

THE PFIZER INOCULATIONS FOR COVID-19

# MORE HARM THAN GOOD



Contact us  
[info@canadiancovidcarealliance.org](mailto:info@canadiancovidcarealliance.org)  
[www.canadiancovidcarealliance.org](http://www.canadiancovidcarealliance.org)

PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



## WHO WE ARE

Our alliance of **over 500 independent Canadian doctors, scientists, and health care practitioners** is committed to providing quality, balanced, evidence-based information to the Canadian public about COVID-19 so that hospitalizations can be reduced, lives saved, and our country safely restored to normal as quickly as possible.

PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



## **WE SUPPORT**

**The doctor/patient relationship** and personalized care

**Informed consent** and treatment options

Free and open **scientific discourse**

**Safe & effective** vaccines



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



## FIRST, DO NO HARM

The federal, provincial and municipal governments in Canada have a **responsibility to protect the health of Canadians as well as our Charter Rights and Freedoms. Any medical interventions approved by Health Canada must first be PROVEN SAFE.**

**Due diligence** in research, as well as **adherence to established protocols of the doctor/patient relationship, informed consent and scientific inquiry** are essential to carrying out that responsibility.

**Deviating from those practices, causing harm and failing to disclose risks of harm is negligent at best.**





# OVERVIEW

## Hierarchy of evidence

### Pfizer's 2 month data report, Dec 31 2020

- ARR vs RRR explained - VIDEO
- Early unblinding of Pfizer's randomized control trial

### Pfizer's 6 month data report, Sep 15 2021

- Increased risk of illness
- Increased risk of death

### The Pfizer Trials - What went wrong

- Pfizer did not follow established protocols
- Misleading demographics - Wrong age
- Misleading demographics - Tested on healthy, given to sick
- Inadequate control groups
- Did not track biomarkers
- Wrong clinical endpoints
- Not tested for spread reduction
- Subjective testing
- Missing data - lost to follow up and Suspected, but unconfirmed

- Failure to test - Why it matters
- 12 - 15 trial - All risk, no benefit
- 12 - 15 trial - Failure to report serious adverse events
- 5 - 11 year olds - Risking their health
- Myocarditis is serious
- The FDA abandons "First, do no harm"
- 5 - 11 year olds - No informed consent
- The BMJ Pfizer trial whistleblower article

### A critical eye on the Sep 15 2020 report

- 6 month data manipulation - Mixed cohorts proved harm
- The Pfizer trials did not prove safety - they proved harm

### How this is playing out in the real world

- Roll out surveillance - You don't find what you don't look for
- Rising incidents of heart issues in young people (Ontario Public Health Report)
- This is not normal - High incidences of deaths in athletes (German, Israeli news articles)

- This is supposed to be rare - VIDEO of athletes collapsing
- Pfizer's post marketing pharmacovigilance report

### Considerable evidence of conflict of interest

- Pfizer is making billions
- The public record of Pfizer's corporate culture
- Links to articles on Pfizer's past behaviour
- Conflicts of interest among Pfizer report authors
- The CDC has redefined "vaccing"
- The media has been captured - VIDEO

### This is no way to manage a supplier

The inoculations should be withdrawn immediately

### Recommended reading & viewing



# THE HIERARCHY OF EVIDENCE

## Levels of Scientific Evidence

Level	Example of Evidence
<b>Level 1</b>	Meta-analysis of Homogenous RCTs <b>Randomized Control Trial</b>
<b>Level 2</b>	Meta-analysis of Level 2 or Heterogenous Level 1 Evidence Prospective Comparative Study
<b>Level 3</b>	Review of Level 3 Evidence Case-control Study Retrospective Cohort Study
<b>Level 4</b>	Uncontrolled Cohort Studies Case Series
<b>Level 5</b>	Expert Opinion Case Report Personal Observation
<b>Foundational Evidence</b>	Animal Research <i>In Vitro</i> Research Ideas, Speculation

- **A randomized control trial is LEVEL 1 Evidence**, the highest form of evidence there is. It is considered the Gold Standard and is the only way to prove something is true.
- **Models are LEVEL 5 or lower** as they are expert opinion/speculation.
- **Policy should be determined by the highest level of evidence available, LEVEL 1.**



# PFIZER'S ORIGINAL TRIAL REPORT

## DECEMBER 31 2020

- Published in New England Journal of Medicine
- Showed **2 months worth of safety & efficacy data**
- Described starting with 43,548 people divided into:
  1. **Treatment group** (received inoculation)
  2. **Control group** (received saline) for 2 months to see who developed COVID-19
- The claim was that the inoculations were safe and showed **95% efficacy 7 days after the 2nd dose**. But that 95% was actually **Relative Risk Reduction**. **Absolute Risk Reduction** was only **0.84%**.

THE NEW ENGLAND JOURNAL OF MEDICINE  
RESEARCH SUMMARY

### Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine

F. P. Polack, et al. DOI: 10.1056/NEJMoa2026437

**CLINICAL PROBLEM**  
Safe and effective vaccines to prevent severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infections and hospitalizations are needed. Existing vaccines against SARS-CoV-2 and other coronaviruses are currently available, and mRNA-based vaccines have not been widely tested.

**CLINICAL TRIAL**  
A double-blind study of an mRNA vaccine encoding the SARS-CoV-2 spike protein.

**DESIGN**  
A 3,548 participants, 316 years old were assigned to receive the vaccine or placebo by intramuscular injection on day 0 and day 21. Participants were followed for 90 days for the development of symptomatic Covid-19 for a median of 2 months.

**RESULTS**  
Vaccine recipients had local reactions (pain, erythema, swelling) at higher rates than placebo recipients, with more reactions following the second dose. Most were mild to moderate and resolved rapidly.

**CONCLUSIONS**  
Two months after inoculation, the mRNA vaccine provided 95% protection against asymptomatic Covid-19 in persons 16 years of age or older.

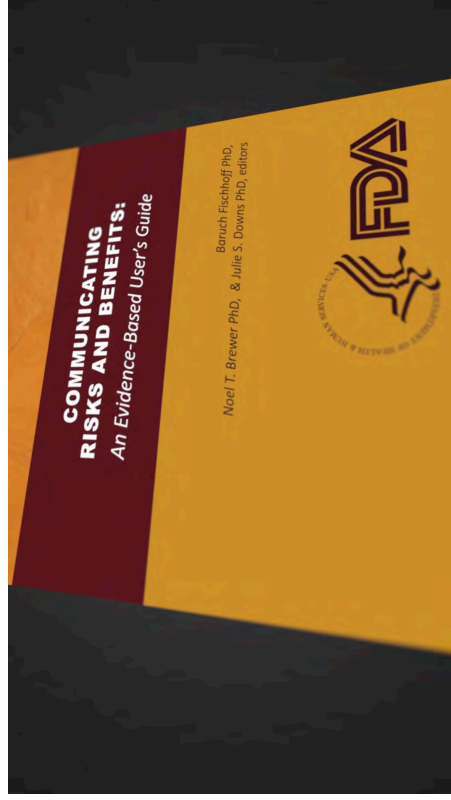
Vaccine efficacy of 95% (95% credible interval, 90.1–97.6%)

Copyright © 2020 Massachusetts Medical Society  
Tables: Full article | Quick Take | Internal



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# ABSOLUTE RISK REDUCTION VS RELATIVE RISK REDUCTION



<https://rumbled.com/vobcg5-relative-vs-absolute-risk-reduction.html>





# EARLY UNBLINDING OF RANDOMIZED CONTROL TRIAL = NO LONG TERM SAFETY DATA

## WHAT WAS SUPPOSED TO HAPPEN

	INOCULATED GROUP	PLACEBO GROUP	
2020			<p><b>July 27 2020</b>  <b>Phase III Begins</b>                      The participants are evenly divided into the inoculated and placebo groups of about 21,000 each. The study is <b>blinded</b>, so participants don't know which group they are in.</p>
2021	↓	↑	
2022	↓	↑	
2023	↓	↑	<p><b>May 2 2023</b>  <b>End of Phase III Clinical Trial</b>                      This is the point where the trial can be <b>unblinded</b> and the Placebo group offered the intervention if it's indicated and they consent.</p>

