to 32 days, median 1 day.
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https://www.phmpt.org/wp-content/uploads/2022/04/reissue_5.3.6-postmarketexperience.pdf

The category of musculoskeletal adverse events of special interest (AESI) is made up of these different diagnoses: arthralgia (joint pain), arthritis (joint inflammation), arthritis/bacterial, chronic fatigue syndrome, polyarthritides (inflammation of multiple joints), post viral fatigue syndrome, and rheumatoid arthritis.

Fibromyalgia is not included in this report but was listed under the category of "Neurological AESIs." Polymyopathy (symmetrical damage to peripheral nerves) is included in this report rather than under "Neurological AESIs."

3,600 cases were reported, **8.5% of the post-marketing data set** of 42,086 cases. These 3,600 individuals reported 3,640 events. **1,614 (44%) were classified as serious.** The most common adverse event was **arthralgia (3,525 or 97%)**, followed by 70 arthritis (2%), 26 rheumatoid arthritis (<1%) and 5 (<1%) polyarthritides.

- Adverse Events were reported to Pfizer during a 90-day period, following the December 1, 2020, public rollout of its COVID-19 experimental "vaccine" product.
- In the Pfizer 5.3.6 document, these AEs were categorized by System Organ Classes (SOC) – in other words, by systems in the body.

The time from administration to adverse event ranged from < 24 hours to 32 days. **50% of the events started within the first 24 hours.**

Of the cases where gender was reported, 2,760 individuals were female, and 711 were male. Once again, the pattern from 5.3.6 SOCs continues, with an almost **4:1 ratio of female to male.**

Age distribution was **predominantly adult (2,850, 85%)**, with 515 (15%) elderly, **two adolescents, four children, and one infant.** During these dates, **BNT162b2 was not approved for use in children or infants.** Approval was only for adolescents 16 and above (not clarified by Pfizer).

Outcomes were reported for 3,662 events: 1,801 (49%) resolved or resolving, **959 (26%) not resolved, 49 (1%) resolved with sequelae, and 853 (23%) were unknown.**

Pfizer's Conclusion:

"This cumulative case review does not raise new safety issues. Surveillance will continue."

Note: As of the date of this team report, no follow-up safety surveillance data have been publicly released.



RECALL
this unsafe "vaccine."

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