## WHAT ACTUALLY HAPPENED

	INOCULATED GROUP	PLACEBO GROUP	
2020			<p><b>July 27 2020</b>  <b>Phase III Begins</b>                      The participants are evenly divided into inoculated and Placebo groups of about 21,000 each. The study is <b>blinded</b>.</p> <p><b>Dec 31 2020</b>                      Release 2 month data report. The trial is unblinded early.</p>
2021		NO DATA	<p><b>Crossover Occurs</b>                      The participants from the <b>Placebo Group</b> are given the opportunity to take the inoculation and by early 2021, the majority of them have crossed over to the inoculated group. It's <b>no longer a randomized control trial, as control group is gone.</b></p>
2022	↓ ↓	NO DATA	
2023	↓ ↓	NO DATA	<p><b>May 2 2023</b>  <b>End of Phase III Clinical Trial</b>                      The long term safety data that was supposed to be assessed at this point is <b>no longer possible to ascertain as the placebo group crossed over two years previously.</b></p>



# PFIZER'S 6 MONTH REPORT DATA LEVEL 1 EVIDENCE OF HARM

- Pfizer's most recent report indicates an **Efficacy of 91.3%**.  
(Which means a **reduction in positive cases** compared to placebo group.)
- **But it also showed**, compared to the placebo group, **an increase in illness and deaths.**
- There is **no benefit to a reduction in cases** if it comes at the cost of **increased sickness and death.**

THE NEW ENGLAND JOURNAL OF MEDICINE

ORIGINAL ARTICLE

### Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine through 6 Months

S. J. Thomas, E. D. Moreira Jr., N. Reichlin, J. Abubakar, A. Gurtman, S. Lockhart, J. L. Peters, C. Peters-Mac, J. P. Polack, C. Zelen, R. B. Bailey, K. A. Swanson, R. W. Frerking, Jr., L. L. Hammitt, O. Turndorf, H. Neff, A. Schaefer, S. Khand, Q. Yan, P. Liberman, D. E. Thornton, S. Mather, P. S. Dominant, U. S. Jain, W. C. Gierber, and G. J. Jensen, for the COVID-19 Clinical Trial Group

---

**ABSTRACT**

**CONCLUSIONS** BNT162b2 is a lipid nanoparticle-formulated, nucleoside-modified RNA vaccine encoding a prefusion-stabilized, membrane-anchored severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike protein. It was granted emergency use authorization for use against coronavirus disease 2019 (COVID-19) and is currently approved, conditionally approved, or authorized for emergency use worldwide. At the time of initial authorization, data beyond 2 months after vaccination were unavailable in an ongoing, placebo-controlled, observer-blinded, multinationa, pivotal efficacy trial, we randomly assigned 4,415 participants 18 years of age or older and 2,094 participants 16 years of age or older to receive either two doses of BNT162b2 or placebo. The trial and sites were selected for efficacy against laboratory-confirmed COVID-19 and safety, which were both evaluated through 6 months after vaccination.

BNT162b2 continued to be safe and have an acceptably adverse-event profile. Few participants had adverse events leading to withdrawal from the trial. Vaccine effectiveness against laboratory-confirmed COVID-19 was 91.3% through 6 months of follow-up among the participants without evidence of previous SARS-CoV-2 infection who could be evaluated. There was a gradual decline in vaccine effectiveness against COVID-19 over time, and the effectiveness was lower in populations with diverse ages, sexes, or ethnic groups, and risk factors for COVID-19 among participants without evidence of previous infection with SARS-CoV-2. The effectiveness against COVID-19 was lower in participants from South Africa, where the SARS-CoV-2 variant of concern B.1.351 (or beta) was predominant, a vaccine efficacy of 100% (95% CI, 53.5 to 100) was observed.

**CONCLUSIONS** The safety and efficacy results of this vaccine efficacy trial suggest that BNT162b2 had a favorable safety profile and was highly efficacious in preventing COVID-19. (Funded by BiNTTech and Pfizer; ClinicalTrials.gov number, NCT04383934.)

DOI: 10.1056/NEJMoa2110345

Copyright © 2022 Massachusetts Medical Society. All rights reserved.

<https://www.nejm.org/doi/pdf/10.1056/NEJMoa2110345?article=fulltext>

PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# INCREASED RISK OF ILLNESS

A **significant increase in illness**, which the Pfizer inoculations were supposed to reduce.

Screen capture from Pfizer 6 Month Supplementary Appendix

Adverse Event	BNT162b2 n, n (%)	Placebo n, n (%)
Any event	6617 (30.2)	3048 (13.9)
Related <sup>a</sup>	5241 (23.9)	1311 (6.0)
Life-threatening	282 (1.2)	150 (0.7)
Any serious adverse event	127 (0.6)	116 (0.5)
Related <sup>a</sup>	3 (0.0)	0
Severe	71 (0.3)	66 (0.3)
Life-threatening	21 (0.1)	26 (0.1)
Any serious event leading to withdrawal	13 (0.1)	11 (0.1)
Related <sup>a</sup>	13 (0.1)	11 (0.1)
Severe	10 (0.0)	10 (0.0)
Life-threatening	3 (0.0)	7 (0.0)
Death	3 (0.0)	5 (0.0)

**Table S1 Participants Reporting at Least 1 Adverse Event from Dose 1 to 1 Month After Dose 2 During the Blinded Follow-up Period.** The population included all ≥16-year-old participants who received ≥1 dose of vaccine irrespective of follow-up time. a. N=number of participants in the specified group. This value is the denominator for the percentage calculations. b. n=Number of participants reporting ≥1 occurrence of the specified event category. The numerator for the percentage calculation is the number of participants reporting ≥1 occurrence of the specified event category. The denominator is the number of participants who received ≥1 dose of vaccine. c. Serious adverse events are defined as death, ventricular arrhythmia, or ventricular fibrillation. d. Shoulder injury related to vaccine administration, right axillary lymphadenopathy, and paroxysmal ventricular arrhythmia (as previously reported). Adverse events for 12-15-year-old participants were reported previously.<sup>11</sup>

Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine Through 6 Months... Supplementary Appendix

	BNT162b2	Placebo	Risk Change
<b>Efficacy</b> (Meaning number of people diagnosed with COVID-19.)	77	850	<b>-91%</b>
<b>Related Adverse Event</b> (Meaning an investigator has assessed it as related to the BNT162b2 injection.)	5,241	1,311	<b>+300%</b>
<b>Any Severe Adverse Event</b> (Interferes significantly with normal function.)	262	150	<b>+75%</b>
<b>Any Serious Adverse Event</b> (Involves visit to ER or hospitalization.)	127	116	<b>+10%</b>



# INCREASED RISK OF DEATH

Screen capture from Pfizer 6 Month Supplementary Appendix

Reported Cause of Death*	BNT162b2 (N=21,929)	Placebo (N=21,921)
<b>Deaths</b>	<b>15</b>	<b>14</b>
Acute respiratory failure	0	1
Aortic rupture	0	1
Atherosclerosis	0	0
Blurred or medicated vision	0	0
<b>COVID-19</b>	<b>0</b>	<b>2</b>
<b>COVID-19 pneumonia</b>	<b>0</b>	<b>2</b>
Cardiac arrest	4	1
Cardiac failure congestive	1	0
Cardiorespiratory arrest	1	1
Chronic obstructive pulmonary disease	1	1
Death	0	1
Dementia	0	1
Empyematous cholecystitis	1	0
Hemorrhagic stroke	0	1
<b>Ischemic heart disease</b>	<b>1</b>	<b>0</b>
Lung cancer metastatic	1	0
Metastases to liver	0	1
Meningeal	0	1
Multiple organ dysfunction syndrome	0	1
<b>Myocardial infarction</b>	<b>0</b>	<b>2</b>
Myocarditis	0	1
Pneumonia	0	2
Sepsis	1	0
Sepsis shock	1	0
Septic arthritis	1	0
Unrecoverable event	1	0

**Table S4 | Causes of Death from Dose 1 to Unblinding (Safety Population, ≥16 Years Old), n.**  
Multiple causes of death could be reported for each participant. There were no deaths among 12- to 15-year-old participants.

\*As used in Pfizer's COVID-19 Clinical Trial Protocol, Version 1.0, 12/2020. See Pfizer's COVID-19 Clinical Trial Protocol, Version 1.0, 12/2020, for details.

## BNT162b2 Placebo

Deaths before unblinding <small>(In Table S4 of Supplementary Appendix)</small>	BNT162b2	Placebo
<b>Deaths before unblinding</b> <small>(Not in table, but mentioned in text of 6 month report. See quote below.)</small>	<b>15</b>	<b>14</b>
<b>Deaths after unblinding</b> <small>(Not in table, but mentioned in text of 6 month report. See quote below.)</small>	<b>5</b>	
<b>Total Deaths</b>	<b>20</b>	<b>14</b>

"After unblinding" means when the Placebo participants were given the opportunity to "cross over" and take the BNT162b2 inoculation. \*

**"...3 participants in the BNT162b2 group and 2 in the original placebo group who received BNT162b2 after unblinding died."**  
*Safety and Efficacy of the BNT162b2 mRNA COVID-19 Vaccine Through 6 Months*

### Concerning Causes of Death

	BNT162b2	Placebo
<b>Total COVID-19 Related Deaths</b>	<b>1</b>	<b>2</b>
<b>Deaths Related to Cardiovascular Events</b>	<b>9</b>	<b>5</b>

\*As used in Pfizer's COVID-19 Clinical Trial Protocol, Version 1.0, 12/2020. See Pfizer's COVID-19 Clinical Trial Protocol, Version 1.0, 12/2020, for details.

**THE PFIZER TRIALS**  
**WHAT WENT WRONG**





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# PFIZER DID NOT FOLLOW ESTABLISHED PROTOCOLS

Regarding the persistent claim that the COVID-19 inoculation products do not need to be tested, because mRNA technology has already undergone testing: mRNA technology is the delivery mechanism, not the inoculation. That's like saying that since we've used syringes safely before, anything injected via syringe is safe. (And in fact, there are still a lot of unknowns about the effects of the mRNA delivery mechanism.)

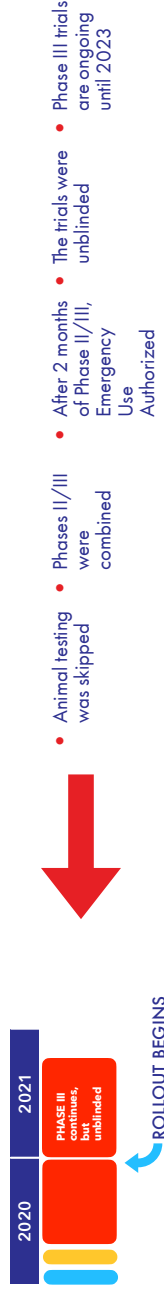
NORMALLY, VACCINE DEVELOPMENT LOOKS LIKE THIS, WITH A TIMELINE OF 5 TO 10 YEARS.



RARELY, IT CAN BE DONE IN AS LITTLE AS 5 YEARS.



FOR THE COVID-19 INOCULATIONS, IT WAS DONE IN 1 YEAR.

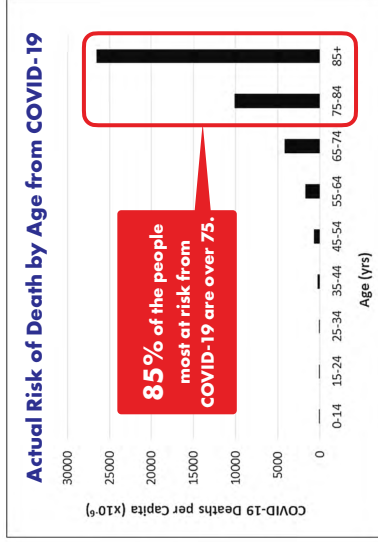




PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# MISLEADING DEMOGRAPHICS WRONG AGE FOR TARGET POPULATION

When designing a trial for the efficacy and safety of a potential treatment, the focus should be on the target population who could most benefit from that treatment. Instead Pfizer chose participants from younger demographic that would be a) less likely to need a vaccine, b) less likely to suffer an adverse event during a trial, c) more likely to respond well to a vaccine, as the elderly have comparatively poor immune responses.



COVID-19 Deaths per capita by age in the United States (as of Jan 5, 2021). Population based on U.S. CDC WONDER Bridge View Population Estimate 2019. Data obtained from [https://wonder.cdc.gov/bridgeview.aspx?\\_lang=en](https://wonder.cdc.gov/bridgeview.aspx?_lang=en)

### Pfizer Trial Demographics

Demographics (population for the primary efficacy endpoint). The number of participants who received vaccine and placebo, stratified by age.

AGE GROUP	Pfizer-BioNTech COVID-19 Vaccine (N = 18,242) n (%)	Placebo (N = 18,379) n (%)
≥12 through 15 years <sup>b</sup>	46 (0.3 %)	42 (0.2 %)
≥16 through 17 years	66 (0.4 %)	68 (0.4 %)
≥18 through 64 years	14,216 (77.9 %)	14,299 (77.8 %)
≥65 through 74 years	3176 (17.4 %)	3226 (17.6 %)
≥75 years	804 (4.4 %)	812 (4.4 %)

Yet 75+ year olds represent only 4% of trial subjects.

FACT SHEET FOR HEALTHCARE PROVIDERS ADMINISTERING VACCINE (VACCINATION PROVIDERS) EMERGENCY USE AUTHORITY FOR COVID-19 VACCINE TO PREVENT CORONAVIRUS DISEASE 2019 (COVID-19)  
<https://labeling.fda.gov/2020/06/24/labeling/ucm746444.htm>







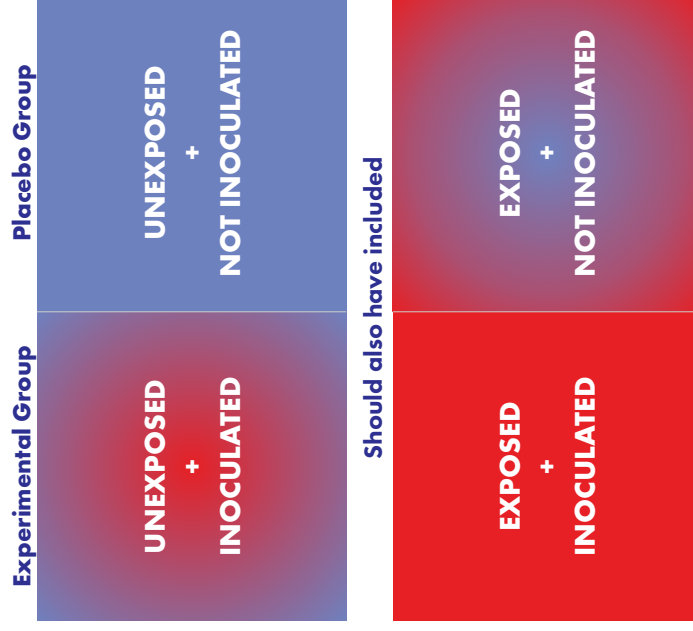
## INADEQUATE CONTROL GROUPS

Pfizer only observed 2 groups:

- **UNEXPOSED & INOCULATED**
- **UNEXPOSED & NOT INOCULATED**

They should have included two more groups:

- **EXPOSED & INOCULATED**, people who had recovered, then got the inoculation, to see if the inoculation was safe for them
- **EXPOSED & NOT INOCULATED** people who were recovered and not inoculated to see how the inoculations stacked up against natural immunity





## LOW QUALITY SAFETY SCIENCE DIDN'T TRACK BIOMARKERS

As Kostoff et al. highlighted in a recent paper, "Why are we vaccinating children against COVID-19?" (highly recommended), that while the Pfizer trials tested for antibodies and tracked adverse events in terms of symptoms, **they didn't test for adverse events at the subclinical (pre-symptom) level.**

This was extremely unsafe, because **symptoms/ diseases are typically end points of processes** that can take months, years, or decades to surface. By the time you get to symptoms, things can have gone pretty wrong. (Think diabetes or high blood pressure, where the disease can be quite advanced before any symptoms occur.) **Pfizer should have been tracking biomarkers that would have been early warning indicators for disease caused by the inoculations.**

**High quality safety science would have meant they should have tested before & after inoculation for:**

- d-dimers for evidence of enhanced **coagulation/ clotting** (several of our doctors have noticed increased levels of d-dimers in inoculated patients presenting with stroke like symptoms - video available [here](#))
- C-reactive protein for evidence of enhanced **inflammation**
- troponins for evidence of **cardiac damage**
- occludin and claudin for evidence of enhanced **barrier permeability**
- blood oxygen levels for evidence of enhanced **hypoxia**
- amyloid-beta and phosphorylated tau for evidence of increased **predisposition to Alzheimer's disease**
- Serum HMGB1, CXCL13, Dickkopf-1 for evidence of an **increased disposition to autoimmune disease, etc.**



**Micro-clots** resulting from the inoculation that were insufficient to cause observable symptoms **could raise the baseline for thrombotic disease.**

RONALD N. KOSTOFF, A. \* DANIELA CAJINA, B. DARA KANIK, C. MICHAEL BRIGGS, D. RANWOTIS VACHONWANNIKULDEE, ANDREW A. SVETIMOV, F. ANISIOS SAKSIS, \*[WITZKATHE@WACHONWANNIKULDEE.COM](mailto:WITZKATHE@WACHONWANNIKULDEE.COM)



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## WRONG CLINICAL ENDPOINTS SHOULD HAVE FOCUSED ON ALL CAUSE MORTALITY & ILLNESS

The fear with COVID-19, was that it was going to a) kill people, b) make them sick.

So any COVID-19 vaccine clinical trial should set out to ask the question "Do people who take the vaccines have less illness and death than those who don't?"

Illness + Death should be the CLINICAL ENDPOINTS. And not just illness + death with COVID-19, but any and all illness and death, in order to make sure that the vaccines are not causing harm.

This is well known. It was learned decades ago with cancer drug trials. At first, they used a clinical endpoint of "Did the drug shrink the cancer?" If it did, they called it effective. But it turned out the drugs were not only killing cancer, they were killing patients. They were forced to change the design of their trials and switch to "all cause mortality" as the primary endpoint instead and show that people receiving the drug actually live longer than those who don't. (J.Bart Classen has written an excellent research article on the subject. Read [here](#).)

### WHAT SHOULD HAVE HAPPENED

(After the proper early safety phases of development were completed.)

"Do people who take the vaccines have less illness and death than those who don't?"

YES. Proceed to long terms safety studies.

NO. Go back to the drawing board.

### WHAT ACTUALLY HAPPENED

(Without the proper early safety phases of development having been completed.)

"Do people who take the vaccines test positive for COVID-19 less often?"

YES. Proceed to world wide roll out.

NO. (The trial set up made this result unlikely).



## NOT TESTED FOR SPREAD REDUCTION VACCINE PASSPORTS UNJUSTIFIED

Although vaccine passports are now being used to ostensibly prevent or reduce transmission of COVID-19, this outcome was never studied in the trial and it is inappropriate to assign that capability to these inoculations. **There is no evidence at all that they reduce the spread of disease and transmission was never one of the study's endpoints.**

### LIMITATIONS AND REMAINING QUESTIONS

Further study is required to understand the following:

- Safety and efficacy beyond 2 months and in groups not included in this trial (e.g., children, pregnant women, and immunocompromised persons).
- Whether the vaccine protects against asymptomatic infection and transmission to unvaccinated persons.
- How to deal with those who miss the second vaccine dose.

**Verify Ontario:**  
Ontario's official app for verifying COVID-19 vaccine certificates.

When a business or organization scans a visitor's digital or paper QR code, this app will:

- protect user privacy by only reading certificates that are trusted and secure
- check if a certificate is valid and the visitor can enter
- show a visitor's name and date of birth so their identity can be verified
- work offline (without an internet connection)

Download the Verify Ontario app at:  
[ontario.ca/verify](https://ontario.ca/verify)

Ontario



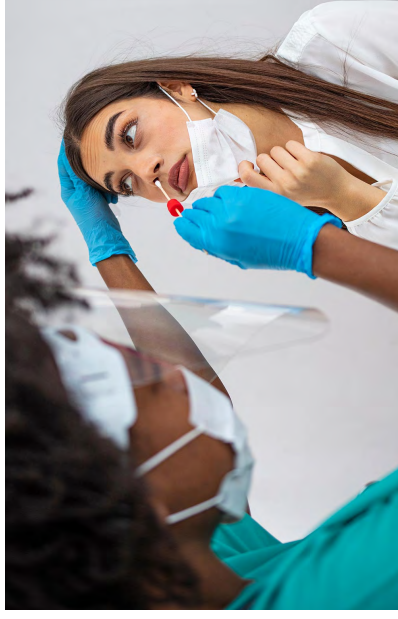
PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## TESTING FAILURES SUBJECTIVE TESTING

**The Pfizer trials DID NOT test all participants for COVID-19.** Instead, they instructed their investigators to test only those with a COVID-19 symptom and **left it up to their discretion** to decide what those were.

This means that:

- ♦ **Asymptomatic infection would be missed entirely**
- ♦ **A high level of subjectivity was introduced to the study - an investigator had the ability to sway the results**
- ♦ **The lack of objective systematic testing makes results unreliable**



All participants should have been tested.



# MISSING DATA

- ◆ **LOST TO FOLLOW UP**
- ◆ **SUSPECTED, BUT UNCONFIRMED**

	INOCULATED GROUP	PLACEBO GROUP
ENDPOINT DATA - Confirmed COVID Cases	8	162
Participants Lost to Follow Up	80	86
Suspected, but Unconfirmed Cases	1,594	1,816

The basis for the Emergency Use Authorization was the Confirmed COVID cases of 8 vs 162, which meant a Relative Risk Reduction of 95%. But **when dealing with such a small number of cases, any change can impact the results significantly.**

**Lost to follow up** means **they lost touch with those subjects** and can't confirm whether they got sick or not. They don't know.

**Suspected, but unconfirmed** means these people were **symptomatic for COVID-19**, but were **never tested**. (Discretion for testing was left up to the investigator.)

The fact that the Lost to Follow Up and Suspected but Unconfirmed numbers are higher - and here they are even significantly higher - than the End Point numbers means that **this data is unreliable. The study should not have been accepted in this state.** In normal scientific practice they should have returned to investigate further.

## Confirmed Cases

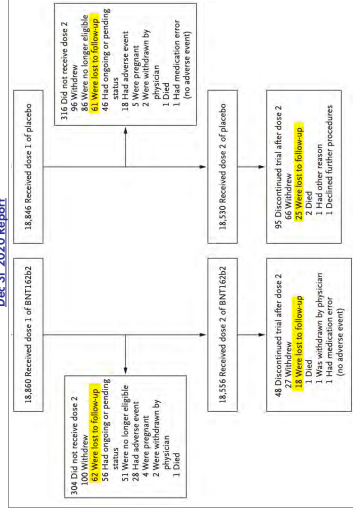
Dec. 31, 2020 Report

**Table 3. Vaccine Efficacy Overall and by Subgroup in Participants without Evidence of Infection before 7 Days after Dose 2.**

Subgroup	RNT16243 (N=18,198)	Placebo (N=18,225)	Vaccine Efficacy, % (95% CI)
Overall	No. of Cases (No. at Risk)* 8 (2,214 (17,411))	No. of Cases (No. at Risk)* 162 (2,222 (17,511))	95.0 (90.4-97.9)

## Lost to Follow Up

Dec. 31, 2020 Report



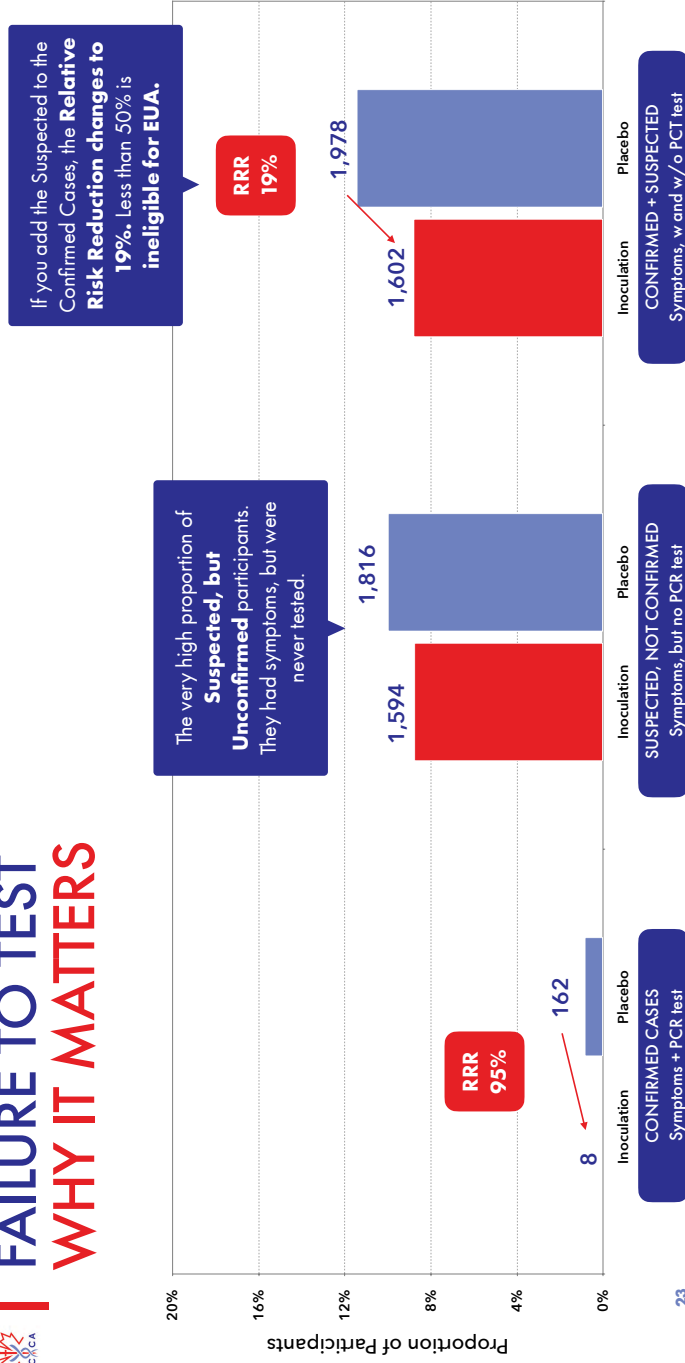
**Suspected but Unconfirmed**  
FDA Briefing Document: Pfizer-BioNTech COVID-19 Vaccine  
Vaccines and Related Biological Products Advisory Committee Meeting December 10, 2020

Among 3410 total cases of suspected but unconfirmed COVID-19 in the overall study population, 1594 occurred in the vaccine group vs. 1816 in the placebo group. Suspected COVID-19 cases that occurred within 7 days after any vaccination were 409 in the vaccine group vs. 287 in the placebo group. It is possible that the imbalance in suspected COVID-19 cases occurring in the study population is related to the study design, which requires that over 90% of those with suspected COVID-19 be confirmed by PCR testing. It is also possible that some of the suspected but unconfirmed COVID-19 cases were not reported to the investigators. It is also possible that some of the suspected but unconfirmed COVID-19 cases could have been clinically significant adverse events that would not have otherwise been detected.

PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



# FAILURE TO TEST WHY IT MATTERS





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## 12-15 ADOLESCENT TRIAL ALL RISK, NO BENEFIT

- This study was severely underpowered, as **a study this small will not show up risk.**
  - Inoculated group - **1,005 (0)** tested positive for COVID-19)
  - Placebo group - **978 (18)** tested positive for COVID-19)
- Pfizer claimed these were great results, but since adolescents are at statistically 0% risk of death from COVID-19, and very low risk of severe illness, **the inoculation is of little benefit to them.** Instead, it presents a very real risk of adverse events.
- But the adolescent Pfizer study wasn't actually designed to find those. **A serious adverse event**, including death, that occurred at a 1/800 rate **might not even show up in a sample of 1,005** people.
- But in this case, it did. **Among the 1,005 adolescents, there WAS at least one serious adverse event - Maddie de Garay.**



"For children without a serious medical condition, the danger of severe Covid is so low as to be difficult to quantify."  
-COVID AND AGE, Oct. 12, 2021, New York Times





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## 12-15 ADOLESCENT TRIAL FAILURE TO REPORT SERIOUS ADVERSE EVENTS

**Maddie de Garay** is a 12 year old trial participant who developed a serious reaction after her second dose and was hospitalized within 24 hours.

Maddie developed gastroparesis, nausea and vomiting, erratic blood pressure, memory loss, brain fog, headaches, dizziness, fainting, seizures, verbal and motor tics, menstrual cycle issues, lost feeling from the waist down, lost bowel and bladder control and had an nasogastric tube placed because she lost her ability to eat. She has been hospitalized many times, and for the past **10 months she has been wheelchair bound and fed via tube.**

In their report to the FDA, **Pfizer described her injuries as "functional abdominal pain."**

- One participant experienced an SAE reported as generalized neuralgia, and also reported a treatment emergent SAE (abdominal pain, ataxia, gait issues) and another SAE (nausea, vomiting, dizziness). The SAE (nausea, vomiting, dizziness) was **disregarded with functional abdominal pain.** The event was reported as ongoing at the time of the cutoff date.

Emancipate.Uk.Aufhorst@doz.Amendment





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



# MYOCARDITIS IS SERIOUS

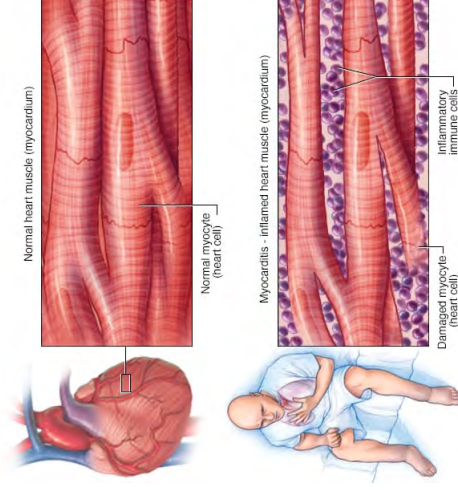
## MYOCARDITIS

"Myocarditis is an inflammatory process of the myocardium. (Heart muscle.) **Severe myocarditis weakens your heart** so that the rest of your body doesn't get enough blood. Clots can form in your heart, **leading to a stroke or heart attack.**"

THE U.S. NATIONAL CENTRE FOR BIOTECHNOLOGY INFORMATION

**"The mortality rate is up to 20% at 6.5 years."**

<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4926492/>



© IMVIO FOUNDATION FOR MEDICAL EDUCATION AND RESEARCH. ALL RIGHTS RESERVED.



# THE FDA ABANDONS FIRST, DO NO HARM

Medical interventions are supposed to be **PROVEN SAFE BEFORE** the are rolled out in the population.

Yet **Dr. Eric Rubin**, one of the 18 members of the **FDA advisory panel** who voted, to approve the inoculations for children 5 - 11, actually said the opposite, and suggested that **a population level roll out was an appropriate way to test for adverse events.**

It's worth noting that Dr. Eric Rubin is the **editor-in-chief of the New England Journal of Medicine, which publishes the Pfizer trial reports.**



**"We're never going to learn about how safe this vaccine is unless we start giving it. That's just the way it goes. That's how we found out about rare complications of other vaccines like the rotavirus vaccine. And I do think we should vote to approve it."**

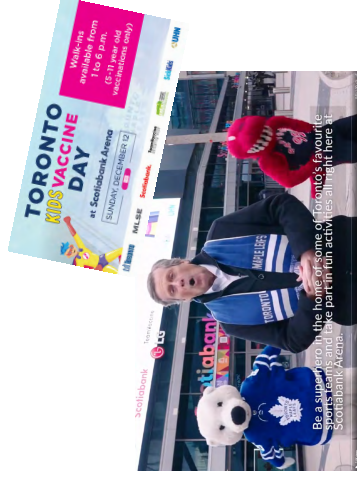
*Dr. Eric Rubin, FDA advisory panel member,  
Harvard professor & editor-in-chief of the New England Journal of Medicine  
Vaccines and Related Biological Products Advisory Committee — 10/26/2021*



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## 5 - 11 YEAR OLDS NO INFORMED CONSENT

- **Direct-to-consumer advertising of prescription drugs is illegal in Canada**, yet politicians from all levels of government are marketing inoculations to children, using cartoons and mascots.
- **They are proclaiming the inoculations to be safe, yet the data is not there to back that up.** In addition to admitting that their inoculations can cause myocarditis, Pfizer also admits, right in their report, that **their long term immune response, efficacy & safety data is limited and that their studies weren't powered to find "rare" side effects** as only 1,517 kids got the inoculation.
- How many parents would take their kids to get this shot if they were informed of this? **The law of informed consent says they should be, but it's not happening.**



Be a superstar in the home of some of Toronto's favourite sports teams and take part in fun activities all right here at Scotiabank Arena.

of a Covid-19 vaccine in this population; trials of other vaccines are under way. Limitations of the study include the lack of longer-term follow-up to assess the duration of immune responses, efficacy, and safety. However, longer-term follow-up from this study, which will continue for 2 years, should provide clarification. This study was also not powered to detect potential rare side effects of BNT162b2 in 5-to-11-year-olds. However, the safety of BNT162b2 observed in the study com-





# A CRITICAL EYE BACK ON THE SEP 15 2021 REPORT





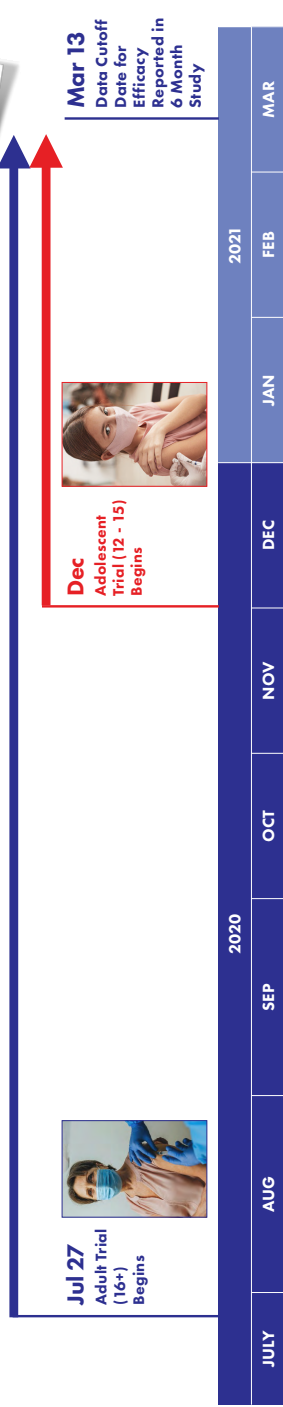


PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# 6 MONTH DATA MANIPULATION MIXED COHORTS

Pfizer took the results from their adult trial, which started July 27, 2020, and then added the results from the 12 - 15 year olds' trial, **despite the fact that the adolescent trial started four months later.**

Since it's well known that the efficacy of the inoculations wanes over time, **this gives a false boost to the efficacy numbers.** The efficacy for these two cohorts should have been reported separately, not presented as one combined result. Without this boost, their efficacy number would likely have fallen.







# PFIZER TRIALS DID NOT PROVE SAFETY THEY PROVED HARM

## ILLNESS

	BNT162b2	Placebo	Risk Change
<b>Efficacy</b> (Meaning number of people diagnosed with COVID-19.)	77	850	<b>-91%</b>
<b>Related Adverse Event</b> (Meaning an investigator has assessed it as related to the BNT162b2 injection.)	5,241	1,311	<b>+300%</b>
<b>Any Severe Adverse Event</b> (Interferes significantly with normal function.)	262	150	<b>+75%</b>
<b>Any Serious Adverse Event</b> (Involves visit to ER or hospitalization.)	127	116	<b>+10%</b>

## DEATHS

BNT162b2	Placebo
20	14

**These are the results of Pfizer's own randomized control trial.  
LEVEL 1 EVIDENCE OF HARM.**

# HOW THIS IS PLAYING OUT IN THE REAL WORLD





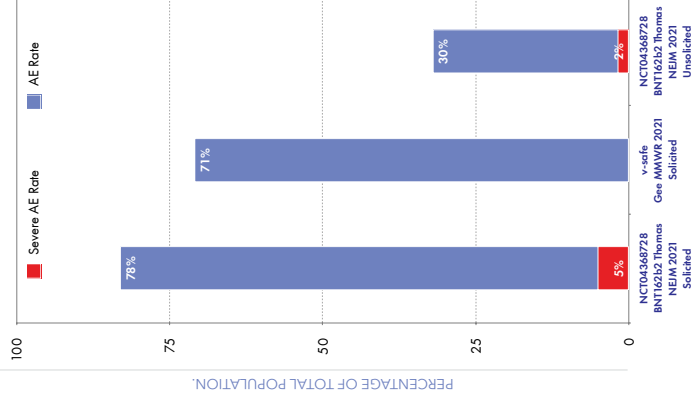
## ROLL OUT SURVEILLANCE YOU DON'T FIND WHAT YOU DON'T LOOK FOR

There is a dramatic difference between passive vs active monitoring of adverse events

1. When participants were **actively** followed for adverse events (AEs) in the trials, high percentages of adverse events were reported.
2. Once the vaccine was rolled out at the population level, **passive** surveillance was used with Health Canada, VAERS or the European Yellow Card system.

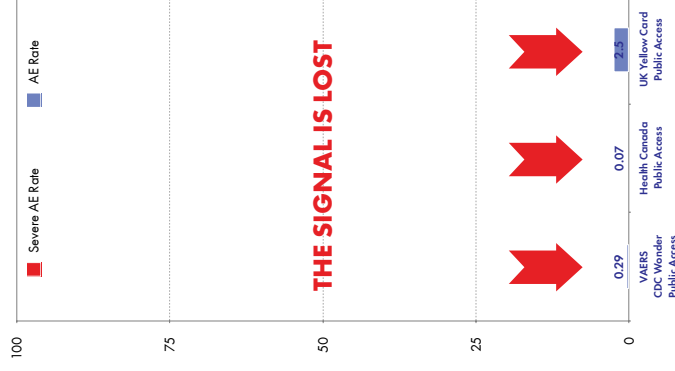
When that happened, the **signal was completely lost**.

### ACTIVE SURVEILLANCE OF TRIAL PARTICIPANTS



VS

### PASSIVE SURVEILLANCE OF POPULATION ROLL OUT



THE SIGNAL IS LOST





# RISING INCIDENTS OF HEART ISSUES IN YOUNG PEOPLE

Ontario Public Health is well aware of this, as they published a [report](#) on it, but they seem inconsistent in their concerns.

- On Sep 29, 2021, Ontario Public Health recommended **young men 18-24** not take the **Moderna shot**, because of a **1 in 5,000 risk of myocarditis**. They suggested **Pfizer shot** instead, which has a **1 in 28,000 risk of myocarditis**.
- But as recently as May 8, 2021, **Ontario had stopped the Astra Zeneca shot because of a 1 in 60,000 risk of clotting side effects**, which was considered too high.
- **Their priorities are inconsistent.**

Ontario

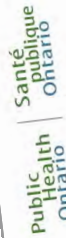
## More than 100 Ontario youth sent to hospital for vaccine-related heart problems: Report

There were 54 persons aged 25-39 included in the tally and 44 persons aged 40 and over

Anthony Furey  
Sep 03, 2021 · September 3, 2021 · 2 minute read · 314 Comments



Moderna coronavirus disease (COVID-19) vaccine labels are seen in this photo. PHOTO BY DAVID ROVIC/REUTERS



### ENHANCED EPIDEMIOLOGICAL SUMMARY

Myocarditis and Pericarditis Following Vaccination with COVID-19 mRNA Vaccines in Ontario: December 13, 2020 to September 4, 2021

#### Purpose

This report summarises reports of myocarditis/pericarditis that have been reported as adverse events following receipt of a COVID-19 mRNA vaccine. Data on myocarditis/pericarditis following COVID-19 mRNA vaccines are

Pfizer's inoculations for COVID-19 / MORE HARM THAN GOOD

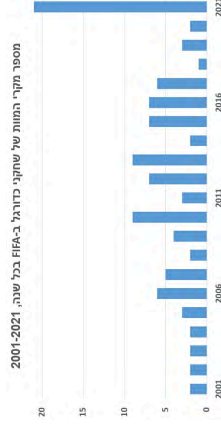
# THIS IS NOT NORMAL

A German news site put together a list of over 75 known cases of athletes collapsing - and even dying - in the last 5 months.

<https://report24.news/deb-13/afirm-langue-lite-ploberlich-wartbarbame-codetz-schwendt-omke-csp-afthar/>

An Israeli news site analyzed the number of sudden deaths "on the pitch" of members of the International Football Association (FIFA) over the past 20 years.

The average number of FIFA sudden deaths between 2000 - 2020 was 4.2. In 2021, it was 21.



<https://www.ynet.co.il/2021/05/22/1.1374900>

**The Guardian**  
for 20th years

Barcelona  
Sergio Agüero out for three months following 'cardiological evaluation'

- Striker admitted to hospital after draw with Alavés
- 33-year-old to undergo 'diagnostic and therapeutic process'

**PROFESIONTECHNICA**  
Isaiah Harris – Pfizer Severe Adverse Reaction

Isaiah Harris Aged 18 – Pfizer May 2021  
Severe Adverse Reaction: Myocarditis (Inflammation of the Heart) Resulting in a Heart Attack

**Health News** / Vaccine Injury Stories / Vaccines  
Ernest Ramirez

Grieving Father Ernest Ramirez Shares Heartbreaking Story of His Teen Son's Death 5 Days After Pfizer Vaccine

**SN**  
US SPORTS / Men's Basketball / Women's Basketball / Men's Hockey / Women's Hockey

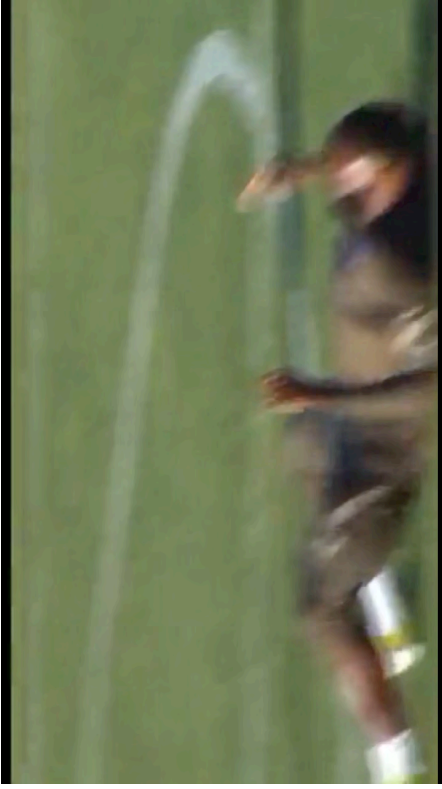
Gees football player Francis Perron dies shortly after season opener

EN MÉMOIRE DE  
IN MEMORY OF  
**FRANCIS PERRON**  
1996 - 2021



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## THIS IS SUPPOSED TO BE RARE



<https://rumble.com/vpnxkr-are-these-side-effects-extremely-rare.html>



# PFIZER'S POST MARKETING PHARMACOVIGILANCE REPORT

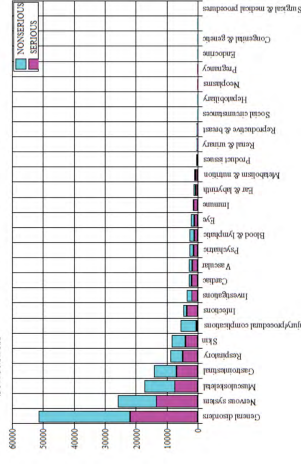
- On Nov 17, 2021, the FDA released the first batch of what will ultimately be **329,000 pages** they were ordered by a court to provide to satisfy a Freedom of Information request by a group called [Public Health and Medical Professionals for Transparency](#) who want access to the **data used by the FDA to approve Pfizer's COVID-19 inoculations**. (The FDA asked in court to have over 50 years to release the documents.)
- One **post marketing pharmacovigilance report** submitted to the FDA, where Pfizer tracked real world adverse events occurring in the first 2.5 months after Emergency Use Authorization, was particularly disturbing.
  - + **Over 1,200 deaths**
  - + **Over 25,000 nervous system adverse events**
  - + Under "Safety concerns" Pfizer listed **Anaphylaxis** and **Vaccine-Associated Enhanced Disease**
- **This document should be incriminating for any agency who saw it and called these inoculations "safe."**

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

Characteristic	Number of Cases	Percent of Total (N=2386)
Gender		
Male	9182	
Female	2914	
Age (years)		
18-30	179	
31-50	4953	
51-60	1386	
61-70	3998	
71-80	2314	
81-90	6816	
Unknown	15582	
Case outcome		
Recovered/Recovering	11261	
Not recovered at the time of report	1223	
Unknown	9460	

a. in 46 cases reported age was <10-year-old and in 34 cases <12-year-old.

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



3.1.2. Summary of Safety Concerns in the US Pharmacovigilance Plan

Table 3. Safety concerns

Important identifier risks	Amplifiers
Important potential risks	Vaccine-Associated Enhanced Disease (VAED), Including Vaccine-associated Enhanced Respiratory Disease (VAERD)
Missing information	Use in Pregnancy and Lactation Vaccine Effectiveness Vaccine Efficacy

# CONSIDERABLE EVIDENCE OF CONFLICT OF INTEREST





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



**PFIZER IS  
MAKING  
BILLIONS  
\$33.5B+ in 2021 alone.**

When the incentive is such an astronomical sum of money, it only makes sense to **ensure rigorous oversight** of the process and to **ensure as many safeguards as possible** are in place.

Their agenda is **their shareholders and their bottom line**, not public health.

Forbes

# Pfizer Expects \$33.5 Billion In Vaccine Revenue In 2021



Albert Bourla, CEO of Pfizer, photographed in June 2020. JANEL TOPPIN FOR FORBES

**B**ioTech giant Pfizer expects to generate \$33.5 billion in Covid-19 vaccine sales in 2021, up from previous estimates of \$26 billion, according to its second quarter earnings reports. These projections are based on the 2.1 billion doses of the Pfizer/BioNTech vaccine which the company expects to manufacture and deliver by the end of the year.

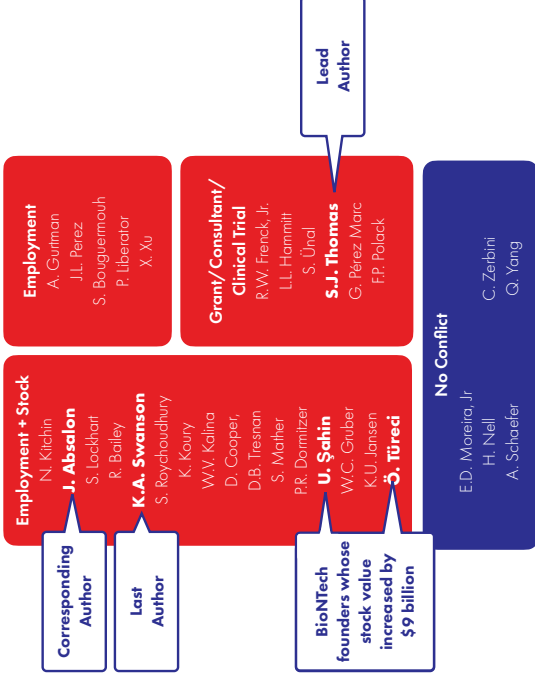






# CONFLICTS OF INTEREST AMONG PFIZER REPORT AUTHORS


## 6 MONTH REPORT AUTHORS



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD



# THE CDC HAS REDEFINED "VACCINE" TO SUIT POLITICAL & PHARMACEUTICAL INTERESTS

For many years	Jul 27, 2021	Aug 18, 2021	Starting Sep 2, 2021
<p><b>CDC Definition of VACCINE</b></p> <p>"A product that stimulates a person's immune system to produce immunity to a specific disease, protecting the person from that disease."</p>	<p>Head of CDC Rochelle Walensky went on CNN and admitted the <u>COVID-19 vaccines do not provide immunity</u> - they don't stop people from catching or transmitting COVID-19.</p> 	<p>Joe Biden announced booster shots for all Americans.</p> 	<p><b>CDC Definition of VACCINE CHANGED</b></p> <p><i>"A preparation that is used to stimulate the body's immune response against diseases."</i></p> <p><b>This looks like fraud.</b></p>



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# THE MEDIA HAS BEEN CAPTURED



<https://rumble.com/voz64j-brought-to-you-by-pfizer.html>



PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## **THIS IS NO WAY TO MANAGE A SUPPLIER**

Pfizer has been **indemnified for damages** in case their inoculations hurt and kill people, and Pfizer **profits to the tune of billions** if the trials are successful.

**No reasonable, responsible person would have given Pfizer carte blanche in such a situation.**

**Instead, you would engage in rigorous oversight and hold them to the highest scientific standards.** This was not done.





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

## THE INOCULATIONS SHOULD BE WITHDRAWN IMMEDIATELY

- It's clear that Pfizer - and the agencies overseeing their trials - failed to follow established, high quality safety and efficacy protocols right from the beginning.
- We have presented **Level 1 evidence of harm from Pfizer's own trial data**. Any government which has approved these inoculations, much less mandated them, **knew or should have known from the available data that harm would be caused to its citizens**.
- Any government that approved this medical intervention for its citizens should have ensured that the trial had used the **appropriate clinical endpoints and high quality safety science**.
- **Any government official who possesses this evidence and continues to allow its citizens to be inoculated with a toxic agent is, at the very least, negligent.**





PFIZER'S INOCULATIONS FOR COVID-19 / MORE HARM THAN GOOD

# RECOMMENDED READING/VIEWING

## PUBLISHED PAPERS REFUTING PFIZER INOCULATIONS

- **Why Are We Vaccinating Children Against COVID-19?** <https://www.sciencedirect.com/science/article/pii/S221475002100161X>
- **US COVID-19 Vaccines Proven to Cause More Harm than Good Based on Pivotal Clinical Trial Data Analyzed Using the Proper Scientific Endpoint, "All Cause Severe Morbidity"** <https://www.scisourcepub.com/pdfs/us-covid-19-vaccines-proven-to-cause-more-harm-than-good-based-on-pivotal-clinical-trial-data-analyzed-using-the-proper-scientific-1811.pdf>
- **Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine** <https://www.nejm.org/doi/full/10.1056/NEJMoa2034572>
- **FDA Briefing Document, Dec 10, 2020** <https://www.fda.gov/media/144245/download>
- **Safety and Efficacy of the BNT162b2 mRNA Covid-19 Vaccine through 6 Months** <https://www.nejm.org/doi/full/10.1056/NEJMoa210345>
- **The 6 Month Supplementary Appendix** [https://www.nejm.org/doi/suppl/10.1056/NEJMoa210345/suppl\\_file](https://www.nejm.org/doi/suppl/10.1056/NEJMoa210345/suppl_file)

## BRITISH MEDICAL JOURNAL

- **Covid-19: Researcher blows the whistle on data integrity issues in Pfizer's vaccine trial** <https://www.bmj.com/content/375/bmj.n2635>
- **Ontario Public Health Epidemiological Summary**
- **Myocarditis and Pericarditis Following Vaccination with COVID-19 mRNA Vaccines in Ontario: December 13, 2020 to September 4, 2021** [https://www.publichealthontario.ca/\\_/media/documents/new/epi/covid-19-myocarditis-pericarditis-vaccines-epi.pdf?sc\\_lang=en](https://www.publichealthontario.ca/_/media/documents/new/epi/covid-19-myocarditis-pericarditis-vaccines-epi.pdf?sc_lang=en)

## SHORT VIDEOS

- **Informed Consent - It's Your Right (3 minutes)** <https://vimeo.com/469434304>
- **Brought to You by Pfizer (1 minute)** <https://vimeo.com/469644brought-to-you-by-pfizer.html>
- **Why Do We Need Vaccine Passports? (2 minutes)** <https://vimeo.com/469644why-do-we-need-vaccine-passports.html>
- **COVID-19 Vaccines and D-Dimer levels (9 minutes)** <https://vimeo.com/469644-rochagan-bilal-on-blowing-the-whistle-on-covid-19-vaccines-and-d-dimer-level.html>
- **How Reliable is the PCR Test? (2 minutes)** <https://youtu.be/gLZ75JmRIM4>



# WE NEED YOU TO HOLD THEM ACCOUNTABLE

- This evidence is a tool you can use. It represents a real opportunity to hold our leaders accountable as it is not opinion, or modelling, or real world evidence that can be dismissed or manipulated, but LEVEL 1 EVIDENCE from a randomized control trial. As such, it has high evidentiary value.
- We're asking that you call your MP and MPP and that you ask for a 1 hour meeting. Preferably in person, but Zoom will work too.
- During the meeting, play them the video and provide them with the PDF version. Ask them questions, like whether or not they were aware of all the issues with the Pfizer trial. Or what they plan to do now that they are. Get them to agree to a follow up meeting where they will provide you with answers.
- Share this video with friends and family. Have group viewing sessions on Zoom and discuss it.
- Share this video and the PDF on social media. When you do, please use the hashtags #CCCA and #MoreHarmThanGood
- Please join our mailing list at [www.canadiancovidcarealliance.org](http://www.canadiancovidcarealliance.org) and we will update you with additional evidence as we have it.
- Follow us on social media. This linktree has all our social accounts.
- This presentation is available in PDF and video format on our website at [www.canadiancovidcarealliance.org](http://www.canadiancovidcarealliance.org)

THE PFIZER INOCULATIONS FOR COVID-19

# MORE HARM THAN GOOD



Contact us  
[info@canadiancovidcarealliance.org](mailto:info@canadiancovidcarealliance.org)  
[www.canadiancovidcarealliance.org](http://www.canadiancovidcarealliance.org)